QUANTITATIVE ECONOMICS with Julia

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Note: You are currently viewing an automatically generated PDF version of our online lectures, which are located at

http://quant-econ.net

Please visit the website for more information on the aims and scope of the lectures and the two language options (Julia or Python). This PDF is generated from a set of source files that are orientated towards the website and to HTML output. As a result, the presentation quality can be less consistent than the website.
This first part of the course provides a relatively fast-paced introduction to the Julia programming language

1.1 Setting up Your Julia Environment

Contents

- Setting up Your Julia Environment
  - Overview
  - First Steps
  - IJulia
  - The QuantEcon Library
  - Exercises

Overview

In this lecture we will cover how to get up and running with Julia

Topics:

1. Installation
2. Interactive Julia sessions
3. Running sample programs
4. Installation of libraries, including the Julia code that underpins these lectures

First Steps

Installation  The first thing you will want to do is install Julia

The best option is probably to install the current release from the download page

  - Read through any download and installation instructions specific to your OS on that page
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- Unless you have good reason to do otherwise, choose the current release rather than nightly build and the platform specific binary rather than source

Assuming there were no problems, you should now be able to start Julia either by

- navigating to Julia through your menus or desktop icons (Windows, OSX), or
- opening a terminal and typing `julia` (Linux)

Either way you should now be looking at something like this (modulo your operating system — this is a Linux machine)

The program that’s running here is called the Julia REPL (Read Eval Print Loop) or Julia interpreter

Let’s try some basic commands:

The Julia interpreter has the kind of nice features you expect from a modern REPL

For example,
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- Pushing the up arrow key retrieves the previously typed command
- If you type `?` the prompt will change to `help?>` and give you access to online documentation

You can also type `;` to get a shell prompt, at which you can enter shell commands

(Here `ls` is a UNIX style command that lists directory contents — your shell commands depend on your operating system)

From now on instead of showing terminal images we’ll show interactions with the interpreter as follows

```julia
julia> x = 10
10

julia> 2 * x
20
```

**Installing Packages** Julia includes many useful tools in the base installation
1.1. SETTING UP YOUR JULIA ENVIRONMENT

However, you’ll quickly find that you also have need for at least some of the many external Julia code libraries.

Fortunately these are very easy to install using Julia’s excellent package management system. For example, let’s install DataFrames, which provides useful functions and data types for manipulating data sets.

```julia
julia> Pkg.add("DataFrames")
```

Assuming you have a working Internet connection this should install the DataFrames package. If you now type Pkg.status() you’ll see DataFrames and its version number. To pull the functionality from DataFrames into the current session we type using DataFrames.

```julia
julia> using DataFrames
```

Now let’s use one of its functions to create a data frame object (something like an R data frame, or a spreadsheet).

```julia
julia> df = DataFrame(x1=[1, 2], x2=["foo", "bar"])
2x2 DataFrame
<table>
<thead>
<tr>
<th>Row</th>
<th>x1</th>
<th>x2</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>1</td>
<td>&quot;foo&quot;</td>
</tr>
<tr>
<td>2</td>
<td>2</td>
<td>&quot;bar&quot;</td>
</tr>
</tbody>
</table>
```

One quick point before we move on: Running

```julia
julia> Pkg.update()
```

will update your installed packages and also update local information on the set of available packages.

It’s a good idea to make a habit of this.

Running Julia Scripts

Julia programs (or “scripts”) are text files containing Julia code, typically with the file extension .jl.

Suppose we have a Julia script called test_script.jl that we wish to run.

The contents of the file is as follows

```julia
for i in 1:3
    println("i = \$i")
end
```

If that file exists in the present working directory we can run it with include("test_script.jl")

(To see what your present working directory is in a Julia session type pwd().)

Here’s an example, where test_script.jl sits in directory /home/john/temp.
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```julia
julia> pwd()
"/home/john/temp"

julia> include("test_script.jl")
i = 1
i = 2
i = 3
```

(Of course paths to files will look different on different operating systems)

If the file is not in your `pwd` you can run it by giving the full path — in the present case

```julia
julia> include("/home/john/temp/test_script.jl")
```

Alternatively you can change your `pwd` to the location of the script

```julia
julia> cd("/home/john/temp")
```

and then run using `include("test_script.jl")` as before

**Editing Julia Scripts**  
Hopefully you can now run Julia scripts  
You also need to know how to edit them

**Text Editors**  
Nothing beats the power and efficiency of a good text editor for working with program text

At a minimum, such an editor should provide

- syntax highlighting for the languages you want to work with
- automatic indentation
- text manipulation basics such as search and replace, copy and paste, etc.

There are many text editors that speak Julia, and a lot of them are free

Suggestions:

**Sublime Text** is a modern, popular and highly regarded text editor with a relatively moderate learning curve (not free but trial period is unlimited)

**Emacs** is a high quality free editor with a sharper learning curve

Finally, if you want an outstanding free text editor and don’t mind a seemingly vertical learning curve plus long days of pain and suffering while all your neural pathways are rewired, try **Vim**

**IDEs**  
IDEs are Integrated Development Environments — they combine an interpreter and text editing facilities in the one application

For Julia one nice option is **Juno**

Alternatively there’s **IJulia**, which is a little bit different again but has some great features that we now discuss

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IJulia

To work with Julia in a scientific context we need at a minimum

1. An environment for editing and running Julia code
2. The ability to generate figures and graphics

A very nice option that provides these features is IJulia

As a bonus, IJulia also provides

- Nicely formatted output in the browser, including tables, figures, animation, video, etc.
- The ability to mix in formatted text and mathematical expressions between cells
- Functions to generate PDF slides, static html, etc.

Whether you end up using IJulia as your primary work environment or not, you’ll find learning about it an excellent investment

Installing IJulia  IJulia is built on top of the IPython notebook

The IPython notebook started off as a Python tool but is in the process of being re-born as a language agnostic scientific programming environment (see Jupyter)

The IPython notebook in turn has a range of dependencies that it needs to work properly

At present the easiest way to install all of these at once is to install the Anaconda Python distribution

Installing Anaconda  Installing Anaconda is straightforward: download the binary and follow the instructions

If you are asked during the installation process whether you’d like to make Anaconda your default Python installation, say yes — you can always remove it later

Otherwise you can accept all of the defaults

Note that the packages in Anaconda update regularly — you can keep up to date by typing conda update anaconda in a terminal

Installing IJulia  Just run

    julia> Pkg.add("IJulia")

Other Requirements  We’ll be wanting to produce plots and while there are several options we’ll start with PyPlot

    julia> Pkg.add("PyPlot")
Finally, since IJulia runs in the browser it might now be a good idea to update your browser. One good option is to install a free modern browser such as Chrome or Firefox. In our experience Chrome plays well with IJulia.

**Getting Started**  To start IJulia in the browser, open up a terminal (or cmd in Windows) and type

```
ipython notebook --profile=julia
```

Here’s an example of the kind of thing you should see.

In this case the address is `localhost:8998/tree`, which indicates that the browser is communicating with a Julia session via port 8998 of the local machine.

The page you are looking at is called the “dashboard.” From here you can now click on New Notebook and see something like this.

The notebook displays an **active cell**, into which you can type Julia commands.

**Notebook Basics**  Notice that in the previous figure the cell is surrounded by a green border. This means that the cell is in **edit mode**.
As a result, you can type in Julia code and it will appear in the cell

When you’re ready to execute these commands, hit Shift-Enter instead of the usual Enter

---

Modal Editing  The next thing to understand about the IPython notebook is that it uses a modal editing system

This means that the effect of typing at the keyboard depends on which mode you are in

The two modes are

1. Edit mode
   - Indicated by a green border around one cell, as in the pictures above
   - Whatever you type appears as is in that cell

2. Command mode
   - The green border is replaced by a grey border
   - Key strokes are interpreted as commands — for example, typing b adds a new cell below the current one
Switching modes

- To switch to command mode from edit mode, hit the Esc key
- To switch to edit mode from command mode, hit Enter or click in a cell

The modal behavior of the IPython notebook is a little tricky at first but very efficient when you get used to it

For more details on the mechanics of using the notebook, see here

Plots As discussed above, IJulia integrates nicely with the plotting package PyPlot.jl

PyPlot in turn relies on the excellent Python graphics library Matplotlib

Once you have PyPlot installed you can load it via using PyPlot

We’ll discuss plotting in detail later on but for now let’s just make sure that it works

Here’s a sample program you can run in IJulia

```julia
using PyPlot

n = 50
srand(1)
x = rand(n)
y = rand(n)
area = pi .* (15 .* rand(n)).^2 # 0 to 15 point radiuses
scatter(x, y, s=area, alpha=0.5)
```

Don’t worry about the details for now — let’s just run it and see what happens

The easiest way to run this code is to copy and paste into a cell in the notebook and Shift-Enter

This is what you should see

Working with the Notebook In this section we’ll run you quickly through some more IPython notebook essentials — just enough so that we can press ahead with programming

Tab Completion A simple but useful feature of IJulia is tab completion

For example if you type `rep` and hit the tab key you’ll get a list of all commands that start with `rep`

IJulia offers up the possible completions

This helps remind you of what’s available and saves a bit of typing

On-Line Help To get help on the Julia function such as `repmat`, enter `help(repmat)`

Documentation should now appear in the browser

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Other Content  In addition to executing code, the IPython notebook allows you to embed text, equations, figures and even videos in the page.

For example, here we enter a mixture of plain text and LaTeX instead of code.

Next we \texttt{Esc} to enter command mode and then type \texttt{m} to indicate that we are writing \texttt{Markdown}, a mark-up language similar to (but simpler than) \texttt{LaTeX}.

(You can also use your mouse to select \texttt{Markdown} from the Code drop-down box just below the list of menu items.)

Now we \texttt{Shift + Enter} to produce this:

\textbf{Shell Commands}  You can execute shell commands (system commands) in IJulia by prepending a semicolon.

For example, \texttt{;ls} will execute the UNIX style shell command \texttt{ls}, which — at least for UNIX style operating systems — lists the contents of the present working directory.

These shell commands are handled by your default system shell and hence are platform specific.

\textbf{Working with Files}  To run an existing Julia file using the notebook we can either
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Out[1]: PyObject <matplotlib.collections.PathCollection object at 0x7f1a00005d90>

In [4]: help(repmat)

Base.repmat(A, n, n)

Construct a matrix by repeating the given matrix \( n \) times in
dimension 1 and \( n \) times in dimension 2.

Given subsets \( A_1, A_2, \ldots \) of \( X \), the set \( \bigcup_n A_n \) is defined by

\[
\bigcup_n A_n := \{ x \in X : x \in A_m \text{ for some } n \}
\]

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1. copy and paste the contents into a cell in the notebook, or
2. use `include("filename")` in the same manner as for the Julia interpreter discussed above

More sophisticated methods for working with files are under active development and should be on-line soon

Sharing Notebooks  Notebook files are just text files structured in JSON and typically ending with .ipynb

A notebook can easily be saved and shared between users — you just need to pass around the ipynb file

To open an existing ipynb file, import it from the dashboard (the first browser page that opens when you start IPython notebook) and run the cells or edit as discussed above

nbviewer  The IPython organization has a site for sharing notebooks called nbviewer

The notebooks you see there are static HTML representations of notebooks

However, each notebook can be downloaded as an ipynb file by clicking on the download icon at the top right of its page

Once downloaded you can open it as a notebook, as we discussed just above

The QuantEcon Library

The QuantEcon library is a community based code library containing open source code for quantitative economic modeling

Thanks to the heroic efforts of Spencer Lyon and some of his collaborators, this now includes a Julia version

You can install this package through the usual Julia package manager:

```
julia> Pkg.add("QuantEcon")
```

For example, the following code creates a discrete approximation to an AR(1) process

```
julia> using QuantEcon; tauchen

julia> tauchen(4, 0.9, 1)
([-6.88247,-2.29416,2.29416,6.88247],
4x4 Array{Float64,2}:
0.945853 0.0541468 2.92863e-10 0.0
0.00580845 0.974718 0.0194737 1.43534e-11
1.43534e-11 0.0194737 0.974718 0.00580845
2.08117e-27 2.92863e-10 0.0541468 0.945853)
```

We’ll learn much more about the library as we go along
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Installing via GitHub  You can also grab a copy of the files in the QuantEcon library directly by downloading the zip file — try clicking the “Download ZIP” button on the main page. Alternatively, you can get a copy of the repo using Git.

For more information see Exercise 1.

Exercises

Exercise 1  If you haven’t heard, Git is a version control system — a piece of software used to manage digital projects such as code libraries.

In many cases the associated collections of files — called repositories — are stored on GitHub. GitHub is a wonderland of collaborative coding projects.

Git is the underlying software used to manage these projects.

Git is an extremely powerful tool for distributed collaboration — for example, we use it to share and synchronize all the source files for these lectures.

There are two main flavors of Git:

1. the plain vanilla command line version
2. the point-and-click GUI versions
   - GUI style Git for Windows
   - GUI style Git for Mac

As an exercise, try getting a copy of the QuantEcon repository using Git.

You can try the GUI options above or install the plain command line Git.

If you’ve installed the command line version, open up a terminal and enter:

```
git clone https://github.com/QuantEcon/QuantEcon.jl
```

This is just `git clone` in front of the URL for the repository.

Even better, sign up to GitHub — it’s free.

Look into ‘forking’ GitHub repositories.

(Loosely speaking, forking means making your own copy of a GitHub repository, stored on GitHub.)

Try forking the QuantEcon repository for the course.

Now try cloning it to some local directory, making edits, adding and committing them, and pushing them back up to your forked GitHub repo.

For reading on these and other topics, try:

- The official Git documentation
- Reading through the docs on GitHub
1.2 An Introductory Example

Overview

We’re now ready to start learning the Julia language itself. Our approach is aimed at those who already have at least some knowledge of programming — perhaps experience with Python, MATLAB, R, C or similar. In particular, we assume you have some familiarity with fundamental programming concepts such as:

- variables
- loops
- conditionals (if/else)

If you have no such programming experience we humbly suggest you try Python first. Python is a great first language and, more importantly, there are many, many introductory treatments. In fact our treatment of Python is much slower than our treatment of Julia, especially at the start. Once you are comfortable with Python you’ll find the leap to Julia is easy.

Approach  In this lecture we will write and then pick apart small Julia programs. At this stage the objective is to introduce you to basic syntax and data structures. Deeper concepts—how things work—will be covered in later lectures. Since we are looking for simplicity the examples are a little contrived.

Other References  The definitive reference is Julia’s own documentation. The manual is thoughtfully written but also quite dense (and somewhat evangelical). The presentation in this and our remaining lectures is more of a tutorial style based around examples.
Example: Plotting a White Noise Process

To begin, let’s suppose that we want to simulate and plot the white noise process $\epsilon_0, \epsilon_1, \ldots, \epsilon_T$, where each draw $\epsilon_t$ is independent standard normal.

In other words, we want to generate figures that look something like this:

This is straightforward using the PyPlot library we installed earlier.

```julia
using PyPlot
ts_length = 100
epsilon_values = randn(ts_length)
plot(epsilon_values, "b-")
```

You should be able to run that code either in IJulia or in the standard REPL (the basic interpreter).

In brief,

- `using PyPlot` makes the functionality in PyPlot available for use
- In particular, it pulls the names exported by the `PyPlot` module into the global scope
- One of these is `plot()`, which in turn calls the `plot` function from Matplotlib
- `randn()` is a Julia function from the standard library for generating standard normals

**Importing Functions** The effect of the statement `using PyPlot` is to make all the names exported by the `PyPlot` module available in the global scope.

If you prefer to be more selective you can replace `using PyPlot` with `import PyPlot: plot`.

Now only the `plot` function is accessible.
Since our program uses only the plot function from this module, either would have worked in the previous example.

**Arrays**  The function call `epsilon_values = randn(ts_length)` creates one of the most fundamental Julia data types: an array

```
 julia> typeof(epsilon_values)
 Array{Float64,1}

 julia> epsilon_values
 100-element Array{Float64,1}:
  -0.908823
  -0.759142
  -1.42078
   0.792799
   0.577181
   1.74219
  -0.912529
   1.06259
   0.5766
  -0.0172788
  -0.591671
  -1.02792
  ...
  -1.29412
  -1.12475
   0.437858
  -0.709243
  -1.96053
   1.31092
   1.19819
   1.54028
  -0.246204
  -1.23305
  -1.16484
```

The information from `typeof()` tells us that `epsilon_values` is an array of 64 bit floating point values, of dimension 1.

Julia arrays are quite flexible — they can store heterogeneous data for example

```
 julia> x = [10, "foo", false]
 3-element Array{Any,1}:
   10
   "foo"
  false
```

Notice now that the data type is recorded as `Any`, since the array contains mixed data.

The first element of `x` is an integer

```
 julia> typeof(x[1])
 Int64
```
The second is a string

```julia
typeof(x[2])
ASCIIString (constructor with 2 methods)
```

The third is the boolean value `false`

```julia
typeof(x[3])
Bool
```

Notice from the above that
- array indices start at 1 (unlike Python, where arrays are zero-based)
- array elements are referenced using square brackets (unlike MATLAB and Fortran)

Julia contains many functions for acting on arrays — we’ll review them later.

For now here’s several examples, applied to the same list `x = [10, "foo", false]`

```julia
length(x)
3
```

```julia
pop!(x)
false
```

```julia
x
2-element Array{Any,1}:
10
"foo"
```

```julia
push!(x, "bar")
3-element Array{Any,1}:
10
"foo"
"bar"
```

```julia
x
3-element Array{Any,1}:
10
"foo"
"bar"
```

The first example just returns the length of the list.

The second, `pop!()`, pops the last element off the list and returns it.

In doing so it changes the list (by dropping the last element).

Because of this we call `pop!` a **mutating method**.

It’s conventional in Julia that mutating methods end in `!` to remind the user that the function has other effects beyond just returning a value.

The function `push!()` is similar, expect that it appends its second argument to the array.
For Loops  Although there’s no need in terms of what we wanted to achieve with our program, for the sake of learning syntax let’s rewrite our program to use a for loop.

```julia
using PyPlot
ts_length = 100
epsilon_values = Array(Float64, ts_length)
for i in 1:ts_length
    epsilon_values[i] = randn()
end
plot(epsilon_values, "b-")
```

Here we first declared `epsilon_values` to be an empty array for storing 64 bit floating point numbers. The for loop then populates this array by successive calls to `randn()`.

- Called without an argument, `randn()` returns a single float.

Like all code blocks in Julia, the end of the for loop code block (which is just one line here) is indicated by the keyword `end`.

The word `in` from the for loop can be replaced by symbol `= `

The expression `1:ts_length` creates an iterator that is looped over — in this case the integers from 1 to `ts_length`.

Iterators are memory efficient because the elements are generated on the fly rather than stored in memory.

In Julia you can also loop directly over arrays themselves, like so:

```julia
words = ["foo", "bar"]
for word in words
    println("Hello ", word)
end
```

The output is:

Hello foo
Hello bar

While Loops  The syntax for the while loop contains no surprises.

```julia
using PyPlot
ts_length = 100
epsilon_values = Array(Float64, ts_length)
i = 1
while i <= ts_length
    epsilon_values[i] = randn()
i = i + 1
end
plot(epsilon_values, "b-")
```

The next example does the same thing with a condition and the `break` statement.
using PyPlot

ts_length = 100
epsilon_values = Array(Float64, ts_length)
i = 1
while true
    epsilon_values[i] = randn()
i = i + 1
    if i == ts_length
        break
    end
end
plot(epsilon_values, "b-")

User-Defined Functions  For the sake of the exercise, let’s now go back to the for loop but restructure our program so that generation of random variables takes place within a user-defined function.

using PyPlot

function generate_data(n)
    epsilon_values = Array(Float64, n)
    for i = 1:n
        epsilon_values[i] = randn()
    end
    return epsilon_values
end

A Slightly More Useful Function  Of course the function generate_data is completely contrived

We could just write the following and be done

ts_length = 100
data = generate_data(ts_length)
plot(data, "b-")

Let’s make a slightly more useful function

This function will be passed a choice of probability distribution and respond by plotting a histogram of observations.

In doing so we’ll make use of the Distributions package.  

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Here’s the code

```julia
using PyPlot
using Distributions

function plot_histogram(distribution, n)
    epsilon_values = rand(distribution, n)  # n draws from distribution
    PyPlot.plt.hist(epsilon_values)
end

lp = Laplace()
plot_histogram(lp, 500)
```

The resulting figure looks like this

Let’s have a casual discussion of how all this works while leaving technical details for later in the lectures

First, `lp = Laplace()` creates an instance of a data type defined in the Distributions module that represents the Laplace distribution

The name `lp` is bound to this object

When we make the function call `plot_histogram(lp, 500)` the code in the body of the function `plot_histogram` is run with

- the name `distribution` bound to the same object as `lp`
- the name `n` bound to the integer 500
A Mystery  Now consider the function call \texttt{rand(distribution, n)}

This looks like something of a mystery.

The function \texttt{rand()} is defined in the base library such that \texttt{rand(n)} returns \(n\) uniform random variables on \([0, 1)\).

\begin{verbatim}
 julia> rand(3)
 3-element Array{Float64,1}:
 0.856817
 0.981502
 0.510947
\end{verbatim}

On the other hand, \texttt{distribution} points to a data type representing the Laplace distribution that has been defined in a third party package.

So how can it be that \texttt{rand()} is able to take this kind of object as an argument and return the output that we want?

The answer in a nutshell is \textbf{multiple dispatch}.

This refers to the idea that functions in Julia can have different behavior depending on the particular arguments that they’re passed.

Hence in Julia we can take an existing function and give it a new behavior by defining how it acts on a new type of object.

The interpreter knows which function definition to apply in a given setting by looking at the types of the objects the function is called on.

In Julia these alternative versions of a function are called \textbf{methods}.

A Small Problem  In many situations multiple dispatch provides a clean solution for resolving the correct action for a given function in a given setting.

You can see however that caution is sometimes required by looking at the line \texttt{PyPlot.plt.hist(epsilon_values)} from the code above.

A function called \texttt{hist()} exists in the standard library and is always available.

\begin{verbatim}
 julia> hist([5, 10, 15, 20])
 (0.0:5.0:20.0,[1,1,1,1])
\end{verbatim}

In addition, to maintain unified syntax with Matplotlib, the library \texttt{PyPlot} also defines its own version of \texttt{hist()}, for plotting.

Because both versions act on arrays, if we simply write \texttt{hist(epsilon_values)} the interpreter can’t tell which version to invoke.

In fact in this case it falls back to the first one defined, which is not the one defined by \texttt{PyPlot}.

This is the reason we need to be more specific, writing \texttt{PyPlot.plt.hist(epsilon_values)} instead of just \texttt{hist(epsilon_values)}.
Exercises

Exercise 1  Recall that $n!$ is read as “$n$ factorial” and defined as $n! = n \times (n - 1) \times \cdots \times 2 \times 1$

In Julia you can compute this value with `factorial(n)`

Write your own version of this function, called `factorial2`, using a for loop

Exercise 2  The binomial random variable $Y \sim Bin(n, p)$ represents

- number of successes in $n$ binary trials
- each trial succeeds with probability $p$

Using only `rand()` from the set of Julia’s built in random number generators (not the Distributions package), write a function `binomial_rv` such that `binomial_rv(n, p)` generates one draw of $Y$

Hint: If $U$ is uniform on $(0, 1)$ and $p \in (0, 1)$, then the expression $U < p$ evaluates to `true` with probability $p$

Exercise 3  Compute an approximation to $\pi$ using Monte Carlo

For random number generation use only `rand()`

Your hints are as follows:

- If $U$ is a bivariate uniform random variable on the unit square $(0, 1)^2$, then the probability that $U$ lies in a subset $B$ of $(0, 1)^2$ is equal to the area of $B$
- If $U_1, \ldots, U_n$ are iid copies of $U$, then, as $n$ gets large, the fraction that falls in $B$ converges to the probability of landing in $B$
- For a circle, area = $\pi \times \text{radius}^2$

Exercise 4  Write a program that prints one realization of the following random device:

- Flip an unbiased coin 10 times
- If 3 consecutive heads occur one or more times within this sequence, pay one dollar
- If not, pay nothing

Once again use only `rand()` as your random number generator

Exercise 5  Simulate and plot the correlated time series

$$x_{t+1} = \alpha x_t + \epsilon_{t+1} \quad \text{where} \quad x_0 = 0 \quad \text{and} \quad t = 0, \ldots, T$$

The sequence of shocks $\{\epsilon_t\}$ is assumed to be iid and standard normal

Set $T = 200$ and $\alpha = 0.9$
Exercise 6  To do the next exercise, you will need to know how to produce a plot legend
The following example should be sufficient to convey the idea

```python
using PyPlot

x = randn(100)
plot(x, "b-", label="white noise")
legend()

```

Running it produces a figure like so

![Plot example](image)

Now, plot three simulated time series, one for each of the cases $\alpha = 0$, $\alpha = 0.8$ and $\alpha = 0.98$

In particular, you should produce (modulo randomness) a figure that looks as follows

(The figure illustrates how time series with the same one-step-ahead conditional volatilities, as these three processes have, can have very different unconditional volatilities.)

Hints:

- If you call the `plot()` function multiple times before calling `show()`, all of the lines you produce will end up on the same figure
- If you omit the argument "b-" to the plot function, PyPlot will automatically select different colors for each line

Solutions

Solution notebook
1.3 Julia Essentials

Contents

- Julia Essentials
  - Overview
  - Common Data Types
  - Input and Output
  - Iterating
  - Comparisons and Logical Operators
  - User Defined Functions
  - Exercises
  - Solutions

Having covered a few examples, let’s now turn to a more systematic exposition of the essential features of the language.

Overview

Topics:

- Common data types
- Basic file I/O
- Iteration
- More on user-defined functions
• Comparisons and logic

Common Data Types

Like most languages, Julia language defines and provides functions for operating on standard data types such as

• integers
• floats
• strings
• arrays, etc...

Let’s learn a bit more about them

Primitive Data Types  A particularly simple data type is a Boolean value, which can be either true or false

```
julia> x = true
true

julia> typeof(x)
Bool

julia> y = 1 > 2  # Now y = false
false
```

Under addition, true is converted to 1 and false is converted to 0

```
julia> true + false
1

julia> sum([true, false, false, true])
2
```

The two most common data types used to represent numbers are integers and floats
(Computers distinguish between floats and integers because arithmetic is handled in a different way)

```
julia> typeof(1.0)
Float64

julia> typeof(1)
Int64
```

If you’re running a 32 bit system you’ll still see Float64, but you will see Int32 instead of Int64 (see the section on Integer types from the Julia manual)

Arithmetic operations are fairly standard
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```julia
julia> x = 2; y = 1.0
1.0

julia> x * y
2.0

julia> x^2
4

julia> y / x
0.5

Although the * can be omitted for multiplication between variables and numeric literals

```
```
julia> 2x - 3y
1.0
```

Also, you can use function (instead of infix) notation if you so desire

```julia
julia> +(10, 20)
30

julia> *(10, 20)
200
```

Complex numbers are another primitive data type, with the imaginary part being specified by `im`

```julia
julia> x = 1 + 2im
1 + 2im

julia> y = 1 - 2im
1 - 2im

julia> x * y  # Complex multiplication
5 + 0im
```

There are several more primitive data types that we’ll introduce as necessary

**Strings** A string is a data type for storing a sequence of characters

```julia
julia> x = "foobar"
"foobar"

```
```
```
julia> typeof(x)
ASCIIString (constructor with 2 methods)
```

You’ve already seen examples of Julia’s simple string formatting operations

```julia
julia> x = 10; y = 20
20

julia> "x = $x"
"x = 10"
```

---

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```julia
julia> "x + y = $(x + y)"
"x + y = 30"
```

To concatenate strings use *

```julia
julia> "foo" * "bar"
"foobar"
```

Julia provides many functions for working with strings

```julia
julia> s = "Charlie don't surf"
"Charlie don't surf"

julia> split(s)
3-element Array{SubString{ASCIIString},1}:
  "Charlie"
  "don't"
  "surf"

julia> replace(s, "surf", "ski")
"Charlie don't ski"

julia> split("fee,fi,fo", ",")
3-element Array{SubString{ASCIIString},1}:
  "fee"
  "fi"
  "fo"

julia> strip(" foobar ")  # Remove whitespace
"foobar"
```

Julia can also find and replace using regular expressions (see the documentation on regular expressions for more info)

```julia
julia> match(r"(\d+)", "Top 10")  # Find numerals in string
RegexMatch("10", 1="10")
```

Containers  Julia has several basic types for storing collections of data

We have already discussed arrays

A related data type is tuples, which can act like “immutable” arrays

```julia
julia> x = ("foo", "bar")
("foo", "bar")

julia> typeof(x)
(ASCIIString,ASCIIString)
```

An immutable object is one that cannot be altered once it resides in memory

In particular, tuples do not support item assignment:
This is similar to Python, as is the fact that the parenthesis can be omitted

```julia
julia> x = "foo", "bar"
("foo","bar")
```

Another similarity with Python is tuple unpacking, which means that the following convenient syntax is valid

```julia
julia> x = ("foo", "bar")
("foo","bar")
```

```julia
julia> word1, word2 = x
("foo","bar")
```

```julia
julia> word1
"foo"
```

```julia
julia> word2
"bar"
```

### Referencing Items

The last element of a sequence type can be accessed with the keyword `end`

```julia
julia> x = [10, 20, 30, 40]
4-element Array{Int64,1}:
  10
  20
  30
  40

julia> x[end]
40
```

```julia
julia> x[end-1]
30
```

To access multiple elements of an array or tuple, you can use slice notation

```julia
julia> x[1:3]
3-element Array{Int64,1}:
  10
  20
  30
```

```julia
julia> x[2:end]
3-element Array{Int64,1}:
  20
  30
  40
```

The same slice notation works on strings
julia> "foobar"[3:end]
"obar"

**Dictionaries**  Another container type worth mentioning is dictionaries.

Dictionaries are like arrays except that the items are named instead of numbered.

julia> d = "name" => "Frodo", "age" => 33
Dict{Any,Any} with 2 entries:
  "name" => "Frodo"
  "age" => 33

julia> d["age"]
33

The strings name and age are called the keys.

The objects that the keys are mapped to ("Frodo" and 33) are called the values.

They can be accessed via keys(d) and values(d) respectively.

**Input and Output**

Let’s have a quick look at reading from and writing to text files.

We’ll start with writing.

julia> f = open("newfile.txt", "w")  # "w" for writing
IOStream(<file newfile.txt>)

julia> write(f, "testing\n")  # \n for newline
7

julia> write(f, "more testing\n")
12

julia> close(f)

The effect of this is to create a file called newfile.txt in your present working directory with contents:

testing
more testing

We can read the contents of newline.txt as follows.

julia> f = open("newfile.txt", "r")  # Open for reading
IOStream(<file newfile.txt>)

julia> print(readall(f))
testing
more testing
Often when reading from a file we want to step through the lines of a file, performing an action on each one.

There’s a neat interface to this in Julia, which takes us to our next topic.

**Iterating**

One of the most important tasks in computing is stepping through a sequence of data and performing a given action.

Julia’s provides neat, flexible tools for iteration as we now discuss.

**Iterables**  An iterable is something you can put on the right hand side of `for` and loop over.

These include sequence data types like arrays:

```julia
actions = ["surf", "ski"]
for action in actions
    println("Charlie don't $action")
end
```

They also include so-called *iterators*.

You’ve already come across these types of objects:

```julia
julia> for i in 1:3 print(i) end
123
```

If you ask for the keys of dictionary you get an iterator:

```julia
julia> d = {"name" => "Frodo", "age" => 33}
Dict{Any, Any} with 2 entries:
  "name" => "Frodo"
  "age" => 33

julia> keys(d)
KeyIterator for a Dict{Any, Any} with 2 entries. Keys:
  "name"
  "age"
```

This makes sense, since the most common thing you want to do with keys is loop over them.

The benefit of providing an iterator rather than an array, say, is that the former is more memory efficient.

Should you need to transform an iterator into an array you can always use `collect()`:

```julia
julia> collect(keys(d))
2-element Array{Any, 1}:
  "name"
  "age"
```
**Looping without Indices** You can loop over sequences without explicit indexing, which often leads to neater code.

For example compare

```julia
for x in x_values
    println(x * x)
end
```

with

```julia
for i in 1:length(x_values)
    println(x_values[i] * x_values[i])
end
```

Julia provides some functional-style helper functions (similar to Python) to facilitate looping without indices.

One is `zip()`, which is used for stepping through pairs from two sequences.

For example, try running the following code

```julia
countries = ("Japan", "Korea", "China")
cities = ("Tokyo", "Seoul", "Beijing")
for (country, city) in zip(countries, cities)
    println("The capital of $country is $city")
end
```

If we happen to need the index as well as the value, one option is to use `enumerate()`

The following snippet will give you the idea

```julia
countries = ("Japan", "Korea", "China")
cities = ("Tokyo", "Seoul", "Beijing")
for (i, country) in enumerate(countries)
    city = cities[i]
    println("The capital of $country is $city")
end
```

**Comprehensions** Comprehensions are an elegant tool for creating new arrays or dictionaries from iterables.

Here’s some examples

```julia
julia> doubles = [2i for i in 1:4]
4-element Array{Int64,1}:
 2
 4
 6
 8
```

```julia
julia> animals = ["dog", "cat", "bird"]
3-element Array{ASCIIString,1}:
"dog"
```

```
```
"cat"
"bird"

```julia
julia> plurals = [animal * "s" for animal in animals]
3-element Array{Union{ASCIIString,UTF8String},1}:
  "dogs"
  "cats"
  "birds"
```

```julia
julia> [i + j for i=1:3, j=4:6]  # can specify multiple parameters
3x3 Array{Int64,2}:
  5  6  7
  6  7  8
  7  8  9
```

```julia
julia> [i + j for i=1:3, j=4:6, k=7:9]
3x3x3 Array{Int64,3}:
[:, :, 1] =
  5  6  7
  6  7  8
  7  8  9
[:, :, 2] =
  5  6  7
  6  7  8
  7  8  9
[:, :, 3] =
  5  6  7
  6  7  8
  7  8  9
```

The same kind of expression works for dictionaries

```julia
julia> d = {"$i" => i for i in 1:3}
Dict{Any,Any} with 3 entries:
  "1" => 1
  "2" => 2
  "3" => 3
```

**Comparisons and Logical Operators**

**Comparisons**  As we saw earlier, when testing for equality we use ==

```julia
julia> x = 1
1
julia> x == 2
false
```

For “not equal” use !=
In Julia we can *chain* inequalities

```julia
julia> 1 < 2 < 3
true

julia> 1 <= 2 <= 3
true
```

In many languages you can use integers or other values when testing conditions but Julia is more fussy

```julia
julia> while 0 println("foo") end
ERROR: type: non-boolean (Int64) used in boolean context
in anonymous at no file

julia> if 1 println("foo") end
ERROR: type: non-boolean (Int64) used in boolean context
```

**Combining Expressions**  Here are the standard logical connectives (conjunction, disjunction)

```julia
julia> true && false
false

julia> true || false
true
```

Remember

- $P \land Q$ is true if both are true, otherwise it’s false
- $P \lor Q$ is false if both are false, otherwise it’s true

**User Defined Functions**

Let’s talk a little more about user defined functions

User defined functions are important for improving the clarity of your code by

- separating different strands of logic
- facilitating code reuse (writing the same thing twice is always a bad idea)

Julia functions are convenient:

- Any number of functions can be defined in a given file
- Any “value” can be passed to a function as an argument, including other functions
- Functions can be (and often are) defined inside other functions
- A function can return any kind of value, including functions
We’ll see many examples of these structures in the following lectures

For now let’s just cover some of the different ways of defining functions

**Return Statement**  In Julia, the `return` statement is optional, so that the following functions have identical behavior

```julia
function f1(a, b)
    return a * b
end

function f2(a, b)
    a * b
end
```

When no return statement is present, the last value obtained when executing the code block is returned.

Although some prefer the second option, we often favor the former on the basis that explicit is better than implicit.

A function can have arbitrarily many `return` statements, with execution terminating when the first return is hit.

You can see this in action when experimenting with the following function

```julia
function foo(x)
    if x > 0
        return "positive"
    end
    return "nonpositive"
end
```

**Other Syntax for Defining Functions**  For short function definitions Julia offers some attractive simplified syntax.

First, when the function body is a simple expression, it can be defined without the `function` keyword or `end`

```julia
julia> f(x) = sin(1 / x)
f (generic function with 2 methods)
```

Let’s check that it works

```julia
julia> f(1 / pi)
1.2246467991473532e-16
```

Julia also allows for you to define anonymous functions

For example, to define `f(x) = sin(1 / x)` you can use `x -> sin(1 / x)`.

The difference is that the second function has no name bound to it.

How can you use a function with no name?
Typically it’s as an argument to another function

```julia
julia> map(x -> sin(1 / x), randn(3))  # Apply function to each element
3-element Array{Float64,1}:
    0.744193
   -0.370506
   -0.458826
```

### Optional and Keyword Arguments

Function arguments can be given default values

```julia
function f(x, a=1)
    return exp(cos(a * x))
end
```

If the argument is not supplied the default value is substituted

```julia
julia> f(pi)
0.36787944117144233
julia> f(pi, 2)
2.718281828459045
```

Another option is to use **keyword** arguments

The difference between keyword and standard (positional) arguments is that they are parsed and bound by name rather than order in the function call

For example, in the call

```julia
simulate(param1, param2, max_iterations=100, error_tolerance=0.01)
```

the last two arguments are keyword arguments and their order is irrelevant (as long as they come after the positional arguments)

To define a function with keyword arguments you need to use `;` like so

```julia
function simulate(param1, param2; max_iterations=100, error_tolerance=0.01)
    # Function body here
end
```

### Exercises

**Exercise 1**

Part 1: Given two numeric arrays or tuples `x_vals` and `y_vals` of equal length, compute their inner product using `zip()`

Part 2: Using a comprehension, count the number of even numbers in 0,...,99

- Hint: `x % 2` returns 0 if `x` is even, 1 otherwise

Part 3: Using a comprehension, take `pairs = ((2, 5), (4, 2), (9, 8), (12, 10))` and count the number of pairs `(a, b)` such that both `a` and `b` are even
Exercise 2  Consider the polynomial

\[ p(x) = a_0 + a_1x + a_2x^2 + \cdots + a_nx^n = \sum_{i=0}^{n} a_i x^i \]  \hspace{1cm} (1.1)

Using `enumerate()` in your loop, write a function \( p \) such that \( p(x, \text{coeff}) \) computes the value in (1.1) given a point \( x \) and an array of coefficients \( \text{coeff} \).

Exercise 3  Write a function that takes a string as an argument and returns the number of capital letters in the string.

Hint: `uppercase("foo")` returns "FOO"

Exercise 4  Write a function that takes two sequences \( \text{seq\_a} \) and \( \text{seq\_b} \) as arguments and returns `true` if every element in \( \text{seq\_a} \) is also an element of \( \text{seq\_b} \), else `false`.

- By “sequence” we mean an array, tuple or string.

Exercise 5  The Julia libraries include functions for interpolation and approximation. Nevertheless, let’s write our own function approximation routine as an exercise.

In particular, write a function \( \text{linapprox} \) that takes as arguments:

- A function \( f \) mapping some interval \([a, b]\) into \( \mathbb{R} \)
- Two scalars \( a \) and \( b \) providing the limits of this interval
- An integer \( n \) determining the number of grid points
- A number \( x \) satisfying \( a \leq x \leq b \)

and returns the piecewise linear interpolation of \( f \) at \( x \), based on \( n \) evenly spaced grid points \( a = \text{point}[1] < \text{point}[2] < \ldots < \text{point}[n] = b \).

Aim for clarity, not efficiency.

Exercise 6  The following data lists US cities and their populations:

- New York: 8,244,910
- Los Angeles: 3,819,702
- Chicago: 2,707,120
- Houston: 2,145,146
- Philadelphia: 1,536,471
- Phoenix: 1,469,471
- San Antonio: 1,359,758
- San Diego: 1,326,179
- Dallas: 1,223,229

Copy this text into a text file called \( \text{us\_cities.txt} \) and save it in your present working directory.

- That is, save it in the location Julia returns when you call `pwd()`.
Write a program to calculate total population across these cities

Hints:

- If \( f \) is a file object then \( \text{eachline}(f) \) provides an iterable that steps you through the lines in the file
- \( \text{int}("100") \) converts the string "100" into an integer

Solutions

Solution notebook

1.4 Vectors, Arrays and Matrices

Contents

- Vectors, Arrays and Matrices
  - Overview
  - Array Basics
  - Operations on Arrays
  - Linear Algebra
  - Exercises
  - Solutions

“Let’s be clear: the work of science has nothing whatever to do with consensus. Consensus is the business of politics. Science, on the contrary, requires only one investigator who happens to be right, which means that he or she has results that are verifiable by reference to the real world. In science consensus is irrelevant. What is relevant is reproducible results.” – Michael Crichton

Overview

In Julia, arrays are the most important data type for working with collections of numerical data

In this lecture we give more details on

- creating and manipulating Julia arrays
- fundamental array processing operations
- basic matrix algebra

Array Basics

Shape and Dimension  We’ve already seen some Julia arrays in action
1.4. VECTORS, ARRAYS AND MATRICES

\[ \texttt{julia> a = [10, 20, 30]} \]
3-element Array{Int64,1}:
  10
  20
  30

\[ \texttt{julia> a = ["foo", "bar", 10]} \]
3-element Array{Any,1}:
  "foo"
  "bar"
  10

The REPL tells us that the arrays are of types Array\{Int64,1\} and Array\{Any,1\} respectively.

Here Int64 and Any are types for the elements inferred by the compiler.

We’ll talk more about types later on.

The 1 in Array\{Int64,1\} and Array\{Any,1\} indicates that the array is \textit{one dimensional}.

This is the default for many Julia functions that create arrays.

\[ \texttt{julia> typeof(linspace(0, 1, 100))} \]
Array\{Float64,1\}

\[ \texttt{julia> typeof(randn(100))} \]
Array\{Float64,1\}

To say that an array is one dimensional is to say that it is flat — neither a row nor a column vector.

We can also confirm that \( a \) is flat using the size() or ndims() functions.

\[ \texttt{julia> size(a)} \]
(3,)

\[ \texttt{julia> ndims(a)} \]
1

The syntax (3,) displays a tuple containing one element.

Here it gives the size along the one dimension that exists.

Here’s a function that creates a two-dimensional array.

\[ \texttt{julia> eye(3)} \]
3x3 Array\{Float64,2\}:
  1.0  0.0  0.0
  0.0  1.0  0.0
  0.0  0.0  1.0

\[ \texttt{julia> diagm([2, 4])} \]
2x2 Array\{Int64,2\}:
  2  0
  0  4
Array vs Vector vs Matrix  In Julia, in addition to arrays you will see the types Vector and Matrix. However, these are just aliases for one- and two-dimensional arrays respectively.

julia> Array{Int64, 1} == Vector{Int64}
true

julia> Array{Int64, 2} == Matrix{Int64}
true

julia> Array{Int64, 1} == Matrix{Int64}
false

julia> Array{Int64, 3} == Matrix{Int64}
false

The only slightly disturbing thing here is that the common mathematical terms “row vector” and “column vector” don’t make sense in Julia.

By definition, a Vector in Julia is flat and hence neither row nor column.

Changing Dimensions  The primary function for changing the dimension of an array is reshape()

julia> a = [10, 20, 30, 40]
4-element Array{Int64,1}:
  10
  20
  30
  40

julia> reshape(a, 2, 2)
2x2 Array{Int64,2}:
  10  30
  20  40

julia> reshape(a, 1, 4)
1x4 Array{Int64,2}:
  10  20  30  40

Notice that this function returns a new copy of the reshaped array rather than modifying the existing one.

To collapse an array along one dimension you can use squeeze()
1.4. VECTORS, ARRAYS AND MATRICES

```
julia> squeeze(a, 1)
4-element Array{Int64,1}:
 1
 2
 3
 4
```

The return value is an Array with the specified dimension “flattened”

**Why Vectors?** As we’ve seen, in Julia we have both

- one-dimensional arrays (flat arrays, or vectors)
- two-dimensional arrays of dimension \((1, n)\) or \((n, 1)\) containing the same elements

Why do we need both?

On one hand, dimension matters when we come to matrix algebra

- Multiplying by a row vector is different to multiplication by a column vector

However, if our vectors are not multiplying matrices, their dimensions don’t matter, and hence are an unnecessary complication

This is why many Julia functions return flat arrays by default

**Creating Arrays**

**Functions that Return Arrays** We’ve already seen some functions for creating arrays

```
julia> eye(2)
2x2 Array{Float64,2}:
 1.0  0.0
 0.0  1.0

julia> zeros(3)
3-element Array{Float64,1}:
 0.0
 0.0
 0.0
```

You can create an empty array using the `Array()` constructor

```
julia> x = Array(Float64, 2, 2)
2x2 Array{Float64,2}:
 0.0  2.82622e-316
 2.76235e-318  2.82622e-316
```

The printed values you see here are just garbage values

(the existing contents of the allocated memory slots being interpreted as 64 bit floats)

Other important functions that return arrays are
1.4. VECTORS, ARRAYS AND MATRICES

```julia
julia> ones(2, 2)
2x2 Array{Float64,2}:
  1.0  1.0
  1.0  1.0
```

```julia
julia> fill("foo", 2, 2)
2x2 Array{ASCIIString,2}:
  "foo"  "foo"  
  "foo"  "foo"
```

**Manual Array Definitions**  As we’ve seen, you can create one dimensional arrays from manually specified data like so

```julia
julia> a = [10, 20, 30, 40]
4-element Array{Int64,1}:
  10
  20
  30
  40
```

In two dimensions we can proceed as follows

```julia
julia> a = [10 20 30 40]  # Two dimensional, shape is 1 x n
1x4 Array{Int64,2}:
  10  20  30  40
```

```julia
julia> ndims(a)
2
```

```julia
julia> a = [10 20; 30 40]  # 2 x 2
2x2 Array{Int64,2}:
  10  20
  30  40
```

You might then assume that `a = [10; 20; 30; 40]` creates a two dimensional column vector by unfortunately this isn’t the case

```julia
julia> a = [10; 20; 30; 40]
4-element Array{Int64,1}:
  10
  20
  30
  40
```

```julia
julia> ndims(a)
1
```

Instead transpose the row vector

```julia
julia> a = [10 20 30 40]'
4x1 Array{Int64,2}:
  10
```

---

**THOMAS SARGENT AND JOHN STACHURSKI**  March 21, 2015
Array Indexing  We’ve already seen the basics of array indexing

julia> a = collect(10:10:40)
4-element Array{Int64,1}:
10
20
30
40

julia> a[end-1]
30

julia> a[1:3]
3-element Array{Int64,1}:
10
20
30

For 2D arrays the index syntax is straightforward

julia> a = randn(2, 2)
2x2 Array{Float64,2}:
1.37556 0.924224
1.52899 0.815694

julia> a[1, 1]
1.375559922478634

julia> a[1, :]  # First row
1x2 Array{Float64,2}:
1.37556 0.924224

julia> a[:, 1]  # First column
2-element Array{Float64,1}:
1.37556
1.52899

Booleans can be used to extract elements

julia> a = randn(2, 2)
2x2 Array{Float64,2}:
-0.121311 0.654559
-0.297859 0.89208

julia> b = [true false; false true]
2x2 Array{Bool,2}:
true  false
false  true

Julia> a[b]
2-element Array{Float64,1}:
-0.121311
0.89208

This is useful for conditional extraction, as we’ll see below

An aside: some or all elements of an array can be set equal to one number using slice notation

Julia> a = Array(Float64, 4)
4-element Array{Float64,1}:
1.30822e-282
1.2732e-313
4.48229e-316
1.30824e-282

Julia> a[2:end] = 42
42

Julia> a
4-element Array{Float64,1}:
1.30822e-282
42.0
42.0
42.0

Passing Arrays  As in Python, all arrays are passed by reference

What this means is that if a is an array and we set b = a then a and b point to exactly the same data

Hence any change in b is reflected in a

Julia> a = ones(3)
3-element Array{Float64,1}:
1.0
1.0
1.0

Julia> b = a
3-element Array{Float64,1}:
1.0
1.0
1.0

Julia> b[3] = 44
44

Julia> a
3-element Array{Float64,1}:
1.0
1.0
44
If you are a MATLAB programmer perhaps you are recoiling in horror at this idea. But this is actually the more sensible default – after all, it’s very inefficient to copy arrays unnecessarily.

If you do need an actual copy in Julia, just use `copy()`

```
julia> a = ones(3)
3-element Array{Float64,1}:
  1.0
  1.0
  1.0

julia> b = copy(a)
3-element Array{Float64,1}:
  1.0
  1.0
  1.0

julia> b[3] = 44
44

julia> a
3-element Array{Float64,1}:
  1.0
  1.0
  1.0
```

### Operations on Arrays

**Array Methods** Julia provides standard functions for acting on arrays, some of which we’ve already seen

```
julia> a = [-1, 0, 1]
3-element Array{Int64,1}:
 -1
  0
  1

julia> length(a)
3

julia> sum(a)
0

julia> mean(a)
0.0
```
julia> std(a)
1.0

julia> var(a)
1.0

julia> maximum(a)
1

julia> minimum(a)
-1

julia> b = sort(a, rev=true)  # Returns new array, original not modified
3-element Array{Int64,1}:
  1
  0
-1

julia> b === a  # === tests if arrays are identical (i.e share same memory)
false

julia> b = sort!(a, rev=true)  # Returns *modified original* array
3-element Array{Int64,1}:
  1
  0
-1

julia> b === a
true

Matrix Algebra  For two dimensional arrays, * means matrix multiplication

julia> a = ones(1, 2)
1x2 Array{Float64,2}:
1.0 1.0

julia> b = ones(2, 2)
2x2 Array{Float64,2}:
1.0 1.0
1.0 1.0

julia> a * b
1x2 Array{Float64,2}:
2.0 2.0

julia> b * a'
2x1 Array{Float64,2}:
2.0
2.0

To solve the linear system $A X = B$ for $X$ use $A \ \backslash \ B$
julia> A = [1 2; 2 3]
2x2 Array{Int64,2}:
  1  2
  2  3

julia> B = ones(2, 2)
2x2 Array{Float64,2}:
  1.0  1.0
  1.0  1.0

julia> A \ B
2x2 Array{Float64,2}:
 -1.0  -1.0
  1.0   1.0

julia> inv(A) * B
2x2 Array{Float64,2}:
 -1.0  -1.0
  1.0   1.0

Although the last two operations give the same result, the first one is numerically more stable and
should be preferred in most cases

Multiplying two one dimensional vectors gives an error — which is reasonable since the meaning
is ambiguous

julia> ones(2) * ones(2)
ERROR: `*` has no method matching *(::Array{Float64,1}, ::Array{Float64,1})

If you want an inner product in this setting use dot()

julia> dot(ones(2), ones(2))
2.0

Matrix multiplication using one dimensional vectors is a bit inconsistent — pre-multiplication by
the matrix is OK, but post-multiplication gives an error

julia> b = ones(2, 2)
2x2 Array{Float64,2}:
  1.0  1.0
  1.0  1.0

julia> b * ones(2)
2-element Array{Float64,1}:
  2.0
  2.0

julia> ones(2) * b
ERROR: DimensionMismatch("*")
in gemm_wrapper! at linalg/matmul.jl:275
in * at linalg/matmul.jl:74

It’s probably best to give your vectors dimension before you multiply them against matrices
Elementwise Operations

**Algebraic Operations** Suppose that we wish to multiply every element of matrix $A$ with the corresponding element of matrix $B$

In that case we need to replace $*$ (matrix multiplication) with .* (elementwise multiplication)

For example, compare

```julia
julia> ones(2, 2) * ones(2, 2)  # Matrix multiplication
2x2 Array{Float64,2}:
  2.0  2.0
  2.0  2.0

julia> ones(2, 2) .* ones(2, 2)  # Element by element multiplication
2x2 Array{Float64,2}:
  1.0  1.0
  1.0  1.0
```

This is a general principle: .x means apply operator x elementwise

```julia
julia> A = -ones(2, 2)
2x2 Array{Float64,2}:
 -1.0  -1.0
 -1.0  -1.0

julia> A.^2  # Square every element
2x2 Array{Float64,2}:
  1.0  1.0
  1.0  1.0
```

However in practice some operations are unambiguous and hence the . can be omitted

```julia
julia> ones(2, 2) + ones(2, 2)  # Same as ones(2, 2) .+ ones(2, 2)
2x2 Array{Float64,2}:
  2.0  2.0
  2.0  2.0
```

Scalar multiplication is similar

```julia
julia> A = ones(2, 2)
2x2 Array{Float64,2}:
  1.0  1.0
  1.0  1.0

julia> 2 * A  # Same as 2 .* A
2x2 Array{Float64,2}:
  2.0  2.0
  2.0  2.0
```

In fact you can omit the $*$ altogether and just write $2A$

**Elementwise Comparisons** Elementwise comparisons also use the .x style notation
We can also do comparisons against scalars with parallel syntax

This is particularly useful for *conditional extraction* — extracting the elements of an array that satisfy a condition
Vectorized Functions Julia provides standard mathematical functions such as \( \log, \exp, \sin \), etc.

```julia
julia> log(1.0)
0.0
```

By default, these functions act *elementwise* on arrays

```julia
julia> log(ones(4))
4-element Array{Float64,1}:
0.0
0.0
0.0
0.0
```

Functions that act elementwise on arrays in this manner are called **vectorized functions**

Note that we can get the same result as with a comprehension or more explicit loop

```julia
julia> [log(x) for x in ones(4)]
4-element Array{Float64,1}:
0.0
0.0
0.0
0.0
```

In Julia loops are typically fast and hence the need for vectorized functions is less intense than for some other high level languages

Nonetheless the syntax is convenient

Linear Algebra

Julia provides some a great deal of additional functionality related to linear operations

```julia
julia> A = [1 2; 3 4]
2x2 Array{Int64,2}:
 1 2
 3 4

julia> det(A)
-2.0

julia> trace(A)
5

julia> eigvals(A)
2-element Array{Float64,1}:
-0.372281
```

---

**THOMAS SARGENT AND JOHN STACHURSKI**  
March 21, 2015
Exercise 1 Consider the stochastic difference equation

\[ X_{t+1} = AX_t + b + \Sigma W_t \]  

(1.2)

Here \( \{W_t\} \) is an iid vector of shocks with variance-covariance matrix equal to the identity matrix. Letting \( S_t \) denote the variance of \( X_t \) and using the rules for computing variances in matrix expressions, it can be shown from (1.2) that \( \{S_t\} \) obeys

\[ S_{t+1} = AS_t A' + \Sigma \Sigma' \]  

(1.3)

Provided that all eigenvalues of \( A \) lie within the unit circle, the sequence \( \{S_t\} \) converges to a unique limit \( S \).

This is the **unconditional variance** or **asymptotic variance** of the stochastic difference equation. As an exercise, try writing a simple function that solves for the limit \( S \) by iterating on (1.3) given \( A \) and \( \Sigma \).

To test your solution, observe that the limit \( S \) is a solution to the matrix equation

\[ S = ASA' + Q \quad \text{where} \quad Q := \Sigma \Sigma' \]  

(1.4)

This kind of equation is known as a **discrete time Lyapunov equation**.

The **QuantEcon package** provides a function called `solve_discrete_lyapunov` that implements a fast “doubling” algorithm to solve this equation.

Test your iterative method against `solve_discrete_lyapunov` using matrices

\[
A = \begin{bmatrix} 0.8 & -0.2 \\ -0.1 & 0.7 \end{bmatrix} \quad \Sigma = \begin{bmatrix} 0.5 & 0.4 \\ 0.4 & 0.6 \end{bmatrix}
\]
Overview

In this lecture we delve more deeply into the structure of Julia, and in particular into

- the concept of types
- building user defined types
- methods and multiple dispatch

These concepts relate to the way that Julia stores and acts on data

While they might be thought of as advanced topics, some understanding is necessary to

1. Read Julia code written by other programmers
2. Write “well organized” Julia code that’s easy to maintain and debug
3. Improve the speed at which your code runs

At the same time, don’t worry about following all the nuances on your first pass.

If you return to these topics after doing some programming in Julia they will make more sense

Types

In Julia all objects (all “values” in memory) have a type, which can be queried using the `typeof()` function

```
julia> x = 42
42

julia> typeof(x)
Int64
```

Note here that the type resides with the object itself, not with the name `x`.

The name `x` is just a symbol bound to an object of type `Int64`.

Here we rebind it to another object, and now `typeof(x)` gives the type of that new object.
**Common Types**  We’ve already met many of the types defined in the core Julia language and its standard library.

For numerical data, the most common types are integers and floats.

For those working on a 64 bit machine, the default integers and floats are 64 bits, and are called `Int64` and `Float64` respectively (they would be `Int32` and `Float64` on a 32 bit machine).

There are many other important types, used for arrays, strings, iterators and so on.

```julia
julia> typeof(1 + 1im)
Complex{Int64} (constructor with 1 method)

julia> typeof(linspace(0, 1, 100))
Array{Float64,1}

julia> typeof(eye(2))
Array{Float64,2}

julia> typeof("foo")
ASCIIString (constructor with 2 methods)

julia> typeof(1:10)
UnitRange{Int64} (constructor with 1 method)

julia> typeof('c')  # Single character is a *Char*
Char
```

Type is important in Julia because it determines what operations will be performed on the data in a given situation.

Moreover, if you try to perform an action that is unexpected for a given type the function call will usually fail.

```julia
julia> 100 + "100"
ERROR: `+` has no method matching +(::Int64, ::ASCIIString)
```

Some languages will try to guess what the programmer wants here and return 200.

Julia doesn’t — in this sense, Julia is a “strongly typed” language.

Type is important and it’s up to the user to supply data in the correct form (as specified by type).

**Methods and Multiple Dispatch**  Looking more closely at how this works brings us to a very important topic concerning Julia’s data model — methods and multiple dispatch.

Let’s look again at the error message.
As discussed earlier, the operator + is just a function, and we can rewrite that call using functional notation to obtain exactly the same result

```
 julia> 100 + "100"
 ERROR: `+` has no method matching +(::Int64, ::ASCIIString)
```

Multiplication is similar

```
 julia> 100 * "100"
 ERROR: `*` has no method matching *(::Int64, ::ASCIIString)
```

What the message tells us is that *(a, b) doesn’t work when a is an integer and b is a string

In particular, the function * has no matching method

In essence, a method in Julia is a version of a function that acts on a particular tuple of data types

For example, if a and b are integers then a method for multiplying integers is invoked

```
 julia> *(100, 100)
 10000
```

On the other hand, if a and b are strings then a method for string concatenation is invoked

```
 julia> *("foo", "bar")
 "foobar"
```

In fact we can see the precise methods being invoked by applying @which

```
 julia> @which *(100, 100)
 *(x::Int64,y::Int64) at int.jl:47
```

```
 julia> @which *("foo", "bar")
 *(s::String...) at string.jl:76
```

We can see the same process with other functions and their methods

```
 julia> isfinite(1.0)  # Call isfinite on a float
 true
```

```
 julia> @which(isfinite(1.0))
 isfinite(x::FloatingPoint) at float.jl:213
```

```
 julia> isfinite(1)  # Call isfinite on an integer
 true
```

```
 julia> @which(isfinite(1))
 isfinite(x::Integer) at float.jl:215
```


March 21, 2015
Here `isfinite()` is a function with multiple methods

It has a method for acting on floating points and another method for acting on integers

In fact it has quite a few methods

```
 julia> methods(isfinite)  # List the methods of isfinite
 # 9 methods for generic function "isfinite":
 isfinite(x::Float16) at float16.jl:115
 isfinite(x::BigFloat) at mpfr.jl:717
 isfinite(x::FloatingPoint) at float.jl:213
 isfinite(x::Integer) at float.jl:215
 isfinite(x::Real) at float.jl:214
 isfinite(x::Complex{T<:Real}) at complex.jl:41
 isfinite{T<:Number}(::AbstractArray{T<:Number,1}) at operators.jl:359
 isfinite{T<:Number}(::AbstractArray{T<:Number,2}) at operators.jl:360
 isfinite{T<:Number}(::AbstractArray{T<:Number,N}) at operators.jl:362
```

The particular method being invoked depends on the data type on which the function is called

We’ll discuss some of the more complicated data types you see towards the end of this list as we go along

**Abstract Types**  Looking at the list of methods above you can see references to types that we haven’t met before, such as `Real` and `Number`

These two types are examples of what are known in Julia as **abstract types**

Abstract types serve a different purpose to **concrete types** such as `Int64` and `Float64`

To understand what that purpose is, imagine that you want to write a function with two methods, one to handle real numbers and the other for complex numbers

As we know, there are multiple types for real numbers, such as integers and floats

There are even multiple integer and float types, such as `Int32`, `Int64`, `Float32`, `Float64`, etc.

If we want to handle all of these types for real numbers in the same way, it’s useful to have a “parent” type called `Real`

Rather than writing a separate method for each concrete type, we can just write one for the abstract `Real` type

In this way, the purpose of abstract types is to serve as a unifying parent type that concrete types can “inherit” from

Indeed, we can see that `Float64`, `Int64`, etc. are **subtypes** of `Real` as follows

```
 julia> Float64 <: Real
 true

 julia> Int64 <: Real
 true
```

On the other hand, 64 bit complex numbers are not reals
They are, however, subtypes of \texttt{Number}

\begin{verbatim}
 julia> Complex64 <: Number
    true
\end{verbatim}

\texttt{Number} in turn is a subtype of \texttt{Any}, which is a parent of all types

\begin{verbatim}
 julia> Number <: Any
    true
\end{verbatim}

\textbf{Type Hierarchy} \hspace{1em} In fact the types form a hierarchy, with \texttt{Any} at the top of the tree and the concrete types at the bottom.

Note that we never see \textit{instances} of abstract types.

For example, if \(x\) is a value then \texttt{typeof(x)} will never return an abstract type.

The point of abstract types is simply to categorize the concrete types (as well as other abstract types that sit below them in the hierarchy).

On the other hand, we cannot subtype concrete types.

While we can build subtypes of abstract types we cannot do the same for concrete types.

\textbf{Multiple Dispatch} \hspace{1em} We can now be a little bit clearer about what happens when you call a function on given types.

Suppose we execute the function call \(f(a, b)\) where \(a\) and \(b\) are of concrete types \(S\) and \(T\) respectively.

The Julia interpreter first queries the types of \(a\) and \(b\) to obtain the tuple \((S, T)\).

It then parses the list of methods belonging to \(f\), searching for a match.

If it finds a method matching \((S, T)\) it calls that method.

If not, it looks to see whether the pair \((S, T)\) matches any method defined for \textit{immediate parent types}.

For example, if \(S\) is \texttt{Float64} and \(T\) is \texttt{Complex64} then the immediate parents are \texttt{FloatingPoint} and \texttt{Number} respectively.

\begin{verbatim}
 julia> super(Float64)
    FloatingPoint

 julia> super(Complex64)
    Number
\end{verbatim}

Hence the interpreter looks next for a method of the form \(f(x::\texttt{FloatingPoint}, y::\texttt{Number})\).

If the interpreter can’t find a match in immediate parents (supertypes) it proceeds up the tree, looking at the parents of the last type it checked at each iteration.
1.5. TYPES, METHODS AND PERFORMANCE

- If it eventually finds a matching method it invokes that method
- If not, we get an error

This is the process that leads to the error that we saw above:

```julia
julia> *(100, "100")
ERROR: `*` has no method matching *(::Int64, ::ASCIIString)
```

The procedure of matching data to appropriate methods is called **dispatch**

Because the procedure starts from the concrete types and works upwards, dispatch always invokes the *most specific method* that is available

For example, if you have methods for function `f` that handle

1. `(Float64, Int64)` pairs
2. `(Number, Number)` pairs

and you call `f` with `f(0.5, 1)` then the first method will be invoked

This makes sense because (hopefully) the first method is designed to work well with exactly this kind of data

The second method is probably more of a “catch all” method that handles other data in a less optimal way

**Defining Types and Methods**

Let’s look at defining our own methods and data types, including composite data types

**User Defined Methods**  It’s straightforward to add methods to existing functions or functions you’ve defined

In either case the process is the same:

- use the standard syntax to define a function of the same name
- but specify the data type for the method in the function signature

For example, we saw above that `+` is just a function with various methods

- recall that `a + b` and `+(a, b)` are equivalent

We saw also that the following call fails because it lacks a matching method

```julia
julia> +(100, "100")
ERROR: `+` has no method matching +(::Int64, ::ASCIIString)
```

This is sensible behavior, but if you want to change it by defining a method to handle the case in question there’s nothing to stop you:
1.5. TYPES, METHODS AND PERFORMANCE

```
julia> +(x::Integer, y::ASCIIString) = x + int(y)
+(generic function with 126 methods)

julia> +(100, "100")
200

julia> 100 + "100"
200
```

Here’s another example, involving a user defined function

We begin with a file called `test.jl` in the present working directory with the following content

```
function f(x)
    println("Generic function invoked")
end

function f(x::Number)
    println("Number method invoked")
end

function f(x::Integer)
    println("Integer method invoked")
end
```

Clearly these methods do nothing more than tell you which method is being invoked

Let’s now run this and see how it relates to our discussion of method dispatch above

```
julia> include("test.jl")
f (generic function with 3 methods)

julia> f(3)
Integer method invoked

julia> f(3.0)
Number method invoked

julia> f("foo")
Generic function invoked
```

Since 3 is an `Int64` and `Int64 <: Integer <: Number`, the call `f(3)` proceeds up the tree to `Integer` and invokes `f(x::Integer)`

On the other hand, `3.0` is a `Float64`, which is not a subtype of `Integer`

Hence the call `f(3.0)` continues up to `f(x::Number)`

Finally, `f("foo")` is handled by the generic function, since it is not a subtype of `Number`

**User Defined Types** Most languages have facilities for creating new data types and Julia is no exception
Julia> type Foo end

Julia> foo = Foo()
Foo()

Julia> typeof(foo)
Foo (constructor with 1 method)

Let’s make some observations about this code

First note that to create a new data type we use the keyword `type` followed by the name

- By convention, type names use CamelCase (e.g., `FloatingPoint`, `Array`, `AbstractArray`)

When a new data type is created in this way, the interpreter simultaneously creates a default constructor for the data type

This constructor is a function for generating new instances of the data type in question

It has the same name as the data type but uses function call notion — in this case `Foo()`

In the code above, `foo = Foo()` is a call to the default constructor

A new instance of type `Foo` is created and the name `foo` is bound to that instance

Now if we want to we can create methods that act on instances of `Foo`

Just for fun, let’s define how to add one `Foo` to another

Julia> +(x::Foo, y::Foo) = "twofoos"
+ (generic function with 126 methods)

Julia> foo1, foo2 = Foo(), Foo()  # Create two Foos
(Foo(), Foo())

Julia> +(foo1, foo2)
"twofoos"

Julia> foo1 + foo2
"twofoos"

We can also create new functions to handle `Foo` data

Julia> foofunc(x::Foo) = "onefoo"
foofunc (generic function with 1 method)

Julia> foofunc(foo)
"onefoo"

This example isn’t of much use but more useful examples follow

**Composite Data Types**  Since the common primitive data types are already built in, most new user-defined data types are composite data types

Composite data types are data types that contain distinct fields of data as attributes
For example, let’s say we are doing a lot of work with AR(1) processes, which are random sequences \{X_t\} that follow a law of motion of the form

\[ X_{t+1} = aX_t + b + \sigma W_{t+1} \] (1.5)

Here \(a, b\) and \(\sigma\) are scalars and \(\{W_t\}\) is an iid sequence of shocks with some given distribution \(\phi\). At times it might be convenient to take these primitives \(a, b, \sigma\) and \(\phi\) and organize them into a single entity like so

```julia
type AR1
    a
    b
    sigma
    phi
end
```

For the distribution \(\phi\) we’ll assign a `Distribution` from the Distributions package. After reading in the `AR1` definition above we can do the following

```julia
julia> using Distributions

julia> m = AR1(0.9, 1, 1, Beta(5, 5))
AR1(0.9,1,1,Beta( alpha=5.0 beta=5.0 ))
```

In this call to the constructor we’ve created an instance of `AR1` and bound the name `m` to it. We can access the fields of `m` using their names and “dotted attribute” notation

```julia
julia> m.a
0.9

julia> m.b
1

julia> m.sigma
1

julia> m.phi
Beta( alpha=5.0 beta=5.0 )
```

For example, the attribute `m.phi` points to an instance of `Beta`, which is in turn a subtype of `Distribution` as defined in the Distributions package

```julia
julia> typeof(m.phi)
Beta (constructor with 3 methods)

julia> typeof(m.phi) <: Distribution
true
```

We can reach into `m` and change this if we want to

```julia
julia> m.phi = Exponential(0.5)
Exponential( scale=0.5 )
```
In our type definition we can be explicit that we want \( \phi \) to be a \texttt{Distribution}, and the other elements to be real scalars

```julia
type AR1
    a::Real
    b::Real
    sigma::Real
    phi::Distribution
end
```

(Before reading this in you might need to restart your REPL session in order to clear the old definition of \texttt{AR1} from memory)

Now the constructor will complain if we try to use the wrong data type

```julia
julia> m = AR1(0.9, 1, "foo", Beta(5, 5))
ERROR: `convert` has no method matching convert(::Type{Real}, ::ASCIIString) in AR1 at no file
```

This is useful if we’re going to have functions that act on instances of \texttt{AR1}

- e.g., simulate time series, compute variances, generate histograms, etc.

If those functions only work with \texttt{AR1} instances built from the specified data types then it’s probably best if we get an error as soon we try to make an instance that doesn’t fit the pattern

Better to fail early rather than deeper into our code where errors are harder to debug

### Type Parameters
Consider the following output

```julia
julia> typeof([10, 20, 30])
Array{Int64,1}
```

Here \texttt{Array} is one of Julia’s predefined types (\texttt{Array <: DenseArray <: AbstractArray <: Any})

The \texttt{Int64,1} in curly brackets are \textbf{type parameters}

In this case they are the element type and the dimension

Many other types have type parameters too

```julia
julia> typeof(1.0 + 1.0im)
Complex{Float64} (constructor with 1 method)

julia> typeof(1 + 1im)
Complex{Int64} (constructor with 1 method)
```

Types with parameters are therefore in fact an indexed family of types, one for each possible value of the parameter

### Defining Parametric Types
We can use parametric types in our own type definitions

Let’s say we’re defining a type called \texttt{FooBar} with attributes \texttt{foo} and \texttt{bar}
Suppose we now decide that we want `foo` and `bar` to have the same type, although we don’t much care what that type is.

We can achieve this with the syntax

```julia
type FooBar{T}
    foo::T
    bar::T
end
```

Now our constructor is happy provided that the arguments do in fact have the same type.

```julia
julia> fb = FooBar(1.0, 2.0)
FooBar{Float64}(1.0,2.0)

julia> fb = FooBar(1, 2)
FooBar{Int64}(1,2)

julia> fb = FooBar(1, 2.0)
ERROR: `FooBar{T}` has no method matching FooBar{T}(::Int64, ::Float64)
```

Now let’s say we want the data to be of the same type *and* that type must be a subtype of `Number`.

We can achieve this as follows.

```julia
type FooBar{T <: Number}
    foo::T
    bar::T
end
```

Let’s try it.

```julia
julia> fb = FooBar(1, 2)
FooBar{Int64}(1,2)

julia> fb = FooBar("fee", "fi")
ERROR: `FooBar{T<:Number}` has no method matching FooBar{T<:Number}(:,:,ASCIIString, :ASCIIString)
```

In the second instance we get an error because `ASCIIString` is not a subtype of `Number`.

**Writing Fast Code**

Let’s briefly discuss how to write Julia code that executes quickly (for a given hardware configuration).

For now our focus is on generating more efficient machine code from essentially the same program (i.e., without parallelization or other more significant changes to the way the program runs).
Basic Concepts  The benchmark for performance is well written compiled code, expressed in languages such as C and Fortran.

This is because computer programs are essentially operations on data, and the details of the operations implemented by the CPU depend on the nature of the data.

When code is written in a language like C and compiled, the compiler has access to sufficient information to build machine code that will organize the data optimally in memory and implement efficient operations as required for the task in hand.

To approach this benchmark, Julia needs to know about the type of data it’s processing as early as possible.

An Example  Consider the following function, which essentially does the same job as Julia’s `sum()` function but acts only on floating point data.

```julia
function sum_float_array(x::Array{Float64, 1})
    sum = 0.0
    for i in 1:length(x)
        sum += x[i]
    end
    return sum
end
```

Calls to this function run very quickly.

```julia
julia> x_float = linspace(0, 1, int(1e6))
julia> @time sum_float_array(x_float)
elapsed time: 0.002731878 seconds (96 bytes allocated)
```

One reason is that data types are fully specified.

When Julia compiles this function via its just-in-time compiler, it knows that the data passed in as `x` will be an array of 64 bit floats.

Hence it’s known to the compiler that the relevant method for `+` is always addition of floating point numbers.

Moreover, the data can be arranged into continuous 64 bit blocks of memory to simplify memory access.

Finally, data types are stable — for example, the local variable `sum` starts off as a float and remains a float throughout.

Type Inferences  What happens if we don’t supply type information?

Here’s the same function minus the type annotation in the function signature.

```julia
function sum_array(x)
    sum = 0.0
    for i in 1:length(x)
        sum += x[i]
    end
end
```
When we run it with the same array of floating point numbers it executes just as fast as before

```julia
julia> @time sum_array(x_float)
elapsed time: 0.002720878 seconds (96 bytes allocated)
```

The reason is that when `sum_array()` is first called on a vector of a given data type, a newly compiled version of the function is compiled to handle that type.

In this case, since we're calling the function on a vector of floats, we get a compiled version of the function with essentially the same internal representation as `sum_float_array()`.

Things get tougher for the interpreter when the data type within the array is imprecise.

For example, the following snippet creates an array where the element type is `Any`.

```julia
julia> n = int(1e6)  
   1000000
julia> x_any = {1/i for i in 1:n}
julia> eltype(x_any)
Any
```

Now summation is much slower and memory management is less efficient.

```julia
julia> @time sum_array(x_any)
elapsed time: 0.051313847 seconds (16000096 bytes allocated)
```

Summary and Tips
To write efficient code use functions to segregate operations into logically distinct blocks.

Data types will be determined at function boundaries.
If types are not supplied then they will be inferred.
If types are stable and can be inferred effectively your functions will run fast.

Further Reading
There are many other aspects to writing fast Julia code.
A good next stop for further reading is the relevant part of the Julia documentation.

Exercise 1
Write a function with the signature `simulate(m::AR1, n::Integer, x0::Real)` that takes as arguments
- an instance `m` of `AR1`
- an integer `n`
• a real number $x_0$

and returns an array containing a time series of length $n$ generated according to (1.5) where

• the primitives of the AR(1) process are as specified in $m$
• the initial condition $X_0$ is set equal to $x_0$

Here $AR1$ is as defined above:

```plaintext
type AR1
    a::Real
    b::Real
    sigma::Real
    phi::Distribution
end
```

Hint: If $d$ is an instance of Distribution then $\text{rand}(d)$ generates one random draw from the distribution specified in $d$

**Exercise 2**  The term **universal function** is sometimes applied to functions which

• when called on a scalar return a scalar
• when called on an array of scalars return an array of the same length by acting elementwise on the scalars in the array

For example, $\sin()$ has this property in Julia

```
julia> sin(pi)
1.2246467991473532e-16
```

```
julia> sin([pi, 2pi])
2-element Array{Float64,1}:
  1.22465e-16
 -2.44929e-16
```

Write a universal function $f$ such that

• $f(k)$ returns a chi-squared random variable with $k$ degrees of freedom when $k$ is an integer
• $f(k\_\text{vec})$ returns a vector where $f(k\_\text{vec})[i]$ is chi-squared with $k\_\text{vec}[i]$ degrees of freedom

Hint: If we take $k$ independent standard normals, square them all and sum we get a chi-squared with $k$ degrees of freedom

**Solutions**

Solution notebook

### 1.6 Useful Libraries
Overview

While Julia lacks the massive scientific ecosystem of Python, it has successfully attracted a small army of enthusiastic and talented developers. As a result, its package system is moving towards a critical mass of useful, well written libraries. In addition, a major advantage of Julia libraries is that, because Julia itself is sufficiently fast, there is less need to mix in low level languages like C and Fortran. As a result, most Julia libraries are written exclusively in Julia. Not only does this make the libraries more portable, it makes them much easier to dive into, read, learn from and modify. In this lecture we introduce a few of the Julia libraries that we’ve found particularly useful for quantitative work in economics.

Plotting

There are already several libraries for generating figures in Julia.

- Winston
- Gadfly
- PyPlot

Of these, the most mature from the point of view of the end user is PyPlot. In fact PyPlot is just a Julia front end to the excellent Python plotting library Matplotlib. In the following we provide some basic information on how to install and work with this library.

Installing PyPlot  The one disadvantage of PyPlot is that it not only requires Python but also a lot of the scientific Python back end. However, this has become less of a hassle with the advent of the Anaconda Python distribution.
Moreover, the scientific Python tools are extremely useful and easily accessible from Julia via PyCall.

We discussed installing Anaconda and PyPlot here.

**Usage** The most important source of information about PyPlot is the documentation for Matplotlib itself.

There are also many useful examples available on the Matplotlib website and elsewhere.

**The Procedural API** Matplotlib has a straightforward plotting API that essentially replicates the plotting routines in MATLAB.

These plotting routines can be expressed in Julia with almost identical syntax.

We’ve already seen some examples of this in earlier lectures.

Here’s another example:

```julia
using PyPlot
x = linspace(0, 10, 200)
y = sin(x)
plot(x, y, "b-", linewidth=2)
```

The resulting figure looks as follows:

![Plot](image)

**The Object Oriented API** Matplotlib also has a more “Pythonic” object orientated API that power users will prefer.

Since Julia doesn’t bundle objects with methods in the same way that Python does, plots based on this API don’t follow exactly the same syntax that they do in Matplotlib.

Fortunately the differences are consistent and after seeing some examples you will find it easy to translate one into the other.
Here’s an example of the syntax we’re discussing, which in this case generates exactly the same plot:

```python
using PyPlot
x = linspace(0, 10, 200)
y = sin(x)
fig, ax = subplots()
ax[0]:plot(x, y, "b-", linewidth=2)
```

In this case we get nothing extra and have to accept more complexity and a less attractive syntax. However, it is a little more explicit and this turns out to be convenient as we move to more sophisticated plots.

Here’s a similar plot with a bit more customization:

```python
using PyPlot
x = linspace(0, 10, 200)
y = sin(x)
fig, ax = subplots()
ax[0]:plot(x, y, "r-", linewidth=2, label="sine function", alpha=0.6)
ax[0]:legend(loc="upper center")
```

The resulting figure has a legend at the top center.

![Example plot](image)

We can render the legend in LaTeX by changing the `ax[0]:plot` line to

```python
ax[0]:plot(x, y, "r-", linewidth=2, label=L"$y = \sin(x)$", alpha=0.6)
```

Note the `L` in front of the string to indicate LaTeX mark up.

The result looks as follows:

Here’s another example, which helps illustrate how to put multiple plots on one figure:

```python
using PyPlot
using Distributions
```
1.6. USEFUL LIBRARIES

```python
u = Uniform()

fig, ax = subplots()
x = linspace(-4, 4, 150)
for i in 1:3
    # == Compute normal pdf from randomly generated mean and std == #
    m, s = rand(u) * 2 - 1, rand(u) + 1
    d = Normal(m, s)
    y = pdf(d, x)
    # == Plot current pdf == #
    ax[:plot](x, y, linewidth=2, alpha=0.6, label="draw $i")
end
ax[:legend]()
```

It generates the following plot
Multiple Subplots  A figure containing \( n \) rows and \( m \) columns of subplots can be created by the call

```python
def, axes = subplots(num_rows, num_cols)
```

Here’s an example

```python
using PyPlot
using Distributions

u = Uniform()
num_rows, num_cols = 3, 2
fig, axes = plt.subplots(num_rows, num_cols, figsize=(8, 12))
subplot_num = 0

for i in 1:num_rows
    for j in 1:num_cols
        ax = axes[i, j]
        subplot_num += 1
        # == Generate a normal sample with random mean and std == #
        m, s = rand(u) * 2 - 1, rand(u) + 1
        d = Normal(m, s)
        x = rand(d, 100)
        # == Histogram the sample == #
        ax[:hist](x, alpha=0.6, bins=20)
        ax[:set_title]("histogram \$\text{subplot_num}\$")
        ax[:set_xticks]([-4, 0, 4])
        ax[:set_yticks]([])
    end
end
```

The resulting figure is as follows

3D Plots  Here’s an example of how to create a 3D plot

```python
using PyPlot
using Distributions
using QuantEcon: meshgrid

n = 50
x = linspace(-3, 3, n)
y = x

z = Array(Float64, n, n)
f(x, y) = \( \cos(x^2 + y^2) / (1 + x^2 + y^2) \)
for i in 1:n
    for j in 1:n
        z[j, i] = f(x[i], y[j])
    end
end

fig = figure(figsize=(8,6))
ax = fig[:gca](projection="3d")
```
1.6. USEFUL LIBRARIES

```python
ax[:set_zlim](-0.5, 1.0)
xgrid, ygrid = meshgrid(x, y)
ax[:plot_surface](xgrid, ygrid, z, rstride=2, cstride=2, cmap=ColorMap("jet"), alpha=0.7, linewidth=0.25)
```

It creates this figure

![Figure](image)

**Probability**

Functions for manipulating probability distributions and generating random variables are supplied by the excellent Distributions package.

This package has first-rate documentation so we'll restrict ourselves to a few comments.

The calls to create instances of various random variables take the form `d = DistributionName(params)`.

Here are several examples:

- Normal with mean $m$ and standard deviation $s$
  - $d = \text{Normal}(m, s)$ with defaults $m = 0$ and $s = 1$

- Uniform on interval $[a, b]$
  - $d = \text{Uniform}(a, b)$ with defaults $a = 0$ and $b = 1$

- Binomial over $n$ trials with success probability $p$
  - $d = \text{Binomial}(n, p)$ with defaults $n = 1$ and $p = 0.5$

The Distributions package defines many methods for acting on these instances in order to obtain...
1.6. USEFUL LIBRARIES

- random draws
- pdf (density), cdf (distribution function), quantiles, etc.
- mean, variance, kurtosis, etc.

For example,
- To generate $k$ draws from the instance $d$ use `rand(d, k)`
- To obtain the mean of the distribution use `mean(d)`
- To evaluate the probability density function of $d$ at $x$ use `pdf(d, x)`

Further details on the interface can be found [here](#)
Several multivariate distributions are also implemented

**Working with Data**

A useful package for working with data is **DataFrames**

The most important data type provided is a DataFrame, a two dimensional array for storing heterogeneous data

Although data can be heterogeneous within a DataFrame, the contents of the columns must be homogeneous

This is analogous to a `data.frame` in R, a DataFrame in Pandas (Python) or, more loosely, a spreadsheet in Excel

The DataFrames package also supplies a DataArray type, which is like a one dimensional DataFrame

In terms of working with data, the advantage of a DataArray over a standard numerical array is that it can handle missing values

Here’s an example

```julia
julia> using DataFrames

julia> commodities = ["crude", "gas", "gold", "silver"]
4-element Array{ASCIIString,1}:
"crude"
"gas"
"gold"
"silver"

julia> last = @data([4.2, 11.3, 12.1, NA])  # Create DataArray
4-element DataArray{Float64,1}:
  4.2
  11.3
  12.1
  NA

julia> df = DataFrame(commod = commodities, price = last)
```

THOMAS SARGENT AND JOHN STACHURSKI

March 21, 2015
4x2 DataFrame

<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Row #</td>
<td>commod</td>
<td>price</td>
</tr>
<tr>
<td>1</td>
<td>&quot;crude&quot;</td>
<td>4.2</td>
</tr>
<tr>
<td>2</td>
<td>&quot;gas&quot;</td>
<td>11.3</td>
</tr>
<tr>
<td>3</td>
<td>&quot;gold&quot;</td>
<td>12.1</td>
</tr>
<tr>
<td>4</td>
<td>&quot;silver&quot;</td>
<td>NA</td>
</tr>
</tbody>
</table>

Columns of the DataFrame can be accessed by name

```julia
df[:price]
df[:commod]
```

The DataFrames package provides a number of methods for acting on DataFrames

A simple one is `describe()`

```julia
describe(df)
```

There are also functions for splitting, merging and other data munging operations

Data can be read from and written to CSV files using syntax `df = readtable("data_file.csv")` and `writetable("data_file.csv", df)` respectively

Other packages for working with data can be found at JuliaStats and JuliaQuant
Optimization, Roots and Fixed Points

Let’s look briefly at the optimization and root finding algorithms

**Roots** A root of a real function \( f \) on \([a, b]\) is an \( x \in [a, b] \) such that \( f(x) = 0 \)

For example, if we plot the function

\[
f(x) = \sin(4(x - 1/4)) + x + x^{20} - 1
\]

with \( x \in [0, 1] \) we get

The unique root is approximately 0.408

One common root-finding algorithm is the Newton-Raphson method

This is implemented as `newton()` in the Roots package and is called with the function and an initial guess

```julia
julia> using Roots
julia> f(x) = sin(4 * (x - 1/4)) + x + x^20 - 1
f (generic function with 1 method)

julia> newton(f, 0.2)
0.40829350427936706
```

The Newton-Raphson method uses local slope information, which can lead to failure of convergence for some initial conditions
For this reason most modern solvers use more robust “hybrid methods”, as does Roots’s \texttt{fzero()} function

```julia
julia> fzero(f, 0, 1)
0.40829350427936706
```

\textbf{Optimization}  For constrained, univariate minimization a useful option is \texttt{optimize()} from the \texttt{Optim} package

This function defaults to a robust hybrid optimization routine called Brent’s method

```julia
julia> using Optim

julia> optimize(x -> x^2, -1.0, 1.0)
Results of Optimization Algorithm
* Algorithm: Brent's Method
* Search Interval: [-1.000000, 1.000000]
* Minimum: -0.000000
* Value of Function at Minimum: 0.000000
* Iterations: 5
* Convergence: \( \max(|x - x_{\text{upper}}|, |x - x_{\text{lower}}|) \leq 2 \times (1.5 \times 10^{-8} \times |x| + 2.2 \times 10^{-16}) \): true
* Objective Function Calls: 6

For other optimization routines, including least squares and multivariate optimization, see the documentation

A number of alternative packages for optimization can be found at \texttt{JuliaOpt}

\textbf{Others Topics}

\textbf{Numerical Integration}  The base library contains a function called \texttt{quadgk()} that performs Gaussian quadrature

```julia
julia> quadgk(x -> cos(x), -2pi, 2pi)
(5.644749237155177e-15,4.696156369056425e-22)
```

This is an adaptive Gauss-Kronrod integration technique that’s relatively accurate for smooth functions

However, its adaptive implementation makes it slow and not well suited to inner loops

For this kind of integration you can use the quadrature routines from QuantEcon

```julia
julia> using QuantEcon

julia> nodes, weights = qnwlege(65, -2pi, 2pi);

julia> integral = do_quad(x -> cos(x), nodes, weights)
-2.912600716165059e-15
```
Let’s time the two implementations

```julia
julia> @time quadgk(x -> cos(x), -2pi, 2pi)
elapsed time: 2.732162971 seconds (984420160 bytes allocated, 40.55% gc time)

julia> @time do_quad(x -> cos(x), nodes, weights)
elapsed time: 0.002805691 seconds (1424 bytes allocated)
```

We get similar accuracy with a speed up factor approaching three orders of magnitude

More numerical integration (and differentiation) routines can be found in the package Calculus

**Linear Algebra**  The standard library contains many useful routines for linear algebra, in addition to standard functions such as \( \text{det}(\cdot) \), \( \text{inv}(\cdot) \), \( \text{eye}(\cdot) \), etc.

Routines are available for

- Cholesky factorization
- LU decomposition
- Singular value decomposition,
- Schur factorization, etc.

See here for further details

**Further Reading**

The full set of libraries available under the Julia packaging system can be browsed at pkg.julialang.org
CHAPTER TWO

INTRODUCTORY APPLICATIONS

This section of the course contains relatively simple applications, one purpose of which is to teach you more about the Python programming environment.

2.1 Linear Algebra

Contents

- Linear Algebra
  - Overview
  - Vectors
  - Matrices
  - Solving Systems of Equations
  - Eigenvalues and Eigenvectors
  - Further Topics

Overview

One of the single most useful branches of mathematics you can learn is linear algebra. For example, many applied problems in economics, finance, operations research and other fields of science require the solution of a linear system of equations, such as

\[
\begin{align*}
y_1 &= ax_1 + bx_2 \\
y_2 &= cx_1 + dx_2
\end{align*}
\]

or, more generally,

\[
\begin{align*}
y_1 &= a_{11}x_1 + a_{12}x_2 + \cdots + a_{1k}x_k \\
\vdots \\
y_n &= a_{n1}x_1 + a_{n2}x_2 + \cdots + a_{nk}x_k
\end{align*}
\]

The objective here is to solve for the “unknowns” \(x_1, \ldots, x_k\) given \(a_{11}, \ldots, a_{nk}\) and \(y_1, \ldots, y_n\). When considering such problems, it is essential that we first consider at least some of the following questions.
• Does a solution actually exist?
• Are there in fact many solutions, and if so how should we interpret them?
• If no solution exists, is there a best “approximate” solution?
• If a solution exists, how should we compute it?

These are the kinds of topics addressed by linear algebra

In this lecture we will cover the basics of linear and matrix algebra, treating both theory and computation.

We admit some overlap with this lecture, where operations on Julia arrays were first explained.

Note that this lecture is more theoretical than most, and contains background material that will be used in applications as we go along.

**Vectors**

A vector of length $n$ is just a sequence (or array, or tuple) of $n$ numbers, which we write as $x = (x_1, \ldots, x_n)$ or $x = [x_1, \ldots, x_n]$

We will write these sequences either horizontally or vertically as we please.

(Later, when we wish to perform certain matrix operations, it will become necessary to distinguish between the two)

The set of all $n$-vectors is denoted by $\mathbb{R}^n$

For example, $\mathbb{R}^2$ is the plane, and a vector in $\mathbb{R}^2$ is just a point in the plane.

Traditionally, vectors are represented visually as arrows from the origin to the point.

The following figure represents three vectors in this manner.

If you’re interested, the Julia code for producing this figure is here.

**Vector Operations** The two most common operators for vectors are addition and scalar multiplication, which we now describe.

As a matter of definition, when we add two vectors, we add them element by element.

$$x + y = \begin{bmatrix} x_1 \\ x_2 \\ \vdots \\ x_n \end{bmatrix} + \begin{bmatrix} y_1 \\ y_2 \\ \vdots \\ y_n \end{bmatrix} := \begin{bmatrix} x_1 + y_1 \\ x_2 + y_2 \\ \vdots \\ x_n + y_n \end{bmatrix}$$

Scalar multiplication is an operation that takes a number $\gamma$ and a vector $x$ and produces

$$\gamma x := \begin{bmatrix} \gamma x_1 \\ \gamma x_2 \\ \vdots \\ \gamma x_n \end{bmatrix}$$
Scalar multiplication is illustrated in the next figure

In Julia, a vector can be represented as a one dimensional *Array*

Julia *Arrays* allow us to express scalar multiplication and addition with a very natural syntax

```julia
julia> x = ones(3)
3-element Array{Float64,1}:
  1.0
  1.0
  1.0

julia> y = [2, 4, 6]
3-element Array{Int64,1}:
  2
  4
  6

julia> x + y
3-element Array{Float64,1}:
  3.0
  5.0
  7.0

julia> 4x  # equivalent to 4 * x and 4 .* x
3-element Array{Float64,1}:
  4.0
  4.0
  4.0
```
2.1. LINEAR ALGEBRA

Inner Product and Norm

The inner product of vectors $x, y \in \mathbb{R}^n$ is defined as

$$x' y := \sum_{i=1}^{n} x_i y_i$$

Two vectors are called orthogonal if their inner product is zero.

The norm of a vector $x$ represents its “length” (i.e., its distance from the zero vector) and is defined as

$$\|x\| := \sqrt{x' x} := \left(\sum_{i=1}^{n} x_i^2\right)^{1/2}$$

The expression $\|x - y\|$ is thought of as the distance between $x$ and $y$.

Continuing on from the previous example, the inner product and norm can be computed as follows:

```julia
julia> dot(x, y)            # Inner product of x and y
12.0
julia> sum(x .* y)         # Gives the same result
12.0
julia> norm(x)             # Norm of x
1.7320508075688772
julia> sqrt(sum(x.^2))     # Gives the same result
1.7320508075688772
```

Span

Given a set of vectors $A := \{a_1, \ldots, a_k\}$ in $\mathbb{R}^n$, it’s natural to think about the new vectors we can create by performing linear operations.
New vectors created in this manner are called linear combinations of $A$

In particular, $y \in \mathbb{R}^n$ is a linear combination of $A := \{a_1, \ldots, a_k\}$ if

$$y = \beta_1 a_1 + \cdots + \beta_k a_k$$

for some scalars $\beta_1, \ldots, \beta_k$

In this context, the values $\beta_1, \ldots, \beta_k$ are called the coefficients of the linear combination.

The set of linear combinations of $A$ is called the span of $A$.

The next figure shows the span of $A = \{a_1, a_2\}$ in $\mathbb{R}^3$.

The span is a 2 dimensional plane passing through these two points and the origin.

The code for producing this figure can be found [here](#).

**Examples**

If $A$ contains only one vector $a_1 \in \mathbb{R}^2$, then its span is just the scalar multiples of $a_1$, which is the unique line passing through both $a_1$ and the origin.

If $A = \{e_1, e_2, e_3\}$ consists of the canonical basis vectors of $\mathbb{R}^3$, that is

$$e_1 := \begin{bmatrix} 1 \\ 0 \\ 0 \end{bmatrix}, \quad e_2 := \begin{bmatrix} 0 \\ 1 \\ 0 \end{bmatrix}, \quad e_3 := \begin{bmatrix} 0 \\ 0 \\ 1 \end{bmatrix}$$

then the span of $A$ is all of $\mathbb{R}^3$, because, for any $x = (x_1, x_2, x_3) \in \mathbb{R}^3$, we can write

$$x = x_1 e_1 + x_2 e_2 + x_3 e_3$$

Now consider $A_0 = \{e_1, e_2, e_1 + e_2\}$.

If $y = (y_1, y_2, y_3)$ is any linear combination of these vectors, then $y_3 = 0$ (check it).

Hence $A_0$ fails to span all of $\mathbb{R}^3$. 

---

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Linear Independence  As we’ll see, it’s often desirable to find families of vectors with relatively large span, so that many vectors can be described by linear operators on a few vectors.

The condition we need for a set of vectors to have a large span is what’s called linear independence. In particular, a collection of vectors $A := \{a_1, \ldots, a_k\}$ in $\mathbb{R}^n$ is said to be

- *linearly dependent* if some strict subset of $A$ has the same span as $A$;
- *linearly independent* if it is not linearly dependent.

Put differently, a set of vectors is linearly independent if no vector is redundant to the span, and linearly dependent otherwise.

To illustrate the idea, recall the figure that showed the span of vectors $\{a_1, a_2\}$ in $\mathbb{R}^3$ as a plane through the origin.

If we take a third vector $a_3$ and form the set $\{a_1, a_2, a_3\}$, this set will be

- linearly dependent if $a_3$ lies in the plane;
- linearly independent otherwise.

As another illustration of the concept, since $\mathbb{R}^n$ can be spanned by $n$ vectors (see the discussion of canonical basis vectors above), any collection of $m > n$ vectors in $\mathbb{R}^n$ must be linearly dependent.

The following statements are equivalent to linear independence of $A := \{a_1, \ldots, a_k\} \subset \mathbb{R}^n$:

1. No vector in $A$ can be formed as a linear combination of the other elements.
2. If $\beta_1a_1 + \cdots + \beta_ka_k = 0$ for scalars $\beta_1, \ldots, \beta_k$, then $\beta_1 = \cdots = \beta_k = 0$.

(The zero in the first expression is the origin of $\mathbb{R}^n$.)

Unique Representations  Another nice thing about sets of linearly independent vectors is that each element in the span has a unique representation as a linear combination of these vectors.

In other words, if $A := \{a_1, \ldots, a_k\} \subset \mathbb{R}^n$ is linearly independent and

$$y = \beta_1a_1 + \cdots + \beta_ka_k$$

then no other coefficient sequence $\gamma_1, \ldots, \gamma_k$ will produce the same vector $y$.

Indeed, if we also have $y = \gamma_1a_1 + \cdots + \gamma_ka_k$, then

$$(\beta_1 - \gamma_1)a_1 + \cdots + (\beta_k - \gamma_k)a_k = 0$$

Linear independence now implies $\gamma_i = \beta_i$ for all $i$.

Matrices  Matrices are a neat way of organizing data for use in linear operations.
2.1. LINEAR ALGEBRA

An \( n \times k \) matrix is a rectangular array \( A \) of numbers with \( n \) rows and \( k \) columns:

\[
A = \begin{bmatrix}
a_{11} & a_{12} & \cdots & a_{1k} \\
a_{21} & a_{22} & \cdots & a_{2k} \\
\vdots & \vdots & \ddots & \vdots \\
a_{n1} & a_{n2} & \cdots & a_{nk}
\end{bmatrix}
\]

Often, the numbers in the matrix represent coefficients in a system of linear equations, as discussed at the start of this lecture.

For obvious reasons, the matrix \( A \) is also called a vector if either \( n = 1 \) or \( k = 1 \). In the former case, \( A \) is called a **row vector**, while in the latter it is called a **column vector**.

If \( n = k \), then \( A \) is called **square**.

The matrix formed by replacing \( a_{ij} \) by \( a_{ji} \) for every \( i \) and \( j \) is called the **transpose** of \( A \), and denoted \( A' \) or \( A^\top \).

If \( A = A' \), then \( A \) is called **symmetric**.

For a square matrix \( A \), the \( i \) elements of the form \( a_{ii} \) for \( i = 1, \ldots, n \) are called the **principal diagonal**.

\( A \) is called **diagonal** if the only nonzero entries are on the principal diagonal.

If, in addition to being diagonal, each element along the principal diagonal is equal to 1, then \( A \) is called the **identity matrix**, and denoted by \( I \).

**Matrix Operations**  Just as was the case for vectors, a number of algebraic operations are defined for matrices.

Scalar multiplication and addition are immediate generalizations of the vector case:

\[
\gamma A = \gamma \begin{bmatrix} a_{11} & \cdots & a_{1k} \\
\vdots & \ddots & \vdots \\
a_{n1} & \cdots & a_{nk} \end{bmatrix} = \begin{bmatrix} \gamma a_{11} & \cdots & \gamma a_{1k} \\
\vdots & \ddots & \vdots \\
\gamma a_{n1} & \cdots & \gamma a_{nk} \end{bmatrix}
\]

and

\[
A + B = \begin{bmatrix} a_{11} & \cdots & a_{1k} \\
\vdots & \ddots & \vdots \\
a_{n1} & \cdots & a_{nk} \end{bmatrix} + \begin{bmatrix} b_{11} & \cdots & b_{1k} \\
\vdots & \ddots & \vdots \\
b_{n1} & \cdots & b_{nk} \end{bmatrix} = \begin{bmatrix} a_{11} + b_{11} & \cdots & a_{1k} + b_{1k} \\
\vdots & \ddots & \vdots \\
a_{n1} + b_{n1} & \cdots & a_{nk} + b_{nk} \end{bmatrix}
\]

In the latter case, the matrices must have the same shape in order for the definition to make sense.

We also have a convention for **multiplying** two matrices.

The rule for matrix multiplication generalizes the idea of inner products discussed above, and is designed to make multiplication play well with basic linear operations.

If \( A \) and \( B \) are two matrices, then their product \( AB \) is formed by taking as its \( i, j \)-th element the inner product of the \( i \)-th row of \( A \) and the \( j \)-th column of \( B \).

There are many tutorials to help you visualize this operation, such as this one, or the discussion on the Wikipedia page.
If \( A \) is \( n \times k \) and \( B \) is \( j \times m \), then to multiply \( A \) and \( B \) we require \( k = j \), and the resulting matrix \( AB \) is \( n \times m \).

As perhaps the most important special case, consider multiplying \( n \times k \) matrix \( A \) and \( k \times 1 \) column vector \( x \).

According to the preceding rule, this gives us an \( n \times 1 \) column vector:

\[
Ax = \begin{bmatrix}
  a_{11} & \cdots & a_{1k} \\
  \vdots & \ddots & \vdots \\
  a_{n1} & \cdots & a_{nk}
\end{bmatrix}
\begin{bmatrix}
  x_1 \\
  \vdots \\
  x_k
\end{bmatrix}
= \begin{bmatrix}
  a_{11}x_1 + \cdots + a_{1k}x_k \\
  \vdots \\
  a_{n1}x_1 + \cdots + a_{nk}x_k
\end{bmatrix}
\tag{2.2}
\]

**Note:** \( AB \) and \( BA \) are not generally the same thing.

Another important special case is the identity matrix.

You should check that if \( A \) is \( n \times k \) and \( I \) is the \( k \times k \) identity matrix, then \( AI = A \).

If \( I \) is the \( n \times n \) identity matrix, then \( IA = A \).

**Matrices in Julia** Julia arrays are also used as matrices, and have fast, efficient functions and methods for all the standard matrix operations.

You can create them as follows:

```
julia> A = [1 2
       3 4]
2x2 Array{Int64,2}:
1  2
3  4

julia> typeof(A)
Array{Int64,2}

julia> size(A)
(2,2)
```

The `size` function returns a tuple giving the number of rows and columns — see [here](#) for more discussion.

To get the transpose of \( A \), use `transpose(A)` or, more simply, \( A' \).

There are many convenient functions for creating common matrices (matrices of zeros, ones, etc.) — see [here](#).

Since operations are performed elementwise by default, scalar multiplication and addition have very natural syntax:

```
julia> A = eye(3)
3x3 Array{Float64,2}:
1.0  0.0  0.0
```
To multiply matrices we use the \( \ast \) operator

In particular, \( A \ast B \) is matrix multiplication, whereas \( A \ast B \) is element by element multiplication

See [here](#) for more discussion

**Matrices as Maps** Each \( n \times k \) matrix \( A \) can be identified with a function \( f(x) = Ax \) that maps \( x \in \mathbb{R}^k \) into \( y = Ax \in \mathbb{R}^n \)

These kinds of functions have a special property: they are *linear*

A function \( f: \mathbb{R}^k \to \mathbb{R}^n \) is called *linear* if, for all \( x, y \in \mathbb{R}^k \) and all scalars \( \alpha, \beta \), we have

\[
f(\alpha x + \beta y) = \alpha f(x) + \beta f(y)
\]

You can check that this holds for the function \( f(x) = Ax + b \) when \( b \) is the zero vector, and fails when \( b \) is nonzero

In fact, it’s known that \( f \) is linear if and only if there exists a matrix \( A \) such that \( f(x) = Ax \) for all \( x \).

**Solving Systems of Equations**

Recall again the system of equations (2.1)

If we compare (2.1) and (2.2), we see that (2.1) can now be written more conveniently as

\[
y = Ax
\]

(2.3)

The problem we face is to determine a vector \( x \in \mathbb{R}^k \) that solves (2.3), taking \( y \) and \( A \) as given

This is a special case of a more general problem: Find an \( x \) such that \( y = f(x) \)
Given an arbitrary function $f$ and a $y$, is there always an $x$ such that $y = f(x)$?

If so, is it always unique?

The answer to both these questions is negative, as the next figure shows

In the first plot there are multiple solutions, as the function is not one-to-one, while in the second there are no solutions, since $y$ lies outside the range of $f$.

Can we impose conditions on $A$ in (2.3) that rule out these problems?

In this context, the most important thing to recognize about the expression $Ax$ is that it corresponds to a linear combination of the columns of $A$.

In particular, if $a_1, \ldots, a_k$ are the columns of $A$, then

$$Ax = x_1a_1 + \cdots + x_ka_k$$

Hence the range of $f(x) = Ax$ is exactly the span of the columns of $A$.

We want the range to be large, so that it contains arbitrary $y$.

As you might recall, the condition that we want for the span to be large is linear independence.

A happy fact is that linear independence of the columns of $A$ also gives us uniqueness.

Indeed, it follows from our earlier discussion that if $\{a_1, \ldots, a_k\}$ are linearly independent and $y = Ax = x_1a_1 + \cdots + x_ka_k$, then no $z \neq x$ satisfies $y = Az$. 
The $n \times n$ Case  
Let's discuss some more details, starting with the case where $A$ is $n \times n$ 
This is the familiar case where the number of unknowns equals the number of equations 
For arbitrary $y \in \mathbb{R}^n$, we hope to find a unique $x \in \mathbb{R}^n$ such that $y = Ax$ 
In view of the observations immediately above, if the columns of $A$ are linearly independent, then 
their span, and hence the range of $f(x) = Ax$, is all of $\mathbb{R}^n$ 
Hence there always exists an $x$ such that $y = Ax$ 
Moreover, the solution is unique 
In particular, the following are equivalent 
1. The columns of $A$ are linearly independent 
2. For any $y \in \mathbb{R}^n$, the equation $y = Ax$ has a unique solution 
The property of having linearly independent columns is sometimes expressed as having full column rank 

Inverse Matrices  
Can we give some sort of expression for the solution? 
If $y$ and $A$ are scalar with $A \neq 0$, then the solution is $x = A^{-1}y$ 
A similar expression is available in the matrix case 
In particular, if square matrix $A$ has full column rank, then it possesses a multiplicative inverse 
matrix $A^{-1}$, with the property that $AA^{-1} = A^{-1}A = I$ 
As a consequence, if we pre-multiply both sides of $y = Ax$ by $A^{-1}$, we get $x = A^{-1}y$ 
This is the solution that we’re looking for 

Determinants  
Another quick comment about square matrices is that to every such matrix we 
assign a unique number called the determinant of the matrix — you can find the expression for it here 
If the determinant of $A$ is not zero, then we say that $A$ is nonsingular 
Perhaps the most important fact about determinants is that $A$ is nonsingular if and only if $A$ is of 
full column rank 
This gives us a useful one-number summary of whether or not a square matrix can be inverted 

More Rows than Columns  
This is the $n \times k$ case with $n > k$ 
This case is very important in many settings, not least in the setting of linear regression (where $n$ 
is the number of observations, and $k$ is the number of explanatory variables) 
Given arbitrary $y \in \mathbb{R}^n$, we seek an $x \in \mathbb{R}^k$ such that $y = Ax$ 
In this setting, existence of a solution is highly unlikely
Without much loss of generality, let’s go over the intuition focusing on the case where the columns of \( A \) are linearly independent.

It follows that the span of the columns of \( A \) is a \( k \)-dimensional subspace of \( \mathbb{R}^n \).

This span is very “unlikely” to contain arbitrary \( y \in \mathbb{R}^n \).

To see why, recall the figure above, where \( k = 2 \) and \( n = 3 \).

Imagine an arbitrarily chosen \( y \in \mathbb{R}^3 \), located somewhere in that three dimensional space.

What’s the likelihood that \( y \) lies in the span of \( \{a_1, a_2\} \) (i.e., the two dimensional plane through these points)?

In a sense it must be very small, since this plane has zero “thickness”.

As a result, in the \( n > k \) case we usually give up on existence.

However, we can still seek a best approximation, for example an \( x \) that makes the distance \( \|y - Ax\| \) as small as possible.

To solve this problem, one can use either calculus or the theory of orthogonal projections.

The solution is known to be \( \hat{x} = (A' A)^{-1} A' y \) — see for example chapter 3 of these notes.

**More Columns than Rows**  This is the \( n \times k \) case with \( n < k \), so there are fewer equations than unknowns.

In this case there are either no solutions or infinitely many — in other words, uniqueness never holds.

For example, consider the case where \( k = 3 \) and \( n = 2 \).

Thus, the columns of \( A \) consists of 3 vectors in \( \mathbb{R}^2 \).

This set can never be linearly independent, since 2 vectors are enough to span \( \mathbb{R}^2 \).

(For example, use the canonical basis vectors.)

It follows that one column is a linear combination of the other two.

For example, let’s say that \( a_1 = aa_2 + \beta a_3 \).

Then if \( y = Ax = x_1a_1 + x_2a_2 + x_3a_3 \), we can also write

\[
y = x_1(aa_2 + \beta a_3) + x_2a_2 + x_3a_3 = (x_1a + x_2)a_2 + (x_1\beta + x_3)a_3
\]

In other words, uniqueness fails.

**Linear Equations with Julia**  Here’s an illustration of how to solve linear equations with Julia’s built-in linear algebra facilities.

```julia
julia> A = [1.0 2.0; 3.0 4.0];

julia> y = ones(2, 1); # A column vector
```
2.1. LINEAR ALGEBRA

```
 julia> det(A)
 -2.0

 julia> A_inv = inv(A)
 2x2 Array{Float64,2}:
  -2.0  1.0
   1.5 -0.5

 julia> x = A_inv * y  # solution
 2x1 Array{Float64,2}:
   -1.0
    1.0

 julia> A * x  # should equal y (a vector of ones)
 2x1 Array{Float64,2}:
   1.0
    1.0

 julia> A\y  # produces the same solution
 2x1 Array{Float64,2}:
   -1.0
    1.0
```

Observe how we can solve for $x = A^{-1}y$ by either via $\text{inv}(A) \times y$, or using $A \ \backslash \ y$

The latter method is preferred because it automatically selects the best algorithm for the problem based on the values of $A$ and $y$

If $A$ is not square then $A \ \backslash \ y$ returns the least squares solution $\hat{x} = (A' A)^{-1}A' y$

### Eigenvalues and Eigenvectors

Let $A$ be an $n \times n$ square matrix

If $\lambda$ is scalar and $v$ is a non-zero vector in $\mathbb{R}^n$ such that

$$Av = \lambda v$$

then we say that $\lambda$ is an eigenvalue of $A$, and $v$ is an eigenvector

Thus, an eigenvector of $A$ is a vector such that when the map $f(x) = Ax$ is applied, $v$ is merely scaled

The next figure shows two eigenvectors (blue arrows) and their images under $A$ (red arrows)

As expected, the image $Av$ of each $v$ is just a scaled version of the original

The eigenvalue equation is equivalent to $(A - \lambda I)v = 0$, and this has a nonzero solution $v$ only when the columns of $A - \lambda I$ are linearly dependent

This in turn is equivalent to stating that the determinant is zero

Hence to find all eigenvalues, we can look for $\lambda$ such that the determinant of $A - \lambda I$ is zero

This problem can be expressed as one of solving for the roots of a polynomial in $\lambda$ of degree $n$
This in turn implies the existence of $n$ solutions in the complex plane, although some might be repeated

Some nice facts about the eigenvalues of a square matrix $A$ are as follows

1. The determinant of $A$ equals the product of the eigenvalues
2. The trace of $A$ (the sum of the elements on the principal diagonal) equals the sum of the eigenvalues
3. If $A$ is symmetric, then all of its eigenvalues are real
4. If $A$ is invertible and $\lambda_1, \ldots, \lambda_n$ are its eigenvalues, then the eigenvalues of $A^{-1}$ are $1/\lambda_1, \ldots, 1/\lambda_n$

A corollary of the first statement is that a matrix is invertible if and only if all its eigenvalues are nonzero

Using Julia, we can solve for the eigenvalues and eigenvectors of a matrix as follows

```
 julia> A = [1.0 2.0; 2.0 1.0];
 julia> evals, evecs = eig(A);
 julia> evals
 2-element Array{Float64,1}:
  -1.0
  3.0
 julia> evecs
 2x2 Array{Float64,2}:
 -0.707107  0.707107
  0.707107  0.707107
```
Note that the columns of \texttt{evecs} are the eigenvectors.

Since any scalar multiple of an eigenvector is an eigenvector with the same eigenvalue (check it), the \texttt{eig} routine normalizes the length of each eigenvector to one.

**Generalized Eigenvalues**  
It is sometimes useful to consider the generalized eigenvalue problem, which, for given matrices \(A\) and \(B\), seeks generalized eigenvalues \(\lambda\) and eigenvectors \(v\) such that

\[
Av = \lambda Bv
\]

This can be solved in Julia via \texttt{eig}(\(A\), \(B\)).

Of course, if \(B\) is square and invertible, then we can treat the generalized eigenvalue problem as an ordinary eigenvalue problem \(B^{-1}Av = \lambda v\), but this is not always the case.

**Further Topics**

We round out our discussion by briefly mentioning several other important topics.

**Series Expansions**  
Recall the usual summation formula for a geometric progression, which states that if \(|a| < 1\), then \(\sum_{k=0}^{\infty} a^k = (1 - a)^{-1}\).

A generalization of this idea exists in the matrix setting.

**Matrix Norms**  
Let \(A\) be a square matrix, and let

\[
\|A\| := \max_{\|x\|=1} \|Ax\|
\]

The norms on the right-hand side are ordinary vector norms, while the norm on the left-hand side is a matrix norm — in this case, the so-called spectral norm.

For example, for a square matrix \(S\), the condition \(\|S\| < 1\) means that \(S\) is contractive, in the sense that it pulls all vectors towards the origin.\(^1\)

**Neumann’s Theorem**  
Let \(A\) be a square matrix and let \(A^k := AA^{k-1}\) with \(A^1 := A\).

In other words, \(A^k\) is the \(k\)-th power of \(A\).

Neumann’s theorem states the following: If \(\|A^k\| < 1\) for some \(k \in \mathbb{N}\), then \(I - A\) is invertible, and

\[
(I - A)^{-1} = \sum_{k=0}^{\infty} A^k
\]

\(^1\) Suppose that \(\|S\| < 1\). Take any nonzero vector \(x\), and let \(r := \|x\|\). We have \(\|Sx\| = r\|S(x/r)\| \leq r\|S\| < r = \|x\|\). Hence every point is pulled towards the origin.
2.1. LINEAR ALGEBRA

Spectral Radius  A result known as Gelfand’s formula tells us that, for any square matrix $A$,  
\[ \rho(A) = \lim_{k \to \infty} \| A^k \|^{1/k} \]
Here $\rho(A)$ is the spectral radius, defined as $\max_{\lambda_i} |\lambda_i|$, where \{\lambda_i\} is the set of eigenvalues of $A$.
As a consequence of Gelfand’s formula, if all eigenvalues are strictly less than one in modulus, there exists a $k$ with $\| A^k \| < 1$.
In which case (2.4) is valid.

Positive Definite Matrices  Let $A$ be a symmetric $n \times n$ matrix.
We say that $A$ is

1. positive definite if $x'Ax > 0$ for every $x \in \mathbb{R}^n \setminus \{0\}$
2. positive semi-definite or nonnegative definite if $x'Ax \geq 0$ for every $x \in \mathbb{R}^n$

Analogous definitions exist for negative definite and negative semi-definite matrices.
It is notable that if $A$ is positive definite, then all of its eigenvalues are strictly positive, and hence $A$ is invertible (with positive definite inverse).

Differentiating Linear and Quadratic forms  The following formulas are useful in many economic contexts. Let

- $z$, $x$ and $a$ all be $n \times 1$ vectors
- $A$ be an $n \times n$ matrix
- $B$ be an $m \times n$ matrix and $y$ be an $m \times 1$ vector

Then

1. $\frac{\partial x'x}{\partial x} = a$
2. $\frac{\partial Ax}{\partial x} = A'$
3. $\frac{\partial x'Ax}{\partial z} = (A + A')x$
4. $\frac{\partial y'Bz}{\partial y} = Bz$
5. $\frac{\partial y'Bz}{\partial B} = yz'$

An Example  Let $x$ be a given $n \times 1$ vector and consider the problem  
\[ v(x) = \max_{y,u} \{ -y'Py - u'Qu \} \]
subject to the linear constraint  
\[ y = Ax + Bu \]
Here
• $P$ is an $n \times n$ matrix and $Q$ is an $m \times m$ matrix
• $A$ is an $n \times n$ matrix and $B$ is an $n \times m$ matrix
• both $P$ and $Q$ are symmetric and positive semidefinite

Question: what must the dimensions of $y$ and $u$ be to make this a well-posed problem?

One way to solve the problem is to form the Lagrangian

$$
\mathcal{L} = -y'Py - u'Qu + \lambda' [Ax + Bu - y]
$$

where $\lambda$ is an $n \times 1$ vector of Lagrange multipliers

Try applying the above formulas for differentiating quadratic and linear forms to obtain the first-order conditions for maximizing $\mathcal{L}$ with respect to $y$, $u$ and minimizing it with respect to $\lambda$

Show that these conditions imply that

1. $\lambda = -2Py$
2. The optimizing choice of $u$ satisfies $u = -(Q + B'PB)^{-1}B'PAx$
3. The function $v$ satisfies $v(x) = -x'\tilde{P}x$ where $\tilde{P} = A'PA - A'PB(Q + B'PB)^{-1}B'PA$

As we will see, in economic contexts Lagrange multipliers often are shadow prices

**Note:** If we don’t care about the Lagrange multipliers, we can substitute the constraint into the objective function, and then just maximize $-(Ax + Bu)'P(Ax + Bu) - u'Qu$ with respect to $u$. You can verify that this leads to the same maximizer.

**Further Reading** The documentation of the linear algebra features built into Julia can be found here

Chapter 2 of these notes contains a discussion of linear algebra along the same lines as above, with solved exercises

If you don’t mind a slightly abstract approach, a nice intermediate-level read on linear algebra is [Janich94]

2.2 Finite Markov Chains
2.2. FINITE MARKOV CHAINS

Contents

• Finite Markov Chains
  – Overview
  – Definitions
  – Simulation
  – Marginal Distributions
  – Stationary Distributions
  – Ergodicity
  – Forecasting Future Values
  – Exercises
  – Solutions

Overview

Markov chains are one of the most useful classes of stochastic processes

Attributes:

• simple, flexible and supported by many elegant theoretical results
• valuable for building intuition about random dynamic models
• very useful in their own right

You will find them in many of the workhorse models of economics and finance

In this lecture we review some of the theory of Markov chains, with a focus on numerical methods

Prerequisite knowledge is basic probability and linear algebra

Definitions

The following concepts are fundamental

Stochastic Matrices  A stochastic matrix (or Markov matrix) is an $n \times n$ square matrix \( P = P[i,j] \) such that

1. each element $P[i,j]$ is nonnegative, and
2. each row $P[i,:]$ sums to one

Let $S := \{1, \ldots, n\}$

Evidently, each row $P[i,:]$ can be regarded as a distribution (probability mass function) on $S$

It is not difficult to check\(^2\) that if $P$ is a stochastic matrix, then so is the $k$-th power $P^k$ for all $k \in \mathbb{N}$

\(^2\) Hint: First show that if $P$ and $Q$ are stochastic matrices then so is their product — to check the row sums, try postmultiplying by a column vector of ones. Finally, argue that $P^n$ is a stochastic matrix using induction.
Markov Chains  A stochastic matrix describes the dynamics of a Markov chain \( \{X_t\} \) that takes values in the state space \( S \).

Formally, we say that a discrete time stochastic process \( \{X_t\} \) taking values in \( S \) is a Markov chain with stochastic matrix \( P \) if

\[
\Pr\{X_{t+1} = j \mid X_t = i\} = P[i,j]
\]

for any \( t \geq 0 \) and \( i, j \in S \); here \( \Pr \) means probability.

Remark: This definition implies that \( \{X_t\} \) has the Markov property, which is to say that, for any \( t \),

\[
\Pr\{X_{t+1} \mid X_t\} = \Pr\{X_{t+1} \mid X_t, X_{t-1}, \ldots\}
\]

Thus the state \( X_t \) is a complete description of the current position of the system.

Thus, by construction,

- \( P[i,j] \) is the probability of going from \( i \) to \( j \) in one unit of time (one step).
- \( P[i,\cdot] \) is the conditional distribution of \( X_{t+1} \) given \( X_t = i \).

Another way to think about this process is to imagine that, when \( X_t = i \), the next value \( X_{t+1} \) is drawn from the \( i \)-th row \( P[i,\cdot] \).

Rephrasing this using more algorithmic language

- At each \( t \), the new state \( X_{t+1} \) is drawn from \( P[X_t,\cdot] \).

**Example 1**  Consider a worker who, at any given time \( t \), is either unemployed (state 1) or employed (state 2).

Let’s write this mathematically as \( X_t = 1 \) or \( X_t = 2 \).

Suppose that, over a one month period,

1. An employed worker loses her job and becomes unemployed with probability \( \beta \in (0,1) \)
2. An unemployed worker finds a job with probability \( \alpha \in (0,1) \)

In terms of a stochastic matrix, this tells us that \( P[1,2] = \alpha \) and \( P[2,1] = \beta \), or

\[
P = \begin{pmatrix} 1 - \alpha & \alpha \\ \beta & 1 - \beta \end{pmatrix}
\]

Once we have the values \( \alpha \) and \( \beta \), we can address a range of questions, such as

- What is the average duration of unemployment?
- Over the long-run, what fraction of time does a worker find herself unemployed?
- Conditional on employment, what is the probability of becoming unemployed at least once over the next 12 months?
- Etc.

We’ll cover such applications below.
Example 2  Using US unemployment data, Hamilton [Ham05] estimated the stochastic matrix

\[ P := \begin{pmatrix}
0.971 & 0.029 & 0 \\
0.145 & 0.778 & 0.077 \\
0 & 0.508 & 0.492
\end{pmatrix} \]

where
- the frequency is monthly
- the first state represents “normal growth”
- the second state represents “mild recession”
- the third state represents “severe recession”

For example, the matrix tells us that when the state is normal growth, the state will again be normal growth next month with probability 0.97.

In general, large values on the main diagonal indicate persistence in the process \( \{X_t\} \).

This Markov process can also be represented as a directed graph, with edges labeled by transition probabilities.

![Directed Graph of Markov Chain]

Here “ng” is normal growth, “mr” is mild recession, etc.

Simulation

One of the most natural ways to answer questions about Markov chains is to simulate them.

(As usual, to approximate the probability of event \( E \), we can simulate many times and count the fraction of times that \( E \) occurs.)

To simulate a Markov chain, we need its stochastic matrix \( P \) and a probability distribution \( \psi \) for the initial state.

Here \( \psi \) is a probability distribution on \( S \) with the interpretation that \( X_0 \) is drawn from \( \psi \).

The Markov chain is then constructed via the following two rules:

1. At time \( t = 0 \), the initial state \( X_0 \) is drawn from \( \psi \).
2. At each subsequent time \( t \), the new state \( X_{t+1} \) is drawn from \( P[X_t, \cdot] \).

In order to implement this simulation procedure, we need a function for generating draws from a given discrete distribution.
We already have this functionality in hand—in the file `discrete_rv.jl`

The module is part of the `QuantEcon` package, and defines a type `DiscreteRV` that can be used as follows

```julia
julia> using QuantEcon

julia> psi = [0.1, 0.9];

julia> d = DiscreteRV(psi);

julia> draw(d, 5)
5-element Array{Int64,1}:
    1
    2
    2
    1
    2
```

Here

- `psi` is understood to be a discrete distribution on the set of outcomes `1, ..., length(psi)`
- `draw(d, 5)` generates 5 independent draws from this distribution

Let's now write a function that generates time series from a specified pair `P, ψ`

Our function will take the following three arguments

- A stochastic matrix `P`,
- An initial state or distribution `init`
- A positive integer `sample_size` representing the length of the time series the function should return

Let's allow `init` to either be

- an integer in `1, ..., n` providing a fixed starting value for `X_0`, or
- a discrete distribution on this same set that corresponds to the initial distribution `ψ`

In the latter case, a random starting value for `X_0` is drawn from the distribution `init`

The function should return a time series (sample path) of length `sample_size`

One solution to this problem can be found in file `mc_tools.jl` from the `QuantEcon` package

The relevant function is `mc_sample_path`

Let's see how it works using the small matrix

\[
P := \begin{pmatrix} 0.4 & 0.6 \\ 0.2 & 0.8 \end{pmatrix}
\] (2.5)

It happens to be true that, for a long series drawn from `P`, the fraction of the sample that takes value 1 will be about 0.25 — we'll see why later on

If you run the following code you should get roughly that answer
using QuantEcon
P = [.4 .6
     .2 .8];
s = mc_sample_path(P, [0.5, 0.5], 100000);
println(mean(s .== 1))  # Should be about 0.25

Marginal Distributions

Suppose that

1. \( \{X_t\} \) is a Markov chain with stochastic matrix \( P \)
2. the distribution of \( X_t \) is known to be \( \psi_t \)

What then is the distribution of \( X_{t+1} \), or, more generally, of \( X_{t+m} \)?
(Motivation for these questions is given below)

Solution  Let’s consider how to solve for the distribution \( \psi_{t+m} \) of \( X_{t+m} \), beginning with the case \( m = 1 \)

Throughout, \( \psi_t \) will refer to the distribution of \( X_t \) for all \( t \)
Hence our first aim is to find \( \psi_{t+1} \) given \( \psi_t \) and \( P \)
To begin, pick any \( j \in S \).
Using the law of total probability, we can decompose the probability that \( X_{t+1} = j \) as follows:

\[
P\{X_{t+1} = j\} = \sum_{i \in S} P\{X_{t+1} = j \mid X_t = i\} \cdot P\{X_t = i\}
\]
(In words, to get the probability of being at \( j \) tomorrow, we account for all ways this can happen and sum their probabilities)

Rewriting this statement in terms of marginal and conditional probabilities gives

\[
\psi_{t+1}[j] = \sum_{i \in S} P[i, j] \psi_t[i]
\]

There are \( n \) such equations, one for each \( j \in S \)
If we think of \( \psi_{t+1} \) and \( \psi_t \) as row vectors, these \( n \) equations are summarized by the matrix expression

\[
\psi_{t+1} = \psi_t P
\]
In other words, to move the distribution forward one unit of time, we postmultiply by \( P \)
By repeating this \( m \) times we move forward \( m \) steps into the future
Hence \( \psi_{t+m} = \psi_t P^m \) is also valid — here \( P^m \) is the \( m \)-th power of \( P \) As a special case, we see that if \( \psi_0 \) is the initial distribution from which \( X_0 \) is drawn, then \( \psi_0 P^m \) is the distribution of \( X_m \)
This is very important, so let’s repeat it

\[
X_0 \sim \psi_0 \implies X_m \sim \psi_0 P^m \tag{2.6}
\]
and, more generally,

\[ X_t \sim \psi_t \implies X_{t+m} \sim \psi_t P^m \]  \hspace{1cm} (2.7)

**Note:** Unless stated otherwise, we follow the common convention in the Markov chain literature that distributions are row vectors.

**Example: Powers of a Markov Matrix** We know that the probability of transitioning from \( i \) to \( j \) in one step is \( P[i, j] \).

It turns out that the probability of transitioning from \( i \) to \( j \) in \( m \) steps is \( P^m[i, j] \), the \([i, j]\)-th element of the \( m \)-th power of \( P \).

To see why, consider again (2.7), but now with \( \psi_t \) put all probability on state \( i \).

If we regard \( \psi_t \) as a vector, it is a vector with 1 in the \( i \)-th position and zero elsewhere.

Inserting this into (2.7), we see that, conditional on \( X_t = i \), the distribution of \( X_{t+m} \) is the \( i \)-th row of \( P^m \).

In particular

\[ P\{X_{t+m} = j\} = P^m[i, j] = [i, j]\text{-th element of } P^m \]

**Example: Future Probabilities** Recall the stochastic matrix \( P \) for recession and growth considered above.

Suppose that the current state is unknown — perhaps statistics are available only at the end of the current month.

We estimate the probability that the economy is in state \( i \) to be \( \psi[i] \).

The probability of being in recession (state 1 or state 2) in 6 months time is given by the inner product

\[ \psi P^6 \cdot \begin{pmatrix} 0 \\ 1 \\ 1 \end{pmatrix} \]

**Example 2: Cross-Sectional Distributions** Recall our model of employment / unemployment dynamics for a given worker discussed above.

Consider a large (i.e., tending to infinite) population of workers, each of whose lifetime experiences are described by the specified dynamics, independently of one another.

Let \( \psi \) be the current cross-sectional distribution over \( \{1, 2\} \).

- For example, \( \psi[1] \) is the unemployment rate.

The cross-sectional distribution records the fractions of workers employed and unemployed at a given moment.

The same distribution also describes the fractions of a particular worker’s career spent being employed and unemployed, respectively.
Stationary Distributions

As stated in the previous section, we can shift probabilities forward one unit of time via postmultiplication by $P$

Some distributions are invariant under this updating process — for example,

```julia
julia> P = [.4 .6; .2 .8];

julia> psi = [0.25, 0.75];

julia> psi'*P
1x2 Array{Float64,2}:
0.25 0.75
```

Such distributions are called stationary, or invariant. Formally, a distribution $\psi^*$ on $S$ is called stationary for $P$ if $\psi^* = \psi^* P$

From this equality we immediately get $\psi^* = \psi^* P^t$ for all $t$

This tells us an important fact: If the distribution of $X_0$ is a stationary distribution, then $X_t$ will have this same distribution for all $t$

Hence stationary distributions have a natural interpretation as stochastic steady states — we’ll discuss this more in just a moment

Mathematically, a stationary distribution is just a fixed point of $P$ when $P$ is thought of as the map $\psi \mapsto \psi P$ from (row) vectors to (row) vectors

At least one such distribution exists for each stochastic matrix $P$ — apply Brouwer’s fixed point theorem, or see EDTC, theorem 4.3.5

There may in fact be many stationary distributions corresponding to a given stochastic matrix $P$

For example, if $P$ is the identity matrix, then all distributions are stationary

One sufficient condition for uniqueness is uniform ergodicity:

**Def.** Stochastic matrix $P$ is called *uniformly ergodic* if there exists a positive integer $m$ such that all elements of $P^m$ are strictly positive

For further details on uniqueness and uniform ergodicity, see, for example, EDTC, theorem 4.3.18

**Example** Recall our model of employment / unemployment dynamics for a given worker discussed above

Assuming $\alpha \in (0, 1)$ and $\beta \in (0, 1)$, the uniform ergodicity condition is satisfied

Let $\psi^* = (p, 1 - p)$ be the stationary distribution, so that $p$ corresponds to unemployment (state 1)

Using $\psi^* = \psi^* P$ and a bit of algebra yields

$$p = \frac{\beta}{\alpha + \beta}$$
This is, in some sense, a steady state probability of unemployment — more on interpretation below. Not surprisingly it tends to zero as $\beta \to 0$, and to one as $\alpha \to 0$.

**Calculating Stationary Distribution**  
As discussed above, a given Markov matrix $P$ can have many stationary distributions. That is, there can be many row vectors $\psi$ such that $\psi = \psi P$.

In fact if $P$ has two distinct stationary distributions $\psi_1, \psi_2$ then it has infinitely many, since in this case, as you can verify,

$$
\psi_3 := \lambda \psi_1 + (1 - \lambda) \psi_2
$$

is a stationary distribution for $P$ for any $\lambda \in [0, 1]$.

If we restrict attention to the case where only one stationary distribution exists, one option for finding it is to try to solve the linear system $\psi (I_n - P) = 0$ for $\psi$, where $I_n$ is the $n \times n$ identity.

But the zero vector solves this equation. Hence we need to impose the restriction that the solution must be a probability distribution.

One function that will do this for us and implement a suitable algorithm is `mc_compute_stationary` from `mc_tools.jl`.

Let’s test it using the matrix (2.5):

```julia
using QuantEcon
P = [.4 .6; .2 .8]
println(mc_compute_stationary(P))
```

If you run this you should find that the unique stationary distribution is (0.25, 0.75).

**Convergence to Stationarity**  
Let $P$ be a stochastic matrix such that the uniform ergodicity assumption is valid.

We know that under this condition there is a unique stationary distribution $\psi^*$.

In fact, under the same condition, we have another important result: for any nonnegative row vector $\psi$ summing to one (i.e., distribution),

$$
\psi P^t \to \psi^* \quad \text{as} \quad t \to \infty
$$

(2.8)

In view of our preceding discussion, this states that the distribution of $X_t$ converges to $\psi^*$, regardless of the distribution of $X_0$.

This adds considerable weight to our interpretation of $\psi^*$ as a stochastic steady state.

For one of several well-known proofs, see EDTC, theorem 4.3.18.

The convergence in (2.8) is illustrated in the next figure.

Here

- $P$ is the stochastic matrix for recession and growth considered above.
2.2. FINITE MARKOV CHAINS

- The highest red dot is an arbitrarily chosen initial probability distribution $\psi$, represented as a vector in $\mathbb{R}^3$.
- The other red dots are the distributions $\psi P^t$ for $t = 1, 2, \ldots$.
- The black dot is $\psi^*$.

The code for the figure can be found in the file `examples/mc_convergence_plot.jl` in the main repository — you might like to try experimenting with different initial conditions.

**Ergodicity**

Under the very same condition of uniform ergodicity, yet another important result obtains: If

1. $\{X_t\}$ is a Markov chain with stochastic matrix $P$.
2. $P$ is uniformly ergodic with stationary distribution $\psi^*$.

then, $\forall j \in S$, 

$$\frac{1}{n} \sum_{t=1}^{n} 1\{X_t = j\} \to \psi^*[j] \quad \text{as } n \to \infty \quad (2.9)$$

Here

- $1\{X_t = j\} = 1$ if $X_t = j$ and zero otherwise.
- convergence is with probability one.
- the result does not depend on the distribution (or value) of $X_0$.

The result tells us that the fraction of time the chain spends at state $j$ converges to $\psi^*[j]$ as time goes to infinity. This gives us another way to interpret the stationary distribution — provided that the convergence result in (2.9) is valid.
Technically, the convergence in (2.9) is a special case of a law of large numbers result for Markov chains — see EDTC, section 4.3.4 for details.

**Example**  Recall our cross-sectional interpretation of the employment / unemployment model discussed above. Assume that $\alpha \in (0, 1)$ and $\beta \in (0, 1)$, so the uniform ergodicity condition is satisfied. We saw that the stationary distribution is $(p, 1 - p)$, where

$$p = \frac{\beta}{\alpha + \beta}$$

In the cross-sectional interpretation, this is the fraction of people unemployed. In view of our latest (ergodicity) result, it is also the fraction of time that a worker can expect to spend unemployed. Thus, in the long-run, cross-sectional averages for a population and time-series averages for a given person coincide. This is one interpretation of the notion of ergodicity.

**Forecasting Future Values**

Let $P$ be an $n \times n$ stochastic matrix with

$$P_{ij} = \mathbb{P}\{x_{t+1} = e_j \mid x_t = e_i\}$$

where $e_i$ is the $i$-th unit vector in $\mathbb{R}^n$. We are said to be “in state $i$” when $x_t = e_i$. Let $\bar{y}$ be an $n \times 1$ vector and let $y_t = \bar{y}'x_t$. In other words, $y_t = \bar{y}_i$ if $x_t = e_i$.

Here are some useful prediction formulas:

$$\mathbb{E} [y_{t+k} \mid x_t = e_i] = \sum_j (P^k)_{ij} \bar{y}_j = (P^k \bar{y})_i$$

for $k = 0, 1, 2, \ldots$, and

$$\mathbb{E} \left[ \sum_{j=0}^{\infty} \beta^j y_{t+j} \mid x_t = e_i \right] = [(I - \beta P)^{-1} \bar{y}]_i$$

where $(P^k)_{ij}$ is the $ij$-th element of $P^k$ and

$$(I - \beta P)^{-1} = I + \beta P + \beta^2 P^2 + \cdots$$

Premultiplication by $(I - \beta P)^{-1}$ amounts to “applying the resolvent operator”.
Exercises

Exercise 1  According to the discussion immediately above, if a worker’s employment dynamics obey the stochastic matrix

\[ P = \begin{pmatrix} 1 - \alpha & \alpha \\ \beta & 1 - \beta \end{pmatrix} \]

with \( \alpha \in (0, 1) \) and \( \beta \in (0, 1) \), then, in the long-run, the fraction of time spent unemployed will be

\[ p := \frac{\beta}{\alpha + \beta} \]

In other words, if \( \{X_t\} \) represents the Markov chain for employment, then \( \bar{X}_n \to p \) as \( n \to \infty \), where

\[ \bar{X}_n := \frac{1}{n} \sum_{t=1}^{n} 1\{X_t = 1\} \]

Your exercise is to illustrate this convergence

First,

- generate one simulated time series \( \{X_t\} \) of length 10,000, starting at \( X_0 = 1 \)
- plot \( \bar{X}_n - p \) against \( n \), where \( p \) is as defined above

Second, repeat the first step, but this time taking \( X_0 = 2 \)

In both cases, set \( \alpha = \beta = 0.1 \)

The result should look something like the following — modulo randomness, of course

(You don’t need to add the fancy touches to the graph—see the solution if you’re interested)
Exercise 2  A topic of interest for economics and many other disciplines is ranking
Let’s now consider one of the most practical and important ranking problems — the rank assigned to web pages by search engines
(Although the problem is motivated from outside of economics, there is in fact a deep connection between search ranking systems and prices in certain competitive equilibria — see [DLP13])
To understand the issue, consider the set of results returned by a query to a web search engine
For the user, it is desirable to
1. receive a large set of accurate matches
2. have the matches returned in order, where the order corresponds to some measure of “importance”
Ranking according to a measure of importance is the problem we now consider
The methodology developed to solve this problem by Google founders Larry Page and Sergey Brin is known as PageRank
To illustrate the idea, consider the following diagram

Imagine that this is a miniature version of the WWW, with
- each node representing a web page
- each arrow representing the existence of a link from one page to another
Now let’s think about which pages are likely to be important, in the sense of being valuable to a search engine user
One possible criterion for importance of a page is the number of inbound links — an indication of popularity
By this measure, $m$ and $j$ are the most important pages, with 5 inbound links each
However, what if the pages linking to $m$, say, are not themselves important?
Thinking this way, it seems appropriate to weight the inbound nodes by relative importance
The PageRank algorithm does precisely this
A slightly simplified presentation that captures the basic idea is as follows

Letting \( j \) be (the integer index of) a typical page and \( r_j \) be its ranking, we set

\[
r_j = \sum_{i \in L_j} \frac{r_i}{\ell_i}
\]

where

- \( \ell_i \) is the total number of outbound links from \( i \)
- \( L_j \) is the set of all pages \( i \) such that \( i \) has a link to \( j \)

This is a measure of the number of inbound links, weighted by their own ranking (and normalized by \( 1/\ell_i \))

There is, however, another interpretation, and it brings us back to Markov chains

Let \( P \) be the matrix given by \( P[i, j] = 1\{i \rightarrow j\} / \ell_i \) where \( 1\{i \rightarrow j\} = 1 \) if \( i \) has a link to \( j \) and zero otherwise

The matrix \( P \) is a stochastic matrix provided that each page has at least one link

With this definition of \( P \) we have

\[
r_j = \sum_{i \in L_j} \frac{r_i}{\ell_i} = \sum_{i} 1\{i \rightarrow j\} \frac{r_i}{\ell_i} = \sum_{i} P[i, j] r_i
\]

Writing \( r \) for the row vector of rankings, this becomes \( r = rP \)

Hence \( r \) is the stationary distribution of the stochastic matrix \( P \)

Let’s think of \( P[i, j] \) as the probability of “moving” from page \( i \) to page \( j \)

The value \( P[i, j] \) has the interpretation

- \( P[i, j] = 1/k \) if \( i \) has \( k \) outbound links, and \( j \) is one of them
- \( P[i, j] = 0 \) if \( i \) has no direct link to \( j \)

Thus, motion from page to page is that of a web surfer who moves from one page to another by randomly clicking on one of the links on that page

Here “random” means that each link is selected with equal probability

Since \( r \) is the stationary distribution of \( P \), assuming that the uniform ergodicity condition is valid, we can interpret \( r_j \) as the fraction of time that a (very persistent) random surfer spends at page \( j \)

Your exercise is to apply this ranking algorithm to the graph pictured above, and return the list of pages ordered by rank

The data for this graph is in the `web_graph_data.txt` file from the main repository — you can also view it here

There is a total of 14 nodes (i.e., web pages), the first named \( a \) and the last named \( n \)

A typical line from the file has the form
This should be interpreted as meaning that there exists a link from \( d \) to \( h \).

To parse this file and extract the relevant information, you can use regular expressions.

The following code snippet provides a hint as to how you can go about this:

```julia
julia> matchall(r"\w", "x +++ y ***** z")
3-element Array{SubString{UTF8String},1}:
    "x"
    "y"
    "z"

julia> matchall(r"\w", "a ^\^ b && \$ \$ c")
3-element Array{SubString{UTF8String},1}:
    "a"
    "b"
    "c"
```

When you solve for the ranking, you will find that the highest ranked node is in fact \( g \), while the lowest is \( a \).

**Exercise 3** In numerical work it is sometimes convenient to replace a continuous model with a discrete one.

In particular, Markov chains are routinely generated as discrete approximations to AR(1) processes of the form

\[
y_{t+1} = \rho y_t + u_{t+1}
\]

Here \( u_t \) is assumed to be iid and \( N(0, \sigma_u^2) \).

The variance of the stationary probability distribution of \( \{y_t\} \) is

\[
\sigma_y^2 := \frac{\sigma_u^2}{1 - \rho^2}
\]

Tauchen’s method [Tau86] is the most common method for approximating this continuous state process with a finite state Markov chain.

As a first step we choose

- \( n \), the number of states for the discrete approximation
- \( m \), an integer that parameterizes the width of the state space

Next we create a state space \( \{x_0, \ldots, x_{n-1}\} \subseteq \mathbb{R} \) and a stochastic \( n \times n \) matrix \( P \) such that

- \( x_0 = -m \sigma_y \)
- \( x_{n-1} = m \sigma_y \)
- \( x_{i+1} = x_i + s \) where \( s = (x_{n-1} - x_0) / (n - 1) \)
- \( P[i, j] \) represents the probability of transitioning from \( x_i \) to \( x_j \)
Let $F$ be the cumulative distribution function of the normal distribution $N(0, \sigma^2_u)$.

The values $P[i, j]$ are computed to approximate the AR(1) process — omitting the derivation, the rules are as follows:

1. If $j = 0$, then set
   
   $$P[i, j] = P[i, 0] = F(x_0 - \rho x_i + s/2)$$

2. If $j = n - 1$, then set
   
   $$P[i, j] = P[i, n - 1] = 1 - F(x_{n-1} - \rho x_i - s/2)$$

3. Otherwise, set
   
   $$P[i, j] = F(x_j - \rho x_i + s/2) - F(x_j - \rho x_i - s/2)$$

The exercise is to write a function `approx_markov(rho, sigma_u, m=3, n=7)` that returns \( \{x_0, \ldots, x_{n-1}\} \subset \mathbb{R} \) and an $n \times n$ matrix $P$ as described above.

Solutions

Solution notebook

### 2.3 Shortest Paths

**Contents**

- Shortest Paths
  - Overview
  - Outline of the Problem
  - Finding Least-Cost Paths
  - Solving for $J$
  - Exercises
  - Solutions

**Overview**

The shortest path problem is a classic problem in mathematics and computer science with applications in

- Economics (sequential decision making, analysis of social networks, etc.)
- Operations research and transportation
- Robotics and artificial intelligence
- Telecommunication network design and routing
- Etc., etc.
For us, the shortest path problem also provides a simple introduction to the logic of dynamic programming, which is one of our key topics.

Variations of the methods we discuss are used millions of times every day, in applications such as Google Maps.

Outline of the Problem

The shortest path problem is one of finding how to traverse a graph from one specified node to another at minimum cost.

Consider the following graph:

We wish to travel from node (vertex) A to node G at minimum cost:

- Arrows (edges) indicate the movements we can take
- Numbers next to edges indicate the cost of traveling that edge

Possible interpretations of the graph include:

- Minimum cost for supplier to reach a destination
- Routing of packets on the internet (minimize time)
- Etc., etc.

For this simple graph, a quick scan of the edges shows that the optimal paths are:

- A, C, F, G at cost 8
- A, D, F, G at cost 8
Finding Least-Cost Paths

For large graphs we need a systematic solution.

Let $J(v)$ denote the minimum cost-to-go from node $v$, understood as the total cost from $v$ if we take the best route.

Suppose that we know $J(v)$ for each node $v$, as shown below for the graph from the preceding example.

\[
\begin{align*}
J(G) &= 0
\end{align*}
\]

Intuitively, the best path can now be found as follows:

- Start at $A$.
- From node $v$, move to any node that solves

\[
\min_{w \in F_v} \{c(v, w) + J(w)\}
\]

(2.10)

where

- $F_v$ is the set of nodes that can be reached from $v$ in one step.
- $c(v, w)$ is the cost of traveling from $v$ to $w$.

Hence, if we know the function $J$, then finding the best path is almost trivial.

But how to find $J$?

Some thought will convince you that, for every node $v$, the function $J$ satisfies

\[
J(v) = \min_{w \in F_v} \{c(v, w) + J(w)\}
\]

(2.11)
This is known as the Bellman equation

- That is, $J$ is the solution to the Bellman equation
- There are algorithms for computing the minimum cost-to-go function $J$

**Solving for $J$**

The standard algorithm for finding $J$ is to start with

$$J_0(v) = M \text{ if } v \neq \text{destination}, \text{ else } J_0(v) = 0$$

where $M$ is some large number

Now we use the following algorithm

1. Set $n = 0$
2. Set $J_{n+1}(v) = \min_{w \in F_v \{c(v, w) + J_n(w)\}}$ for all $v$
3. If $J_{n+1}$ and $J_n$ are not equal then increment $n$, go to 2

In general, this sequence converges to $J$—the proof is omitted

**Exercises**

**Exercise 1** Use the algorithm given above to find the optimal path (and its cost) for this graph

Here the line `node0, node1 0.04, node8 11.11, node14 72.21` means that from node0 we can go to

- node1 at cost 0.04
- node8 at cost 11.11
- node14 at cost 72.21

and so on

According to our calculations, the optimal path and its cost are like this

Your code should replicate this result

**Solutions**

Solution notebook

# 2.4 Schelling’s Segregation Model
2.4. SCHELLING’S SEGREGATION MODEL

Outline

In 1969, Thomas C. Schelling developed a simple but striking model of racial segregation [Sch69]. His model studies the dynamics of racially mixed neighborhoods. Like much of Schelling’s work, the model shows how local interactions can lead to surprising aggregate structure. In particular, it shows that relatively mild preference for neighbors of similar race can lead in aggregate to the collapse of mixed neighborhoods, and high levels of segregation. In recognition of this and other research, Schelling was awarded the 2005 Nobel Prize in Economic Sciences (joint with Robert Aumann).

In this lecture we (in fact you) will build and run a version of Schelling’s model.

The Model

We will cover a variation of Schelling’s model that is easy to program and captures the main idea.

Set Up

Suppose we have two types of people: orange people and green people. For the purpose of this lecture, we will assume there are 250 of each type. These agents all live on a single unit square. The location of an agent is just a point \((x, y)\), where \(0 < x, y < 1\).

Preferences

We will say that an agent is happy if half or more of her 10 nearest neighbors are of the same type. Here ‘nearest’ is in terms of Euclidean distance.

An agent who is not happy is called unhappy. An important point here is that agents are not averse to living in mixed areas. They are perfectly happy if half their neighbors are of the other color.
2.4. SCHELLING’S SEGREGATION MODEL

**Behavior** Initially, agents are mixed together (integrated)

In particular, the initial location of each agent is an independent draw from a bivariate uniform distribution on $S = (0, 1)^2$

Now, cycling through the set of all agents, each agent is now given the chance to stay or move

We assume that each agent will stay put if they are happy and move if unhappy

The algorithm for moving is as follows

1. Draw a random location in $S$
2. If happy at new location, move there
3. Else, go to step 1

In this way, we cycle continuously through the agents, moving as required

We continue to cycle until no one wishes to move

**Results**

Let’s have a look at the results we got when we coded and ran this model

As discussed above, agents are initially mixed randomly together

But after several cycles they become segregated into distinct regions

In this instance, the program terminated after 4 cycles through the set of agents, indicating that all agents had reached a state of happiness

What is striking about the pictures is how rapidly racial integration breaks down
2.4. SCHELLING’S SEGREGATION MODEL
2.4. SCHELLING’S SEGREGATION MODEL

This is despite the fact that people in the model don’t actually mind living mixed with the other type. Even with these preferences, the outcome is a high degree of segregation.

**Exercises**

Rather than show you the program that generated these figures, we’ll now ask you to write your own version. You can see our program at the end, when you look at the solution.

**Exercise 1**  Implement and run this simulation for yourself. Consider the following structure for your program:

Agents are modeled as objects.

(Have a look at *this lecture* if you’ve forgotten how to build your own objects)

Here’s an indication of how they might look:

* Data:

  * type (green or orange)
  * location

* Methods:

  * Determine whether happy or not given locations of other agents
2.5. LLN AND CLT

* If not happy, move
  * find a new location where happy

And here's some pseudocode for the main loop

```python
while agents are still moving
  for agent in agents
    give agent the opportunity to move
  end
end
```

Use 250 agents of each type

Solutions

Solution notebook

2.5 LLN and CLT

Contents

- LLN and CLT
  - Overview
  - Relationships
  - LLN
  - CLT
  - Exercises
  - Solutions

Overview

This lecture illustrates two of the most important theorems of probability and statistics: The law of large numbers (LLN) and the central limit theorem (CLT)

These beautiful theorems lie behind many of the most fundamental results in econometrics and quantitative economic modeling

The lecture is based around simulations that show the LLN and CLT in action

We also demonstrate how the LLN and CLT break down when the assumptions they are based on do not hold

In addition, we examine several useful extensions of the classical theorems, such as

- The delta method, for smooth functions of random variables
- The multivariate case
Some of these extensions are presented as exercises

**Relationships**

The CLT refines the LLN.

The LLN gives conditions under which sample moments converge to population moments as sample size increases.

The CLT provides information about the rate at which sample moments converge to population moments as sample size increases.

**LLN**

We begin with the law of large numbers, which tells us when sample averages will converge to their population means.

**The Classical LLN**

The classical law of large numbers concerns independent and identically distributed (IID) random variables.

Here is the strongest version of the classical LLN, known as Kolmogorov’s strong law.

Let $X_1, \ldots, X_n$ be independent and identically distributed scalar random variables, with common distribution $F$.

When it exists, let $\mu$ denote the common mean of this sample:

$$
\mu := \mathbb{E}X = \int xF(dx)
$$

In addition, let

$$
\bar{X}_n := \frac{1}{n} \sum_{i=1}^{n} X_i
$$

Kolmogorov’s strong law states that, if $\mathbb{E}|X|$ is finite, then

$$
P \{ \bar{X}_n \to \mu \text{ as } n \to \infty \} = 1
$$

What does this last expression mean?

Let’s think about it from a simulation perspective, imagining for a moment that our computer can generate perfect random samples (which of course it can’t).

Let’s also imagine that we can generate infinite sequences, so that the statement $\bar{X}_n \to \mu$ can be evaluated.

In this setting, (2.13) should be interpreted as meaning that the probability of the computer producing a sequence where $\bar{X}_n \to \mu$ fails to occur is zero.
The proof of Kolmogorov’s strong law is nontrivial – see, for example, theorem 8.3.5 of [Dud02].

On the other hand, we can prove a weaker version of the LLN very easily and still get most of the intuition.

The version we prove is as follows: If \( X_1, \ldots, X_n \) is IID with \( \mathbb{E}X_i^2 < \infty \), then, for any \( \epsilon > 0 \), we have

\[
P \{ |\bar{X}_n - \mu| \geq \epsilon \} \to 0 \quad \text{as} \quad n \to \infty
\]  

(2.14)

(This version is weaker because we claim only convergence in probability rather than almost sure convergence, and assume a finite second moment.)

To see that this is so, fix \( \epsilon > 0 \), and let \( \sigma^2 \) be the variance of each \( X_i \).

Recall the Chebyshev inequality, which tells us that

\[
P \{ |\bar{X}_n - \mu| \geq \epsilon \} \leq \frac{\mathbb{E}[(\bar{X}_n - \mu)^2]}{\epsilon^2}
\]  

(2.15)

Now observe that

\[
\mathbb{E}[(\bar{X}_n - \mu)^2] = \mathbb{E} \left\{ \left[ \frac{1}{n} \sum_{i=1}^{n} (X_i - \mu) \right]^2 \right\}
\]

\[
= \frac{1}{n^2} \sum_{i=1}^{n} \sum_{j=1}^{n} \mathbb{E}(X_i - \mu)(X_j - \mu)
\]

\[
= \frac{1}{n^2} \sum_{i=1}^{n} \mathbb{E}(X_i - \mu)^2
\]

\[
= \frac{\sigma^2}{n}
\]

Here the crucial step is at the third equality, which follows from independence.

Independence means that if \( i \neq j \), then the covariance term \( \mathbb{E}(X_i - \mu)(X_j - \mu) \) drops out.

As a result, \( n^2 - n \) terms vanish, leading us to a final expression that goes to zero in \( n \).

Combining our last result with (2.15), we come to the estimate

\[
P \{ |\bar{X}_n - \mu| \geq \epsilon \} \leq \frac{\sigma^2}{n\epsilon^2}
\]  

(2.16)

The claim in (2.14) is now clear.

Of course, if the sequence \( X_1, \ldots, X_n \) is correlated, then the cross-product terms \( \mathbb{E}(X_i - \mu)(X_j - \mu) \) are not necessarily zero.

While this doesn’t mean that the same line of argument is impossible, it does mean that if we want a similar result then the covariances should be “almost zero” for “most” of these terms.

In a long sequence, this would be true if, for example, \( \mathbb{E}(X_i - \mu)(X_j - \mu) \) approached zero when the difference between \( i \) and \( j \) became large.
In other words, the LLN can still work if the sequence $X_1, \ldots, X_n$ has a kind of “asymptotic independence”, in the sense that correlation falls to zero as variables become further apart in the sequence.

This idea is very important in time series analysis, and we’ll come across it again soon enough.

**Illustration**  Let’s now illustrate the classical IID law of large numbers using simulation.

In particular, we aim to generate some sequences of IID random variables and plot the evolution of $\bar{X}_n$ as $n$ increases.

Below is a figure that does just this (as usual, you can click on it to expand it).

It shows IID observations from three different distributions and plots $\bar{X}_n$ against $n$ in each case. The dots represent the underlying observations $X_i$ for $i = 1, \ldots, 100$.

In each of the three cases, convergence of $\bar{X}_n$ to $\mu$ occurs as predicted.

The figure was produced by `illustrates_lln.jl`, which is shown below (and can be found in the `examples` directory of the main repository).
The three distributions are chosen at random from a selection stored in the dictionary `distributions`.

#=
Visual illustration of the law of large numbers.

@author: Spencer Lyon <spencer.lyon@nyu.edu>

References
---------
Based off the original python file illustrates_lln.py

```
# using PyPlot
using Distributions

n = 100
srand(42)  # reproducible results

# == Arbitrary collection of distributions == #
distributions = {"student's t with 10 degrees of freedom" => TDist(10),
    "beta(2, 2)" => Beta(2.0, 2.0),
    "lognormal LN(0, 1/2)" => LogNormal(0.5),
    "gamma(5, 1/2)" => Gamma(5.0, 2.0),
    "poisson(4)" => Poisson(4),
    "exponential with lambda = 1" => Exponential(1)}

# == Create a figure and some axes == #
um_plots = 3
fig, axes = plt.subplots(num_plots, 1, figsize=(10, 10))
bbox = [0., 1.02, 1., .102]
legend_args = {:ncol => 2,
    :bbox_to_anchor => bbox,
    :loc => 3,
    :mode => "expand"}
subplots_adjust(hspace=0.5)

for ax in axes
    dist_names = collect(keys(distributions))
    # == Choose a randomly selected distribution == #
    name = dist_names[rand(1:length(dist_names))]
    dist = pop!(distributions, name)

    # == Generate n draws from the distribution == #
    data = rand(dist, n)

    # == Compute sample mean at each n == #
    sample_mean = Array(Float64, n)
    for i=1:n
        sample_mean[i] = mean(data[1:i])
    end
```
Infinite Mean  What happens if the condition $E|X| < \infty$ in the statement of the LLN is not satisfied?

This might be the case if the underlying distribution is heavy tailed — the best known example is the Cauchy distribution, which has density

$$f(x) = \frac{1}{\pi(1+x^2)} \quad (x \in \mathbb{R})$$

The next figure shows 100 independent draws from this distribution

Notice how extreme observations are far more prevalent here than the previous figure

Let’s now have a look at the behavior of the sample mean

Here we’ve increased $n$ to 1000, but the sequence still shows no sign of converging

Will convergence become visible if we take $n$ even larger?

The answer is no

To see this, recall that the characteristic function of the Cauchy distribution is

$$\phi(t) = \mathbb{E}e^{itX} = \int e^{itx} f(x) dx = e^{-|t|} \quad (2.17)$$
2.5. LLN AND CLT

Using independence, the characteristic function of the sample mean becomes
\[
E e^{it\bar{X}_n} = E \exp \left\{ i \frac{t}{n} \sum_{j=1}^{n} X_j \right\}
= E \prod_{j=1}^{n} \exp \left\{ i \frac{t}{n} X_j \right\}
= \prod_{j=1}^{n} E \exp \left\{ i \frac{t}{n} X_j \right\} = [\phi(t/n)]^n
\]

In view of (2.17), this is just \( e^{-|t|} \)

Thus, in the case of the Cauchy distribution, the sample mean itself has the very same Cauchy distribution, regardless of \( n \)

In particular, the sequence \( \bar{X}_n \) does not converge to a point

CLT

Next we turn to the central limit theorem, which tells us about the distribution of the deviation between sample averages and population means

**Statement of the Theorem**  The central limit theorem is one of the most remarkable results in all of mathematics

In the classical IID setting, it tells us the following: If the sequence \( X_1, \ldots, X_n \) is IID, with common mean \( \mu \) and common variance \( \sigma^2 \in (0, \infty) \), then
\[
\sqrt{n}(\bar{X}_n - \mu) \overset{d}{\to} N(0, \sigma^2) \quad \text{as} \quad n \to \infty
\] (2.18)

Here \( \overset{d}{\to} N(0, \sigma^2) \) indicates convergence in distribution to a centered (i.e, zero mean) normal with standard deviation \( \sigma \).
**Intuition**  The striking implication of the CLT is that for any distribution with finite second moment, the simple operation of adding independent copies always leads to a Gaussian curve.

A relatively simple proof of the central limit theorem can be obtained by working with characteristic functions (see, e.g., theorem 9.5.6 of [Dud02]).

The proof is elegant but almost anticlimactic, and it provides surprisingly little intuition.

In fact all of the proofs of the CLT that we know are similar in this respect.

Why does adding independent copies produce a bell-shaped distribution?

Part of the answer can be obtained by investigating addition of independent Bernoulli random variables.

In particular, let $X_i$ be binary, with $P\{X_i = 0\} = P\{X_i = 1\} = 0.5$, and let $X_1, \ldots, X_n$ be independent.

Think of $X_i = 1$ as a “success”, so that $Y_n = \sum_{i=1}^n X_i$ is the number of successes in $n$ trials.

The next figure plots the probability mass function of $Y_n$ for $n = 1, 2, 4, 8$.

When $n = 1$, the distribution is flat — one success or no successes have the same probability.

When $n = 2$ we can either have 0, 1 or 2 successes.

Notice the peak in probability mass at the mid-point $k = 1$.

The reason is that there are more ways to get 1 success (“fail then succeed” or “succeed then fail”) than to get zero or two successes.

Moreover, the two trials are independent, so the outcomes “fail then succeed” and “succeed then fail” are just as likely as the outcomes “fail then fail” and “succeed then succeed.”
(If there was positive correlation, say, then “succeed then fail” would be less likely than “succeed then succeed”)

Here, already we have the essence of the CLT: addition under independence leads probability mass to pile up in the middle and thin out at the tails.

For \( n = 4 \) and \( n = 8 \) we again get a peak at the “middle” value (halfway between the minimum and the maximum possible value).

The intuition is the same — there are simply more ways to get these middle outcomes.

If we continue, the bell-shaped curve becomes ever more pronounced.

We are witnessing the binomial approximation of the normal distribution.

**Simulation 1**  Since the CLT seems almost magical, running simulations that verify its implications is one good way to build intuition.

To this end, we now perform the following simulation:

1. Choose an arbitrary distribution \( F \) for the underlying observations \( X_i \)
2. Generate independent draws of \( Y_n := \sqrt{n}(\bar{X}_n - \mu) \)
3. Use these draws to compute some measure of their distribution — such as a histogram
4. Compare the latter to \( N(0, \sigma^2) \)

Here’s some code that does exactly this for the exponential distribution \( F(x) = 1 - e^{-\lambda x} \)

(Please experiment with other choices of \( F \), but remember that, to conform with the conditions of the CLT, the distribution must have finite second moment.)
# == Compute mean of each row, producing k draws of \( \bar X_n \) == #

```python
sample_means = mean(data, 2)
```

# == Generate observations of \( Y_n \) == #

```python
Y = sqrt(n) * (sample_means .- mu)
```

# == Plot == #

```python
fig, ax = subplots()
xmin, xmax = -3 * s, 3 * s
ax[:set_xlim](xmin, xmax)
ax[:hist](Y, bins=60, alpha=0.5, normed=true)
xgrid = linspace(xmin, xmax, 200)
ax[:plot](xgrid, pdf(Normal(0.0, s), xgrid), "k-", lw=2, label=LaTeXString("\( \mathcal{N}(0, \sigma^2=\( s^2 \)) \))")
ax[:legend]()
```

The file is `illustrates_clt.jl`, from the main repository.

The program produces figures such as the one below.

The fit to the normal density is already tight, and can be further improved by increasing \( n \)

You can also experiment with other specifications of \( F \)

**Simulation 2**  Our next simulation is somewhat like the first, except that we aim to track the distribution of \( Y_n := \sqrt{n}(\bar{X}_n - \mu) \) as \( n \) increases.

In the simulation we’ll be working with random variables having \( \mu = 0 \)

Thus, when \( n = 1 \), we have \( Y_1 = X_1 \), so the first distribution is just the distribution of the underlying random variable.

For \( n = 2 \), the distribution of \( Y_2 \) is that of \( (X_1 + X_2)/\sqrt{2} \), and so on.
What we expect is that, regardless of the distribution of the underlying random variable, the distribution of $Y_n$ will smooth out into a bell shaped curve.

The next figure shows this process for $X_i \sim f$, where $f$ was specified as the convex combination of three different beta densities.

(Taking a convex combination is an easy way to produce an irregular shape for $f$)

In the figure, the closest density is that of $Y_1$, while the furthest is that of $Y_5$

As expected, the distribution smooths out into a bell curve as $n$ increases.

The figure is generated by file examples/clt3d.jl, which is available from the main repository.

We leave you to investigate its contents if you wish to know more.

If you run the file from the ordinary Julia or IJulia shell, the figure should pop up in a window that you can rotate with your mouse, giving different views on the density sequence.

**The Multivariate Case**  The law of large numbers and central limit theorem work just as nicely in multidimensional settings.

To state the results, let’s recall some elementary facts about random vectors.

A random vector $X$ is just a sequence of $k$ random variables $(X_1, \ldots, X_k)$.

Each realization of $X$ is an element of $\mathbb{R}^k$.

A collection of random vectors $X_1, \ldots, X_n$ is called independent if, given any $n$ vectors $x_1, \ldots, x_n$ in $\mathbb{R}^k$, we have

$$
\mathbb{P}\{X_1 \leq x_1, \ldots, X_n \leq x_n\} = \mathbb{P}\{X_1 \leq x_1\} \times \cdots \times \mathbb{P}\{X_n \leq x_n\}
$$
2.5. LLN AND CLT

(The vector inequality $X \leq x$ means that $X_j \leq x_j$ for $j = 1, \ldots, k$)

Let $\mu_j := E[X_j]$ for all $j = 1, \ldots, k$

The expectation $E[X]$ of $X$ is defined to be the vector of expectations:

$$E[X] := \begin{pmatrix} E[X_1] \\ E[X_2] \\ \vdots \\ E[X_k] \end{pmatrix} = \begin{pmatrix} \mu_1 \\ \mu_2 \\ \vdots \\ \mu_k \end{pmatrix} =: \mu$$

The variance-covariance matrix of random vector $X$ is defined as

$$\text{Var}[X] := E[(X - \mu)(X - \mu)']$$

Expanding this out, we get

$$\text{Var}[X] = \begin{pmatrix} E[(X_1 - \mu_1)(X_1 - \mu_1)] & \cdots & E[(X_1 - \mu_1)(X_k - \mu_k)] \\ E[(X_2 - \mu_2)(X_1 - \mu_1)] & \cdots & E[(X_2 - \mu_2)(X_k - \mu_k)] \\ \vdots & \vdots & \vdots \\ E[(X_k - \mu_k)(X_1 - \mu_1)] & \cdots & E[(X_k - \mu_k)(X_k - \mu_k)] \end{pmatrix}$$

The $j, k$-th term is the scalar covariance between $X_j$ and $X_k$

With this notation we can proceed to the multivariate LLN and CLT

Let $X_1, \ldots, X_n$ be a sequence of independent and identically distributed random vectors, each one taking values in $\mathbb{R}^k$

Let $\mu$ be the vector $E[X_i]$, and let $\Sigma$ be the variance-covariance matrix of $X_i$

Interpreting vector addition and scalar multiplication in the usual way (i.e., pointwise), let

$$\bar{X}_n := \frac{1}{n} \sum_{i=1}^n X_i$$

In this setting, the LLN tells us that

$$\mathbb{P}\{\bar{X}_n \to \mu \text{ as } n \to \infty\} = 1$$

(2.19)

Here $\bar{X}_n \to \mu$ means that $||\bar{X}_n - \mu|| \to 0$, where $|| \cdot ||$ is the standard Euclidean norm

The CLT tells us that, provided $\Sigma$ is finite,

$$\sqrt{n}(\bar{X}_n - \mu) \xrightarrow{d} N(0, \Sigma) \text{ as } n \to \infty$$

(2.20)

Exercises

Exercise 1 One very useful consequence of the central limit theorem is as follows

Assume the conditions of the CLT as stated above
If \( g : \mathbb{R} \rightarrow \mathbb{R} \) is differentiable at \( \mu \) and \( g'(\mu) \neq 0 \), then
\[
\sqrt{n}\{g(\bar{X}_n) - g(\mu)\} \xrightarrow{d} N(0, g'(\mu)^2 \sigma^2) \quad \text{as} \quad n \to \infty
\] (2.21)

This theorem is used frequently in statistics to obtain the asymptotic distribution of estimators — many of which can be expressed as functions of sample means

(These kinds of results are often said to use the “delta method”)

The proof is based on a Taylor expansion of \( g \) around the point \( \mu \)

Taking the result as given, let the distribution \( F \) of each \( X_i \) be uniform on \([0, \pi/2]\) and let \( g(x) = \sin(x) \)

Derive the asymptotic distribution of \( \sqrt{n}\{g(\bar{X}_n) - g(\mu)\} \) and illustrate convergence in the same spirit as the program illustrate_clt.jl discussed above

What happens when you replace \([0, \pi/2]\) with \([0, \pi]\)?

What is the source of the problem?

Exercise 2 Here’s a result that’s often used in developing statistical tests, and is connected to the multivariate central limit theorem

If you study econometric theory, you will see this result used again and again

Assume the setting of the multivariate CLT discussed above, so that

1. \( X_1, \ldots, X_n \) is a sequence of IID random vectors, each taking values in \( \mathbb{R}^k \)
2. \( \mu := \mathbb{E}[X_i] \), and \( \Sigma \) is the variance-covariance matrix of \( X_i \)
3. The convergence
\[
\sqrt{n}(\bar{X}_n - \mu) \xrightarrow{d} N(0, \Sigma)
\] (2.22)

is valid

In a statistical setting, one often wants the right hand side to be standard normal, so that confidence intervals are easily computed

This normalization can be achieved on the basis of three observations

First, if \( X \) is a random vector in \( \mathbb{R}^k \) and \( A \) is constant and \( k \times k \), then
\[
\text{Var}[AX] = A \text{Var}[X]A'
\]

Second, by the continuous mapping theorem, if \( Z_n \xrightarrow{d} Z \) in \( \mathbb{R}^k \) and \( A \) is constant and \( k \times k \), then
\[
AZ_n \xrightarrow{d} AZ
\]

Third, if \( S \) is a \( k \times k \) symmetric positive definite matrix, then there exists a symmetric positive definite matrix \( Q \), called the inverse square root of \( S \), such that
\[
QSQ' = I
\]
Here \( \mathbf{I} \) is the \( k \times k \) identity matrix.

Putting these things together, your first exercise is to show that if \( \mathbf{Q} \) is the inverse square root of \( \Sigma \), then

\[
\mathbf{Z}_n := \sqrt{n} \mathbf{Q} (\bar{\mathbf{X}}_n - \mu) \xrightarrow{d} \mathbf{Z} \sim N(0, \mathbf{I})
\]

Applying the continuous mapping theorem one more time tells us that

\[
\|\mathbf{Z}_n\|^2 \xrightarrow{d} \|\mathbf{Z}\|^2
\]

Given the distribution of \( \mathbf{Z} \), we conclude that

\[
n \|\mathbf{Q} (\bar{\mathbf{X}}_n - \mu)\|^2 \xrightarrow{d} \chi^2(k)
\]

(2.23)

where \( \chi^2(k) \) is the chi-squared distribution with \( k \) degrees of freedom.

(Recall that \( k \) is the dimension of \( \mathbf{X}_i \), the underlying random vectors).

Your second exercise is to illustrate the convergence in (2.23) with a simulation.

In doing so, let

\[
\mathbf{X}_i := \left( \begin{array}{c} \mathbf{W}_i \\ \mathbf{U}_i + \mathbf{W}_i \end{array} \right)
\]

where

- each \( \mathbf{W}_i \) is an IID draw from the uniform distribution on \([-1, 1]\)
- each \( \mathbf{U}_i \) is an IID draw from the uniform distribution on \([-2, 2]\)
- \( \mathbf{U}_i \) and \( \mathbf{W}_i \) are independent of each other

Hints:

1. \( \sqrt{\mathbf{m}}(\mathbf{A}) \) computes the square root of \( \mathbf{A} \). You still need to invert it.
2. You should be able to work out \( \Sigma \) from the preceding information.

Solutions

Solution notebook

2.6 Linear State Space Models
2.6. LINEAR STATE SPACE MODELS

Contents

• Linear State Space Models
  – Overview
  – The Linear State Space Model
  – Distributions and Moments
  – Stationarity and Ergodicity
  – Prediction
  – Code
  – Exercises
  – Solutions

“We may regard the present state of the universe as the effect of its past and the cause of its future” – Marquis de Laplace

Overview

This lecture introduces the linear state space dynamic system

Easy to use and carries a powerful theory of prediction

A workhorse with many applications

• representing dynamics of higher-order linear systems
• predicting the position of a system \( j \) steps into the future
• predicting a geometric sum of future values of a variable like
  – non financial income
  – dividends on a stock
  – the money supply
  – a government deficit or surplus
  – etc., etc., . . .

• key ingredient of useful models
  – Friedman’s permanent income model of consumption smoothing
  – Barro’s model of smoothing total tax collections
  – Rational expectations version of Cagan’s model of hyperinflation
  – Sargent and Wallace’s “unpleasant monetarist arithmetic”
  – etc., etc., . . .
The Linear State Space Model

Objects in play

- An $n \times 1$ vector $x_t$ denoting the state at time $t = 0, 1, 2, \ldots$
- An $m \times 1$ vector of iid shocks $w_{t+1} \sim N(0, I)$
- A $k \times 1$ vector of observations $y_t$ at time $t = 0, 1, 2, \ldots$
- An $n \times n$ matrix $A$ called the transition matrix
- An $n \times m$ matrix $C$ called the volatility matrix
- A $k \times n$ matrix $G$ sometimes called the output matrix

Here is the linear state-space system

$$
\begin{align*}
x_{t+1} &= Ax_t + Cw_{t+1} \\
y_t &= Gx_t \\
x_0 &\sim N(\mu_0, \Sigma_0)
\end{align*}
$$

Primitives

The primitives of the model are

1. the matrices $A, C, G$
2. shock distribution, which we have specialized to $N(0, I)$
3. the distribution of the initial condition $x_0$, which we have set to $N(\mu_0, \Sigma_0)$

Given $A, C, G$ and draws of $x_0$ and $w_1, w_2, \ldots$, the model (2.24) pins down the values of the sequences $\{x_t\}$ and $\{y_t\}$

Even without these draws, the primitives 1–3 pin down the probability distributions of $\{x_t\}$ and $\{y_t\}$

Later we’ll see how to compute these distributions and their moments

Martingale difference shocks

We’ve made the common assumption that the shocks are independent standardized normal vectors

But some of what we say will go through under the assumption that $\{w_{t+1}\}$ is a martingale difference sequence

A martingale difference sequence is a sequence that is zero mean when conditioned on past information

In the present case, since $\{x_t\}$ is our state sequence, this means that it satisfies

$$
\mathbb{E} [w_{t+1} | x_t, x_{t-1}, \ldots] = 0
$$

This is a weaker condition than that $\{w_t\}$ is iid with $w_{t+1} \sim N(0, I)$
Examples  By appropriate choice of the primitives, a variety of dynamics can be represented in terms of the linear state space model

The following examples help to highlight this point
They also illustrate the wise dictum *finding the state is an art*

**Second-order difference equation**  Let \( \{y_t\} \) be a deterministic sequence that satisfies

\[
y_{t+1} = \phi_0 + \phi_1 y_t + \phi_2 y_{t-1} \quad \text{s.t.} \quad y_0, y_{-1} \text{ given} \tag{2.25}
\]

To map (2.25) into our state space system (2.24), we set

\[
x_t = \begin{bmatrix} 1 \\ y_t \\ y_{t-1} \end{bmatrix}, \quad A = \begin{bmatrix} 1 & 0 & 0 \\ \phi_0 & \phi_1 & \phi_2 \\ 0 & 1 & 0 \end{bmatrix}, \quad C = \begin{bmatrix} 0 \\ 0 \\ 0 \end{bmatrix}, \quad G = \begin{bmatrix} 0 & 1 & 0 \end{bmatrix}
\]

You can confirm that under these definitions, (2.24) and (2.25) agree

The next figure shows dynamics of this process when \( \phi_0 = 1.1, \phi_1 = 0.8, \phi_2 = -0.8, y_0 = y_{-1} = 1 \)

![Graph showing dynamics of a process](image)

Later you’ll be asked to recreate this figure

**Univariate Autoregressive Processes**  We can use (2.24) to represent the model

\[
y_{t+1} = \phi_1 y_t + \phi_2 y_{t-1} + \phi_3 y_{t-2} + \phi_4 y_{t-3} + \sigma w_{t+1} \tag{2.26}
\]

where \( \{w_t\} \) is iid and standard normal

To put this in the linear state space format we take \( x_t = [y_t \ y_{t-1} \ y_{t-2} \ y_{t-3}]' \) and

\[
A = \begin{bmatrix} \phi_1 & \phi_2 & \phi_3 & \phi_4 \\ 1 & 0 & 0 & 0 \\ 0 & 1 & 0 & 0 \\ 0 & 0 & 1 & 0 \end{bmatrix}, \quad C = \begin{bmatrix} \sigma \\ 0 \\ 0 \\ 0 \end{bmatrix}, \quad G = \begin{bmatrix} 1 & 0 & 0 & 0 \end{bmatrix}
\]
The matrix $A$ has the form of the companion matrix to the vector $[\phi_1 \ \phi_2 \ \phi_3 \ \phi_4]$.

The next figure shows dynamics of this process when

$$\phi_1 = 0.5, \phi_2 = -0.2, \phi_3 = 0, \phi_4 = 0.5, \sigma = 0.2, y_0 = y_{-1} = y_{-2} = y_{-3} = 1$$

Vector Autoregressions

Now suppose that

- $y_t$ is a $k \times 1$ vector
- $\phi_j$ is a $k \times k$ matrix and
- $w_t$ is $k \times 1$

Then (2.26) is termed a vector autoregression

To map this into (2.24), we set

$$x_t = \begin{bmatrix} y_t \\ y_{t-1} \\ y_{t-2} \\ y_{t-3} \end{bmatrix}, \quad A = \begin{bmatrix} \phi_1 & \phi_2 & \phi_3 & \phi_4 \\ I & 0 & 0 & 0 \\ 0 & I & 0 & 0 \\ 0 & 0 & I & 0 \end{bmatrix}, \quad C = \begin{bmatrix} \sigma \\ 0 \\ 0 \end{bmatrix}, \quad G = \begin{bmatrix} I & 0 & 0 & 0 \end{bmatrix}$$

where $I$ is the $k \times k$ identity matrix and $\sigma$ is a $k \times k$ matrix

Seasonals

We can use (2.24) to represent

1. the deterministic seasonal $y_t = y_{t-4}$
2. the indeterministic seasonal $y_t = \phi_4 y_{t-4} + w_t$
In fact both are special cases of (2.26)
With the deterministic seasonal, the transition matrix becomes
\[
A = \begin{bmatrix}
0 & 0 & 0 & 1 \\
1 & 0 & 0 & 0 \\
0 & 1 & 0 & 0 \\
0 & 0 & 1 & 0
\end{bmatrix}
\]
The eigenvalues are \((1, -1, i, -i)\), and so have period four.

The resulting sequence oscillates deterministically with period four, and can be used to model deterministic seasonals in quarterly time series.

The indeterministic seasonal produces recurrent, but aperiodic, seasonal fluctuations.

**Time Trends**

The model \(y_t = at + b\) is known as a *linear time trend*.

We can represent this model in the linear state space form by taking
\[
A = \begin{bmatrix} 1 & 1 \\ 0 & 1 \end{bmatrix} \quad C = \begin{bmatrix} 0 \\ 0 \end{bmatrix} \quad G = \begin{bmatrix} a \\ b \end{bmatrix}
\] (2.27)

and starting at initial condition \(x_0 = \begin{bmatrix} 0 \\ 1 \end{bmatrix}'\).

In fact it’s possible to use the state-space system to represent polynomial trends of any order.

For instance, let
\[
x_0 = \begin{bmatrix} 0 \\ 1 \\ 1 \end{bmatrix}, \quad A = \begin{bmatrix} 1 & 1 & 0 \\ 0 & 1 & 1 \\ 0 & 0 & 1 \end{bmatrix} \quad C = \begin{bmatrix} 0 \\ 0 \end{bmatrix}
\]

It follows that
\[
A^t = \begin{bmatrix} 1 & t & t(t-1)/2 \\ 0 & 1 & t \\ 0 & 0 & 1 \end{bmatrix}
\]

Then \(x_t' = \begin{bmatrix} t(t-1)/2 & t & 1 \end{bmatrix}\), so that \(x_t\) contains linear and quadratic time trends.

As a variation on the linear time trend model, consider \(y_t = t + b + \sum_{j=0}^t w_j\) with \(w_0 = 0\).

To modify (2.27) accordingly, we set
\[
A = \begin{bmatrix} 1 & 1 \\ 0 & 1 \end{bmatrix} \quad C = \begin{bmatrix} 1 \\ 0 \end{bmatrix} \quad G = \begin{bmatrix} 1 & b \end{bmatrix}
\] (2.28)

For reasons explained below, \(y_t\) is called a *martingale with drift*.

---

3 For example, note that \(i = \cos(\pi/2) + i \sin(\pi/2)\), so the period associated with \(i\) is \(\frac{2\pi}{\pi/2} = 4\).
Moving Average Representations  A nonrecursive expression for $x_t$ as a function of $x_0, w_1, w_2, \ldots, w_t$ can be found by using (2.24) repeatedly to obtain

$$
x_t = Ax_{t-1} + Cw_t
= A^2x_{t-2} + ACw_{t-1} + Cw_t
\vdots
= \sum_{j=0}^{t-1} A^jCw_{t-j} + A^tx_0
$$

Representation (2.29) is a moving average representation

It expresses \{x_t\} as a linear function of

1. current and past values of the process \{w_t\} and
2. the initial condition $x_0$

As an example of a moving average representation, recall the model (2.28)

You will be able to show that $A^t = \begin{bmatrix} 1 & t \\ 0 & 1 \end{bmatrix}$ and $A^tC = \begin{bmatrix} 1 & 0 \end{bmatrix}$

Substituting into the moving average representation (2.29), we obtain

$$
x_{1t} = \sum_{j=0}^{t-1} w_{t-j} + \begin{bmatrix} 1 & t \end{bmatrix} x_0
$$

where $x_{1t}$ is the first entry of $x_t$

The first term on the right is a cumulated sum of martingale differences, and is therefore a martingale

The second term is a translated linear function of time

For this reason, $x_{1t}$ is called a martingale with drift

Distributions and Moments

Unconditional Moments  Using (2.24), it’s easy to obtain expressions for the (unconditional) mean of $x_t$ and $y_t$

We’ll explain what unconditional and conditional mean soon

Letting $\mu_t := \mathbb{E}[x_t]$ and using linearity of expectations, we find that

$$
\mu_{t+1} = A\mu_t
$$

The initial condition for (2.30) is the primitive $\mu_0$ from (2.24)

The expectation $\mathbb{E}[y_t]$ of $y_t$ is $G\mu_t$

The variance-covariance matrix of $x_t$ is $\Sigma_t := \mathbb{E}[(x_t - \mu_t)(x_t - \mu_t)']$
Using $x_{t+1} - \mu_{t+1} = A(x_t - \mu_t) + Cw_{t+1}$, we can determine this matrix recursively via

$$
\Sigma_{t+1} = A\Sigma_t A' + CC' \quad \text{with} \quad \Sigma_0 \text{ given}
$$

(2.31)

The initial condition is $\Sigma_0$ from the initial distribution of $x_0$.

As a matter of terminology, we will sometimes call

- $\mu_t$ the *unconditional mean* of $x_t$
- $\Sigma_t$ the *unconditional variance-covariance matrix* of $x_t$

This is to distinguish $\mu_t$ and $\Sigma_t$ from related objects that use conditioning information, to be defined below.

However, you should be aware that these "unconditional" moments do depend on the initial distribution $N(\mu_0, \Sigma_0)$.

**Distributions** In general, knowing the mean and variance-covariance matrix of a random vector is not quite as good as knowing the full distribution.

However, there are some situations where these moments alone tell us all we need to know.

One such situation is when the vector in question is Gaussian (i.e., normally distributed).

This is the case here, given

1. our Gaussian assumptions on the primitives
2. the fact that normality is preserved under linear operations

In fact, it’s well-known that

$$
u \sim N(\bar{u}, S) \quad \text{and} \quad v = a + Bu \implies v \sim N(a + B\bar{u}, BSB')
$$

(2.32)

In particular, given our Gaussian assumptions on the primitives and the linearity of (2.24) we can see immediately that both $x_t$ and $y_t$ are Gaussian for all $t \geq 0$.

Since $x_t$ is Gaussian, to find the distribution, all we need to do is find its mean and variance-covariance matrix.

But in fact we’ve already done this, in (2.30) and (2.31).

Letting $\mu_t$ and $\Sigma_t$ be as defined by these equations, we have

$$
x_t \sim N(\mu_t, \Sigma_t) \quad \text{and} \quad y_t \sim N(G\mu_t, GS\Sigma_t G')
$$

(2.33)

**Ensemble Interpretations** How should we interpret the distributions defined by (2.33)?

Intuitively, the probabilities in a distribution correspond to relative frequencies in a large population drawn from that distribution.

Let’s apply this idea to our setting, focusing on the distribution of $y_T$ for fixed $T$.

---

4 The correct way to argue this is by induction. Suppose that $x_t$ is Gaussian. Then (2.24) and (2.32) imply that $x_{t+1}$ is Gaussian. Since $x_0$ is assumed to be Gaussian, it follows that every $x_t$ is Gaussian. Evidently this implies that each $y_t$ is Gaussian.
We can generate independent draws of $y_T$ by repeatedly simulating the evolution of the system up to time $T$, using an independent set of shocks each time.

The next figure shows 20 simulations, producing 20 time series for $\{y_t\}$, and hence 20 draws of $y_T$.

The system in question is the univariate autoregressive model (2.26).

The values of $y_T$ are represented by black dots in the left-hand figure.

![Graph showing the simulation of $y_T$](image1)

In the right-hand figure, these values are converted into a rotated histogram that shows relative frequencies from our sample of 20 $y_T$'s.

(The parameters and source code for the figures can be found in file examples/paths_and_hist.jl from the main repository.)

Here is another figure, this time with 100 observations.

![Graph showing the simulation of $y_T$](image2)

Let's now try with 500,000 observations, showing only the histogram (without rotation).

The black line is the density of $y_T$ calculated analytically, using (2.33).

The histogram and analytical distribution are close, as expected.
By looking at the figures and experimenting with parameters, you will gain a feel for how the distribution depends on the model primitives listed above.

**Ensemble means** In the preceding figure we recovered the distribution of $y_T$ by

1. generating $I$ sample paths (i.e., time series) where $I$ is a large number
2. recording each observation $y_T^i$
3. histogramming this sample

Just as the histogram corresponds to the distribution, the *ensemble* or *cross-sectional average* $\bar{y}_T := \frac{1}{I} \sum_{i=1}^{I} y_T^i$

approximates the expectation $\mathbb{E}[y_T] = G \mu_t$ (as implied by the law of large numbers)

Here’s a simulation comparing the ensemble average and true mean at time points $t = 0, \ldots, 50$

The parameters are the same as for the preceding figures, and the sample size is relatively small ($I = 20$)

The ensemble mean for $x_t$ is

$$\bar{x}_T := \frac{1}{I} \sum_{i=1}^{I} x_T^i \rightarrow \mu_T \quad (I \rightarrow \infty)$$

The right-hand side $\mu_T$ can be thought of as a “population average”

(By *population average* we mean the average for an infinite ($I = \infty$) number of sample $x_T$’s)
Another application of the law of large numbers assures us that
\[
\frac{1}{I} \sum_{i=1}^{I} (x_T^i - \bar{x}_T)(x_T^i - \bar{x}_T)' \to \Sigma_T \quad (I \to \infty)
\]

Joint Distributions In the preceding discussion we looked at the distributions of \(x_t\) and \(y_t\) in isolation. This gives us useful information, but doesn’t allow us to answer questions like

- what’s the probability that \(x_t \geq 0\) for all \(t\)?
- what’s the probability that the process \(\{y_t\}\) exceeds some value \(a\) before falling below \(b\)?
- etc., etc.

Such questions concern the joint distributions of these sequences. To compute the joint distribution of \(x_0, x_1, \ldots, x_T\), recall that in general joint and conditional densities are linked by the rule
\[
p(x, y) = p(y \mid x)p(x) \quad \text{(joint = conditional \times marginal)}
\]

From this rule we get
\[
p(x_0, x_1) = p(x_1 \mid x_0)p(x_0)
\]

Repeated applications of the same rule lead us to
\[
p(x_0, x_1, \ldots, x_T) = p(x_0) \prod_{t=0}^{T-1} p(x_{t+1} \mid x_t)
\]

The marginal \(p(x_0)\) is just the primitive \(N(\mu_0, \Sigma_0)\). In view of (2.24), the conditional densities are
\[
p(x_{t+1} \mid x_t) = N(Ax_t, CC')
\]
2.6. LINEAR STATE SPACE MODELS

**Autocovariance functions** An important object related to the joint distribution is the autocovariance function

\[ \Sigma_{t+j,t} := \mathbb{E} \left[ (x_{t+j} - \mu_{t+j})(x_t - \mu_t) \right] \]  

(2.34)

Elementary calculations show that

\[ \Sigma_{t+j,t} = A^j \Sigma_t \]  

(2.35)

Notice that \( \Sigma_{t+j,t} \) in general depends on both \( j \), the gap between the two dates, and \( t \), the earlier date.

**Stationarity and Ergodicity**

Two properties that greatly aid analysis of linear state space models when they hold are stationarity and ergodicity.

Let’s start with the intuition.

**Visualizing Stability** Let’s look at some more time series from the same model that we analyzed above.

This picture shows cross-sectional distributions for \( y \) at times \( T, T', T'' \)

Note how the time series “settle down” in the sense that the distributions at \( T' \) and \( T'' \) are relatively similar to each other — but unlike the distribution at \( T \).

In essence, the distributions of \( y_t \) are converging to a fixed long-run distribution as \( t \to \infty \).

When such a distribution exists it is called a stationary distribution.

THOMAS SARGENT AND JOHN STACHURSKI  March 21, 2015
2.6. LINEAR STATE SPACE MODELS

Stationary Distributions  In our setting, a distribution $\psi_\infty$ is said to be stationary for $x_t$ if

$$x_t \sim \psi_\infty \text{ and } x_{t+1} = Ax_t + Cw_{t+1} \implies x_{t+1} \sim \psi_\infty$$

Since

1. in the present case all distributions are Gaussian
2. a Gaussian distribution is pinned down by its mean and variance-covariance matrix

we can restate the definition as follows: $\psi_\infty$ is stationary for $x_t$ if

$$\psi_\infty = N(\mu_\infty, \Sigma_\infty)$$

where $\mu_\infty$ and $\Sigma_\infty$ are fixed points of (2.30) and (2.31) respectively

Covariance Stationary Processes  Let’s see what happens to the preceding figure if we start $x_0$ at the stationary distribution

Now the differences in the observed distributions at $T, T'$ and $T''$ come entirely from random fluctuations due to the finite sample size

By

- our choosing $x_0 \sim N(\mu_\infty, \Sigma_\infty)$
- the definitions of $\mu_\infty$ and $\Sigma_\infty$ as fixed points of (2.30) and (2.31) respectively

we’ve ensured that

$$\mu_t = \mu_\infty \text{ and } \Sigma_t = \Sigma_\infty \text{ for all } t$$

Moreover, in view of (2.35), the autocovariance function takes the form $\Sigma_{t+j,t} = A^j \Sigma_\infty$, which depends on $j$ but not on $t$
This motivates the following definition
A process \( \{x_t\} \) is said to be covariance stationary if
- both \( \mu_t \) and \( \Sigma_t \) are constant in \( t \)
- \( \Sigma_{t+j,t} \) depends on the time gap \( j \) but not on time \( t \)

In our setting, \( \{x_t\} \) will be covariance stationary if \( \mu_0, \Sigma_0, A, C \) assume values that imply that none of \( \mu_t, \Sigma_t, \Sigma_{t+j,t} \) depends on \( t \)

Conditions for Stationarity

The globally stable case  The difference equation \( \mu_{t+1} = A \mu_t \) is known to have unique fixed point \( \mu_\infty = 0 \) if all eigenvalues of \( A \) have moduli strictly less than unity
That is, if \( \text{all(abs(eigvals(A)) < 1)} \Rightarrow \text{true} \)

The difference equation (2.31) also has a unique fixed point in this case, and, moreover
\[
\mu_t \to \mu_\infty = 0 \quad \text{and} \quad \Sigma_t \to \Sigma_\infty \quad \text{as} \quad t \to \infty
\]
regardless of the initial conditions \( \mu_0 \) and \( \Sigma_0 \)
This is the globally stable case — see these notes for more a theoretical treatment
However, global stability is more than we need for stationary solutions, and often more than we want
To illustrate, consider our second order difference equation example
Here the state is \( x_t = [1 \ y_t \ y_{t-1}]' \)
Because of the constant first component in the state vector, we will never have \( \mu_t \to 0 \)
How can we find stationary solutions that respect a constant state component?

Processes with a constant state component  To investigate such a process, suppose that \( A \) and \( C \) take the form
\[
A = \begin{bmatrix} A_1 & a \\ 0 & 1 \end{bmatrix} \quad C = \begin{bmatrix} C_1 \\ 0 \end{bmatrix}
\]
where
- \( A_1 \) is an \( (n-1) \times (n-1) \) matrix
- \( a \) is an \( (n-1) \times 1 \) column vector
Let \( x_t = [x_{1t}' \ 1]' \) where \( x_{1t} \) is \( (n-1) \times 1 \)
It follows that
\[
x_{1,t+1} = A_1 x_{1t} + a + C_1 w_{t+1}
\]
Let $\mu_{1t} = E[x_{1t}]$ and take expectations on both sides of this expression to get

$$\mu_{1,t+1} = A_1 \mu_{1t} + a$$

(2.36)

Assume now that the moduli of the eigenvalues of $A_1$ are all strictly less than one. Then (2.36) has a unique stationary solution, namely,

$$\mu_{1\infty} = (I - A_1)^{-1}a$$

The stationary value of $\mu_t$ itself is then $\mu_{\infty} := [\mu'_{1\infty} \ 1]'$

The stationary values of $\Sigma_t$ and $\Sigma_{t+j,t}$ satisfy

$$\Sigma_{\infty} = A \Sigma_{\infty} A' + CC'$$

(2.37)

$$\Sigma_{t+j,t} = A^j \Sigma_{\infty}$$

Notice that $\Sigma_{t+j,t}$ depends on the time gap $j$ but not on calendar time $t$.

In conclusion, if

- $x_0 \sim N(\mu_{\infty}, \Sigma_{\infty})$ and
- the moduli of the eigenvalues of $A_1$ are all strictly less than unity

then the $\{x_t\}$ process is covariance stationary, with constant state component.

**Note:** If the eigenvalues of $A_1$ are less than unity in modulus, then (a) starting from any initial value, the mean and variance-covariance matrix both converge to their stationary values; and (b) iterations on (2.31) converge to the fixed point of the discrete Lyapunov equation in the first line of (2.37).

---

**Ergodicity**

Let’s suppose that we’re working with a covariance stationary process.

In this case we know that the ensemble mean will converge to $\mu_{\infty}$ as the sample size $I$ approaches infinity.

**Averages over time**

Ensemble averages across simulations are interesting theoretically, but in real life we usually observe only a single realization $\{x_t, y_t\}_{t=0}^T$.

So now let’s take a single realization and form the time series averages

$$\bar{x} := \frac{1}{T} \sum_{t=1}^T x_t \quad \text{and} \quad \bar{y} := \frac{1}{T} \sum_{t=1}^T y_t$$

Do these time series averages converge to something interpretable in terms of our basic state-space representation?

To get this desideratum, we require something called *ergodicity*.

Ergodicity is the property that time series and ensemble averages coincide.
More formally, ergodicity implies that time series sample averages converge to their expectation under the stationary distribution.

In particular,

- \( \frac{1}{T} \sum_{t=0}^{T} x_t \to \mu_{\infty} \)
- \( \frac{1}{T} \sum_{t=0}^{T} (x_t - \bar{x}_T)(x_t - \bar{x}_T)' \to \Sigma_{\infty} \)
- \( \frac{1}{T} \sum_{t=0}^{T} (x_{t+j} - \bar{x}_T)(x_t - \bar{x}_T)' \to A^j \Sigma_{\infty} \)

In our linear Gaussian setting, any covariance stationary process is also ergodic.

**Prediction**

The theory of prediction for linear state space systems is elegant and simple.

**Forecasting Formulas – Conditional Means** The natural way to predict variables is to use conditional distributions.

For example, the optimal forecast of \( x_{t+1} \) given information known at time \( t \) is

\[
E_t [x_{t+1}] := E [x_{t+1} \mid x_t, x_{t-1}, \ldots, x_0] = Ax_t
\]

The right-hand side follows from \( x_{t+1} = Ax_t + Cw_{t+1} \) and the fact that \( w_{t+1} \) is zero mean and independent of \( x_t, x_{t-1}, \ldots, x_0 \).

Observe that in the present case, conditioning on the entire history is the same as conditioning on the present.

In other words, \( E_t [x_{t+1}] = E_t [x_{t+1} \mid x_t] \), an implication of \( \{ x_t \} \) having the *Markov property*.

The one-step-ahead forecast error is

\[
x_{t+1} - E_t [x_{t+1}] = Cw_{t+1}
\]

The covariance matrix of the forecast error is

\[
E [ (x_{t+1} - E_t [x_{t+1}]) (x_{t+1} - E_t [x_{t+1}])' ] = CC'
\]

More generally, we’d like to compute

- \( j \)-step ahead forecasts of \( x \): \( E_t [x_{t+j}] := E [x_{t+j} \mid x_t, x_{t-1}, \ldots, x_0] \)
- \( j \)-step ahead forecasts of \( y \): \( E_t [y_{t+j}] := E [y_{t+j} \mid x_t, x_{t-1}, \ldots, x_0] \)

Here are the pertinent formulas

- \( j \)-step ahead forecast of \( x \):
  \[
  E_t [x_{t+j}] = A^j x_t
  \]

- \( j \)-step ahead forecast of \( y \):
  \[
  E_t [y_{t+j}] = GA^j x_t
  \]
**Covariance of Prediction Errors**  It is useful to obtain the covariance matrix of the vector of $j$-step-ahead prediction errors

$$x_{t+j} - \mathbb{E}_t[x_{t+j}] = \sum_{s=0}^{j-1} A^s C w_{t-s+j}$$

(2.38)

Evidently,

$$V_j := \mathbb{E}_t[(x_{t+j} - \mathbb{E}_t[x_{t+j}])(x_{t+j} - \mathbb{E}_t[x_{t+j}])'] = \sum_{k=0}^{j-1} A^k CC' A'^k$$

(2.39)

$V_j$ defined in (2.39) can be calculated recursively via $V_1 = CC'$ and

$$V_j = CC' + AV_{j-1}A', \quad j \geq 2$$

(2.40)

$V_j$ is the *conditional covariance matrix* of the errors in forecasting $x_{t+j}$, conditioned on time $t$ information $x_t$.

Under particular conditions, $V_j$ converges to

$$V_\infty = CC' + AV_\infty A'$$

(2.41)

Equation (2.41) is an example of a *discrete Lyapunov* equation in the covariance matrix $V_\infty$.

A sufficient condition for $V_j$ to converge is that the eigenvalues of $A$ be strictly less than one in modulus.

Weaker sufficient conditions for convergence associate eigenvalues equaling or exceeding one in modulus with elements of $C$ that equal 0.

**Forecasts of Geometric Sums**  In several contexts, we want to compute forecasts of geometric sums of future random variables governed by the linear state-space system (2.24).

We want the following objects

- Forecast of a geometric sum of future $x$’s, or $\mathbb{E} \left[ \sum_{j=0}^{\infty} \beta^j x_{t+j} | x_t \right]$
- Forecast of a geometric sum of future $y$’s, or $\mathbb{E} \left[ \sum_{j=0}^{\infty} \beta^j y_{t+j} | x_t \right]$

These objects are important components of some famous and interesting dynamic models.

For example,

- if $\{y_t\}$ is a stream of dividends, then $\mathbb{E} \left[ \sum_{j=0}^{\infty} \beta^j y_{t+j} | x_t \right]$ is a model of a stock price
- if $\{y_t\}$ is the money supply, then $\mathbb{E} \left[ \sum_{j=0}^{\infty} \beta^j y_{t+j} | x_t \right]$ is a model of the price level

**Formulas**  Fortunately, it is easy to use a little matrix algebra to compute these objects.

Suppose that every eigenvalue of $A$ has modulus strictly less than $\frac{1}{\beta}$.

It then follows that $I + \beta A + \beta^2 A^2 + \cdots = [I - \beta A]^{-1}$

This leads to our formulas:
• Forecast of a geometric sum of future \( x \)'s

\[
E_t \left[ \sum_{j=0}^{\infty} \beta^j x_{t+j} \right] = [I + \beta A + \beta^2 A^2 + \cdots] x_t = [I - \beta A]^{-1} x_t
\]

• Forecast of a geometric sum of future \( y \)'s

\[
E_t \left[ \sum_{j=0}^{\infty} \beta^j y_{t+j} \right] = G[I + \beta A + \beta^2 A^2 + \cdots] x_t = G[I - \beta A]^{-1} x_t
\]

**Code**

Our preceding simulations and calculations are based on code in the file lss.jl from the QuantEcon package. The code implements a type for handling linear state space models (simulations, calculating moments, etc.). We repeat it here for convenience.

```julia
# Computes quantities related to the Gaussian linear state space model

x_{t+1} = A x_t + C w_{t+1}

y_t = G x_t

The shocks \( \{w_t\} \) are iid and \( N(0, I) \)

@author : Spencer Lyon <spencer.lyon@nyu.edu>
@date : 2014-07-28

References
---------

Simple port of the file quantecon.lss

http://quant-econ.net/linear_models.html

# import Distributions: MultivariateNormal, rand

# numpy allows its multivariate_normal function to have a matrix of zeros for the covariance matrix; Stats.jl doesn't. This type just gives a `rand` method when we pass in a matrix of zeros for Sigma_0 so the rest of the api can work, unaffected

The behavior of `rand` is to just pass back the mean vector when the covariance matrix is zero.
```

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March 21, 2015
type FakeMVTNorm{T <: Real}
    mu_0::Array{T}
    Sigma_0::Array{T}
end

Base.rand{T}(d::FakeMVTNorm{T}) = copy(d.mu_0)

type LSS
    A::Matrix
    C::Matrix
    G::Matrix
    k::Int
    n::Int
    m::Int
    mu_0::Vector
    Sigma_0::Matrix
    dist::Union(MultivariateNormal, FakeMVTNorm)
end

function LSS(A::ScalarOrArray, C::ScalarOrArray, G::ScalarOrArray,
             mu_0::ScalarOrArray=zeros(size(G, 2)),
             Sigma_0::Matrix=zeros(size(G, 2), size(G, 2)))
    k = size(G, 1)
    n = size(G, 2)
    m = size(C, 2)

    # coerce shapes
    A = reshape([A], n, n)
    C = reshape([C], n, m)
    G = reshape([G], k, n)

    mu_0 = reshape([mu_0], n)

    # define distribution
    if all(Sigma_0 .== 0.0)  # no variance -- no distribution
        dist = FakeMVTNorm(mu_0, Sigma_0)
    else
        dist = MultivariateNormal(mu_0, Sigma_0)
    end
    LSS(A, C, G, k, n, m, mu_0, Sigma_0, dist)
end

# make kwarg version
function LSS(A::Matrix, C::Matrix, G::Matrix;
             mu_0::Vector=zeros(size(G, 2)),
             Sigma_0::Matrix=zeros(size(G, 2), size(G, 2)))
    return LSS(A, C, G, mu_0, Sigma_0)
end

function simulate(lss::LSS, ts_length=100)
    x = Array(Float64, lss.n, ts_length)
x[:, 1] = rand(lss.dist)
for t=1:ts_length-1
  x[:, t+1] = lss.A * x[:, t] .+ lss.C * w[:, t]
end
y = lss.G * x

return x, y
end

function replicate(lss::LSS, t=10, num_reps=100)
  x = Array(Float64, lss.n, num_reps)
  for j=1:num_reps
    x_t, _ = simulate(lss, t+1)
    x[:, j] = x_t[:, end]
  end
  y = lss.G * x
  return x, y
end

replicate(lss::LSS; t=10, num_reps=100) = replicate(lss, t, num_reps)

function moment_sequence(lss::LSS)
  mu_x, Sigma_x = copy(lss.mu_0), copy(lss.Sigma_0)
  while true
    mu_y, Sigma_y = G * mu_x, G * Sigma_x * G'
    produce((mu_x, mu_y, Sigma_x, Sigma_y))
    # Update moments of x
    mu_x = A * mu_x
    Sigma_x = A * Sigma_x * A' + C * C'
  end
  nothing
end

function stationary_distributions(lss::LSS; max_iter=200, tol=1e-5)
  # Initialize iteration
  m = @task moment_sequence(lss)
  mu_x, mu_y, Sigma_x, Sigma_y = consume(m)
  i = 0
  err = tol + 1.
  while err > tol
    if i > max_iter
      println("Convergence failed after $i iterations")
      break
    else
      # Update moments of x
      mu_x = A * mu_x
      Sigma_x = A * Sigma_x * A' + C * C'
    end
    mu_y, Sigma_y = G * mu_x, G * Sigma_x * G'
    produce((mu_x, mu_y, Sigma_x, Sigma_y))
    # Update moments of x
    mu_x = A * mu_x
    Sigma_x = A * Sigma_x * A' + C * C'
    i += 1
    err = norm(Sigma_y - Sigma_x)
  end
  nothing
end
i += 1
mu_x1, mu_y, Sigma_x1, Sigma_y = consume(m)
err_mu = Base.maxabs(mu_x1 - mu_x)
err_Sigma = Base.maxabs(Sigma_x1 - Sigma_x)
err = max(err_Sigma, err_mu)
mu_x, Sigma_x = mu_x1, Sigma_x1
end
end

return mu_x, mu_y, Sigma_x, Sigma_y
end
def geometric_sums(lss::LSS, bet, x_t):
    I = eye(lss.n)
    S_x = (I - bet .* A) \ x_t
    S_y = lss.G * S_x
    return S_x, S_y
end

Hopefully the code is relatively self explanatory and adequately documented
Examples of usage are given in the solutions to the exercises

Exercises

Exercise 1  Replicate this figure using the LSS type from lss.jl

Exercise 2  Replicate this figure modulo randomness using the same type

Exercise 3  Replicate this figure modulo randomness using the same type
The state space model and parameters are the same as for the preceding exercise

Exercise 4  Replicate this figure modulo randomness using the same type
The state space model and parameters are the same as for the preceding exercise, except that the
initial condition is the stationary distribution
Hint: You can use the stationary_distributions method to get the initial conditions
The number of sample paths is 80, and the time horizon in the figure is 100
Producing the vertical bars and dots is optional, but if you wish to try, the bars are at dates 10, 50
and 75

Solutions

Solution notebook
2.7 A First Look at the Kalman Filter

Overview

This lecture provides a simple and intuitive introduction to the Kalman filter, for those who either
• have heard of the Kalman filter but don’t know how it works, or
• know the Kalman filter equations, but don’t know where they come from

For additional (more advanced) reading on the Kalman filter, see
• [LS12], section 2.7.
• [AM05]

The last reference gives a particularly clear and comprehensive treatment of the Kalman filter

Required knowledge: Familiarity with matrix manipulations, multivariate normal distributions, covariance matrices, etc.

The Basic Idea

The Kalman filter has many applications in economics, but for now let’s pretend that we are rocket scientists

A missile has been launched from country Y and our mission is to track it

Let \( x \in \mathbb{R}^2 \) denote the current location of the missile—a pair indicating latitude-longitude coordinates on a map

At the present moment in time, the precise location \( x \) is unknown, but we do have some beliefs about \( x \)

One way to summarize our knowledge is a point prediction \( \hat{x} \)

• But what if the President wants to know the probability that the missile is currently over the Sea of Japan?
• Better to summarize our initial beliefs with a bivariate probability density \( p \)
\[ \int_E p(x)dx \] indicates the probability that we attach to the missile being in region \( E \)

The density \( p \) is called our *prior* for the random variable \( x \)

To keep things tractable, we will always assume that our prior is Gaussian. In particular, we take

\[ p = N(\hat{x}, \Sigma) \quad (2.42) \]

where \( \hat{x} \) is the mean of the distribution and \( \Sigma \) is a \( 2 \times 2 \) covariance matrix. In our simulations, we will suppose that

\[ \hat{x} = \begin{pmatrix} 0.2 \\ -0.2 \end{pmatrix}, \quad \Sigma = \begin{pmatrix} 0.4 & 0.3 \\ 0.3 & 0.45 \end{pmatrix} \quad (2.43) \]

This density \( p(x) \) is shown below as a contour map, with the center of the red ellipse being equal to \( \hat{x} \)

![Contour map of prior density](image)

**Figure 2.1:** Prior density (Click this or any other figure to enlarge.)

**The Filtering Step**  We are now presented with some good news and some bad news

The good news is that the missile has been located by our sensors, which report that the current location is \( y = (2.3, -1.9) \)

The next figure shows the original prior \( p(x) \) and the new reported location \( y \)

The bad news is that our sensors are imprecise.

In particular, we should interpret the output of our sensor not as \( y = x \), but rather as

\[ y = Gx + v, \quad \text{where} \quad v \sim N(0, R) \quad (2.44) \]

Here \( G \) and \( R \) are \( 2 \times 2 \) matrices with \( R \) positive definite. Both are assumed known, and the noise term \( v \) is assumed to be independent of \( x \)
How then should we combine our prior \( p(x) = N(\hat{x}, \Sigma) \) and this new information \( y \) to improve our understanding of the location of the missile?

As you may have guessed, the answer is to use Bayes’ theorem, which tells us we should update our prior \( p(x) \) to \( p(x | y) \) via

\[
p(x | y) = \frac{p(y | x) p(x)}{p(y)}
\]

where \( p(y) = \int p(y | x) p(x) dx \)

In solving for \( p(x | y) \), we observe that

- \( p(x) = N(\hat{x}, \Sigma) \)
- In view of (2.44), the conditional density \( p(y | x) \) is \( N(Gx, R) \)
- \( p(y) \) does not depend on \( x \), and enters into the calculations only as a normalizing constant

Because we are in a linear and Gaussian framework, the updated density can be computed by calculating population linear regressions.

In particular, the solution is known \(^5\) to be

\[
p(x | y) = N(\hat{x}^F, \Sigma^F)
\]

where

\[
\hat{x}^F := \hat{x} + \Sigma G' (G \Sigma G' + R)^{-1} (y - G \hat{x}) \quad \text{and} \quad \Sigma^F := \Sigma - \Sigma G' (G \Sigma G' + R)^{-1} G \Sigma
\]  

(2.45)

Here \( \Sigma G' (G \Sigma G' + R)^{-1} \) is the matrix of population regression coefficients of the hidden object \( x - \hat{x} \) on the surprise \( y - G \hat{x} \)

---

\(^5\) See, for example, page 93 of [Bis06]. To get from his expressions to the ones used above, you will also need to apply the Woodbury matrix identity.
This new density \( p(x \mid y) = N(\hat{x}^F, \Sigma^F) \) is shown in the next figure via contour lines and the color map.

The original density is left in as contour lines for comparison.

Our new density twists the prior \( p(x) \) in a direction determined by the new information \( y - G\hat{x} \).

In generating the figure, we set \( G \) to the identity matrix and \( R = 0.5\Sigma \) for \( \Sigma \) defined in (2.43).

(The code for generating this and the preceding figures can be found in the file examples/gaussian_contours.jl from the main repository.)

**The Forecast Step**  What have we achieved so far?

We have obtained probabilities for the current location of the state (missile) given prior and current information.

This is called “filtering” rather than forecasting, because we are filtering out noise rather than looking into the future.

- \( p(x \mid y) = N(\hat{x}^F, \Sigma^F) \) is called the *filtering distribution*.

But now let’s suppose that we are given another task: To predict the location of the missile after one unit of time (whatever that may be) has elapsed.

To do this we need a model of how the state evolves.

Let’s suppose that we have one, and that it’s linear and Gaussian: In particular,

\[
x_{t+1} = Ax_t + w_{t+1}, \quad \text{where} \quad w_t \sim N(0, Q)
\]  \hspace{1cm} (2.46)

Our aim is to combine this law of motion and our current distribution \( p(x \mid y) = N(\hat{x}^F, \Sigma^F) \) to come up with a new *predictive* distribution for the location one unit of time hence
In view of (2.46), all we have to do is introduce a random vector $x^F \sim N(\hat{x}^F, \Sigma^F)$ and work out the distribution of $Ax^F + w$ where $w$ is independent of $x^F$ and has distribution $N(0, Q)$.

Since linear combinations of Gaussians are Gaussian, $Ax^F + w$ is Gaussian.

Elementary calculations and the expressions in (2.45) tell us that

$$
E[Ax^F + w] = A\hat{x}^F + E[w] = A\hat{x}^F + A\Sigma G' (G\Sigma G' + R)^{-1}(y - G\hat{x})
$$

and

$$
\text{Var}[Ax^F + w] = A\text{Var}[x^F]A' + Q = A\Sigma F A' + Q = A\Sigma A' - A\Sigma G' (G\Sigma G' + R)^{-1}G\Sigma A' + Q
$$

The matrix $A\Sigma G' (G\Sigma G' + R)^{-1}$ is often written as $K_\Sigma$ and called the Kalman gain.

- the subscript $\Sigma$ has been added to remind us that $K_\Sigma$ depends on $\Sigma$, but not $y$ or $\hat{x}$

Using this notation, we can summarize our results as follows: Our updated prediction is the density $N(\hat{x}_{\text{new}}, \Sigma_{\text{new}})$ where

$$
\begin{align*}
\hat{x}_{\text{new}} &:= A\hat{x} + K_\Sigma (y - G\hat{x}) \\
\Sigma_{\text{new}} &:= A\Sigma A' - K_\Sigma G\Sigma A' + Q
\end{align*}
$$

- The density $p_{\text{new}}(x) = N(\hat{x}_{\text{new}}, \Sigma_{\text{new}})$ is called the predictive distribution.

The predictive distribution is the new density shown in the following figure, where the update has used parameters

$$
A = \begin{pmatrix} 1.2 & 0.0 \\ 0.0 & -0.2 \end{pmatrix}, \quad Q = 0.3 \times \Sigma
$$
The Recursive Procedure  Let’s look back at what we’ve done.

We started the current period with a prior \( p(x) \) for the location \( x \) of the missile

We then used the current measurement \( y \) to update to \( p(x | y) \)

Finally, we used the law of motion (2.46) for \( \{x_t\} \) to update to \( p_{\text{new}}(x) \)

If we now step into the next period, we are ready to go round again, taking \( p_{\text{new}}(x) \) as the current prior

Swapping notation \( p_t(x) \) for \( p(x) \) and \( p_{t+1}(x) \) for \( p_{\text{new}}(x) \), the full recursive procedure is:

1. Start the current period with prior \( p_t(x) = N(\hat{x}_t, \Sigma_t) \)
2. Observe current measurement \( y_t \)
3. Compute the filtering distribution \( p_t(x | y) = N(\hat{x}_F^t, \Sigma_F^t) \) from \( p_t(x) \) and \( y_t \), applying Bayes rule and the conditional distribution (2.44)
4. Compute the predictive distribution \( p_{t+1}(x) = N(\hat{x}_{t+1}, \Sigma_{t+1}) \) from the filtering distribution and (2.46)
5. Increment \( t \) by one and go to step 1

Repeating (2.47), the dynamics for \( \hat{x}_t \) and \( \Sigma_t \) are as follows

\[
\hat{x}_{t+1} = A\hat{x}_t + K\Sigma_t(y_t - G\hat{x}_t) \tag{2.48}
\]

\[
\Sigma_{t+1} = A\Sigma_tA' - K\Sigma_tG\Sigma_tA' + Q
\]

These are the standard dynamic equations for the Kalman filter. See, for example, [LS12], page 58.

Convergence

The matrix \( \Sigma_t \) is a measure of the uncertainty of our prediction \( \hat{x}_t \) of \( x_t \)

Apart from special cases, this uncertainty will never be fully resolved, regardless of how much time elapses

One reason is that our prediction \( \hat{x}_t \) is made based on information available at \( t - 1 \), not \( t \)

Even if we know the precise value of \( x_{t-1} \) (which we don’t), the transition equation (2.46) implies that \( x_t = Ax_{t-1} + w_t \)

Since the shock \( w_t \) is not observable at \( t - 1 \), any time \( t - 1 \) prediction of \( x_t \) will incur some error (unless \( w_t \) is degenerate)

However, it is certainly possible that \( \Sigma_t \) converges to a constant matrix as \( t \to \infty \)

To study this topic, let’s expand the second equation in (2.48):

\[
\Sigma_{t+1} = A\Sigma_tA' - A\Sigma_tG'(G\Sigma_tG' + R)^{-1}G\Sigma_tA' + Q \tag{2.49}
\]

This is a nonlinear difference equation in \( \Sigma_t \)

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A fixed point of (2.49) is a constant matrix \( \Sigma \) such that
\[
\Sigma = A \Sigma A' - A \Sigma G' (G \Sigma G' + R)^{-1} G \Sigma A' + Q
\] (2.50)

Equation (2.49) is known as a discrete time Riccati difference equation
Equation (2.50) is known as a discrete time algebraic Riccati equation

Conditions under which a fixed point exists and the sequence \( \{ \Sigma_t \} \) converges to it are discussed in [AHMS96] and [AM05], chapter 4

One sufficient (but not necessary) condition is that all the eigenvalues \( \lambda_i \) of \( A \) satisfy \( |\lambda_i| < 1 \) (cf. e.g., [AM05], p. 77)

(This strong condition assures that the unconditional distribution of \( x_t \) converges as \( t \to +\infty \))

In this case, for any initial choice of \( \Sigma_0 \) that is both nonnegative and symmetric, the sequence \( \{ \Sigma_t \} \) in (2.49) converges to a nonnegative symmetric matrix \( \Sigma \) that solves (2.50)

**Implementation**

The type `Kalman` from the `QuantEcon` package implements the Kalman filter

The class bundles together
- Instance data:
  - The parameters \( A, G, Q, R \) of a given model
  - the moments \( (\hat{x}_t, \Sigma_t) \) of the current prior
- Methods:
  - a method `prior_to_filtered` to update \( (\hat{x}_t, \Sigma_t) \) to \( (\hat{x}_F^t, \Sigma_F^t) \)
  - a method `filtered_to_forecast` to update the filtering distribution to the predictive distribution – which becomes the new prior \( (\hat{x}_{t+1}, \Sigma_{t+1}) \)
  - an `update` method, which combines the last two methods
  - a `stationary_values` method, which computes the solution to (2.50) and the corresponding (stationary) Kalman gain

You can view the program on GitHub but we repeat it here for convenience

```python
#= Implements the Kalman filter for a linear Gaussian state space model.

@author : Spencer Lyon  <spencer.lyon@nyu.edu>
@date: 2014-07-29

References
---------
Simple port of the file quantecron.kalman
```

---

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http://quant-econ.net/kalman.html

```julia
# type Kalman
A  G  Q  R
k  n
cur_x_hat
cur_sigma

end

# Initializes current mean and cov to zeros
function Kalman(A, G, Q, R)
    k = size(G, 1)
    n = size(G, 2)
    xhat = n == 1 ? zero(eltype(A)) : zeros(n)
    Sigma = n == 1 ? zero(eltype(A)) : zeros(n, n)
    return Kalman(A, G, Q, R, k, n, xhat, Sigma)
end

function set_state!(k::Kalman, x_hat, Sigma)
    k.cur_x_hat = x_hat
    k.cur_sigma = Sigma
    nothing
end

function prior_to_filtered!(k::Kalman, y)
    # simplify notation
    G, R = k.G, k.R
    x_hat, Sigma = k.cur_x_hat, k.cur_sigma
    # and then update
    if k.k > 1
        reshape(y, k.k, 1)
    end
    A = Sigma * G'
    B = G * Sigma' * G' + R
    M = A * inv(B)
    k.cur_x_hat = x_hat + M * (y - G * x_hat)
    k.cur_sigma = Sigma - M * G * Sigma
    nothing
end

function filtered_to_forecast!(k::Kalman)
    # simplify notation
    A, Q = k.A, k.Q
```
x_hat, Sigma = k.cur_x_hat, k.cur_sigma

# and then update
k.cur_x_hat = A * x_hat
k.cur_sigma = A * Sigma * A' + Q
nothing
end

function update!(k::Kalman, y)
prior_to_filtered!(k, y)
filtered_to_forecast!(k)
nothing
end

function stationary_values(k::Kalman)
    # simplify notation

    # solve Riccati equation, obtain Kalman gain
    Sigma_inf = solve_discrete_riccati(A', G', Q, R)
    K_inf = A * Sigma_inf * G' * inv(G * Sigma_inf * G' + R)
    return Sigma_inf, K_inf
end

Exercises

**Exercise 1** Consider the following simple application of the Kalman filter, loosely based on [LS12], section 2.9.2

Suppose that

- all variables are scalars
- the hidden state \( \{ x_t \} \) is in fact constant, equal to some \( \theta \in \mathbb{R} \) unknown to the modeler

State dynamics are therefore given by (2.46) with \( A = 1, Q = 0 \) and \( x_0 = \theta \)

The measurement equation is \( y_t = \theta + v_t \) where \( v_t \) is \( N(0,1) \) and iid

The task of this exercise to simulate the model and, using the code from kalman.jl, plot the first five predictive densities \( p_t(x) = N(\hat{x}_t, \Sigma_t) \)

As shown in [LS12], sections 2.9.1–2.9.2, these distributions asymptotically put all mass on the unknown value \( \theta \)

In the simulation, take \( \theta = 10, \hat{x}_0 = 8 \) and \( \Sigma_0 = 1 \)

Your figure should – modulo randomness – look something like this

**Exercise 2** The preceding figure gives some support to the idea that probability mass converges to \( \theta \)
To get a better idea, choose a small $\epsilon > 0$ and calculate

$$z_t := 1 - \int_{\theta-\epsilon}^{\theta+\epsilon} p_t(x) \, dx$$

for $t = 0, 1, 2, \ldots, T$

Plot $z_t$ against $T$, setting $\epsilon = 0.1$ and $T = 600$

Your figure should show error erratically declining something like this

---

**Exercise 3**  As discussed above, if the shock sequence $\{\hat{w}_t\}$ is not degenerate, then it is not in general possible to predict $x_t$ without error at time $t - 1$ (and this would be the case even if we could observe $x_{t-1}$)

Let’s now compare the prediction $\hat{x}_t$ made by the Kalman filter against a competitor who is allowed to observe $x_{t-1}$

This competitor will use the conditional expectation $E[x_t \mid x_{t-1}]$, which in this case is $Ax_{t-1}$

The conditional expectation is known to be the optimal prediction method in terms of minimizing mean squared error

(More precisely, the minimizer of $E \|x_t - g(x_{t-1})\|^2$ with respect to $g$ is $g^*(x_{t-1}) := E[x_t \mid x_{t-1}]$)

Thus we are comparing the Kalman filter against a competitor who has more information (in the sense of being able to observe the latent state) and behaves optimally in terms of minimizing squared error

Our horse race will be assessed in terms of squared error

In particular, your task is to generate a graph plotting observations of both $\|x_t - Ax_{t-1}\|^2$ and $\|x_t - \hat{x}_t\|^2$ against $t$ for $t = 1, \ldots, 50$
For the parameters, set $G = I$, $R = 0.5I$ and $Q = 0.3I$, where $I$ is the $2 \times 2$ identity matrix.

Set

$$A = \begin{pmatrix} 0.5 & 0.4 \\ 0.6 & 0.3 \end{pmatrix}$$

To initialize the prior density, set

$$\Sigma_0 = \begin{pmatrix} 0.9 & 0.3 \\ 0.3 & 0.9 \end{pmatrix}$$

and $\hat{x}_0 = (8, 8)$

Finally, set $x_0 = (0, 0)$

You should end up with a figure similar to the following (modulo randomness)

Observe how, after an initial learning period, the Kalman filter performs quite well, even relative to the competitor who predicts optimally with knowledge of the latent state.

**Exercise 4** Try varying the coefficient 0.3 in $Q = 0.3I$ up and down

Observe how the diagonal values in the stationary solution $\Sigma$ (see (2.50)) increase and decrease in line with this coefficient.

The interpretation is that more randomness in the law of motion for $x_t$ causes more (permanent) uncertainty in prediction.

**Solutions**

Solution notebook
2.8 Infinite Horizon Dynamic Programming

Overview

In a previous lecture we gained some intuition about finite stage dynamic programming by studying the shortest path problem.

The aim of this lecture is to introduce readers to methods for solving simple infinite-horizon dynamic programming problems using Julia.

We will also introduce and motivate some of the modeling choices used throughout the lectures to treat this class of problems.

The particular application we will focus on is solving for consumption in an optimal growth model. Although the model is quite specific, the key ideas extend to many other problems in dynamic optimization.

The model is also very simplistic — we favor ease of exposition over realistic assumptions throughout the current lecture.
2.8. INFINITE HORIZON DYNAMIC PROGRAMMING

Other References  For supplementary reading see

- [LS12], section 3.1
- EDTC, section 6.2 and chapter 10
- [Sun96], chapter 12
- [SLP89], chapters 2–5
- [HLL96], all

An Optimal Growth Model

Consider an agent who owns at time \( t \) capital stock \( k_t \in \mathbb{R}_+ := [0, \infty) \) and produces output

\[
y_t := f(k_t) \in \mathbb{R}_+
\]

This output can either be consumed or saved as capital for next period

For simplicity we assume that depreciation is total, so that next period capital is just output minus consumption:

\[
k_{t+1} = y_t - c_t \tag{2.51}
\]

Taking \( k_0 \) as given, we suppose that the agent wishes to maximize

\[
\sum_{t=0}^{\infty} \beta^t u(c_t) \tag{2.52}
\]

where \( u \) is a given utility function and \( \beta \in (0, 1) \) is a discount factor

More precisely, the agent wishes to select a path \( c_0, c_1, c_2, \ldots \) for consumption that is

1. nonnegative
2. feasible in the sense that the capital path \( \{k_t\} \) determined by \( \{c_t\}, k_0 \) and (2.51) is always nonnegative
3. optimal in the sense that it maximizes (2.52) relative to all other feasible consumption sequences

A well-known result from the standard theory of dynamic programming (cf., e.g., [SLP89], section 4.1) states that, for kind of this problem, any optimal consumption sequence \( \{c_t\} \) must be Markov

That is, there exists a function \( \sigma \) such that

\[
c_t = \sigma(k_t) \quad \text{for all } t
\]

In other words, the current control is a fixed (i.e., time homogeneous) function of the current state.
The Policy Function Approach  As it turns out, we are better off seeking the function $\sigma$ directly, rather than the optimal consumption sequence.

The main reason is that the functional approach — seeking the optimal policy — translates directly over to the stochastic case, whereas the sequential approach does not.

For this model, we will say that function $\sigma$ mapping $\mathbb{R}_+$ into $\mathbb{R}_+$ is a feasible consumption policy if it satisfies

$$\sigma(k) \leq f(k) \quad \text{for all} \quad k \in \mathbb{R}_+ \quad (2.53)$$

The set of all such policies will be denoted by $\Sigma$.

Using this notation, the agent’s decision problem can be rewritten as

$$\max_{\sigma \in \Sigma} \left\{ \sum_{t=0}^{\infty} \beta^t u(\sigma(k_t)) \right\} \quad (2.54)$$

where the sequence $\{k_t\}$ in (2.54) is given by

$$k_{t+1} = f(k_t) - \sigma(k_t), \quad k_0 \text{ given} \quad (2.55)$$

In the next section we discuss how to solve this problem for the maximizing $\sigma$.

Dynamic Programming

We will solve for the optimal policy using dynamic programming.

The first step is to define the policy value function $v_\sigma$ associated with a given policy $\sigma$, which is

$$v_\sigma(k_0) := \sum_{t=0}^{\infty} \beta^t u(\sigma(k_t)) \quad (2.56)$$

when $\{k_t\}$ is given by (2.55).

Evidently $v_\sigma(k_0)$ is the total present value of discounted utility associated with following policy $\sigma$ forever, given initial capital $k_0$.

The value function for this optimization problem is then defined as

$$v^*(k_0) := \sup_{\sigma \in \Sigma} v_\sigma(k_0) \quad (2.57)$$

The value function gives the maximal value that can be obtained from state $k_0$, after considering all feasible policies.

A policy $\sigma \in \Sigma$ is called optimal if it attains the supremum in (2.57) for all $k_0 \in \mathbb{R}_+$.

The Bellman equation for this problem takes the form

$$v^*(k) = \max_{0 \leq c \leq f(k)} \left\{ u(c) + \beta v^*(f(k) - c) \right\} \quad \text{for all} \quad k \in \mathbb{R}_+ \quad (2.58)$$

It states that maximal value from a given state can be obtained by trading off current reward from a given action against the (discounted) future value of the state resulting from that action.
(If the intuition behind the Bellman equation is not clear to you, try working through this lecture)

As a matter of notation, given a continuous function \( w \) on \( \mathbb{R}_+ \), we say that policy \( \sigma \in \Sigma \) is \( w \)-greedy if \( \sigma(k) \) is a solution to

\[
\max_{0 \leq c \leq f(k)} \{ u(c) + \beta w(f(k) - c) \} \tag{2.59}
\]

for every \( k \in \mathbb{R}_+ \)

**Theoretical Results**  As with most optimization problems, conditions for existence of a solution typically require some form of continuity and compactness.

In addition, some restrictions are needed to ensure that the sum of discounted utility is always finite.

For example, if we are prepared to assume that \( f \) and \( u \) are continuous and \( u \) is bounded, then

1. The value function \( v^* \) is finite, bounded, continuous and satisfies the Bellman equation
2. At least one optimal policy exists
3. A policy is optimal if and only if it is \( v^*- \)greedy

(For a proof see, for example, proposition 10.1.13 of EDTC)

In view of these results, to find an optimal policy, one option — perhaps the most common — is to

1. compute \( v^* \)
2. solve for a \( v^*- \)greedy policy

The advantage is that, once we get to the second step, we are solving a one-dimensional optimization problem — the problem on the right-hand side of (2.58)

This is much easier than an infinite-dimensional optimization problem, which is what we started out with

(An infinite sequence \( \{c_t\} \) is a point in an infinite-dimensional space)

In fact step 2 is almost trivial once \( v^* \) is obtained

For this reason, most of our focus is on the first step — how to obtain the value function

**Value Function Iteration**  The value function \( v^* \) can be obtained by an iterative technique: Starting with a guess — some initial function \( w \) — and successively improving it.

The improvement step involves applying an “operator” (a mathematical term for a function that takes a function as an input and returns a new function as an output).

The operator in question is the *Bellman operator*

The Bellman operator for this problem is a map \( T \) sending function \( w \) into function \( Tw \) via

\[
Tw(k) := \max_{0 \leq c \leq f(k)} \{ u(c) + \beta w(f(k) - c) \} \tag{2.60}
\]

Now let \( w \) be any continuous bounded function
It is known that iteratively applying $T$ from initial condition $w$ produces a sequence of functions $w, Tw, T(Tw) = T^2w, \ldots$ that converges uniformly to $v^*$

(For a proof see, for example, lemma 10.1.20 of EDTC)

This convergence will be prominent in our numerical experiments

**Unbounded Utility**  The theoretical results stated above assume that the utility function is bounded

In practice economists often work with unbounded utility functions

For utility functions that are bounded below (but possibly unbounded above), a clean and comprehensive theory now exists

(Section 12.2 of EDTC provides one exposition)

For utility functions that are unbounded both below and above the situation is more complicated

For recent work on deterministic problems, see, for example, [Kam12] or [MdRV10]

In this lecture we will use both bounded and unbounded utility functions without dwelling on the theory

**Computation**

Let’s now look at computing the value function and the optimal policy

**Fitted Value Iteration**  The first step is to compute the value function by iterating with the Bellman operator

In theory, the algorithm is as follows

1. Begin with a function $w$ — an initial condition
2. Solving (2.60), obtain the function $Tw$
3. Unless some stopping condition is satisfied, set $w = Tw$ and go to step 2

However, there is a problem we must confront before we implement this procedure: The iterates can neither be calculated exactly nor stored on a computer

To see the issue, consider (2.60)

Even if $w$ is a known function, unless $Tw$ can be shown to have some special structure, the only way to store this function is to record the value $Tw(k)$ for every $k \in \mathbb{R}_+$

Clearly this is impossible

What we will do instead is use fitted value function iteration

The procedure is to record the value of the function $Tw$ at only finitely many “grid” points $\{k_1, \ldots, k_I\} \subset \mathbb{R}_+$, and reconstruct it from this information when required

More precisely, the algorithm will be
1. Begin with an array of values \( \{w_1, \ldots, w_I\} \), typically representing the values of some initial function \( w \) on the grid points \( \{k_1, \ldots, k_I\} \)

2. Build a function \( \hat{w} \) on the state space \( \mathbb{R}_+ \) by interpolating the points \( \{w_1, \ldots, w_I\} \)

3. By repeatedly solving (2.60), obtain and record the value \( T\hat{w}(k_i) \) on each grid point \( k_i \)

4. Unless some stopping condition is satisfied, set \( \{w_1, \ldots, w_I\} = \{T\hat{w}(k_1), \ldots, T\hat{w}(k_I)\} \) and go to step 2

How should we go about step 2?

This is a problem of function approximation, and there are many ways to approach it.

What’s important here is that the function approximation scheme must not only produce a good approximation to \( Tw \), but also combine well with the broader iteration algorithm described above.

One good choice from both respects is continuous piecewise linear interpolation (see this paper for further discussion).

The next figure illustrates piecewise linear interpolation of an arbitrary function on grid points 0, 0.2, 0.4, \ldots, 1

Another advantage of piecewise linear interpolation is that it preserves useful shape properties such as monotonicity and concavity / convexity.

**A First Pass Implementation**  Let’s now look at an implementation of fitted value function iteration using Julia.

In the example below,

- \( f(k) = k^\alpha \) with \( \alpha = 0.65 \)
- \( u(c) = \ln c \) and \( \beta = 0.95 \)
As is well-known (see [LS12], section 3.1.2), for this particular problem an exact analytical solution is available, with

$$v^*(k) = c_1 + c_2 \ln k$$  \hspace{1cm} (2.61)

for

$$c_1 := \frac{\ln(1 - \alpha \beta)}{1 - \beta} + \frac{\ln(\alpha \beta)}{(1 - \alpha \beta)(1 - \beta)} \quad \text{and} \quad c_2 := \frac{\alpha}{1 - \alpha \beta}$$

At this stage, our only aim is to see if we can replicate this solution numerically, using fitted value function iteration.

Here’s a first-pass solution, the details of which are explained below.

The code can be found in file examples/optgrowth_v0.jl from the main repository.

We repeat it here for convenience.

```julia
# A first pass at solving the optimal growth problem via value function iteration. A more general version is provided in optgrowth.py.

@doc

using Optim: optimize
using Grid: CoordInterpGrid, BCnan, InterpLinear
using PyPlot

## Primitives and grid

alpha = 0.65
bet = 0.95
grid_max = 2
grid_size = 150
grid = 1e-6:(grid_max-1e-6)/(grid_size-1):grid_max

## Exact solution

ab = alpha * bet

c1 = (log(1 - ab) + log(ab) * ab / (1 - ab)) / (1 - bet)
c2 = alpha / (1 - ab)
v_star(k) = c1 .+ c2 .* log(k)

function bellman_operator(grid, w)
    Aw = CoordInterpGrid(grid, w, BCnan, InterpLinear)
    Tw = zeros(w)
    for (i, k) in enumerate(grid)
        objective(c) = - log(c) - bet * Aw[k^alpha - c]
        res = optimize(objective, 1e-6, k^alpha)
        Tw[i] = - objective(res.minimum)
    end
    return Tw
end
```

---

**Thomas Sargent and John Stachurski**  
March 21, 2015
function main(n::Int=35)
    w = 5 .* log(grid) .- 25  # An initial condition -- fairly arbitrary
    fig, ax = subplots()
    ax[:set_ylim](-40, -20)
    ax[:set_xlim](minimum(grid), maximum(grid))
    lb = "initial condition"
    jet = ColorMap("jet")[:__call__]
    ax[:plot](grid, w, color=jet(0), lw=2, alpha=0.6, label=lb)

    for i=1:n
        w = bellman_operator(grid, w)
        ax[:plot](grid, w, color=jet(i/n), lw=2, alpha=0.6)
    end
    lb = "true value function"
    ax[:plot](grid, v_star(grid), "k-", lw=2, alpha=0.8, label=lb)
    ax[:legend](loc="upper left")
    nothing
end

Running the code produces the following figure

The curves in this picture represent

1. the first 36 functions generated by the fitted value function iteration algorithm described above, with hotter colors given to higher iterates

2. the true value function as specified in (2.61), drawn in black

The sequence of iterates converges towards $v^*$
If we increase $n$ and run again we see further improvement — the next figure shows $n = 75$

Incidentally, it is true that knowledge of the functional form of $v^*$ for this model has influenced our choice of the initial condition

$$w = 5 \ast \log(\text{grid}) - 25$$

In more realistic problems such information is not available, and convergence will probably take longer

**Comments on the Code**  The function `bellman_operator` implements steps 2–3 of the fitted value function algorithm discussed above.

Linear interpolation is performed by the `getindex` method on the `CoordInterpGrid` from `Grid.jl`.

The numerical solver `optimize` from `Optim.jl` minimizes its objective, so we use the identity $\max_x f(x) = -\min_x -f(x)$ to solve (2.60)

Notice that we wrap the code used to generate the figure in a function named `main`. This allows us to import the functionality of `optgrowth_v0.jl` into a Julia session, without necessarily generating the figure.

**The Policy Function**  To compute an approximate optimal policy, we run the *fitted value function algorithm* until approximate convergence.

Taking the function so produced as an approximation to $v^*$, we then compute the (approximate) $v^*$-greedy policy.
For this particular problem, the optimal consumption policy has the known analytical solution
\[ \sigma(k) = (1 - \alpha \beta) k^\alpha \]

The next figure compares the numerical solution to this exact solution.

In the three figures, the approximation to \( \sigma^* \) is obtained by running the loop in the fitted value function algorithm 2, 4 and 6 times respectively.

Even with as few as 6 iterates, the numerical result is quite close to the true policy.

Exercise 1 asks you to reproduce this figure — although you should read the next section first.

**Writing Reusable Code**

The title of this section might sound uninteresting and a departure from our topic, but it’s equally important if not more so.

It’s understandable that many economists never consider the basic principles of software development, preoccupied as they are with the applied aspects of trying to implement their projects. However, in programming as in many things, success tends to find those who focus on what is important, not just what is urgent.
The Danger of Copy and Paste  For computing the value function of the particular growth model studied above, the code we have already written (in file \texttt{optgrowth\_v0.jl}, shown here) is perfectly fine.

However, suppose that we now want to solve a different growth model, with different technology and preferences.

Probably we want to keep our existing code, so let’s follow our first instinct and

1. copy the contents of \texttt{optgrowth\_v0.jl} to a new file
2. then make the necessary changes

Now let’s suppose that we repeat this process again and again over several years, so we now have many similar files.

(And perhaps we’re doing similar things with other projects, leading to hundreds of specialized and partially related Julia files lying around our file system.)

There are several potential problems here.

Problem 1  First, if we now realize we’ve been making some small but fundamental error with our dynamic programming all this time, we have to modify all these different files.

And now we realize that we don’t quite remember which files they were, and where exactly we put them...

So we fix all the ones we can find — spending a few hours in the process, since each implementation is slightly different and takes time to remember — and leave the ones we can’t.

Now, 6 weeks later, we need to use one of these files.

But is file X one that we have fixed, or is it not?

In this way, our code base becomes a mess, with related functionality scattered across many files, and errors creeping into them.

Problem 2  A second issue here is that since all these files are specialized and might not be used again, there’s little incentive to invest in writing them cleanly and efficiently.

DRY  The preceding discussion leads us to one of the most fundamental principles of code development: don’t repeat yourself.

To the extent that it’s practical,

- always strive to write code that is abstract and generic in order to facilitate reuse
- try to ensure that each distinct logical concept is repeated in your code base as few times as possible

To this end, we are now going to rewrite our solution to the optimal growth problem given in \texttt{optgrowth\_v0.jl} (shown above) with the intention of producing a more generic version.
While some aspects of this exercise might seem like overkill, the principles are important, and easy to illustrate in the context of the current problem.

**Implementation 2** In writing our second implementation, we want our function `bellman_operator` to be able to handle a wider class of models.

In particular, we don’t want model specifics hardwired into this function.

Instead, we would like `bellman_operator` to act in conjunction with a more general description of a model (technology, preferences, etc.)

To do so it’s convenient to wrap the model description up in a type and add the Bellman operator as a method.

(Review *this lecture* if you have forgotten the syntax for type definitions)

This idea is implemented in the code below, in file `optgrowth.jl` from the `QuantEcon` package:

```julia
#=
Solving the optimal growth problem via value function iteration.

@author: Spencer Lyon <spencer.lyon@nyu.edu>
@date: 2014-07-05

References
--------

Simple port of the file quantecon.models.optgrowth

http://quant-econ.net/dp_intro.html
#=

#=
This type defines the primitives representing the growth model. The default values are

\[
f(k) = k^{0.65}, \text{i.e., Cobb-Douglas production function}
\]
\[
u(c) = \log(c), \text{i.e., log utility}
\]

See the constructor below for details
#=

type GrowthModel
    f::Function
    bet::Real
    u::Function
    grid_max::Int
    grid_size::Int
    grid::FloatRange
end

default_f(k) = k^0.65
default_u(c) = log(c)
```
function GrowthModel(f=default_f, bet=0.95, u=default_u,
                        grid_max=2, grid_size=150)
    grid = linspace_range(1e-6, grid_max, grid_size)
    return GrowthModel(f, bet, u, grid_max, grid_size, grid)
end

#=
The approximate Bellman operator, which computes and returns the
updated value function T \(w\) on the grid points. Could also return the
policy function instead if asked.
=#
function bellman_operator!(g::GrowthModel, w::Vector, out::Vector;
                        ret_policy::Bool=false)
    # Apply linear interpolation to \(w\)
    Aw = CoordInterpGrid(g.grid, w, BCnan, InterpLinear)

    for (i, k) in enumerate(g.grid)
        objective(c) = - g.u(c) - g.bet * Aw[g.f(k) - c]
        res = optimize(objective, 1e-6, g.f(k))
        c_star = res.minimum

        if ret_policy
            # set the policy equal to the optimal \(c\)
            out[i] = c_star
        else
            # set \(T_w[i]\) equal to \( \max_c \{ u(c) + \beta w(f(k_i) - c) \}\)
            out[i] = - objective(c_star)
        end
    end

    return out
end

function bellman_operator(g::GrowthModel, w::Vector;
                        ret_policy::Bool=false)
    out = similar(w)
    bellman_operator!(g, w, out, ret_policy=ret_policy)
end

#=
Compute the \(w\)-greedy policy on the grid points.
=#
function get_greedy!(g::GrowthModel, w::Vector, out::Vector)
    bellman_operator!(g, w, out, ret_policy=true)
end

get_greedy(g::GrowthModel, w::Vector) = bellman_operator(g, w, ret_policy=true)

Of course we could omit the type structure and just pass date to bellman_operator and
compute_greedy as a list of separate arguments

For example
Tw = bellman_operator(f, beta, u, grid_max, grid_size, w)

This approach is also fine, and many prefer it.

Our own view is that the type structure is more convenient and a bit less error prone because once an instance is created we can call the methods repeatedly without having to specify a lot of arguments.

**Iteration**  The next thing we need to do is implement iteration of the Bellman operator.

Since iteratively applying an operator is something we’ll do a lot of, let’s write this as generic, reusable code.

Our code is written in the file `compute_fp.jl` from the main repository, and displayed below:

```julia
# Compute the fixed point of a given operator T, starting from specified initial condition v.

@copyright: Spencer Lyon <spencer.lyon@nyu.edu>
@date: 2014-07-05

References
---------

Simple port of the file `quantecon.compute_fp`

http://quant-econ.net/dp_intro.html

#=
# Computes and returns $T^k v$, where $T$ is an operator, $v$ is an initial condition and $k$ is the number of iterates. Provided that $T$ is a contraction mapping or similar, $T^k v$ will be an approximation to the fixed point.
#=  

```
function compute_fixed_point(T::Function, v; err_tol=1e-3, max_iter=50, verbose=true, print_skip=10)
    iterate = 0
    err = err_tol + 1
    while iterate < max_iter && err > err_tol
        new_v = T(v)
        iterate += 1
        err = Base.maxabs(new_v - v)
        if verbose
            if iterate % print_skip == 0
                println("Compute iterate ", iterate, " with error ", err)
            end
        end
        v = new_v
    end
```
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If iterate < max_iter && verbose
    println("Converged in 
$iterate steps")
elseif iterate == max_iter
    warn("max_iter exceeded in compute_fixed_point")
end

return v
end

As currently written, the code continues iteration until one of two stopping conditions holds

1. Successive iterates become sufficiently close together, in the sense that the maximum deviation between them falls below error_tol
2. The number of iterations exceeds max_iter

Examples of usage for all the code above can be found in the solutions to the exercises

Exercises

Exercise 1  Replicate the optimal policy figure shown above
Use the same parameters and initial condition found in optgrowth.jl

Exercise 2  Once an optimal consumption policy $\sigma$ is given, the dynamics for the capital stock follows (2.55)

The next figure shows the first 25 elements of this sequence for three different discount factors (and hence three different policies)

![Graph showing capital sequence over time for different discount factors]

In each sequence, the initial condition is $k_0 = 0.1$

The discount factors are discount_factors = (0.9, 0.94, 0.98)
Otherwise, the parameters and primitives are the same as found in `optgrowth.jl
Replicate the figure

Solutions

Solution notebook

2.9 LQ Control Problems

Overview

Linear quadratic (LQ) control refers to a class of dynamic optimization problems that have found applications in almost every scientific field.

This lecture provides an introduction to LQ control and its economic applications.

As we will see, LQ systems have a simple structure that makes them an excellent workhorse for a wide variety of economic problems.

Moreover, while the linear-quadratic structure is restrictive, it is in fact far more flexible than it may appear initially.

These themes appear repeatedly below.

Mathematically, LQ control problems are closely related to the Kalman filter, although we won’t pursue the deeper connections in this lecture.

In reading what follows, it will be useful to have some familiarity with

- matrix manipulations
- vectors of random variables
- dynamic programming and the Bellman equation (see for example this lecture and this lecture)

For additional reading on LQ control, see, for example,
In order to focus on computation, we leave longer proofs to these sources (while trying to provide as much intuition as possible).

**Introduction**

The “linear” part of LQ is a linear law of motion for the state, while the “quadratic” part refers to preferences.

Let’s begin with the former, move on to the latter, and then put them together into an optimization problem.

**The Law of Motion** Let $x_t$ be a vector describing the state of some economic system.

Suppose that $x_t$ follows a linear law of motion given by

$$x_{t+1} = Ax_t + Bu_t + Cw_{t+1}, \quad t = 0, 1, 2, \ldots$$  \hspace{1cm} (2.62)

Here

- $u_t$ is a “control” vector, incorporating choices available to a decision maker confronting the current state $x_t$.
- $\{w_t\}$ is an uncorrelated zero mean shock process satisfying $\mathbb{E}w_tw'_t = I$, where the right-hand side is the identity matrix.

Regarding the dimensions:

- $x_t$ is $n \times 1$, $A$ is $n \times n$.
- $u_t$ is $k \times 1$, $B$ is $n \times k$.
- $w_t$ is $j \times 1$, $C$ is $n \times j$.

**Example 1** Consider a household budget constraint given by

$$a_{t+1} + c_t = (1 + r)a_t + y_t$$

Here $a_t$ is assets, $r$ is a fixed interest rate, $c_t$ is current consumption, and $y_t$ is current non-financial income.

If we suppose that $\{y_t\}$ is uncorrelated and $N(0, \sigma^2)$, then, taking $\{w_t\}$ to be standard normal, we can write the system as

$$a_{t+1} = (1 + r)a_t - c_t + \sigma w_{t+1}$$

This is clearly a special case of (2.62), with assets being the state and consumption being the control.
Example 2  One unrealistic feature of the previous model is that non-financial income has a zero mean and is often negative.

This can easily be overcome by adding a sufficiently large mean.

Hence in this example we take \( y_t = \sigma w_{t+1} + \mu \) for some positive real number \( \mu \).

Another alteration that’s useful to introduce (we’ll see why soon) is to change the control variable from consumption to the deviation of consumption from some “ideal” quantity \( \bar{c} \).

(Most parameterizations will be such that \( \bar{c} \) is large relative to the amount of consumption that is attainable in each period, and hence the household wants to increase consumption.)

For this reason, we now take our control to be \( u_t := c_t - \bar{c} \).

In terms of these variables, the budget constraint \( a_{t+1} = (1 + r)a_t - c_t + y_t \) becomes

\[
a_{t+1} = (1 + r)a_t - u_t - \bar{c} + \sigma w_{t+1} + \mu
\]  

(2.63)

How can we write this new system in the form of equation (2.62)?

If, as in the previous example, we take \( a_t \) as the state, then we run into a problem: the law of motion contains some constant terms on the right-hand side.

This means that we are dealing with an affine function, not a linear one (recall this discussion).

Fortunately, we can easily circumvent this problem by adding an extra state variable.

In particular, if we write

\[
\begin{pmatrix}
  a_{t+1} \\
  1
\end{pmatrix}
= 
\begin{pmatrix}
  1 + r & -\bar{c} + \mu \\
  0 & 1
\end{pmatrix}
\begin{pmatrix}
  a_t \\
  1
\end{pmatrix}
+ 
\begin{pmatrix}
  -1 \\
  0
\end{pmatrix}
u_t + 
\begin{pmatrix}
  \sigma \\
  0
\end{pmatrix}w_{t+1}
\]

(2.64)

then the first row is equivalent to (2.63).

Moreover, the model is now linear, and can be written in the form of (2.62) by setting

\[
x_t := \begin{pmatrix}
  a_t \\
  1
\end{pmatrix}, \quad A := \begin{pmatrix}
  1 + r & -\bar{c} + \mu \\
  0 & 1
\end{pmatrix}, \quad B := \begin{pmatrix}
  -1 \\
  0
\end{pmatrix}, \quad C := \begin{pmatrix}
  \sigma \\
  0
\end{pmatrix}
\]

(2.65)

In effect, we’ve bought ourselves linearity by adding another state.

Preferences  In the LQ model, the aim is to minimize a flow of losses, where time-\( t \) loss is given by the quadratic expression

\[
x_t'Rx_t + u_t'Qu_t
\]  

(2.66)

Here

- \( R \) is assumed to be \( n \times n \), symmetric and nonnegative definite.
- \( Q \) is assumed to be \( k \times k \), symmetric and positive definite.

Note: In fact, for many economic problems, the definiteness conditions on \( R \) and \( Q \) can be relaxed. It is sufficient that certain submatrices of \( R \) and \( Q \) be nonnegative definite. See [HS08] for details.
Example 1  A very simple example that satisfies these assumptions is to take $R$ and $Q$ to be identity matrices, so that current loss is

$$x_t^I I x_t + u_t^I I u_t = \|x_t\|^2 + \|u_t\|^2$$

Thus, for both the state and the control, loss is measured as squared distance from the origin

(In fact the general case (2.66) can also be understood in this way, but with $R$ and $Q$ identifying other – non-Euclidean – notions of “distance” from the zero vector)

Intuitively, we can often think of the state $x_t$ as representing deviation from a target, such as

- deviation of inflation from some target level
- deviation of a firm’s capital stock from some desired quantity

The aim is to put the state close to the target, while using controls parsimoniously

Example 2  In the household problem studied above, setting $R = 0$ and $Q = 1$ yields preferences

$$x_t^I R x_t + u_t^I Q u_t = u_t^2 = (c_t - \bar{c})^2$$

Under this specification, the household’s current loss is the squared deviation of consumption from the ideal level $\bar{c}$

Optimality – Finite Horizon

Let’s now be precise about the optimization problem we wish to consider, and look at how to solve it

The Objective  We will begin with the finite horizon case, with terminal time $T \in \mathbb{N}$

In this case, the aim is to choose a sequence of controls $\{u_0, \ldots, u_{T-1}\}$ to minimize the objective

$$E \left\{ \sum_{t=0}^{T-1} \beta^t (x_t^I R x_t + u_t^I Q u_t) + \beta^T x_T^I R_f x_T \right\}$$

subject to the law of motion (2.62) and initial state $x_0$

The new objects introduced here are $\beta$ and the matrix $R_f$

The scalar $\beta$ is the discount factor, while $x^I R_f x$ gives terminal loss associated with state $x$

Comments:

- We assume $R_f$ to be $n \times n$, symmetric and nonnegative definite
- We allow $\beta = 1$, and hence include the undiscounted case
- $x_0$ may itself be random, in which case we require it to be independent of the shock sequence $w_1, \ldots, w_T$
2.9. LQ CONTROL PROBLEMS

Information  There’s one constraint we’ve neglected to mention so far, which is that the decision maker who solves this LQ problem knows only the present and the past, not the future.

To clarify this point, consider the sequence of controls \( \{u_0, \ldots, u_{T-1}\} \)

When choosing these controls, the decision maker is permitted to take into account the effects of the shocks \( \{w_1, \ldots, w_T\} \) on the system.

However, it is typically assumed — and will be assumed here — that the time-\( t \)-control \( u_t \) can only be made with knowledge of past and present shocks.

The fancy measure-theoretic way of saying this is that \( u_t \) must be measurable with respect to the \( \sigma \)-algebra generated by \( x_0, w_1, w_2, \ldots, w_t \).

This is in fact equivalent to stating that \( u_t \) can be written in the form \( u_t = g_t(x_0, w_1, w_2, \ldots, w_t) \) for some Borel measurable function \( g_t \).

(Just about every function that’s useful for applications is Borel measurable, so, for the purposes of intuition, you can read that last phrase as “for some function \( g_t \)”)

Now note that \( x_t \) will ultimately depend on the realizations of \( x_0, w_1, w_2, \ldots, w_t \).

In fact it turns out that \( x_t \) summarizes all the information about these historical shocks that the decision maker needs to set controls optimally.

More precisely, it can be shown that any optimal control \( u_t \) can always be written as a function of the current state alone.

Hence in what follows we restrict attention to control policies (i.e., functions) of the form \( u_t = g_t(x_t) \).

Actually, the preceding discussion applies to all standard dynamic programming problems.

What’s special about the LQ case is that — as we shall soon see — the optimal \( u_t \) turns out to be a linear function of \( x_t \).

Solution  To solve the finite horizon LQ problem we can use a dynamic programming strategy based on backwards induction that is conceptually similar to the approach adopted in this lecture.

For reasons that will soon become clear, we first introduce the notation \( J_T(x) := x'R_f x \).

Now consider the problem of the decision maker in the second to last period.

In particular, let the time be \( T-1 \), and suppose that the state is \( x_{T-1} \).

The decision maker must trade off current and (discounted) final losses, and hence solves

\[
\min_u \{ x_{T-1}' R x_{T-1} + u' Q u + \beta \mathbb{E} J_T(A x_{T-1} + B u + C w_T) \}
\]

At this stage, it is convenient to define the function

\[
J_{T-1}(x) := \min_u \{ x' R x + u' Q u + \beta \mathbb{E} J_T(A x + B u + C w_T) \} \tag{2.68}
\]

The function \( J_{T-1} \) will be called the \( T-1 \) value function, and \( J_{T-1}(x) \) can be thought of as representing total “loss-to-go” from state \( x \) at time \( T-1 \) when the decision maker behaves optimally.
Now let’s step back to $T - 2$

For a decision maker at $T - 2$, the value $J_{T-1}(x)$ plays a role analogous to that played by the terminal loss $J_T(x) = x'Rf/x$ for the decision maker at $T - 1$

That is, $J_{T-1}(x)$ summarizes the future loss associated with moving to state $x$

The decision maker chooses her control $u$ to trade off current loss against future loss, where

- the next period state is $x_{T-1} = Ax_{T-2} + Bu + Cw_{T-1}$, and hence depends on the choice of current control
- the “cost” of landing in state $x_{T-1}$ is $J_{T-1}(x_{T-1})$

Her problem is therefore

$$\min_u \{x'_{T-2}Rx_{T-2} + u'Qu + \beta \mathbb{E}J_{T-1}(Ax_{T-2} + Bu + Cw_{T-1})\}$$

Letting

$$J_{T-2}(x) := \min_u \{x'Rx + u'Qu + \beta \mathbb{E}J_{T-1}(Ax + Bu + Cw_{T-1})\}$$

the pattern for backwards induction is now clear

In particular, we define a sequence of value functions $\{J_0, \ldots, J_T\}$ via

$$J_t(x) = \min_u \{x'Rx + u'Qu + \beta \mathbb{E}J_{t+1}(Ax + Bu + Cw_t)\} \quad \text{and} \quad J_T(x) = x'Rf/x$$

The first equality is the Bellman equation from dynamic programming theory specialized to the finite horizon LQ problem

Now that we have $\{J_0, \ldots, J_T\}$, we can obtain the optimal controls

As a first step, let’s find out what the value functions look like

It turns out that every $J_t$ has the form $J_t(x) = x'P_tx + d_t$ where $P_t$ is a $n \times n$ matrix and $d_t$ is a constant

We can show this by induction, starting from $P_T := Rf$ and $d_T = 0$

Using this notation, (2.68) becomes

$$J_{T-1}(x) := \min_u \{x'Rx + u'Qu + \beta \mathbb{E}J_{T}(Ax + Bu + Cw_{T})\} \quad (2.69)$$

To obtain the minimizer, we can take the derivative of the r.h.s. with respect to $u$ and set it equal to zero

Applying the relevant rules of matrix calculus, this gives

$$u = -(Q + \beta B'P_TB)^{-1}\beta B'P_TA$$ \quad (2.70)

Plugging this back into (2.69) and rearranging yields

$$J_{T-1}(x) := x'P_{T-1}x + d_{T-1}$$

where

$$P_{T-1} := R - \beta^2A'P_TB(Q + \beta B'P_TB)^{-1}B'P_TA + \beta A'P_TA$$ \quad (2.71)
and
\[ d_{T-1} := \beta \text{trace}(C'P_tC) \] (2.72)
(The algebra is a good exercise — we’ll leave it up to you)
If we continue working backwards in this manner, it soon becomes clear that \( J_t(x) := x'P_tx + d_t \) as claimed, where \( \{P_t\} \) and \( \{d_t\} \) satisfy the recursions
\[ P_{t-1} := R - \beta^2 A'P_tB(Q + \beta B'P_tB)^{-1}B'P_tA + \beta A'P_tA \quad \text{with} \quad P_T = R_f \] (2.73)
and
\[ d_{t-1} := \beta(d_t + \text{trace}(C'P_tC)) \quad \text{with} \quad d_T = 0 \] (2.74)
Recalling (2.70), the minimizers from these backward steps are
\[ u_t = -F_t x_t \quad \text{where} \quad F_t := (Q + \beta B'P_{t+1}B)^{-1}\beta B'P_{t+1}A \] (2.75)
These are the linear optimal control policies we discussed above
In particular, the sequence of controls given by (2.75) and (2.62) solves our finite horizon LQ problem
Rephrasing this more precisely, the sequence \( u_0, \ldots, u_{T-1} \) given by
\[ u_t = -F_t x_t \quad \text{with} \quad x_{t+1} = (A - BF_t)x_t + Cw_{t+1} \] (2.76)
for \( t = 0, \ldots, T - 1 \) attains the minimum of (2.67) subject to our constraints

**An Application**  Early Keynesian models assumed that households have a constant marginal propensity to consume from current income
Data contradicted the constancy of the marginal propensity to consume
In response, Milton Friedman, Franco Modigliani and many others built models based on a consumer’s preference for a stable consumption stream
(See, for example, [Fri56] or [MB54])
One property of those models is that households purchase and sell financial assets to make consumption streams smoother than income streams
The household savings problem outlined above captures these ideas
The optimization problem for the household is to choose a consumption sequence in order to minimize
\[ \mathbb{E} \left\{ \sum_{t=0}^{T-1} \beta^t (c_t - \bar{c})^2 + \beta^T qa_T^2 \right\} \] (2.77)
subject to the sequence of budget constraints \( a_{t+1} = (1+r)a_t - c_t + y_t, \ t \geq 0 \)
Here \( q \) is a large positive constant, the role of which is to induce the consumer to target zero debt at the end of her life
(Without such a constraint, the optimal choice is to choose \( c_t = \bar{c} \) in each period, letting assets adjust accordingly)
As before we set \( y_t = \sigma w_{t+1} + \mu \) and \( u_t := c_t - \overline{c} \), after which the constraint can be written as in (2.63).

We saw how this constraint could be manipulated into the LQ formulation \( x_{t+1} = Ax_t + Bu_t + Cw_{t+1} \) by setting \( x_t = (a_t, 1)' \) and using the definitions in (2.65).

To match with this state and control, the objective function (2.77) can be written in the form of (2.67) by choosing
\[
Q := 1, \quad R := \begin{pmatrix} 0 & 0 \\ 0 & 0 \end{pmatrix}, \quad \text{and} \quad R_f := \begin{pmatrix} q & 0 \\ 0 & 0 \end{pmatrix}
\]

Now that the problem is expressed in LQ form, we can proceed to the solution by applying (2.73) and (2.75).

After generating shocks \( w_1, \ldots, w_T \), the dynamics for assets and consumption can be simulated via (2.76).

We provide code for all these operations below.

The following figure was computed using this code, with \( r = 0.05, \beta = 1/(1+r), \overline{c} = 2, \mu = 1, \sigma = 0.25, T = 45 \) and \( q = 10^6 \).

The shocks \( \{w_t\} \) were taken to be iid and standard normal.

The top panel shows the time path of consumption \( c_t \) and income \( y_t \) in the simulation.

As anticipated by the discussion on consumption smoothing, the time path of consumption is much smoother than that for income.

(But note that consumption becomes more irregular towards the end of life, when the zero final asset requirement impinges more on consumption choices)

The second panel in the figure shows that the time path of assets \( a_t \) is closely correlated with cumulative unanticipated income, where the latter is defined as
\[
z_t := \sum_{j=0}^{t} \sigma w_j
\]

A key message is that unanticipated windfall gains are saved rather than consumed, while unanticipated negative shocks are met by reducing assets.

(Again, this relationship breaks down towards the end of life due to the zero final asset requirement).

These results are relatively robust to changes in parameters.

For example, let’s increase \( \beta \) from \( 1/(1+r) \approx 0.952 \) to 0.96 while keeping other parameters fixed.

This consumer is slightly more patient than the last one, and hence puts relatively more weight on later consumption values.

A simulation is shown below.

We now have a slowly rising consumption stream and a hump-shaped build up of assets in the middle periods to fund rising consumption.

However, the essential features are the same: consumption is smooth relative to income, and assets are strongly positively correlated with cumulative unanticipated income.
2.9. LQ CONTROL PROBLEMS

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Extensions and Comments

Let’s now consider a number of standard extensions to the LQ problem treated above.

Nonstationary Parameters In some settings it can be desirable to allow $A, B, C, R$ and $Q$ to depend on $t$.

For the sake of simplicity, we’ve chosen not to treat this extension in our implementation given below.

However, the loss of generality is not as large as you might first imagine.

In fact, we can tackle many nonstationary models from within our implementation by suitable choice of state variables.

One illustration is given below.

For further examples and a more systematic treatment, see [HS13], section 2.4.

Adding a Cross-Product Term In some LQ problems, preferences include a cross-product term $u_t'N x_t$, so that the objective function becomes

$$
\mathbb{E}\left\{ \sum_{t=0}^{T-1} \beta^t (x_t' Rx_t + u_t' Qu_t + 2u_t' N x_t) + \beta^T x_T' R_f x_T \right\}
$$

(2.78)

Our results extend to this case in a straightforward way.

The sequence $\{P_t\}$ from (2.73) becomes

$$
P_{t-1} := R - (\beta B'P_t A + N)'(Q + \beta B'P_t A + N) + \beta A' P_t A \quad \text{with} \quad P_T = R_f
$$

(2.79)

The policies in (2.75) are modified to

$$
u_t = -F_t x_t \quad \text{where} \quad F_t := (Q + \beta B'P_{t+1} A + N)^{-1}(\beta B'P_{t+1} A + N)
$$

(2.80)

The sequence $\{d_t\}$ is unchanged from (2.74).

We leave interested readers to confirm these results (the calculations are long but not overly difficult).

Infinite Horizon Finally, we consider the infinite horizon case, with cross-product term, unchanged dynamics and objective function given by

$$
\mathbb{E}\left\{ \sum_{t=0}^{\infty} \beta^t (x_t' Rx_t + u_t' Qu_t + 2u_t' N x_t) \right\}
$$

(2.81)

In the infinite horizon case, optimal policies can depend on time only if time itself is a component of the state vector $x_t$.

In other words, there exists a fixed matrix $F$ such that $u_t = -Fx_t$ for all $t$. 

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This stationarity is intuitive — after all, the decision maker faces the same infinite horizon at every stage, with only the current state changing.

Not surprisingly, $P$ and $d$ are also constant.

The stationary matrix $P$ is given by the fixed point of (2.73)

Equivalently, it is the solution $P$ to the discrete time algebraic Riccati equation

$$P := R - (\beta B'PA + N)'(Q + \beta B'PB)^{-1}(\beta B'PA + N) + \beta A'PA$$

Equation (2.82) is also called the LQ Bellman equation, and the map that sends a given $P$ into the right-hand side of (2.82) is called the LQ Bellman operator.

The stationary optimal policy for this model is

$$u = -Fx$$

where

$$F := (Q + \beta B'PB)^{-1}(\beta B'PA + N)$$

The sequence $\{d_t\}$ from (2.74) is replaced by the constant value

$$d := \text{trace}(C'PC)\frac{\beta}{1 - \beta}$$

The state evolves according to the time-homogeneous process $x_{t+1} = (A - BF)x_t + Cw_{t+1}$

An example infinite horizon problem is treated below.

**Certainty Equivalence** Linear quadratic control problems of the class discussed above have the property of certainty equivalence.

By this we mean that the optimal policy $F$ is not affected by the parameters in $C$, which specify the shock process.

This can be confirmed by inspecting (2.83) or (2.80).

It follows that we can ignore uncertainty when solving for optimal behavior, and plug it back in when examining optimal state dynamics.

**Implementation**

We have put together some code for solving finite and infinite horizon linear quadratic control problems.

The code can be found in the file lqcontrol.jl from the QuantEcon package.

You can view the program on GitHub but we repeat it here for convenience.

```julia
#=
Provides a type called LQ for solving linear quadratic control problems.

@author : Spencer Lyon <spencer.lyon@nyu.edu>

@date : 2014-07-05
```

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2.9. LQ CONTROL PROBLEMS

References
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Simple port of the file quantecon.lqcontrol

http://quant-econ.net/lqcontrol.html

#

type LQ
    Q::Matrix
    R::Matrix
    A::Matrix
    B::Matrix
    C::Union(Nothing, Matrix)
    bet::Real
    T::Union(Int, Nothing)
    Rf::Matrix
    k::Int
    n::Int
    j::Int
    P::Matrix
    d::Real
    F::Matrix
end

function LQ(Q::ScalarOrArray,
            R::ScalarOrArray,
            A::ScalarOrArray,
            B::ScalarOrArray,
            C::Union(Nothing, ScalarOrArray)=nothing,
            bet::ScalarOrArray=1.0,
            T::Union(Int, Nothing)=nothing,
            Rf::Union(Nothing, ScalarOrArray)=nothing)
    k = size(Q, 1)
    n = size(R, 1)

    if C == nothing
        j = 1
        C = zeros(n, j)
    else
        j = size(C, 2)
        if j == 1
            C = reshape([C], n, j) # make sure C is a Matrix
        end
    end

    if Rf == nothing
        Rf = fill(NaN, size(R)...)  
    end

    # Reshape arrays to make sure they are Matrix
Q = reshape([Q], k, k)
R = reshape([R], n, n)
A = reshape([A], n, n)
B = reshape([B], n, k)
Rf = reshape([Rf], n, n)
F = zeros(Float64, k, n)
P = copy(Rf)
d = 0.0
LQ(Q, R, A, B, C, bet, T, Rf, k, n, j, P, d, F)
end

# make kwarg version
function LQ(Q::ScalarOrArray,
    R::ScalarOrArray,
    A::ScalarOrArray,
    B::ScalarOrArray;
    C::Union(Nothing, ScalarOrArray)=nothing,
    bet::ScalarOrArray=1.0,
    T::Union(Int, Nothing)=nothing,
    Rf::Union(Nothing, ScalarOrArray)=nothing)
    LQ(Q, R, A, B, C, bet, T, Rf)
end

function update_values!(lq::LQ)
    # Simplify notation

    # Some useful matrices
    S1 = Q .+ lq.bet .* (B' * P * B)
    S2 = lq.bet .* (B' * P * A)
    S3 = lq.bet .* (A' * P * A)

    # Compute F as (Q + B'PB)^{-1} (beta B'PA)
    lq.F = S1 \ S2

    # Shift P back in time one step
    new_P = R - S2' * lq.F + S3

    # Recalling that trace(AB) = trace(BA)
    new_d = lq.bet * (d + trace(P * C * C'))

    # Set new state
    lq.P = new_P
    lq.d = new_d
    return nothing
end

function stationary_values!(lq::LQ)
# simplify notation

# solve Riccati equation, obtain P
A0, B0 = sqrt(lq.bet) .* A, sqrt(lq.bet) .* B
P = solve_discrete_riccati(A0, B0, R, Q)

# Compute F
S1 = Q .+ lq.bet .* (B' * P * B)
S2 = lq.bet .* (B' * P * A)
F = S1 \ S2

# Compute d
d = lq.bet .* trace(P * C * C') / (1 - lq.bet)

# Bind states
nothing

end

function stationary_values(lq::LQ)
    stationary_values!(lq)
    return lq.P, lq.F, lq.d
end

function compute_sequence(lq::LQ, x0::ScalarOrArray, ts_length=100)
    # simplify notation

    # Preliminaries,
    if lq.T != nothing
        # finite horizon case
        T = min(ts_length, lq.T)
        lq.P, lq.d = lq.Rf, 0.0
    else
        # infinite horizon case
        T = ts_length
        stationary_values!(lq)
    end

    # Set up initial condition and arrays to store paths
    x0 = reshape([x0], lq.n, 1)  # make sure x0 is a column vector
    x_path = Array(eltype(x0), lq.n, T+1)
    u_path = Array(eltype(x0), lq.k, T)
    w_path = C * randn(lq.j, T+1)

    # Compute and record the sequence of policies
    policies = Array(typeof(lq.F), T)
    for t=1:T
        if lq.T != nothing
            update_values!(lq)
        end
        # ...
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end

policies[t] = lq.F
end

# Use policy sequence to generate states and controls
F = pop!(policies)
x_path[:, 1] = x0
u_path[:, 1] = -(F * x0)

for t=2:T
    F = pop!(policies)
    Ax, Bu = A * x_path[:, t-1], B * u_path[:, t-1]
    x_path[:, t] = Ax .+ Bu .+ w_path[:, t]
    u_path[:, t] = -(F * x_path[:, t])
end

Ax, Bu = A * x_path[:, T], B * u_path[:, T]
x_path[:, T+1] = Ax .+ Bu .+ w_path[:, T+1]
return x_path, u_path, w_path
end

In the module, the various updating, simulation and fixed point methods are wrapped in a type called LQ, which includes

- Instance data:
  - The required parameters $Q, R, A, B$ and optional parameters $C, \beta, T, R_f, N$ specifying a given LQ model
    * set $T$ and $R_f$ to None in the infinite horizon case
    * set $C = \text{None}$ (or zero) in the deterministic case
  - the value function and policy data
    * $d_t, P_t, F_t$ in the finite horizon case
    * $d, P, F$ in the infinite horizon case

- Methods:
  - update_values — shifts $d_t, P_t, F_t$ to their $t-1$ values via (2.73), (2.74) and (2.75)
  - stationary_values — computes $P, d, F$ in the infinite horizon case
  - compute_sequence — simulates the dynamics of $x_t, u_t, w_t$ given $x_0$ and assuming standard normal shocks

An example of usage is given in lq_permanent_1.jl from the main repository, the contents of which are shown below

This program can be used to replicate the figures shown in our section on the permanent income model

(Some of the plotting techniques are rather fancy and you can ignore those details if you wish)
2.9. LQ CONTROL PROBLEMS

\[
\begin{align*}
\sigma & = 0.25 \\
\mu & = 1.0 \\
q & = 1e6 \\
# == Formulate as an LQ problem == # \\
Q & = 1.0 \\
R & = \text{zeros}(2, 2) \\
Rf & = \text{zeros}(2, 2); Rf[1, 1] = q \\
A & = [1.0+r -c\_bar+\mu; \\
& \quad 0.0 \quad 1.0] \\
B & = [-1.0, 0.0] \\
C & = [\sigma, 0.0] \\
# == Compute solutions and simulate == # \\
lq & = \text{LQ}(Q, R, A, B, C, bet, T, Rf) \\
x0 & = [0.0, 1.0] \\
xp, up, wp & = \text{compute_sequence}(lq, x0) \\
# == Convert back to assets, consumption and income == # \\
assets & = \text{squeeze}(xp[1, :, 1], 1) \quad # a_t \\
c & = \text{squeeze}(up + c\_bar, 1) \quad # c_t \\
income & = \text{squeeze}(wp[1, 2:end] + mu, 1) \quad # y_t \\
# == Plot results == # \\
n\_rows & = 2 \\
fig, axes = \text{subplots}(n\_rows, 1, figsize=(12, 10)) \\
subplots\_adjust(hspace=0.5) \\
for i=1:n\_rows \\
\text{axes[i][:grid]()} \\
\text{axes[i][:set\_xlabel]("Time")} \\
end \\
bbox & = [0.0, 1.02, 1.0, 0.102] \\
axes[1][:plot](2:T+1, income, "g-", label="non-financial income", lw=2, alpha=0.7) \\
axes[1][:plot](t; c, "k-", label="consumption", lw=2, alpha=0.7) \\
axes[1][:legend](ncol=2, bbox_to_anchor=bbox, loc=3, mode="expand") \\
axes[2][:plot](2:T+1, cumsum(income .- mu), "r-", label="cumulative unanticipated income", lw=2, alpha=0.7) \\
axes[2][:plot](t; assets, "b-", label="assets", lw=2, alpha=0.7) \\
axes[2][:plot](t; zeros(T), "k-") \\
axes[2][:legend](ncol=2, bbox_to_anchor=bbox, loc=3, mode="expand") \\

Further Applications

Application 1: Nonstationary Income  \textit{Previously} we studied a permanent income model that generated consumption smoothing.

One unrealistic feature of that model is the assumption that the mean of the random income process does not depend on the consumer’s age.

A more realistic income profile is one that rises in early working life, peaks towards the middle
and maybe declines toward end of working life, and falls more during retirement. 

In this section, we will model this rise and fall as a symmetric inverted “U” using a polynomial in age.

As before, the consumer seeks to minimize

$$\mathbb{E} \left\{ \sum_{t=0}^{T-1} \beta^t (c_t - \bar{c})^2 + \beta^T q a_T^2 \right\}$$

(2.85)

subject to $a_{t+1} = (1 + r)a_t - c_t + y_t$, $t \geq 0$

For income we now take $y_t = p(t) + \sigma w_{t+1}$ where $p(t) := m_0 + m_1 t + m_2 t^2$

(In the next section we employ some tricks to implement a more sophisticated model)

The coefficients $m_0, m_1, m_2$ are chosen such that $p(0) = 0$, $p(T/2) = \mu$, and $p(T) = 0$.

You can confirm that the specification $m_0 = 0, m_1 = T \mu / (T/2)^2, m_2 = -\mu / (T/2)^2$ satisfies these constraints.

To put this into an LQ setting, consider the budget constraint, which becomes

$$a_{t+1} = (1 + r)a_t - c_t + m_1 t + m_2 t^2 + \sigma w_{t+1}$$

(2.86)

The fact that $a_{t+1}$ is a linear function of $(a_t, 1, t, t^2)$ suggests taking these four variables as the state vector $x_t$.

Once a good choice of state and control (recall $u_t = c_t - \bar{c}$) has been made, the remaining specifications fall into place relatively easily.

Thus, for the dynamics we set

$$x_t := \begin{pmatrix} a_t \\ 1 \\ t \\ t^2 \end{pmatrix}, \quad A := \begin{pmatrix} 1 + r & -\bar{c} & m_1 & m_2 \\ 0 & 1 & 0 & 0 \\ 0 & 1 & 1 & 0 \\ 0 & 1 & 2 & 1 \end{pmatrix}, \quad B := \begin{pmatrix} -1 \\ 0 \\ 0 \\ 0 \end{pmatrix}, \quad C := \begin{pmatrix} \sigma \\ 0 \\ 0 \end{pmatrix}$$

(2.87)

If you expand the expression $x_{t+1} = Ax_t + Bu_t + Cw_{t+1}$ using this specification, you will find that assets follow (2.86) as desired, and that the other state variables also update appropriately.

To implement preference specification (2.85) we take

$$Q := 1, \quad R := \begin{pmatrix} 0 & 0 & 0 & 0 \\ 0 & 0 & 0 & 0 \\ 0 & 0 & 0 & 0 \\ 0 & 0 & 0 & 0 \end{pmatrix} \quad \text{and} \quad R_f := \begin{pmatrix} q & 0 & 0 & 0 \\ 0 & 0 & 0 & 0 \\ 0 & 0 & 0 & 0 \\ 0 & 0 & 0 & 0 \end{pmatrix}$$

(2.88)

The next figure shows a simulation of consumption and assets computed using the compute_sequence method of lqcontrol.jl with initial assets set to zero.

Once again, smooth consumption is a dominant feature of the sample paths.

The asset path exhibits dynamics consistent with standard life cycle theory.

Exercise 1 gives the full set of parameters used here and asks you to replicate the figure.
2.9. LQ CONTROL PROBLEMS

![Graph showing non-financial income, consumption, and assets over time.](image)

- Non-financial income (green line)
- Consumption (black line)
- Assets (blue line)

Time axis ranges from 0 to 50.
**Application 2: A Permanent Income Model with Retirement**  
In the *previous application*, we generated income dynamics with an inverted U shape using polynomials, and placed them in an LQ framework.

It is arguably the case that this income process still contains unrealistic features.

A more common earning profile is where

1. income grows over working life, fluctuating around an increasing trend, with growth flattening off in later years
2. retirement follows, with lower but relatively stable (non-financial) income

Letting $K$ be the retirement date, we can express these income dynamics by

$$y_t = \begin{cases} p(t) + \sigma w_{t+1} & \text{if } t \leq K \\ s & \text{otherwise} \end{cases}$$  \hspace{1cm} (2.89)

Here

- $p(t) := m_1 t + m_2 t^2$ with the coefficients $m_1, m_2$ chosen such that $p(K) = \mu$ and $p(0) = p(2K) = 0$
- $s$ is retirement income

We suppose that preferences are unchanged and given by (2.77)

The budget constraint is also unchanged and given by $a_{t+1} = (1 + r)a_t - c_t + y_t$

Our aim is to solve this problem and simulate paths using the LQ techniques described in this lecture.

In fact this is a nontrivial problem, as the kink in the dynamics (2.89) at $K$ makes it very difficult to express the law of motion as a fixed-coefficient linear system.

However, we can still use our LQ methods here by suitably linking two component LQ problems.

These two LQ problems describe the consumer’s behavior during her working life (`lq_working`) and retirement (`lq_retired`).

(This is possible because in the two separate periods of life, the respective income processes [polynomial trend and constant] each fit the LQ framework)

The basic idea is that although the whole problem is not a single time-invariant LQ problem, it is still a dynamic programming problem, and hence we can use appropriate Bellman equations at every stage.

Based on this logic, we can

1. solve `lq_retired` by the usual backwards induction procedure, iterating back to the start of retirement
2. take the start-of-retirement value function generated by this process, and use it as the terminal condition $R_f$ to feed into the `lq_working` specification
3. solve `lq_working` by backwards induction from this choice of $R_f$, iterating back to the start of working life

---

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This process gives the entire life-time sequence of value functions and optimal policies. The next figure shows one simulation based on this procedure.

![Graph showing non-financial income and consumption over time.](image1)

![Graph showing assets over time.](image2)

The full set of parameters used in the simulation is discussed in Exercise 2, where you are asked to replicate the figure.

Once again, the dominant feature observable in the simulation is consumption smoothing.

The asset path fits well with standard life cycle theory, with dissaving early in life followed by later saving.

Assets peak at retirement and subsequently decline.

**Application 3: Monopoly with Adjustment Costs** Consider a monopolist facing stochastic inverse demand function

\[ p_t = a_0 - a_1 q_t + d_t \]

Here \( q_t \) is output, and the demand shock \( d_t \) follows

\[ d_{t+1} = \rho d_t + \sigma w_{t+1} \]
where \( \{w_t\} \) is iid and standard normal.

The monopolist maximizes the expected discounted sum of present and future profits

\[
E \left\{ \sum_{t=0}^{\infty} \beta^t \pi_t \right\}
\]

where \( \pi_t := p_tq_t - cq_t - \gamma(q_{t+1} - q_t)^2 \) (2.90)

Here

- \( \gamma(q_{t+1} - q_t)^2 \) represents adjustment costs
- \( c \) is average cost of production

This can be formulated as an LQ problem and then solved and simulated, but first let’s study the problem and try to get some intuition.

One way to start thinking about the problem is to consider what would happen if \( \gamma = 0 \).

Without adjustment costs there is no intertemporal trade-off, so the monopolist will choose output to maximize current profit in each period.

It’s not difficult to show that profit-maximizing output is

\[
\bar{q}_t := \frac{a_0 - c + d_t}{2a_1}
\]

In light of this discussion, what we might expect for general \( \gamma \) is that

- if \( \gamma \) is close to zero, then \( q_t \) will track the time path of \( \bar{q}_t \) relatively closely
- if \( \gamma \) is larger, then \( q_t \) will be smoother than \( \bar{q}_t \), as the monopolist seeks to avoid adjustment costs

This intuition turns out to be correct.

The following figures show simulations produced by solving the corresponding LQ problem.

The only difference in parameters across the figures is the size of \( \gamma \).

To produce these figures we converted the monopolist problem into an LQ problem.

The key to this conversion is to choose the right state — which can be a bit of an art.

Here we take \( x_t = (\bar{q}_t \ q_t \ 1)' \), while the control is chosen as \( u_t = q_{t+1} - q_t \).

We also manipulated the profit function slightly.

In (2.90), current profits are \( \pi_t := p_tq_t - cq_t - \gamma(q_{t+1} - q_t)^2 \).

Let’s now replace \( \pi_t \) in (2.90) with \( \hat{\pi}_t := \pi_t - a_1\bar{q}_t^2 \).

This makes no difference to the solution, since \( a_1\bar{q}_t^2 \) does not depend on the controls.
2.9. LQ CONTROL PROBLEMS

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(In fact we are just adding a constant term to (2.90), and optimizers are not affected by constant terms)

The reason for making this substitution is that, as you will be able to verify, \( \hat{\pi}_t \) reduces to the simple quadratic

\[
\hat{\pi}_t = -a_1 (q_t - \bar{q}_t)^2 - \gamma u_t^2
\]

After negation to convert to a minimization problem, the objective becomes

\[
\min E \sum_{t=0}^{\infty} \beta^t \{ a_1 (q_t - \bar{q}_t)^2 + \gamma u_t^2 \}
\]  

(2.91)

It’s now relatively straightforward to find \( R \) and \( Q \) such that (2.91) can be written as (2.81)

Furthermore, the matrices \( A, B \) and \( C \) from (2.62) can be found by writing down the dynamics of each element of the state

Exercise 3 asks you to complete this process, and reproduce the preceding figures

Exercises

Exercise 1  Replicate the figure with polynomial income *shown above*

The parameters are \( r = 0.05, \beta = 1/(1 + r), \bar{c} = 1.5, \mu = 2, \sigma = 0.15, T = 50 \) and \( q = 10^4 \)

Exercise 2  Replicate the figure on work and retirement *shown above*

The parameters are \( r = 0.05, \beta = 1/(1 + r), \bar{c} = 4, \mu = 4, \sigma = 0.35, K = 40, T = 60, s = 1 \) and \( q = 10^4 \)

To understand the overall procedure, carefully read the section containing that figure

Some hints are as follows:

First, in order to make our approach work, we must ensure that both LQ problems have the same state variables and control

As with previous applications, the control can be set to \( u_t = c_t - \bar{c} \)

For \( lq\_working \), \( x_t, A, B, C \) can be chosen as in (2.87)

• Recall that \( m_1, m_2 \) are chosen so that \( p(K) = \mu \) and \( p(2K) = 0 \)

For \( lq\_retired \), use the same definition of \( x_t \) and \( u_t \), but modify \( A, B, C \) to correspond to constant income \( y_t = s \)

For \( lq\_retired \), set preferences as in (2.88)

For \( lq\_working \), preferences are the same, except that \( R_f \) should be replaced by the final value function that emerges from iterating \( lq\_retired \) back to the start of retirement

With some careful footwork, the simulation can be generated by patching together the simulations from these two separate models
Exercise 3  Reproduce the figures from the monopolist application *given above*

For parameters, use $a_0 = 5, a_1 = 0.5, \sigma = 0.15, \rho = 0.9, \beta = 0.95$ and $c = 2$, while $\gamma$ varies between 1 and 50 (see figures)

**Solutions**

Solution notebook

## 2.10 Rational Expectations Equilibrium

**Contents**

- Rational Expectations Equilibrium
  - Overview
  - Defining Rational Expectations Equilibrium
  - Computation of the Equilibrium
  - Exercises
  - Solutions

“If you’re so smart, why aren’t you rich?”

**Overview**

This lecture introduces the concept of *rational expectations equilibrium*

To illustrate it, we describe a linear quadratic version of a famous and important model due to Lucas and Prescott [LP71]

This 1971 paper is one of a small number of research articles that kicked off the *rational expectations revolution*

We follow Lucas and Prescott by employing a setting that is readily “Bellmanized” (i.e., capable of being formulated in terms of dynamic programming problems)

Because we use linear quadratic setups for demand and costs, we can adapt the LQ programming techniques described in *this lecture*

We will learn about how a representative agent’s problem differs from a planner’s, and how a planning problem can be used to compute rational expectations quantities

We will also learn about how a rational expectations equilibrium can be characterized as a fixed point of a mapping from a perceived law of motion to an actual law of motion

Equality between a perceived and an actual law of motion for endogenous market-wide objects captures in a nutshell what the rational expectations equilibrium concept is all about

Finally, we will learn about the important “Big $K$, little $k$” trick, a modeling device widely used in macroeconomics

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Except that for us

- Instead of “Big K” it will be “Big Y”
- Instead of “little k” it will be “little y”

**The Big Y, little y trick**  This widely used method applies in contexts in which a “representative firm” or agent is a “price taker” operating within a competitive equilibrium.

We want to impose that

- The representative firm or individual takes *aggregate Y* as given when it chooses individual *y*, but . . .
- At the end of the day, *Y* = *y*, so that the representative firm is indeed representative.

The Big Y, little y trick accomplishes these two goals by

- Taking *Y* as a given “state” variable or process, beyond the control of the representative individual, when posing the problem of the individual firm or agent; but . . .
- Imposing *Y* = *y* after having solved the individual’s optimization problem.

Please watch for how this strategy is applied as the lecture unfolds.

We begin by applying the Big Y, little y trick in a very simple static context.

**A simple static example of the Big Y, little y trick**  Consider a static model in which a collection of *n* firms produce a homogeneous good that is sold in a competitive market.

Each of these *n* firms sells output *y*.

The price *p* of the good lies on an inverse demand curve

\[ p = a_0 - a_1 Y \]  \hspace{1cm} (2.92)

where

- \( a_i > 0 \) for \( i = 0, 1 \)
- \( Y = ny \) is the market-wide level of output

Each firm has total cost function

\[ c(y) = c_1 y + 0.5 c_2 y^2, \quad c_i > 0 \text{ for } i = 1, 2 \]

The profits of a representative firm are \( py - c(y) \).

Using (2.92), we can express the problem of the representative firm as

\[ \max_y \left[ (a_0 - a_1 Y)y - c_1 y - 0.5 c_2 y^2 \right] \]

In posing problem (2.93), we want the firm to be a *price taker*.

We do that by regarding *p* and therefore *Y* as exogenous to the firm.
The essence of the Big $Y$, little $y$ trick is not to set $Y = ny$ before taking the first-order condition with respect to $y$ in problem (2.93)

The first order condition for problem (2.93) is

$$a_0 - a_1 Y - c_1 - c_2 y = 0$$ (2.94)

At this point, but not before, we substitute $Y = ny$ into (2.94) to obtain the following linear equation

$$a_0 - c_1 - (a_1 + n^{-1} c_2) Y = 0$$ (2.95)

to be solved for the competitive equilibrium market wide output $Y$

After solving for $Y$, we can compute the competitive equilibrium price from the inverse demand curve (2.92)

**Further Reading** References for this lecture include

- [LP71]
- [Sar87], chapter XIV
- [LS12], chapter 7

**Defining Rational Expectations Equilibrium**

Our first illustration of rational expectations equilibrium involves a market with $n$ firms, each of whom seeks to maximize profits in the face of adjustment costs

The adjustment costs encourage the firms to make gradual adjustments, which in turn requires consideration of future prices

Individual firms understand that prices are determined by aggregate supply from other firms, and hence each firm must forecast this quantity

In our context, a forecast is expressed as a belief about the law of motion for the aggregate state

Rational expectations equilibrium is obtained when this belief coincides with the actual law of motion generated by production choices made on the basis of this belief

**Competitive Equilibrium with Adjustment Costs** To illustrate, consider a collection of $n$ firms producing a homogeneous good that is sold in a competitive market.

Each of these $n$ firms sells output $y_t$

The price $p_t$ of the good lies on the inverse demand curve

$$p_t = a_0 - a_1 Y_t$$ (2.96)

where

- $a_i > 0$ for $i = 0, 1$
- $Y_t = ny_t$ is the market-wide level of output
2.10. RATIONAL EXPECTATIONS EQUILIBRIUM

The Firm’s Problem  The firm is a price taker
While it faces no uncertainty, it does face adjustment costs
In particular, it chooses a production plan to maximize
\[ \sum_{t=0}^{\infty} \beta^t r_t \]  \hspace{1cm} (2.97)
where
\[ r_t := p_t y_t - \frac{\gamma (y_{t+1} - y_t)^2}{2}, \quad y_0 \text{ given} \]  \hspace{1cm} (2.98)
Regarding the parameters,
- \( \beta \in (0, 1) \) is a discount factor
- \( \gamma > 0 \) measures the cost of adjusting the rate of output
Regarding timing, the firm observes \( p_t \) and \( y_t \) at time \( t \) when it chooses \( y_{t+1} \)
To state the firm’s optimization problem completely requires that we specify dynamics for all state variables
This includes ones like \( p_t \), which the firm cares about but does not control
We turn to this problem now

Prices and Aggregate Output  In view of (2.96), the firm’s incentive to forecast the market price translates into an incentive to forecast the level of aggregate output \( Y_t \)
Aggregate output depends on the choices of other firms
We assume that \( n \) is a large number so that the output of any single firm has a negligible effect on aggregate output
That justifies firms in treating their forecast of aggregate output as being unaffected by their own output decisions

The Firm’s Beliefs  We suppose the firm believes that market-wide output \( Y_t \) follows the law of motion
\[ Y_{t+1} = H(Y_t) \]  \hspace{1cm} (2.99)
where \( Y_0 \) is a known initial condition
The belief function \( H \) is an equilibrium object, and hence remains to be determined

Optimal Behavior Given Beliefs  For now let’s fix a particular belief \( H \) in (2.99) and investigate the firm’s response
Let \( v \) be the corresponding value function for the firm’s problem
The value function satisfies the Bellman equation
\[ v(y, Y) = \max_y \left\{ a_0 y - a_1 y Y - \frac{\gamma (y' - y)^2}{2} + \beta v(y', H(Y)) \right\} \]  \hspace{1cm} (2.100)
Let’s denote the firm’s optimal policy function by $h$, so that

$$y_{t+1} = h(y_t, Y_t)$$ \tag{2.101}$$

where

$$h(y, Y) := \arg \max_{y'} \left\{ a_0 y - a_1 y Y - \frac{\gamma (y' - y)^2}{2} + \beta v(y', H(Y)) \right\}$$ \tag{2.102}$$

Evidently $v$ and $h$ both depend on $H$

**First Order Characterization of $h$** In what follows it will be helpful to have a second characterization of $h$, based on first order conditions

The first-order necessary condition for choosing $y'$ is

$$-\gamma (y' - y) + \beta v_y(y', H(Y)) = 0$$ \tag{2.103}$$

A well-known envelope result [BS79] implies that to differentiate $v$ with respect to $y$ we can naively differentiate the right-hand side of (2.100), giving

$$v_y(y, Y) = a_0 - a_1 Y + \gamma (y' - y)$$

Substituting this equation into (2.103) gives the Euler equation

$$-\gamma (y_{t+1} - y_t) + \beta [a_0 - a_1 Y_{t+1} + \gamma (y_{t+2} - y_{t+1})] = 0$$ \tag{2.104}$$

In the process of solving its Bellman equation, the firm sets an output path that satisfies (2.104), taking (2.99) as given, and subject to

- the initial conditions for $(y_0, Y_0)$
- the terminal condition $\lim_{t \to \infty} \beta^t y_t v_y(y_t, Y_t) = 0$

This last condition is called the transversality condition, and acts as a first-order necessary condition “at infinity”

The firm’s decision rule solves the difference equation (2.104) subject to the given initial condition $y_0$ and the transversality condition

Note that solving the Bellman equation (2.100) for $v$ and then $h$ in (2.102) yields a decision rule that automatically imposes both the Euler equation (2.104) and the transversality condition

**The Actual Law of Motion for $\{Y_t\}$** As we’ve seen, a given belief translates into a particular decision rule $h$

Recalling that $Y_t = ny_t$, the actual law of motion for market-wide output is then

$$Y_{t+1} = nh(Y_t/n, Y_t)$$ \tag{2.105}$$

Thus, when firms believe that the law of motion for market-wide output is (2.99), their optimizing behavior makes the actual law of motion be (2.105)
**Definition of Rational Expectations Equilibrium**  A rational expectations equilibrium or recursive competitive equilibrium of the model with adjustment costs is a decision rule \( h \) and an aggregate law of motion \( H \) such that

1. Given belief \( H \), the map \( h \) is the firm’s optimal policy function
2. The law of motion \( H \) satisfies \( H(Y) = nh(Y/n, Y) \) for all \( Y \)

Thus, a rational expectations equilibrium equates the perceived and actual laws of motion (2.99) and (2.105)

**Fixed point characterization**  As we’ve seen, the firm’s optimum problem induces a mapping \( \Phi \) from a perceived law of motion \( H \) for market-wide output to an actual law of motion \( \Phi(H) \)

The mapping \( \Phi \) is the composition of two operations, taking a perceived law of motion into a decision rule via (2.100)–(2.102), and a decision rule into an actual law via (2.105)

The \( H \) component of a rational expectations equilibrium is a fixed point of \( \Phi \)

**Computation of the Equilibrium**

Now let’s consider the problem of computing the rational expectations equilibrium

**Misbehavior of \( \Phi \)**  Readers accustomed to dynamic programming arguments might try to address this problem by choosing some guess \( H_0 \) for the aggregate law of motion and then iterating with \( \Phi \)

Unfortunately, the mapping \( \Phi \) is not a contraction

In particular, there is no guarantee that direct iterations on \( \Phi \) converge

Fortunately, there is another method that works here

The method exploits a general connection between equilibrium and Pareto optimality expressed in the fundamental theorems of welfare economics (see, e.g, [MCWG95])

Lucas and Prescott [LP71] used this method to construct a rational expectations equilibrium

The details follow

**A Planning Problem Approach**  Our plan of attack is to match the Euler equations of the market problem with those for a single-agent planning problem

As we’ll see, this planning problem can be solved by LQ control

The optimal quantities from the planning problem are then rational expectations equilibrium quantities

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The rational expectations equilibrium price can be obtained as a shadow price in the planning problem.

For convenience, in this section we set $n = 1$.

We first compute a sum of consumer and producer surplus at time $t$

$$s(Y_t, Y_{t+1}) := \int_0^{Y_t} (a_0 - a_1 x) \, dx - \frac{\gamma(Y_{t+1} - Y_t)^2}{2} \quad (2.106)$$

The first term is the area under the demand curve, while the second is the social costs of changing output.

The planning problem is to choose a production plan $\{Y_t\}$ to maximize

$$\sum_{t=0}^{\infty} \beta^ts(Y_t, Y_{t+1})$$

subject to an initial condition for $Y_0$.

**Solution of the Planning Problem** Evaluating the integral in (2.106) yields the quadratic form $a_0 Y_t - a_1 Y_t^2/2$.

As a result, the Bellman equation for the planning problem is

$$V(Y) = \max_{Y'} \left\{ a_0 Y - \frac{a_1 Y^2}{2} - \frac{\gamma(Y' - Y)^2}{2} + \beta V(Y') \right\} \quad (2.107)$$

The associated first order condition is

$$-\gamma(Y' - Y) + \beta V'(Y') = 0 \quad (2.108)$$

Applying the same Benveniste-Scheinkman formula gives

$$V'(Y) = a_0 - a_1 Y + \gamma(Y' - Y)$$

Substituting this into equation (2.108) and rearranging leads to the Euler equation

$$\beta a_0 + \gamma Y_t - [\beta a_1 + \gamma(1 + \beta)]Y_{t+1} + \gamma \beta Y_{t+2} = 0 \quad (2.109)$$

**The Key Insight** Return to equation (2.104) and set $y_t = Y_t$ for all $t$.

(Recall that for this section we’ve set $n = 1$ to simplify the calculations)

A small amount of algebra will convince you that when $y_t = Y_t$, equations (2.109) and (2.104) are identical.

Thus, the Euler equation for the planning problem matches the second-order difference equation that we derived by

1. finding the Euler equation of the representative firm and
2. substituting into it the expression $Y_t = ny_t$ that “makes the representative firm be representative”
If it is appropriate to apply the same terminal conditions for these two difference equations, which it is, then we have verified that a solution of the planning problem also is a rational expectations equilibrium.

It follows that for this example we can compute an equilibrium by forming the optimal linear regulator problem corresponding to the Bellman equation (2.107).

The optimal policy function for the planning problem is the aggregate law of motion $H$ that the representative firm faces within a rational expectations equilibrium.

**Structure of the Law of Motion** As you are asked to show in the exercises, the fact that the planner’s problem is an LQ problem implies an optimal policy — and hence aggregate law of motion — taking the form

$$Y_{t+1} = \kappa_0 + \kappa_1 Y_t$$

(2.110)

for some parameter pair $\kappa_0, \kappa_1$.

Now that we know the aggregate law of motion is linear, we can see from the firm’s Bellman equation (2.100) that the firm’s problem can be framed as an LQ problem.

As you’re asked to show in the exercises, the LQ formulation of the firm’s problem implies a law of motion that looks as follows

$$y_{t+1} = h_0 + h_1 y_t + h_2 Y_t$$

(2.111)

Hence a rational expectations equilibrium will be defined by the parameters $(\kappa_0, \kappa_1, h_0, h_1, h_2)$ in (2.110)–(2.111).

**Exercises**

**Exercise 1** Consider the firm problem described above.

Let the firm’s belief function $H$ be as given in (2.110).

Formulate the firm’s problem as a discounted optimal linear regulator problem, being careful to describe all of the objects needed.

Use the type LQ from the QuantEcon package to solve the firm’s problem for the following parameter values:

$$a_0 = 100, a_1 = 0.05, \beta = 0.95, \gamma = 10, \kappa_0 = 95.5, \kappa_1 = 0.95$$

Express the solution of the firm’s problem in the form (2.111) and give the values for each $h_j$.

If there were $n$ identical competitive firms all behaving according to (2.111), what would (2.111) imply for the actual law of motion (2.99) for market supply?

**Exercise 2** Consider the following $\kappa_0, \kappa_1$ pairs as candidates for the aggregate law of motion component of a rational expectations equilibrium (see (2.110)).

Extending the program that you wrote for exercise 1, determine which if any satisfy the definition of a rational expectations equilibrium.

- $(94.0886298678, 0.923409232937)$
Describe an iterative algorithm that uses the program that you wrote for exercise 1 to compute a rational expectations equilibrium.

(You are not being asked actually to use the algorithm you are suggesting)

**Exercise 3**  
Recall the planner’s problem described above

1. Formulate the planner’s problem as an LQ problem
2. Solve it using the same parameter values in exercise 1
   - $a_0 = 100, a_1 = 0.05, \beta = 0.95, \gamma = 10$
3. Represent the solution in the form $Y_{t+1} = \kappa_0 + \kappa_1 Y_t$
4. Compare your answer with the results from exercise 2

**Exercise 4**  
A monopolist faces the industry demand curve (2.96) and chooses $\{Y_t\}$ to maximize $\sum_{t=0}^{\infty} \beta^t r_t$ where

$$r_t = p_t Y_t - \frac{\gamma(Y_{t+1} - Y_t)^2}{2}$$

Formulate this problem as an LQ problem.

Compute the optimal policy using the same parameters as the previous exercise.

In particular, solve for the parameters in

$$Y_{t+1} = m_0 + m_1 Y_t$$

Compare your results with the previous exercise. Comment.

**Solutions**

**Solution notebook**

**2.11 Markov Asset Pricing**
Overview

An asset is a claim on a stream of prospective payments. The spot price of an asset depends primarily on:

- the anticipated dynamics for the stream of income accruing to the owners
- the pricing model, which determines how prices react to different income streams

In this lecture, we consider some standard pricing models and dividend stream specifications. We study how prices and dividend-price ratios respond in these different scenarios. We also look at creating and pricing derivative assets by repackaging income streams.

Key tools for the lecture are:

- Formulas for predicting future values of functions of a Markov state
- A formula for predicting the discounted sum of future values of a Markov state

Pricing Models

We begin with some notation and then proceed to foundational pricing models.

In what follows, let $d_0, d_1, \ldots$ be a stream of dividends:

- A time-$t$ cum-dividend asset is a claim to the stream $d_t, d_{t+1}, \ldots$
- A time-$t$ ex-dividend asset is a claim to the stream $d_{t+1}, d_{t+2}, \ldots$

Risk Neutral Pricing

Let $\beta = 1/(1 + \rho)$ be an intertemporal discount factor. In other words, $\rho$ is the rate at which agents discount the future.

The basic risk-neutral asset pricing equation for pricing one unit of a cum-dividend asset is

$$p_t = d_t + \beta E_t[p_{t+1}]$$  \hspace{1cm} (2.112)
This is a simple “cost equals expected benefit” relationship.

Here \( \mathbb{E}_t[y] \) denotes the best forecast of \( y \), conditioned on information available at time \( t \).

In the present case this information set consists of observations of dividends up until time \( t \).

For an ex-dividend asset (buy today in exchange for the asset and dividend tomorrow), the basic risk-neutral asset pricing equation is

\[
p_t = \beta \mathbb{E}_t[d_{t+1} + p_{t+1}]
\]  
(2.113)

**Pricing Under Risk Aversion** Let’s now introduce risk aversion by supposing that all agents evaluate payoffs according to strictly concave period utility function \( u \).

In this setting Robert Lucas [Luc78] showed that under certain equilibrium conditions the price of an ex-dividend asset obeys the famous consumption-based asset pricing equation

\[
p_t = \mathbb{E}_t \left[ \beta u'(d_{t+1}) (d_{t+1} + p_{t+1}) \right]
\]  
(2.114)

Comparing (2.113) and (2.114), the difference is that \( \beta \) in (2.113) has been replaced by

\[
\beta \frac{u'(d_{t+1})}{u'(d_t)}
\]

This term is usually called the *stochastic discount factor*.

We give a full derivation of (2.114) in a *later lecture*.

For now we focus more on implications.

For the most part we will assume preferences take the form

\[
u(c) = \frac{1 - \gamma}{c^{1-\gamma}} \text{ with } \gamma > 0 \quad \text{or} \quad u(c) = \ln c
\]

**Simple Examples** What price dynamics result from these models?

The answer to this question depends on the process we specify for dividends.

Let’s look at some examples that illustrate this idea.

**Example 1: Constant dividends, risk neutral pricing** The simplest case is a constant, non-random dividend stream \( d_t = d > 0 \).

Removing the expectation from (2.112) and iterating forward gives

\[
p_t = d + \beta p_{t+1} = d + \beta (d + \beta p_{t+2}) = \cdots = d + \beta d + \beta^2 d + \cdots + \beta^{k-1} d + \beta^k p_{t+k}
\]
Unless prices explode in the future, this sequence converges to
\[ p_t = \frac{1}{1 - \beta} d_t \]
This price is the equilibrium price in the constant dividend case
The ex-dividend equilibrium price is \( p_t = (1 - \beta)^{-1} \beta d_t \)

**Example 2: Deterministic dividends, risk neutral pricing**  Consider a growing, non-random dividend process \( d_t = \lambda t d_0 \) where \( 0 < \lambda \beta < 1 \)
The cum-dividend price under risk neutral pricing is then
\[ p_t = \frac{d_t}{1 - \beta \lambda} = \frac{\lambda t d_0}{1 - \beta \lambda} \] (2.115)
(Hint: Set \( v_t = p_t / d_t \) in (2.112) and then \( v_t = v_{t+1} = v \) to solve for constant \( v \))
The ex-dividend price is \( p_t = (1 - \beta \lambda)^{-1} \beta \lambda d_t \)
If, in this example, we take \( \lambda = 1 + g \), then the ex-dividend price becomes
\[ p_t = \frac{1 + g}{\rho - g} d_t \]
This is called the Gordon formula

**Example 3: Markov growth, risk neutral pricing**  Next we consider a dividend process where the growth rate is Markovian
In particular,
\[ d_{t+1} = \lambda_{t+1} d_t \quad \text{where} \quad \mathbb{P}\{\lambda_{t+1} = s_j | \lambda_t = s_i\} = P_{ij} := P[i,j] \]
This notation means that \( \{\lambda_t\} \) is an \( n \) state Markov chain with transition matrix \( P \) and state space \( s = \{s_1, \ldots, s_n\} \)
To obtain asset prices under risk neutrality, recall that in (2.115) the price dividend ratio \( p_t / d_t \) is constant and depends on \( \lambda \)
This encourages us to guess that, in the current case, \( p_t / d_t \) is constant given \( \lambda_t \)
That is \( p_t = v(\lambda_t) d_t \) for some unknown function \( v \) on the state space
To simplify notation, let \( v_t := v(s_t) \)
For a cum-dividend stock we find that \( v_t = 1 + \beta \sum_{j=1}^{n} P_{ij} s_j v_j \)
Letting \( \mathbf{1} \) be an \( n \times 1 \) vector of ones and \( \tilde{P}_{ij} = P_{ij} s_j \), we can express this in matrix notation as
\[ v = (I - \beta \tilde{P})^{-1} \mathbf{1} \]
Here we are assuming invertibility, which requires that the growth rate of the Markov chain is not too large relative to \( \beta \)
(In particular, that the eigenvalues of $\tilde{P}$ be strictly less than $\beta^{-1}$ in modulus)

Similar reasoning yields the ex-dividend price-dividend ratio $w$, which satisfies

$$w = \beta(I - \beta \tilde{P})^{-1}Ps'$$

**Example 4: Deterministic dividends, risk averse pricing**  Our formula for pricing a cum-dividend claim to a non random stream $d_t = \lambda t d$ then becomes

$$p_t = d_t + \beta \lambda^{-\gamma} p_{t+1}$$

Guessing again that the price obeys $p_t = v d_t$ where $v$ is a constant price-dividend ratio, we have $vd_t = d_t + \beta \lambda^{-\gamma} vd_{t+1}$, or

$$v = \frac{1}{1 - \beta \lambda^{1-\gamma}}$$

If $u'(c) = 1/c$, then the preceding formula for the price-dividend ratio becomes $v = 1/(1 - \beta)$

Here the price-dividend ratio is constant and independent of the dividend growth rate $\lambda$

**Finite Markov Asset Pricing**

For the remainder of the lecture we focus on computing asset prices when

- endowments follow a finite state Markov chain
- agents are risk averse, and prices obey (2.114)

Our finite state Markov setting emulates [MP85]

In particular, we’ll assume that there is an endowment of a consumption good that follows

$$c_{t+1} = \lambda_{t+1} c_t$$  \hspace{1cm} (2.116)

Here $\lambda_t$ is governed by the $n$ state Markov chain discussed above

A *Lucas tree* is a unit claim on this endowment stream

We’ll price several distinct assets, including

- The Lucas tree itself
- A consol (a type of bond issued by the UK government in the 19th century)
- Finite and infinite horizon call options on a consol

**Pricing the Lucas tree**  Using (2.114), the definition of $u$ and (2.116) leads to

$$p_t = E_t \left[ \beta \lambda_{t+1}^{-\gamma} (c_{t+1} + p_{t+1}) \right]$$  \hspace{1cm} (2.117)

Drawing intuition from our earlier discussion on pricing with Markov growth, we guess a pricing function of the form $p_t = v(\lambda_t) c_t$ where $v$ is yet to be determined
If we substitute this guess into (2.117) and rearrange, we obtain

\[ v(\lambda_t) c_t = \mathbb{E}_t \left[ \beta \lambda \lambda_{t+1}^{1-\gamma} (c_{t+1} + c_{t+1} v(\lambda_{t+1})) \right] \]

Using (2.116) again and simplifying gives

\[ v(\lambda_t) = \mathbb{E}_t \left[ \beta \lambda^{1-\gamma} (1 + v(\lambda_{t+1})) \right] \]

As before we let \( v(s_i) = v_i \), so that \( v \) is modeled as an \( n \times 1 \) vector, and

\[ v_i = \beta \sum_{j=1}^{n} P_{ij} s_j^{1-\gamma} (1 + v_j) \quad (2.118) \]

Letting \( \tilde{P}_{ij} = P_{ij} s_j^{1-\gamma} \), we can write (2.118) as \( v = \beta \tilde{P} \mathbf{1} + \beta \tilde{P} v \)

Assuming again that the eigenvalues of \( \tilde{P} \) are strictly less than \( \beta^{-1} \) in modulus, we can solve this to yield

\[ v = \beta (I - \beta \tilde{P})^{-1} \tilde{P} \mathbf{1} \quad (2.119) \]

With log preferences, \( \gamma = 1 \) and hence \( s^{1-\gamma} = 1 \), from which we obtain

\[ v = \frac{\beta}{1 - \beta} \mathbf{1} \]

Thus, with log preferences, the price-dividend ratio for a Lucas tree is constant

**A Risk-Free Consol**  Consider the same pure exchange representative agent economy

A risk-free consol promises to pay a constant amount \( \zeta > 0 \) each period

Recycling notation, let \( p_t \) now be the price of an ex-coupon claim to the consol

An ex-coupon claim to the consol entitles the owner at the end of period \( t \) to

- \( \zeta \) in period \( t+1 \), plus
- the right to sell the claim for \( p_{t+1} \) next period

The price satisfies

\[ u'(c_t) p_t = \beta \mathbb{E}_t \left[ u'(c_{t+1}) (\zeta + p_{t+1}) \right] \]

Substituting \( u'(c) = c^{-\gamma} \) into the above equation yields

\[ c_t^{-\gamma} p_t = \beta \mathbb{E}_t \left[ c_{t+1}^{-\gamma} (\zeta + p_{t+1}) \right] = \beta c_t^{-\gamma} \mathbb{E}_t \left[ \lambda_{t+1}^{-\gamma} (\zeta + p_{t+1}) \right] \]

It follows that

\[ p_t = \beta \mathbb{E}_t \left[ \lambda_{t+1}^{-\gamma} (\zeta + p_{t+1}) \right] \quad (2.120) \]

Now guess that the price takes the form

\[ p_t = p(\lambda_t) = p_i \quad \text{when} \quad \lambda_t = s_i \]
Then (2.120) becomes

\[ p_i = \beta \sum_j P_{ij} s_j^{-\gamma} (\zeta + p_j) \]

which can be expressed as

\[ p = \beta \tilde{P} \zeta \mathbf{1} + \beta \tilde{P} p, \]

or

\[ p = \beta (I - \beta \tilde{P})^{-1} \tilde{P} \zeta \mathbf{1} \]  \hspace{1cm} (2.121)

where \( \tilde{P}_{ij} = P_{ij} s_j^{-\gamma} \)

**Pricing an Option to Purchase the Consol**  
Let’s now price options of varying maturity that give the right to purchase a consol at a price \( p_S \)

**An infinite horizon call option**  
We want to price an infinite horizon option to purchase a consol at a price \( p_S \)

The option entitles the owner at the beginning of a period either to

1. purchase the bond at price \( p_S \) now, or
2. to hold the option until next period

Thus, the owner either *exercises* the option now, or chooses *not to exercise* and wait until next period

This is termed an infinite-horizon *call option* with *strike price* \( p_S \)

The owner of the option is entitled to purchase the consol at the price \( p_S \) at the beginning of any period, after the coupon has been paid to the previous owner of the bond

The economy is identical with the one above

Let \( w(\lambda_t, p_S) \) be the value of the option when the time \( t \) growth state is known to be \( \lambda_t \) but before the owner has decided whether or not to exercise the option at time \( t \) (i.e., today)

Recalling that \( p(\lambda_t) \) is the value of the consol when the initial growth state is \( \lambda_t \), the value of the option satisfies

\[ w(\lambda_t, p_S) = \max \left\{ \beta \mathbb{E}_t \left[ \frac{u'(c_{t+1})}{u'(c_t)} w(\lambda_{t+1}, p_S), \ p(\lambda_t) - p_S \right] \right\} \]

The first term on the right is the value of waiting, while the second is the value of exercising

We can also write this as

\[ w(s_t, p_S) = \max \left\{ \beta \sum_{j=1}^n P_{ij} s_j^{-\gamma} w(s_j, p_S), \ p(s_t) - p_S \right\} \]  \hspace{1cm} (2.122)

Letting \( \tilde{P}_{ij} = P_{ij} s_j^{-\gamma} \) and \( w_t = w(s_t, p_S) \), we can express (2.122) as the nonlinear vector equation

\[ w = \max \{ \beta \tilde{P} w, \ p - p_S \mathbf{1} \} \]  \hspace{1cm} (2.123)

To solve (2.123), form the operator \( T \) mapping vector \( w \) into vector \( Tw \) via

\[ Tw = \max \{ \beta \tilde{P} w, \ p - p_S \mathbf{1} \} \]

Start at some initial \( w \) and iterate to convergence with \( T \)
Finite-horizon options Finite horizon options obey functional equations closely related to (2.122)

A $k$ period option expires after $k$ periods

At time $t$, a $k$ period option gives the owner the right to exercise the option to purchase the risk-free consol at the strike price $p_S$ at $t, t+1, \ldots, t+k-1$

The option expires at time $t+k$

Thus, for $k = 1, 2, \ldots$, let $w(s_i, k)$ be the value of a $k$-period option

It obeys

$$w(s_i, k) = \max \left\{ \beta \sum_{j=1}^{n} P_{ij} s_j^{-\gamma} w(s_j, k-1), p(s_i) - p_S \right\}$$

where $w(s_i, 0) = 0$ for all $i$

We can express the preceding as the sequence of nonlinear vector equations

$$w_i^{(k)} = \max \left\{ \beta \sum_{j=1}^{n} \hat{P}_{ij} w_j^{(k-1)}, p_i - p_S \right\}, \quad k = 1, 2, \ldots \quad \text{with } w^0 = 0$$

Other Prices Let’s look at the pricing of several other assets

The one-period risk-free interest rate For this economy, the stochastic discount factor is

$$m_{t+1} = \beta \frac{c_{t+1}}{c_t} = \beta \lambda_{t+1}^{-\gamma}$$

It follows that the reciprocal $R_t^{-1}$ of the gross risk-free interest rate $R_t$ is

$$\mathbb{E}_t m_{t+1} = \beta \sum_{j=1}^{n} P_{ij} s_j^{-\gamma}$$

or

$$m_1 = \beta P s^{-\gamma}$$

where the $i$-th element of $m_1$ is the reciprocal of the one-period gross risk-free interest rate when $\lambda_t = s_i$

$j$ period risk-free interest rates Let $m_j$ be an $n \times 1$ vector whose $i$ th component is the reciprocal of the $j$ -period gross risk-free interest rate when $\lambda_t = s_i$

Again, let $\hat{P}_{ij} = P_{ij} s_j^{-\gamma}$

Then $m_1 = \beta \hat{P}$, and $m_{j+1} = \hat{P} m_j$ for $j \geq 1$
2.12. THE PERMANENT INCOME MODEL

Implementation

The type AssetPrices from the QuantEcon package provides methods for computing some of the prices described above.

We print the code here for convenience.

Exercises

Exercise 1 Compute the price of the Lucas tree in an economy with the following primitives

\[ n = 5 \]
\[ P = 0.0125 \times \text{ones}(n, n) \]
\[ P +\text{diagm}(0.95 - 0.0125 \times \text{ones}(5)) \]
\[ s = [1.05, 1.025, 1.0, 0.975, 0.95] \]
\[ \text{gamm} = 2.0 \]
\[ \text{bet} = 0.94 \]
\[ \text{zet} = 1.0 \]

Using the same set of primitives, compute the price of the risk-free console when \( \zeta = 1 \)

Do the same for the call option on the console when \( p_S = 150.0 \)

Compute the value of the option at dates \( T = [10, 20, 30] \)

Solutions

Solution notebook

2.12 The Permanent Income Model

Contents

- The Permanent Income Model
  - Overview
  - The Savings Problem
  - Alternative Representations
  - Two Classic Examples
  - Further Reading
  - Appendix: The Euler Equation

Overview

This lecture describes a rational expectations version of the famous permanent income model of Friedman [Fri56]
Hall cast Friedman’s model within a linear-quadratic setting [Hal78].

Like Hall, we formulate an infinite-horizon linear-quadratic savings problem. We use the model as a vehicle for illustrating

- alternative formulations of the state of a dynamic system
- the idea of cointegration
- impulse response functions
- the idea that changes in consumption are useful as predictors of movements in income

**The Savings Problem**

In this section we state and solve the savings and consumption problem faced by the consumer.

**Preliminaries** The discussion below requires a casual familiarity with martingales. A discrete time martingale is a stochastic process (i.e., a sequence of random variables) \( \{X_t\} \) with finite mean and satisfying

\[
E_t[X_{t+1}] = X_t, \quad t = 0, 1, 2, \ldots
\]

Here \( E_t := E [\cdot | F_t] \) is a mathematical expectation conditional on the time \( t \) information set \( F_t \).

The latter is just a collection of random variables that the modeler declares to be visible at \( t \)

- When not explicitly defined, it is usually understood that \( F_t = \{X_t, X_{t-1}, \ldots, X_0\} \)

Martingales have the feature that the history of past outcomes provides no predictive power for changes between current and future outcomes.

For example, the current wealth of a gambler engaged in a “fair game” has this property. One common class of martingales is the family of random walks.

A random walk is a stochastic process \( \{X_t\} \) that satisfies

\[
X_{t+1} = X_t + w_{t+1}
\]

for some iid zero mean innovation sequence \( \{w_t\} \).

Evidently \( X_t \) can also be expressed as

\[
X_t = \sum_{j=1}^{t} w_j + X_0
\]

Not every martingale arises as a random walk (see, for example, Wald’s martingale).
The Decision Problem  A consumer has preferences over consumption streams that are ordered by the utility functional

$$E_0 \left[ \sum_{t=0}^{\infty} \beta^t u(c_t) \right]$$

(2.124)

where

- $E_t$ is the mathematical expectation conditioned on the consumer’s time $t$ information
- $c_t$ is time $t$ consumption
- $u$ is a strictly concave one-period utility function
- $\beta \in (0, 1)$ is a discount factor

The consumer maximizes (2.124) by choosing a consumption, borrowing plan $\{c_t, b_{t+1}\}_{t=0}^{\infty}$ subject to the sequence of budget constraints

$$b_{t+1} = (1 + r)(c_t + b_t - y_t) \quad t \geq 0$$

(2.125)

Here

- $y_t$ is an exogenous endowment process
- $r > 0$ is the risk-free interest rate
- $b_t$ is one-period risk-free debt maturing at $t$
- $b_0$ is a given initial condition

Assumptions  For the remainder of this lecture, we follow Friedman and Hall in assuming that $(1 + r)^{-1} = \beta$

Regarding the endowment process, we assume it has the state-space representation

$$x_{t+1} = Ax_t + Cw_{t+1}$$

$$y_t = Ux_t$$

(2.126) (2.127)

where

- $\{w_t\}$ is an iid vector process with $E w_t = 0$ and $E w_t w'_t = I$
- the spectral radius of $A$ satisfies $\rho(A) < 1/\beta$
- $U$ is a selection vector that pins down $y_t$ as a particular linear combination of the elements of $x_t$.

The restriction on $\rho(A)$ prevents income from growing so fast that certain sums become infinite

We also impose the no Ponzi scheme condition

$$E_0 \left[ \sum_{t=0}^{\infty} \beta^t b_t^2 \right] < \infty$$

(2.128)

This condition rules out an always-borrow scheme that would allow the household to enjoy unbounded or bliss consumption forever
Regarding preferences, we assume the quadratic utility function
\[ u(c_t) = -(c_t - \bar{c})^2 \]
where \( \bar{c} \) is a bliss level of consumption
(Along with this quadratic utility specification, we allow consumption to be negative)

**First Order Conditions**  First-order conditions for maximizing (2.124) subject to (2.125) are
\[ \mathbb{E}_t[u'(c_{t+1})] = u'(c_t), \quad t = 0, 1, \ldots \] (2.129)
These equations are also known as the *Euler equations* for the model
If you’re not sure where they come from, you can find a proof sketch in the *appendix*
With our quadratic preference specification, (2.129) has the striking implication that consumption follows a martingale:
\[ \mathbb{E}_t[c_{t+1}] = c_t \] (2.130)
(In fact quadratic preferences are *necessary* for this conclusion 7)
One way to interpret (2.130) is that consumption will only change when “new information” about permanent income is revealed
These ideas will be clarified below

**The Optimal Decision Rule**  The state vector confronting the household at \( t \) is \([b_t \ x_t]\)
Here
- \( x_t \) is an exogenous component, unaffected by household behavior
- \( b_t \) is an endogenous component (since it depends on the decision rule)
Note that \( x_t \) contains all variables useful for forecasting the household’s future endowment
Now let’s deduce the optimal decision rule 8

---

**Note:** One way to solve the consumer’s problem is to apply *dynamic programming* as in *this lecture*. We do this later. But first we use an alternative approach that is revealing and shows the work that dynamic programming does for us automatically

We want to solve the system of difference equations formed by (2.125) and (2.130) subject to the boundary condition (2.128)
To accomplish this, observe first that (2.128) implies \( \lim_{t \to \infty} \beta^t b_{t+1} = 0 \)

---

7 A linear marginal utility is essential for deriving (2.130) from (2.129). Suppose instead that we had imposed the following more standard assumptions on the utility function: \( u'(c) > 0, u''(c) < 0, u'''(c) > 0 \) and required that \( c \geq 0 \). The Euler equation remains (2.129). But the fact that \( u''' < 0 \) implies via Jensen’s inequality that \( \mathbb{E}_t[u'(c_{t+1})] > u'\left(\mathbb{E}_t[c_{t+1}]\right) \). This inequality together with (2.129) implies that \( \mathbb{E}_t[c_{t+1}] > c_t \) (consumption is said to be a ‘submartingale’), so that consumption stochastically diverges to +\( \infty \). The consumer’s savings also diverge to +\( \infty \).
8 An optimal decision rule is a map from current state into current actions—in this case, consumption
2.12. THE PERMANENT INCOME MODEL

Using this restriction on the debt path and solving (2.125) forward yields

\[ b_t = \sum_{j=0}^{\infty} \beta^j (y_{t+j} - c_{t+j}) \]  

(2.131)

Take conditional expectations on both sides of (2.131) and use the law of iterated expectations to deduce

\[ b_t = \sum_{j=0}^{\infty} \beta^j \mathbb{E}_t [y_{t+j}] - \frac{c_t}{1 - \beta} \]  

(2.132)

Expressed in terms of \( c_t \) we get

\[ c_t = (1 - \beta) \left[ \sum_{j=0}^{\infty} \beta^j \mathbb{E}_t [y_{t+j}] - b_t \right] \]  

(2.133)

If we define the net rate of interest \( r \) by \( \beta = \frac{1}{1+r} \), we can also express this equation as

\[ c_t = \frac{r}{1+r} \left[ \sum_{j=0}^{\infty} \beta^j \mathbb{E}_t [y_{t+j}] - b_t \right] \]

These last two equations assert that consumption equals economic income

- financial wealth equals \( b_t \)
- non-financial wealth equals \( \sum_{j=0}^{\infty} \beta^j \mathbb{E}_t [y_{t+j}] \)
- A marginal propensity to consume out of wealth equals the interest factor \( \frac{r}{1+r} \)
- economic income equals
  - a constant marginal propensity to consume times the sum of nonfinancial wealth and financial wealth
  - the amount the household can consume while leaving its wealth intact

A State-Space Representation The preceding results provide a decision rule and hence the dynamics of both state and control variables

First note that equation (2.133) represents \( c_t \) as a function of the state \([b_t \ x_t]\) confronting the household

If the last statement isn’t clear, recall that \( \mathbb{E}_t [y_{t+j}] \) can be expressed as a function of \( x_t \), since the latter contains all information useful for forecasting the household’s endowment process

In fact, from this discussion we see that

\[ \sum_{j=0}^{\infty} \beta^j \mathbb{E}_t [y_{t+j}] = \mathbb{E}_t \left[ \sum_{j=0}^{\infty} \beta^j y_{t+j} \right] = U(I - \beta A)^{-1} x_t \]
Using this expression, we can obtain a linear state-space system governing consumption, debt and income:

\[
x_{t+1} = Ax_t + Cw_{t+1} \\
b_{t+1} = b_t + U[(I - \beta A)^{-1}(A - I)]x_t \\
y_t = Ux_t \\
c_t = (1 - \beta)[U(I - \beta A)^{-1}x_t - b_t]
\]

(2.134)  (2.135)  (2.136)  (2.137)

**A Simple Example with iid Income**  To gain some preliminary intuition on the implications of (2.134), let’s look at a highly stylized example where income is just iid (Later examples will investigate more realistic income streams)

In particular, let \( \{w_t\}_{t=1}^{\infty} \) be iid and scalar standard normal, and let

\[
x_t = \begin{bmatrix} x_t^1 \\ 1 \end{bmatrix}, \quad A = \begin{bmatrix} 0 & 0 \\ 0 & 1 \end{bmatrix}, \quad U = \begin{bmatrix} 1 & \mu \end{bmatrix}, \quad C = \begin{bmatrix} \sigma \\ 0 \end{bmatrix}
\]

Finally, let \( b_0 = x_0^1 = 0 \)

Under these assumptions we have \( y_t = \mu + \sigma w_t \sim N(\mu, \sigma^2) \)

Further, if you work through the state space representation, you will see that

\[
b_t = -\sigma \sum_{j=1}^{t-1} w_j \\
c_t = \mu + (1 - \beta)\sigma \sum_{j=1}^{t} w_j
\]

Thus income is iid and debt and consumption are both Gaussian random walks

Defining assets as \(-b_t\), we see that assets are just the cumulative sum of unanticipated income prior to the present date

The next figure shows a typical realization with \( r = 0.05, \mu = 1 \) and \( \sigma = 0.15 \)

Observe that consumption is considerably smoother than income

The figure below shows the consumption paths of 250 consumers with independent income streams

The code for these figures can be found in perm_inc_figs.jl

**Alternative Representations**

In this section we shed more light on the evolution of savings, debt and consumption by representing their dynamics in several different ways
2.12. THE PERMANENT INCOME MODEL
Hall’s Representation  Hall [Hal78] suggests a sharp way to summarize the implications of LQ permanent income theory.

First, to represent the solution for $b_t$, shift (2.133) forward one period and eliminate $b_{t+1}$ by using (2.125) to obtain

$$c_{t+1} = (1 - \beta) \sum_{j=0}^{\infty} \beta^j \mathbb{E}_{t+1}[y_{t+j+1}] - (1 - \beta) \left[ \beta^{-1}(c_t + b_t - y_t) \right]$$

If we add and subtract $\beta^{-1}(1 - \beta) \sum_{j=0}^{\infty} \beta^j \mathbb{E}_t[y_{t+j}]$ from the right side of the preceding equation and rearrange, we obtain

$$c_{t+1} - c_t = (1 - \beta) \sum_{j=0}^{\infty} \beta^j \left\{ \mathbb{E}_{t+1}[y_{t+j+1}] - \mathbb{E}_t[y_{t+j+1}] \right\} \tag{2.138}$$

The right side is the time $t+1$ innovation to the expected present value of the endowment process $\{y_t\}$.

We can represent the optimal decision rule for $c_t, b_{t+1}$ in the form of (2.138) and (2.132), which is repeated here:

$$b_t = \sum_{j=0}^{\infty} \beta^j \mathbb{E}_t[y_{t+j}] - \frac{1}{1 - \beta} c_t \tag{2.139}$$

Equation (2.139) asserts that the household’s debt due at $t$ equals the expected present value of its endowment minus the expected present value of its consumption stream.

A high debt thus indicates a large expected present value of surpluses $y_t - c_t$.

Recalling again our discussion on forecasting geometric sums, we have

$$\mathbb{E}_t \sum_{j=0}^{\infty} \beta^j y_{t+j} = U(I - \beta A)^{-1} x_t$$

$$\mathbb{E}_{t+1} \sum_{j=0}^{\infty} \beta^j y_{t+j+1} = U(I - \beta A)^{-1} x_{t+1}$$

$$\mathbb{E}_t \sum_{j=0}^{\infty} \beta^j y_{t+j+1} = U(I - \beta A)^{-1} A x_t$$

Using these formulas together with (2.126) and substituting into (2.138) and (2.139) gives the following representation for the consumer’s optimum decision rule:

$$c_{t+1} = c_t + (1 - \beta) U(I - \beta A)^{-1} C w_{t+1} \tag{2.140}$$

$$b_t = U(I - \beta A)^{-1} x_t - \frac{1}{1 - \beta} c_t \tag{2.141}$$

$$y_t = U x_t \tag{2.142}$$

$$x_{t+1} = A x_t + C w_{t+1} \tag{2.143}$$

Representation (2.140) makes clear that

- The state can be taken as $(c_t, x_t)$
2.12. THE PERMANENT INCOME MODEL

- The endogenous part is $c_t$ and the exogenous part is $x_t$
- Debt $b_t$ has disappeared as a component of the state because it is encoded in $c_t$

- Consumption is a random walk with innovation $(1 - \beta)U(I - \beta A)^{-1}Cw_{t+1}$
- This is a more explicit representation of the martingale result in (2.130)

Cointegration  Representation (2.140) reveals that the joint process $\{c_t, b_t\}$ possesses the property that Engle and Granger [EG87] called cointegration.

Cointegration is a tool that allows us to apply powerful results from the theory of stationary processes to (certain transformations of) nonstationary models.

To clarify cointegration in the present context, suppose that $x_t$ is asymptotically stationary.$^9$

Despite this, both $c_t$ and $b_t$ will be non-stationary because they have unit roots (see (2.134) for $b_t$).

Nevertheless, there is a linear combination of $c_t, b_t$ that is asymptotically stationary.

In particular, from the second equality in (2.140) we have

$$(1 - \beta)b_t + c_t = (1 - \beta)U(I - \beta A)^{-1}x_t$$

(2.144)

Hence the linear combination $(1 - \beta)b_t + c_t$ is asymptotically stationary.

Accordingly, Granger and Engle would call $[(1 - \beta) \ 1]$ a cointegrating vector for the state.

When applied to the nonstationary vector process $[b_t \ c_t]'$, it yields a process that is asymptotically stationary.

Equation (2.144) can be arranged to take the form

$$(1 - \beta)b_t + c_t = (1 - \beta)\mathbb{E}_t \sum_{j=0}^{\infty} \beta^jy_{t+j},$$

(2.145)

Equation (2.145) asserts that the cointegrating residual on the left side equals the conditional expectation of the geometric sum of future incomes on the right.$^{10}$

Cross-Sectional Implications  Consider again (2.140), this time in light of our discussion of distribution dynamics in the lecture on linear systems.

The dynamics of $c_t$ are given by

$$c_{t+1} = c_t + (1 - \beta)U(I - \beta A)^{-1}Cw_{t+1}$$

(2.146)

or

$$c_t = c_0 + \sum_{j=1}^{t} \hat{w}_j \quad \text{for} \quad \hat{w}_{t+1} := (1 - \beta)U(I - \beta A)^{-1}Cw_{t+1}$$

The unit root affecting $c_t$ causes the time $t$ variance of $c_t$ to grow linearly with $t$.

$^9$ This would be the case if, for example, the spectral radius of $A$ is strictly less than one.

In particular, since \( \{ \hat{w}_t \} \) is iid, we have
\[
\text{Var}[c_t] = \text{Var}[c_0] + t \hat{\sigma}^2 \tag{2.147}
\]
when
\[
\hat{\sigma}^2 := (1 - \beta)^2 U(I - \beta A)^{-1}CC'(I - \beta A')^{-1}U'
\]
Assuming that \( \hat{\sigma} > 0 \), this means that \( \{ c_t \} \) has no asymptotic distribution.

Let’s consider what this means for a cross-section of ex ante identical households born at time 0.
Let the distribution of \( c_0 \) represent the cross-section of initial consumption values.
Equation (2.147) tells us that the distribution of \( c_t \) spreads out over time at a rate proportional to \( t \).
A number of different studies have investigated this prediction (see, e.g., [DP94], [STY04]).

**Impulse Response Functions**  
Impulse response functions measure the change in a dynamic system subject to a given impulse (i.e., temporary shock).  
The impulse response function of \( \{ c_t \} \) to the innovation \( \{ w_t \} \) is a box.  
In particular, the response of \( c_{t+j} \) to a unit increase in the innovation \( w_{t+1} \) is \( (1 - \beta) U(I - \beta A)^{-1}C \) for all \( j \geq 1 \).

**Moving Average Representation**  
It’s useful to express the innovation to the expected present value of the endowment process in terms of a moving average representation for income \( y_t \).

The endowment process defined by (2.126) has the moving average representation
\[
y_{t+1} = d(L)w_{t+1} \tag{2.148}
\]
where
- \( d(L) = \sum_{j=0}^{\infty} d_j L^j \) for some sequence \( d_j \), where \( L \) is the lag operator \(^{11}\)
- at time \( t \), the household has an information set \(^{12}\) \( w^t = [w_t, w_{t-1}, \ldots] \)

Notice that
\[
y_{t+j} - \mathbb{E}_t[y_{t+j}] = d_0 w_{t+j} + d_1 w_{t+j-1} + \cdots + d_{j-1} w_{t+1}
\]
It follows that
\[
\mathbb{E}_{t+1}[y_{t+j}] - \mathbb{E}_t[y_{t+j}] = d_{j-1} w_{t+1} \tag{2.149}
\]
Using (2.149) in (2.138) gives
\[
c_{t+1} - c_t = (1 - \beta) d(\beta) w_{t+1} \tag{2.150}
\]
The object \( d(\beta) \) is the *present value of the moving average coefficients* in the representation for the endowment process \( y_t \).

\(^{11}\) Representation (2.126) implies that \( d(L) = U(I - AL)^{-1}C \).

\(^{12}\) A moving average representation for a process \( y_t \) is said to be *fundamental* if the linear space spanned by \( y^t \) is equal to the linear space spanned by \( w^t \). A time-invariant innovations representation, attained via the Kalman filter, is by construction fundamental.
Two Classic Examples

We illustrate some of the preceding ideas with the following two examples. In both examples, the endowment follows the process $y_t = x_{1t} + x_{2t}$ where

$$
\begin{bmatrix}
x_{1t+1} \\
x_{2t+1}
\end{bmatrix}
= 
\begin{bmatrix}
1 & 0 \\
0 & 0
\end{bmatrix}
\begin{bmatrix}
x_{1t} \\
x_{2t}
\end{bmatrix}
+ 
\begin{bmatrix}
\sigma_1 & 0 \\
0 & \sigma_2
\end{bmatrix}
\begin{bmatrix}
w_{1t+1} \\
w_{2t+1}
\end{bmatrix}
$$

Here
- $w_{t+1}$ is an iid $2 \times 1$ process distributed as $N(0, I)$
- $x_{1t}$ is a permanent component of $y_t$
- $x_{2t}$ is a purely transitory component

**Example 1** Assume as before that the consumer observes the state $x_t$ at time $t$

In view of (2.140) we have

$$
c_{t+1} - c_t = \sigma_1 w_{1t+1} + (1 - \beta)\sigma_2 w_{2t+1}
$$

(2.151)

Formula (2.151) shows how an increment $\sigma_1 w_{1t+1}$ to the permanent component of income $x_{1t+1}$ leads to
- a permanent one-for-one increase in consumption and
- no increase in savings $-b_{t+1}$

But the purely transitory component of income $\sigma_2 w_{2t+1}$ leads to a permanent increment in consumption by a fraction $1 - \beta$ of transitory income.

The remaining fraction $\beta$ is saved, leading to a permanent increment in $-b_{t+1}$

Application of the formula for debt in (2.134) to this example shows that

$$
b_{t+1} - b_t = -x_{2t} = -\sigma_2 w_{2t}
$$

(2.152)

This confirms that none of $\sigma_1 w_{1t}$ is saved, while all of $\sigma_2 w_{2t}$ is saved.

The next figure illustrates these very different reactions to transitory and permanent income shocks using impulse-response functions.

The code for generating this figure is in file examples/perm_inf.jl from the main repository, as shown below

```julia
#=
@author: Spencer Lyon
@date: 07/09/2014
#

using PyPlot
```
const r = 0.05
const beta = 1.0 / (1.0 + r)
const T = 20  # Time horizon
const S = 5   # Impulse date
const sigma1 = 0.15
const sigma2 = 0.15

function time_path(permanent=false)
    w1 = zeros(T+1)
    w2 = zeros(T+1)
    b = zeros(T+1)
    c = zeros(T+1)

    if permanent == false
        w2[S+2] = 1.0
    else
        w1[S+2] = 1.0
    end

    for t=2:T
        b[t+1] = b[t] - sigma2 * w2[t]
        c[t+1] = c[t] + sigma1 * w1[t+1] + (1 - beta) * sigma2 * w2[t+1]
    end

    return b, c
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Example 2  Assume now that at time $t$ the consumer observes $y_t$, and its history up to $t$, but not $x_t$

Under this assumption, it is appropriate to use an innovation representation to form $A, C, U$ in (2.140)

The discussion in sections 2.9.1 and 2.11.3 of [LS12] shows that the pertinent state space representation for $y_t$ is

$$
\begin{bmatrix}
    y_{t+1} \\
    a_{t+1}
\end{bmatrix} = \begin{bmatrix}
    1 & -(1-K) \\
    0 & 0
\end{bmatrix} \begin{bmatrix}
    y_t \\
    a_t
\end{bmatrix} + \begin{bmatrix}
    1 \\
    1
\end{bmatrix} a_{t+1}
$$

$$
y_t = \begin{bmatrix}
    1 & 0
\end{bmatrix} \begin{bmatrix}
    y_t \\
    a_t
\end{bmatrix}
$$

where

- $K$ := the stationary Kalman gain
- $a_t := y_t - E[y_t | y_{t-1}, \ldots, y_0]$
2.12. THE PERMANENT INCOME MODEL

In the same discussion in [LS12] it is shown that \( K \in [0, 1] \) and that \( K \) increases as \( \sigma_1/\sigma_2 \) does.

In other words, as the ratio of the standard deviation of the permanent shock to that of the transitory shock increases

Applying formulas (2.140) implies

\[
ct+1 - ct = [1 - \beta(1 - K)]at+1
\]  
(2.153)

where the endowment process can now be represented in terms of the univariate innovation to \( y_t \) as

\[
y_{t+1} - y_t = at+1 - (1 - K)at
\]  
(2.154)

Equation (2.154) indicates that the consumer regards

- fraction \( K \) of an innovation \( at+1 \) to \( y_{t+1} \) as permanent
- fraction \( 1 - K \) as purely transitory

The consumer permanently increases his consumption by the full amount of his estimate of the permanent part of \( at+1 \), but by only \( (1 - \beta) \) times his estimate of the purely transitory part of \( at+1 \).

Therefore, in total he permanently increments his consumption by a fraction \( K + (1 - \beta)(1 - K) = 1 - \beta(1 - K) \) of \( at+1 \).

He saves the remaining fraction \( \beta(1 - K) \).

According to equation (2.154), the first difference of income is a first-order moving average.

Equation (2.153) asserts that the first difference of consumption is iid.

Application of formula to this example shows that

\[
b_{t+1} - b_t = (K - 1)at
\]  
(2.155)

This indicates how the fraction \( K \) of the innovation to \( y_t \) that is regarded as permanent influences the fraction of the innovation that is saved.

**Further Reading**

The model described above significantly changed how economists think about consumption.

At the same time, it’s generally recognized that Hall’s version of the permanent income hypothesis fails to capture all aspects of the consumption/savings data.

For example, liquidity constraints and buffer stock savings appear to be important.

Further discussion can be found in, e.g., [HM82], [Par99], [Dea91], [Car01].

**Appendix: The Euler Equation**

Where does the first order condition (2.129) come from?

Here we’ll give a proof for the two period case, which is representative of the general argument.
The finite horizon equivalent of the no-Ponzi condition is that the agent cannot end her life in debt, so $b_2 = 0$.

From the budget constraint (2.125) we then have

$$c_0 = \frac{b_1}{1 + r} - b_0 + y_0 \quad \text{and} \quad c_1 = y_1 - b_1$$

Here $b_0$ and $y_0$ are given constants.

Substituting these constraints into our two period objective $u(c_0) + \beta \mathbb{E}_0 [u(c_1)]$ gives

$$\max_{b_1} \left\{ u \left( \frac{b_1}{R} - b_0 + y_0 \right) + \beta \mathbb{E}_0 [u(y_1 - b_1)] \right\}$$

You will be able to verify that the first order condition is

$$u'(c_0) = \beta R \mathbb{E}_0 [u'(c_1)]$$

Using $\beta R = 1$ gives (2.129) in the two period case.

The proof for the general case is not dissimilar.
ADVANCED APPLICATIONS

This advanced section of the course contains more complex applications, and can be read selectively, according to your interests.

3.1 Continuous State Markov Chains

Overview

In a previous lecture we learned about finite Markov chains, a relatively elementary class of stochastic dynamic models.

The present lecture extends this analysis to continuous (i.e., uncountable) state Markov chains.

Most stochastic dynamic models studied by economists either fit directly into this class or can be represented as continuous state Markov chains after minor modifications.

In this lecture, our focus will be on continuous Markov models that

- evolve in discrete time
- are often nonlinear

The fact that we accommodate nonlinear models here is significant, because linear stochastic models have their own highly developed tool set, as we’ll see later on.
The question that interests us most is: Given a particular stochastic dynamic model, how will the state of the system evolve over time?

In particular,

- What happens to the distribution of the state variables?
- Is there anything we can say about the “average behavior” of these variables?
- Is there a notion of “steady state” or “long run equilibrium” that’s applicable to the model?
  - If so, how can we compute it?

Answering these questions will lead us to revisit many of the topics that occupied us in the finite state case, such as simulation, distribution dynamics, stability, ergodicity, etc.

**Note:** For some people, the term “Markov chain” always refers to a process with a finite or discrete state space. We follow the mainstream mathematical literature (e.g., [MT09]) in using the term to refer to any discrete time Markov process

---

**The Density Case**

You are probably aware that some distributions can be represented by densities and some cannot (For example, distributions on the real numbers \( \mathbb{R} \) that put positive probability on individual points have no density representation)

We are going to start our analysis by looking at Markov chains where the one step transition probabilities have density representations

The benefit is that the density case offers a very direct parallel to the finite case in terms of notation and intuition

Once we’ve built some intuition we’ll cover the general case

**Definitions and Basic Properties** In our lecture on finite Markov chains, we studied discrete time Markov chains that evolve on a finite state space \( S \)

In this setting, the dynamics of the model are described by a stochastic matrix — a nonnegative square matrix \( P = P[i, j] \) such that each row \( P[i, \cdot] \) sums to one

The interpretation of \( P \) is that \( P[i, j] \) represents the probability of transitioning from state \( i \) to state \( j \) in one unit of time

In symbols,

\[
\mathbb{P}\{X_{t+1} = j \mid X_t = i\} = P[i, j]
\]

Equivalently,

- \( P \) can be thought of as a family of distributions \( P[i, \cdot] \), one for each \( i \in S \)
- \( P[i, \cdot] \) is the distribution of \( X_{t+1} \) given \( X_t = i \)
As you probably recall, when using Julia arrays, $P[i, \cdot]$ is expressed as $P[i, :]$.

In this section, we’ll allow $S$ to be a subset of $\mathbb{R}$, such as

- $\mathbb{R}$ itself
- the positive reals $(0, \infty)$
- a bounded interval $(a, b)$

The family of discrete distributions $P[i, \cdot]$ will be replaced by a family of densities $p(x, \cdot)$, one for each $x \in S$

Analogous to the finite state case, $p(x, \cdot)$ is to be understood as the distribution (density) of $X_{t+1}$ given $X_t = x$

More formally, a stochastic kernel on $S$ is a function $p: S \times S \to \mathbb{R}$ with the property that

1. $p(x, y) \geq 0$ for all $x, y \in S$
2. $\int p(x, y)dy = 1$ for all $x \in S$

(Integrals are over the whole space unless otherwise specified)

For example, let $S = \mathbb{R}$ and consider the particular stochastic kernel $p_w$ defined by

$$p_w(x, y) := \frac{1}{\sqrt{2\pi}} \exp\left\{-\frac{(y-x)^2}{2}\right\}$$

What kind of model does $p_w$ represent?

The answer is, the (normally distributed) random walk

$$X_{t+1} = X_t + \xi_{t+1} \quad \text{where} \quad \{\xi_t\} \overset{\text{iid}}{\sim} \mathcal{N}(0, 1)$$

To see this, let’s find the stochastic kernel $p$ corresponding to (3.2)

Recall that $p(x, \cdot)$ represents the distribution of $X_{t+1}$ given $X_t = x$

Letting $X_t = x$ in (3.2) and considering the distribution of $X_{t+1}$, we see that $p(x, \cdot) = \mathcal{N}(x, 1)$

In other words, $p$ is exactly $p_w$, as defined in (3.1)

**Connection to Stochastic Difference Equations**  In the previous section, we made the connection between stochastic difference equation (3.2) and stochastic kernel (3.1)

In economics and time series analysis we meet stochastic difference equations of all different shapes and sizes

It will be useful for us if we have some systematic methods for converting stochastic difference equations into stochastic kernels

To this end, consider the generic (scalar) stochastic difference equation given by

$$X_{t+1} = \mu(X_t) + \sigma(X_t) \xi_{t+1}$$

Here we assume that
\{\xi_t\} \overset{\text{IID}}{\sim} \phi, \text{ where } \phi \text{ is a given density on } \mathbb{R} \\
\mu \text{ and } \sigma \text{ are given functions on } S, \text{ with } \sigma(x) > 0 \text{ for all } x \\

**Example 1:** The random walk (3.2) is a special case of (3.3), with \(\mu(x) = x\) and \(\sigma(x) = 1\)

**Example 2:** Consider the ARCH model 
\[ X_{t+1} = \alpha X_t + \sigma \xi_{t+1}, \quad \sigma_t^2 = \beta + \gamma X_t^2, \quad \beta, \gamma > 0 \]
Alternatively, we can write the model as 
\[ X_{t+1} = \alpha X_t + (\beta + \gamma X_t^2)^{1/2} \xi_{t+1} \]  
(3.4)
This is a special case of (3.3) with \(\mu(x) = \alpha x\) and \(\sigma(x) = (\beta + \gamma x^2)^{1/2}\)

**Example 3:** With stochastic production and a constant savings rate, the one-sector neoclassical growth model leads to a law of motion for capital per worker such as 
\[ k_{t+1} = sA_{t+1}f(k_t) + (1 - \delta)k_t \]  
(3.5)
Here 
- \(s\) is the rate of savings 
- \(A_{t+1}\) is a production shock 
  - The \((t + 1)\) subscript indicates that \(A_{t+1}\) is not visible at time \(t\) 
- \(\delta\) is a depreciation rate 
- \(f: \mathbb{R}_+ \to \mathbb{R}_+\) is a production function satisfying \(f(k) > 0\) whenever \(k > 0\) 

(The fixed savings rate can be rationalized as the optimal policy for a particular set of technologies and preferences (see [LS12], section 3.1.2), although we omit the details here)

Equation (3.5) is a special case of (3.3) with \(\mu(x) = (1 - \delta)x\) and \(\sigma(x) = sf(x)\)

Now let’s obtain the stochastic kernel corresponding to the generic model (3.3)
To find it, note first that if \(U\) is a random variable with density \(f_U\), and \(V = a + bU\) for some constants \(a, b\) with \(b > 0\), then the density of \(V\) is given by 
\[ f_V(v) = \frac{1}{b} f_U \left( \frac{v - a}{b} \right) \]  
(3.6)
(The proof is below. For a multidimensional version see EDTC, theorem 8.1.3)
 Taking (3.6) as given for the moment, we can obtain the stochastic kernel \(p\) for (3.3) by recalling that \(p(x, \cdot)\) is the conditional density of \(X_{t+1}\) given \(X_t = x\)
In the present case, this is equivalent to stating that \(p(x, \cdot)\) is the density of \(Y := \mu(x) + \sigma(x) \xi_{t+1}\) when \(\xi_{t+1} \sim \phi\)
Hence, by (3.6), 
\[ p(x, y) = \frac{1}{\sigma(x)} \phi \left( \frac{y - \mu(x)}{\sigma(x)} \right) \]  
(3.7)
For example, the growth model in (3.5) has stochastic kernel
\[ p(x, y) = \frac{1}{sf(x)} \varphi \left( \frac{y - (1 - \delta)x}{sf(x)} \right) \]  
(3.8)
where \( \varphi \) is the density of \( A_{t+1} \)
(Regarding the state space \( S \) for this model, a natural choice is \( (0, \infty) \) — in which case \( \sigma(x) = sf(x) \) is strictly positive for all \( s \) as required)

**Distribution Dynamics** In *this section* of our lecture on *finite* Markov chains, we asked the following question: If
1. \( \{X_t\} \) is a Markov chain with stochastic matrix \( P \)
2. the distribution of \( X_t \) is known to be \( \psi_t \)
then what is the distribution of \( X_{t+1} \)?
Letting \( \psi_{t+1} \) denote the distribution of \( X_{t+1} \), the answer we gave was that
\[ \psi_{t+1}[j] = \sum_{i \in S} P[i, j]\psi_t[i] \] 
This intuitive equality states that the probability of being at \( j \) tomorrow is the probability of visiting \( i \) today and then going on to \( j \), summed over all possible \( i \)
In the density case, we just replace the sum with an integral and probability mass functions with densities, yielding
\[ \psi_{t+1}(y) = \int p(x, y)\psi_t(x) \, dx, \quad \forall y \in S \]  
(3.9)
It is convenient to think of this updating process in terms of an operator
(An operator is just a function, but the term is usually reserved for a function that sends functions into functions)
Let \( \mathcal{D} \) be the set of all densities on \( S \), and let \( P \) be the operator from \( \mathcal{D} \) to itself that takes density \( \psi \) and sends it into new density \( \psi P \), where the latter is defined by
\[ (\psi P)(y) = \int p(x, y)\psi(x) \, dx \]  
(3.10)
This operator is usually called the *Markov operator* corresponding to \( p \)

**Note:** Unlike most operators, we write \( P \) to the right of its argument, instead of to the left (i.e., \( \psi P \) instead of \( P\psi \)). This is a common convention, with the intention being to maintain the parallel with the finite case — see here

With this notation, we can write (3.9) more succinctly as \( \psi_{t+1}(y) = (\psi_t P)(y) \) for all \( y \), or, dropping the \( y \) and letting “\( = \)” indicate equality of functions,
\[ \psi_{t+1} = \psi_t P \]  
(3.11)
Equation (3.11) tells us that if we specify a distribution for \( \psi_0 \), then the entire sequence of future distributions can be obtained by iterating with \( P \)
It’s interesting to note that (3.11) is a deterministic difference equation.

Thus, by converting a stochastic difference equation such as (3.3) into a stochastic kernel $p$ and hence an operator $P$, we convert a stochastic difference equation into a deterministic one (albeit in a much higher dimensional space).

**Note:** Some people might be aware that discrete Markov chains are in fact a special case of the continuous Markov chains we have just described. The reason is that probability mass functions are densities with respect to the **counting measure**.

**Computation**  To learn about the dynamics of a given process, it’s useful to compute and study the sequences of densities generated by the model.

One way to do this is to try to implement the iteration described by (3.10) and (3.11) using numerical integration.

However, to produce $\psi P$ from $\psi$ via (3.10), you would need to integrate at every $y$, and there is a continuum of such $y$.

Another possibility is to discretize the model, but this introduces errors of unknown size.

A nicer alternative in the present setting is to combine simulation with an elegant estimator called the **look ahead** estimator.

Let’s go over the ideas with reference to the growth model discussed above, the dynamics of which we repeat here for convenience:

$$k_{t+1} = sA_{t+1}f(k_t) + (1 - \delta)k_t$$  

(3.12)

Our aim is to compute the sequence $\{\psi_t\}$ associated with this model and fixed initial condition $\psi_0$.

To approximate $\psi_t$ by simulation, recall that, by definition, $\psi_t$ is the density of $k_t$ given $k_0 \sim \psi_0$.

If we wish to generate observations of this random variable, all we need to do is

1. draw $k_0$ from the specified initial condition $\psi_0$
2. draw the shocks $A_1, \ldots, A_t$ from their specified density $\phi$
3. compute $k_t$ iteratively via (3.12)

If we repeat this $n$ times, we get $n$ independent observations $k^1_t, \ldots, k^n_t$.

With these draws in hand, the next step is to generate some kind of representation of their distribution $\psi_t$.

A naive approach would be to use a histogram, or perhaps a smoothed histogram using the `kde` function from `KernelDensity.jl`.

However, in the present setting there is a much better way to do this, based on the look-ahead estimator.

With this estimator, to construct an estimate of $\psi_t$, we actually generate $n$ observations of $k_{t-1}$, rather than $k_t$. 

3.1. CONTINUOUS STATE MARKOV CHAINS

Now we take these \( n \) observations \( k_{t-1}^1, \ldots, k_{t-1}^n \) and form the estimate

\[
\psi_t^n(y) = \frac{1}{n} \sum_{i=1}^{n} p(k_{t-1}^i, y)
\]  

(3.13)

where \( p \) is the growth model stochastic kernel in (3.8)

What is the justification for this slightly surprising estimator?

The idea is that, by the strong \textit{law of large numbers},

\[
\frac{1}{n} \sum_{i=1}^{n} p(k_{t-1}^i, y) \to \mathbb{E} p(k_{t-1}^i, y) = \int p(x, y) \psi_{t-1}(x) \, dx = \psi_t(y)
\]

with probability one as \( n \to \infty \)

Here the first equality is by the definition of \( \psi_{t-1} \), and the second is by (3.9)

We have just shown that our estimator \( \psi_t^n(y) \) in (3.13) converges almost surely to \( \psi_t(y) \), which is just what we want to compute

In fact much stronger convergence results are true (see, for example, this paper)

**Implementation** A type called LAE for estimating densities by this technique can be found in QuantEcon

We repeat it here for convenience

```python
# Computes a sequence of marginal densities for a continuous state space
# Markov chain \( X_t \) where the transition probabilities can be represented as densities. The estimate of the marginal density of \( X_t \) is

.. math::
\frac{1}{n} \sum_{i=0}^{n} p(X_{t-1}^i, y)

This is a density in \( y \).

@author: Spencer Lyon <spencer.lyon@nyu.edu>

@date: 2014-08-01

References
----------

Simple port of the file quantecon.lae.py

http://quant-econ.net/stationary_densities.html
```

```python
type LAE
    p::Function
    X::Matrix
```
3.1. CONTINUOUS STATE MARKOV CHAINS

```julia
function LAE(p::Function, X::Array)
    n = length(X)
    new(p, reshape(X, n, 1))
end
end

function lae_est{T}(l::LAE, y::Array{T})
    k = length(y)
    v = l.p(l.X, reshape(y, 1, k))
    psi_vals = mean(v, 1)
    return squeeze(psi_vals, 1)
end
```

This function returns the right-hand side of (3.13) using
- an object of type LAE that stores the stochastic kernel and the observations
- the value y as its second argument

The function is vectorized, in the sense that if psi is such an instance and y is an array, then the call psi(y) acts elementwise

(This is the reason that we reshaped X and y inside the type — to make vectorization work)

**Example** An example of usage for the stochastic growth model described above can be found in examples/stochasticgrowth.jl

When run, the code produces a figure like this

![Density of \( k_1 \) (lighter) to \( k_T \) (darker) for \( T = 30 \)](image)
The figure shows part of the density sequence \( \{ \psi_t \} \), with each density computed via the look ahead estimator.

Notice that the sequence of densities shown in the figure seems to be converging — more on this in just a moment.

Another quick comment is that each of these distributions could be interpreted as a cross sectional distribution (recall this discussion).

**Beyond Densities**

Up until now, we have focused exclusively on continuous state Markov chains where all conditional distributions \( p(x, \cdot) \) are densities.

As discussed above, not all distributions can be represented as densities.

If the conditional distribution of \( X_{t+1} \) given \( X_t = x \) cannot be represented as a density for some \( x \in S \), then we need a slightly different theory.

The ultimate option is to switch from densities to probability measures, but not all readers will be familiar with measure theory.

We can, however, construct a fairly general theory using distribution functions.

**Example and Definitions** To illustrate the issues, recall that Hopenhayn and Rogerson [HR93] study a model of firm dynamics where individual firm productivity follows the exogenous process

\[
X_{t+1} = a + \rho X_t + \xi_{t+1}, \quad \text{where} \quad \{\xi_t\} \overset{\text{iid}}{\sim} N(0, \sigma^2)
\]

As is, this fits into the density case we treated above.

However, the authors wanted this process to take values in \([0, 1]\), so they added boundaries at the end points 0 and 1.

One way to write this is

\[
X_{t+1} = h(a + \rho X_t + \xi_{t+1}) \quad \text{where} \quad h(x) := x 1\{0 \leq x \leq 1\} + 1\{x > 1\}
\]

If you think about it, you will see that for any given \( x \in [0, 1] \), the conditional distribution of \( X_{t+1} \) given \( X_t = x \) puts positive probability mass on 0 and 1.

Hence it cannot be represented as a density.

What we can do instead is use cumulative distribution functions (cdfs).

To this end, set

\[
G(x, y) := P\{h(a + \rho x + \xi_{t+1}) \leq y\} \quad (0 \leq x, y \leq 1)
\]

This family of cdfs \( G(x, \cdot) \) plays a role analogous to the stochastic kernel in the density case.

The distribution dynamics in (3.9) are then replaced by

\[
F_{t+1}(y) = \int G(x, y) F_t(dx)
\]
Here $F_t$ and $F_{t+1}$ are cdfs representing the distribution of the current state and next period state. The intuition behind (3.14) is essentially the same as for (3.9).

**Computation.** If you wish to compute these cdfs, you cannot use the look-ahead estimator as before.

Indeed, you should not use any density estimator, since the objects you are estimating/computing are not densities.

One good option is simulation as before, combined with the *empirical distribution function*.

**Stability.**

In our *lecture* on finite Markov chains we also studied stationarity, stability and ergodicity. Here we will cover the same topics for the continuous case.

We will, however, treat only the density case (as in this section), where the stochastic kernel is a family of densities.

The general case is relatively similar — references are given below.

**Theoretical Results.** Analogous to the finite case, given a stochastic kernel $p$ and corresponding Markov operator as defined in (3.10), a density $\psi^*$ on $S$ is called *stationary* for $P$ if it is a fixed point of the operator $P$.

In other words,

$$\psi^*(y) = \int p(x,y)\psi^*(x) \, dx, \quad \forall y \in S$$

(3.15)

As with the finite case, if $\psi^*$ is stationary for $P$, and the distribution of $X_0$ is $\psi^*$, then, in view of (3.11), $X_t$ will have this same distribution for all $t$.

Hence $\psi^*$ is the stochastic equivalent of a steady state.

In the finite case, we learned that at least one stationary distribution exists, although there may be many.

When the state space is infinite, the situation is more complicated.

Even existence can fail very easily.

For example, the random walk model has no stationary density (see, e.g., *EDTC*, p. 210).

However, there are well-known conditions under which a stationary density $\psi^*$ exists.

With additional conditions, we can also get a unique stationary density ($\psi \in \mathcal{D}$ and $\psi = \psi P \implies \psi = \psi^*$), and also global convergence in the sense that

$$\forall \psi \in \mathcal{D}, \quad \psi P^t \to \psi^* \text{ as } t \to \infty$$

(3.16)

This combination of existence, uniqueness and global convergence in the sense of (3.16) is often referred to as *global stability*.  

---

**Thomas Sargent and John Stachurski**

March 21, 2015
Under very similar conditions, we get \textit{ergodicity}, which means that

\[
\frac{1}{n} \sum_{i=1}^{n} h(X_i) \to \int h(x) \psi^*(x) dx \quad \text{as} \; n \to \infty
\]  

(3.17)

for any (measurable) function \( h: S \to \mathbb{R} \) such that the right-hand side is finite.

Note that the convergence in (3.17) does not depend on the distribution (or value) of \( X_0 \).

This is actually very important for simulation — it means we can learn about \( \psi^* \) (i.e., approximate the right hand side of (3.17) via the left hand side) without requiring any special knowledge about what to do with \( X_0 \).

So what are these conditions we require to get global stability and ergodicity?

In essence, it must be the case that

1. Probability mass does not drift off to the “edges” of the state space
2. Sufficient “mixing” obtains

For one such set of conditions see theorem 8.2.14 of \textit{EDTC}.

In addition

- \textbf{[SLP89]} contains a classic (but slightly outdated) treatment of these topics
- From the mathematical literature, \textbf{[LM94]} and \textbf{[MT09]} give outstanding in depth treatments
- Section 8.1.2 of \textit{EDTC} provides detailed intuition, and section 8.3 gives additional references
- \textit{EDTC}, section 11.3.4 provides a specific treatment for the growth model we considered in this lecture

\textbf{An Example of Stability}  \hspace{1em} As stated above, the \textit{growth model treated here} is stable under mild conditions on the primitives

- See \textit{EDTC}, section 11.3.4 for more details

We can see this stability in action — in particular, the convergence in (3.16) — by simulating the path of densities from various initial conditions.

Here is such a figure.

All sequences are converging towards the same limit, regardless of their initial condition.

The details regarding initial conditions and so on are given in \textit{this exercise}, where you are asked to replicate the figure.

\textbf{Computing Stationary Densities}  \hspace{1em} In the preceding figure, each sequence of densities is converging towards the unique stationary density \( \psi^* \).

Even from this figure we can get a fair idea what \( \psi^* \) looks like, and where its mass is located.

However, there is a much more direct way to estimate the stationary density, and it involves only a slight modification of the look ahead estimator.
Let’s say that we have a model of the form (3.3) that is stable and ergodic.

Let $p$ be the corresponding stochastic kernel, as given in (3.7).

To approximate the stationary density $\psi^*$, we can simply generate a long time series $X_0, X_1, \ldots, X_n$ and estimate $\psi^*$ via

$$ \psi^*_n(y) = \frac{1}{n} \sum_{t=1}^{n} p(X_t, y) $$

(3.18)

This is essentially the same as the look ahead estimator (3.13), except that now the observations we generate are a single time series, rather than a cross section.

The justification for (3.18) is that, with probability one as $n \to \infty$,

$$ \frac{1}{n} \sum_{t=1}^{n} p(X_t, y) \to \int p(x, y) \psi^*(x) \, dx = \psi^*(y) $$

where the convergence is by (3.17) and the equality on the right is by (3.15).

The right hand side is exactly what we want to compute.

On top of this asymptotic result, it turns out that the rate of convergence for the look ahead estimator is very good.

The first exercise helps illustrate this point.

**Exercises**

**Exercise 1** Consider the simple threshold autoregressive model

$$ X_{t+1} = \theta |X_t| + (1 - \theta^2)^{1/2} \xi_{t+1} $$

where $\{\xi_t\} \sim N(0, 1)$

(3.19)
This is one of those rare nonlinear stochastic models where an analytical expression for the stationary density is available.

In particular, provided that $|\theta| < 1$, there is a unique stationary density $\psi^*$ given by

$$
\psi^*(y) = 2 \phi(y) \Phi \left[ \frac{\theta y}{(1 - \theta^2)^{1/2}} \right]
$$

(3.20)

Here $\phi$ is the standard normal density and $\Phi$ is the standard normal cdf.

As an exercise, compute the look ahead estimate of $\psi^*$, as defined in (3.18), and compare it with $\psi^*$ in (3.20) to see whether they are indeed close for large $n$.

In doing so, set $\theta = 0.8$ and $n = 500$.

The next figure shows the result of such a computation.

The additional density (black line) is a nonparametric kernel density estimate, added to the solution for illustration.

(You can try to replicate it before looking at the solution if you want to.)

As you can see, the look ahead estimator is a much tighter fit than the kernel density estimator.

If you repeat the simulation you will see that this is consistently the case.

**Exercise 2**  Replicate the figure on global convergence shown above.

The densities come from the stochastic growth model treated at the start of the lecture.

Begin with the code found in examples/stochasticgrowth.jl.

Use the same parameters.
For the four initial distributions, use the beta distribution and shift the random draws as shown below

\[\psi_0 = \text{Beta}(5.0, 5.0) \quad \# \text{ Initial distribution}\]
\[n = 1000\]
\[\# \text{.... more setup}\]
\[\text{for } i=1:4\]
\[\# \text{.... some code}\]
\[\text{rand\_draws} = (\text{rand}(\psi_0, n) + 2.5i) / 2\]

**Exercise 3**  A common way to compare distributions visually is with boxplots

To illustrate, let’s generate three artificial data sets and compare them with a boxplot

```
using PyPlot

n = 500
n = 500
x = randn(n) \quad \# N(0, 1)
x = \exp(x) \quad \# \text{Map } x \text{ to lognormal}
y = randn(n) + 2.0 \quad \# N(2, 1)
z = randn(n) + 4.0 \quad \# N(4, 1)

fig, ax = subplots()
ax[:boxplot]([x y z])
ax[:set_xticks]((1, 2, 3))
ax[:set_ylim](-2, 14)
ax[:set_xticklabels]((L"X", L"Y", L"Z"), fontsize=16)
plt.show()
```

The three data sets are

\[\{X_1, \ldots, X_n\} \sim LN(0,1), \quad \{Y_1, \ldots, Y_n\} \sim N(2,1), \quad \text{and } \{Z_1, \ldots, Z_n\} \sim N(4,1),\]

The figure looks as follows

Each data set is represented by a box, where the top and bottom of the box are the third and first quartiles of the data, and the red line in the center is the median

The boxes give some indication as to

- the location of probability mass for each sample
- whether the distribution is right-skewed (as is the lognormal distribution), etc

Now let’s put these ideas to use in a simulation

Consider the threshold autoregressive model in (3.19)

We know that the distribution of \(X_t\) will converge to (3.20) whenever \(|\theta| < 1\)

Let’s observe this convergence from different initial conditions using boxplots

In particular, the exercise is to generate \(J\) boxplot figures, one for each initial condition \(X_0\) in
initial_conditions = linspace(8, 0, J)

For each \( X_0 \) in this set,

1. Generate \( k \) time series of length \( n \), each starting at \( X_0 \) and obeying (3.19)
2. Create a boxplot representing \( n \) distributions, where the \( t \)-th distribution shows the \( k \) observations of \( X_t \)

Use \( \theta = 0.9, n = 20, k = 5000, J = 8 \)

**Solutions**

Solution notebook

**Appendix**

Here’s the proof of (3.6)

Let \( F_U \) and \( F_V \) be the cumulative distributions of \( U \) and \( V \) respectively

By the definition of \( V \), we have

\[
F_V(v) = \mathbb{P}\{a + bU \leq v\} = \mathbb{P}\{U \leq (v-a)/b\}
\]

In other words, \( F_V(v) = F_U((v-a)/b) \)

Differentiating with respect to \( v \) yields (3.6)
3.2 The Lucas Asset Pricing Model

Overview

As stated in an earlier lecture, an asset is a claim on a stream of prospective payments. What is the correct price to pay for such a claim?

The elegant asset pricing model of Lucas [Luc78] attempts to answer this question in an equilibrium setting with risk averse agents.

While we mentioned some consequences of Lucas’ model earlier, it is now time to work through the model more carefully, and try to understand where the fundamental asset pricing equation comes from.

A side benefit of studying Lucas’ model is that it provides a beautiful illustration of model building in general and equilibrium pricing in competitive models in particular.

The Lucas Model

Lucas studied a pure exchange economy with a representative consumer (or household), where

- **Pure exchange** means that all endowments are exogenous
- **Representative** consumer means that either
  - there is a single consumer (sometimes also referred to as a household), or
  - all consumers have identical endowments and preferences

Either way, the assumption of a representative agent means that prices adjust to eradicate desires to trade.

This makes it very easy to compute competitive equilibrium prices.

Basic Setup

Let’s review the set up.

Assets

There is a single “productive unit” that costlessly generates a sequence of consumption goods \( \{ y_t \}_{t=0}^{\infty} \).

Another way to view \( \{ y_t \}_{t=0}^{\infty} \) is as a *consumption endowment* for this economy.
We will assume that this endowment is Markovian, following the exogenous process

\[ y_{t+1} = G(y_t, \xi_{t+1}) \]

Here \( \{\xi_t\} \) is an iid shock sequence with known distribution \( \phi \) and \( y_t \geq 0 \)

An asset is a claim on all or part of this endowment stream

The consumption goods \( \{y_t\}_{t=0}^\infty \) are nonstorable, so holding assets is the only way to transfer wealth into the future

For the purposes of intuition, it’s common to think of the productive unit as a “tree” that produces fruit

Based on this idea, a “Lucas tree” is a claim on the consumption endowment

**Consumers**  A representative consumer ranks consumption streams \( \{c_t\} \) according to the time separable utility functional

\[ E \sum_{t=0}^\infty \beta^t u(c_t) \]  

(3.21)

Here

- \( \beta \in (0, 1) \) is a fixed discount factor
- \( u \) is a strictly increasing, strictly concave, continuously differentiable period utility function
- \( E \) is a mathematical expectation

**Pricing a Lucas Tree**  What is an appropriate price for a claim on the consumption endowment?

We’ll price an *ex dividend* claim, meaning that

- the seller retains this period’s dividend
- the buyer pays \( p_t \) today to purchase a claim on
  - \( y_{t+1} \) and
  - the right to sell the claim tomorrow at price \( p_{t+1} \)

Since this is a competitive model, the first step is to pin down consumer behavior, taking prices as given

Next we’ll impose equilibrium constraints and try to back out prices

In the consumer problem, the consumer’s control variable is the share \( \pi_t \) of the claim held in each period

Thus, the consumer problem is to maximize (3.21) subject to

\[ c_t + \pi_{t+1} p_t \leq \pi_t y_t + \pi_t p_t \]

along with \( c_t \geq 0 \) and \( 0 \leq \pi_t \leq 1 \) at each \( t \)

The decision to hold share \( \pi_t \) is actually made at time \( t - 1 \)

But this value is inherited as a state variable at time \( t \), which explains the choice of subscript
The dynamic program  We can write the consumer problem as a dynamic programming problem.

Our first observation is that prices depend on current information, and current information is really just the endowment process up until the current period.

In fact the endowment process is Markovian, so that the only relevant information is the current state \( y \in \mathbb{R}_+ \) (dropping the time subscript).

This leads us to guess an equilibrium where price is a function \( p \) of \( y \).

Remarks on the solution method

- Since this is a competitive (read: price taking) model, the consumer will take this function \( p \) as given.
- In this way we determine consumer behavior given \( p \) and then use equilibrium conditions to recover \( p \).
- This is the standard way to solve competitive equilibrium models.

Using the assumption that price is a given function \( p \) of \( y \), we write the value function and constraint as

\[
 v(\pi, y) = \max_{c, \pi'} \left\{ u(c) + \beta \int v(\pi', G(y, z)) \phi(dz) \right\}
\]

subject to

\[
 c + \pi' p(y) \leq \pi y + \pi p(y) \tag{3.22}
\]

We can invoke the fact that utility is increasing to claim equality in (3.22) and hence eliminate the constraint, obtaining

\[
 v(\pi, y) = \max_{\pi'} \left\{ u[\pi(y + p(y)) - \pi' p(y)] + \beta \int v(\pi', G(y, z)) \phi(dz) \right\} \tag{3.23}
\]

The solution to this dynamic programming problem is an optimal policy expressing either \( \pi' \) or \( c \) as a function of the state \( (\pi, y) \).

- Each one determines the other, since \( c(\pi, y) = \pi(y + p(y)) - \pi'(\pi, y)p(y) \).

Next steps  What we need to do now is determine equilibrium prices.

It seems that to obtain these, we will have to

1. Solve this two dimensional dynamic programming problem for the optimal policy.
2. Impose equilibrium constraints.
3. Solve out for the price function \( p(y) \) directly.

However, as Lucas showed, there is a related but more straightforward way to do this.

Equilibrium constraints  Since the consumption good is not storable, in equilibrium we must have \( c_t = y_t \) for all \( t \).

In addition, since there is one representative consumer (alternatively, since all consumers are identical), there should be no trade in equilibrium.
In particular, the representative consumer owns the whole tree in every period, so $\pi_t = 1$ for all $t$

Prices must adjust to satisfy these two constraints

**The equilibrium price function**  Now observe that the first order condition for (3.23) can be written as

$$u'(c)p(y) = \beta \int v'_1(\pi', G(y, z)) \phi(dz)$$

where $v'_1$ is the derivative of $v$ with respect to its first argument

To obtain $v'_1$ we can simply differentiate the right hand side of (3.23) with respect to $\pi$, yielding

$$v'_1(\pi, y) = u'(c)(y + p(y))$$

Next we impose the equilibrium constraints while combining the last two equations to get

$$p(y) = \beta \int \frac{u'[G(y, z)]}{u'(y)} [G(y, z) + p(G(y, z))] \phi(dz)$$

(3.24)

In sequential rather than functional notation, we can also write this as

$$p_t = E_t \left[ \beta \frac{u'(c_{t+1})}{u'(c_t)} (c_{t+1} + p_{t+1}) \right]$$

(3.25)

This is the famous consumption-based asset pricing equation

Before discussing it further we want to solve out for prices

**Solving the Model**  Equation (3.24) is a *functional equation* in the unknown function $p$

The solution is an equilibrium price function $p^*$

Let’s look at how to obtain it

**Setting up the problem**  Instead of solving for it directly we’ll follow Lucas’ indirect approach, first setting

$$f(y) := u'(y)p(y)$$

(3.26)

so that (3.24) becomes

$$f(y) = h(y) + \beta \int f[G(y, z)] \phi(dz)$$

(3.27)

Here $h(y) := \beta \int u'[G(y, z)]G(y, z) \phi(dz)$ is a function that depends only on the primitives

Equation (3.27) is a functional equation in $f$

The plan is to solve out for $f$ and convert back to $p$ via (3.26)

To solve (3.27) we’ll use a standard method: convert it to a fixed point problem

First we introduce the operator $T$ mapping $f$ into $Tf$ as defined by

$$(Tf)(y) = h(y) + \beta \int f[G(y, z)] \phi(dz)$$

(3.28)
The reason we do this is that a solution to (3.27) now corresponds to a function \( f^* \) satisfying 
\[(Tf^*)(y) = f^*(y)\] for all \( y \)
In other words, a solution is a fixed point of \( T \)
This means that we can use fixed point theory to obtain and compute the solution

A little fixed point theory    Let \( cb\mathbb{R}_+ \) be the set of continuous bounded functions \( f: \mathbb{R}_+ \to \mathbb{R}_+ \)
We now show that
1. \( T \) has exactly one fixed point \( f^* \) in \( cb\mathbb{R}_+ \)
2. For any \( f \in cb\mathbb{R}_+ \), the sequence \( T^k f \) converges uniformly to \( f^* \)
(Note: If you find the mathematics heavy going you can take 1–2 as given and skip to the next section)
Recall the Banach contraction mapping theorem
It tells us that the previous statements will be true if we can find an \( \alpha < 1 \) such that
\[
\| T f - T g \| \leq \alpha \| f - g \|, \quad \forall f, g \in cb\mathbb{R}_+ \tag{3.29}
\]
Here \( \| h \| := \sup_{x \in \mathbb{R}_+} |h(x)| \)
To see that (3.29) is valid, pick any \( f, g \in cb\mathbb{R}_+ \) and any \( y \in \mathbb{R}_+ \)
Observe that, since integrals get larger when absolute values are moved to the inside,
\[
|Tf(y) - Tg(y)| = \left| \beta \int f(G(y, z))\phi(dz) - \beta \int g(G(y, z))\phi(dz) \right|
\leq \beta \int |f(G(y, z)) - g(G(y, z))| \phi(dz)
\leq \beta \int \| f - g \| \phi(dz)
= \beta \| f - g \|
\]
Since the right hand side is an upper bound, taking the sup over all \( y \) on the left hand side gives (3.29) with \( \alpha := \beta \)

Computation – An Example    The preceding discussion tells that we can compute \( f^* \) by picking any arbitrary \( f \in cb\mathbb{R}_+ \) and then iterating with \( T \)
The equilibrium price function \( p^* \) can then be recovered by \( p^*(y) = f^*(y) / u'(y) \)
Let’s try this when \( \ln y_{t+1} = \alpha \ln y_t + \sigma \epsilon_{t+1} \) where \( \{\epsilon_t\} \) is iid and standard normal
Utility will take the isoelastic form \( u(c) = c^{1-\gamma} / (1 - \gamma) \), where \( \gamma > 0 \) is the coefficient of relative risk aversion
Some code to implement the iterative computational procedure can be found in lucastree.jl from the QuantEcon package
We repeat it here for convenience
An example of usage is given in the docstring and repeated here

```
import LucasTree

tree = LucasTree(2, 0.95, 0.90, 0.1)
grid, price_vals = compute_lt_price(tree)
```

Here’s the resulting price function

![Price function graph]

The price is increasing, even if we remove all serial correlation from the endowment process.
The reason is that a larger current endowment reduces current marginal utility.
The price must therefore rise to induce the household to consume the entire endowment (and hence satisfy the resource constraint).

What happens with a more patient consumer?
Here the blue line corresponds to the previous parameters and the green line is price when $\beta = 0.98$.

We see that when consumers are more patient the asset becomes more valuable, and the price of the Lucas tree shifts up.

Exercise 1 asks you to replicate this figure.

**Exercises**

**Exercise 1** Replicate the figure to show how discount rates affect prices.
Next we study a computational problem concerning career and job choices. The model is originally due to Derek Neal [Nea99] and this exposition draws on the presentation in [LS12], section 6.5.

Model features
3.3. MODELING CAREER CHOICE

- career and job within career both chosen to maximize expected discounted wage flow
- infinite horizon dynamic programming with two states variables

Model

In what follows we distinguish between a career and a job, where
- a career is understood to be a general field encompassing many possible jobs, and
- a job is understood to be a position with a particular firm

For workers, wages can be decomposed into the contribution of job and career
- \( w_t = \theta_t + \epsilon_t \), where
  - \( \theta_t \) is contribution of career at time \( t \)
  - \( \epsilon_t \) is contribution of job at time \( t \)

At the start of time \( t \), a worker has the following options
- retain a current (career, job) pair \((\theta_t, \epsilon_t)\) — referred to hereafter as “stay put”
- retain a current career \( \theta_t \) but redraw a job \( \epsilon_t \) — referred to hereafter as “new job”
- redraw both a career \( \theta_t \) and a job \( \epsilon_t \) — referred to hereafter as “new life”

Draws of \( \theta \) and \( \epsilon \) are independent of each other and past values, with
- \( \theta_t \sim F \)
- \( \epsilon_t \sim G \)

Notice that the worker does not have the option to retain a job but redraw a career — starting a new career always requires starting a new job

A young worker aims to maximize the expected sum of discounted wages

\[
E \sum_{t=0}^{\infty} \beta^t w_t
\]

subject to the choice restrictions specified above

Let \( V(\theta, \epsilon) \) denote the value function, which is the maximum of (3.30) over all feasible (career, job) policies, given the initial state \((\theta, \epsilon)\)

The value function obeys

\[
V(\theta, \epsilon) = \max \{ I, II, III \},
\]

where

\[
I = \theta + \epsilon + \beta V(\theta, \epsilon)
\]

\[
II = \theta + \int \epsilon' G(d\epsilon') + \beta \int V(\theta, \epsilon') G(d\epsilon')
\]

\[
III = \int \theta' F(d\theta') + \int \epsilon' G(d\epsilon') + \beta \int \int V(\theta', \epsilon') G(d\epsilon') F(d\theta')
\]

Evidently I, II and III correspond to “stay put”, “new job” and “new life”, respectively
Parameterization   As in [LS12], section 6.5, we will focus on a discrete version of the model, parameterized as follows:

- both $\theta$ and $\epsilon$ take values in the set $\text{linspace}(0, B, N)$ — an even grid of $N$ points between 0 and $B$ inclusive
- $N = 50$
- $B = 5$
- $\beta = 0.95$

The distributions $F$ and $G$ are discrete distributions generating draws from the grid points $\text{linspace}(0, B, N)$

A very useful family of discrete distributions is the Beta-binomial family, with probability mass function

$$p(k \mid n, a, b) = \binom{n}{k} \frac{B(k + a, n - k + b)}{B(a, b)}, \quad k = 0, \ldots, n$$

Interpretation:

- draw $q$ from a Beta distribution with shape parameters $(a, b)$
- run $n$ independent binary trials, each with success probability $q$
- $p(k \mid n, a, b)$ is the probability of $k$ successes in these $n$ trials

Nice properties:

- very flexible class of distributions, including uniform, symmetric unimodal, etc.
- only three parameters

Here’s a figure showing the effect of different shape parameters when $n = 50$

![Figure showing the effect of different shape parameters](image)

The code that generated this figure can be found [here](link)
Implementation: career.jl

The QuantEcon package provides some code for solving the DP problem described above. See in particular this file, which is repeated here for convenience.

```julia
#=
A type to solve the career / job choice model due to Derek Neal.

@author: Spencer Lyon <spencer.lyon@nyu.edu>
@date: 2014-08-05

References
--------
http://quant-econ.net/career.html


=#

type CareerWorkerProblem
    beta::Real
    N::Int
    B::Real
    theta::Vector
    epsilon::Vector
    F_probs::Vector
    G_probs::Vector
    F_mean::Real
    G_mean::Real
end

function CareerWorkerProblem(beta::Real=0.95, B::Real=5.0, N::Real=50, F_a::Real=1, F_b::Real=1, G_a::Real=1, G_b::Real=1)
    theta = linspace(0, B, N)
    epsilon = copy(theta)
    F_probs::Vector{Float64} = pdf(BetaBinomial(N-1, F_a, F_b))
    G_probs::Vector{Float64} = pdf(BetaBinomial(N-1, G_a, G_b))
    F_mean = sum(theta .* F_probs)
    G_mean = sum(epsilon .* G_probs)
    CareerWorkerProblem(beta, N, B, theta, epsilon, F_probs, G_probs, F_mean, G_mean)
end

# create kwarg version
function CareerWorkerProblem(;beta::Real=0.95, B::Real=5.0, N::Real=50, F_a::Real=1, F_b::Real=1, G_a::Real=1, G_b::Real=1)
    CareerWorkerProblem(beta, N, B, F_a, F_b, G_a, G_b)
end
```

THOMAS SARGENT AND JOHN STACHURSKI

March 21, 2015
function bellman_operator!(cp::CareerWorkerProblem, v::Array, out::Array; ret_policy=false)
    # new life. This is a function of the distribution parameters and is always constant. No need to recompute it in the loop
    v3 = (cp.G_mean + cp.F_mean + cp.beta .* cp.F_probs' * v * cp.G_probs)[1]  # don't need 1 element array
    for j=1:cp.N
        for i=1:cp.N
            # stay put
            v1 = cp.theta[i] + cp.epsilon[j] + cp.beta * v[i, j]

            # new job
            v2 = (cp.theta[i] + cp.G_mean + cp.beta .* v[i, :] * cp.G_probs)[1]  # don't need a single element array

            if ret_policy
                if v1 > max(v2, v3)
                    action = 1
                elseif v2 > max(v1, v3)
                    action = 2
                else
                    action = 3
                end
                out[i, j] = action
            else
                out[i, j] = max(v1, v2, v3)
            end
        end
    end
end

function bellman_operator(cp::CareerWorkerProblem, v::Array; ret_policy=false)
    out = similar(v)
    bellman_operator!(cp, v, out, ret_policy=ret_policy)
    return out
end

function get_greedy!(cp::CareerWorkerProblem, v::Array, out::Array)
    bellman_operator!(cp, v, out, ret_policy=true)
end

function get_greedy(cp::CareerWorkerProblem, v::Array)
    bellman_operator(cp, v, ret_policy=true)
end

The code defines
3.3. MODELING CAREER CHOICE

• a type CareerWorkerProblem that
  – encapsulates all the details of a particular parameterization
  – implement the Bellman operator $T$

In this model, $T$ is defined by $Tv(\theta, \epsilon) = \max\{I, II, III\}$, where $I$, $II$ and $III$ are as given in (3.31), replacing $V$ with $v$

The default probability distributions in CareerWorkerProblem correspond to discrete uniform distributions (see the Beta-binomial figure)

In fact all our default settings correspond to the version studied in [LS12], section 6.5.

Hence we can reproduce figures 6.5.1 and 6.5.2 shown there, which exhibit the value function and optimal policy respectively

Here’s the value function

![Value function with uniform probabilities](image)

Figure 3.1: Value function with uniform probabilities

The code used to produce this plot was examples/career_vf_plot.jl

The optimal policy can be represented as follows (see Exercise 3 for code)

Interpretation:

• If both job and career are poor or mediocre, the worker will experiment with new job and new career

• If career is sufficiently good, the worker will hold it and experiment with new jobs until a sufficiently good one is found

• If both job and career are good, the worker will stay put
3.3. MODELING CAREER CHOICE

Notice that the worker will always hold on to a sufficiently good career, but not necessarily hold on to even the best paying job.

The reason is that high lifetime wages require both variables to be large, and the worker cannot change careers without changing jobs.

- Sometimes a good job must be sacrificed in order to change to a better career.

**Exercises**

**Exercise 1** Using the default parameterization in the type `CareerWorkerProblem`, generate and plot typical sample paths for $\theta$ and $\epsilon$ when the worker follows the optimal policy.

In particular, modulo randomness, reproduce the following figure (where the horizontal axis represents time).

Hint: To generate the draws from the distributions $F$ and $G$, use the type `DiscreteRV`.

**Exercise 2** Let’s now consider how long it takes for the worker to settle down to a permanent job, given a starting point of $(\theta, \epsilon) = (0, 0)$.

In other words, we want to study the distribution of the random variable

\[ T^* := \text{the first point in time from which the worker’s job no longer changes} \]
Evidently, the worker’s job becomes permanent if and only if \((\theta_t, \epsilon_t)\) enters the “stay put” region of \((\theta, \epsilon)\) space.

Letting \(S\) denote this region, \(T^*\) can be expressed as the first passage time to \(S\) under the optimal policy:

\[
T^* := \inf\{t \geq 0 \mid (\theta_t, \epsilon_t) \in S\}
\]

Collect 25,000 draws of this random variable and compute the median (which should be about 7).

Repeat the exercise with \(\beta = 0.99\) and interpret the change.

**Exercise 3**  As best you can, reproduce the figure showing the optimal policy.

Hint: The `get_greedy()` function returns a representation of the optimal policy where values 1, 2 and 3 correspond to “stay put”, “new job” and “new life” respectively. Use this and `contourf` from `PyPlot.jl` to produce the different shadings.

Now set \(G_a = G_b = 100\) and generate a new figure with these parameters. Interpret.

**Solutions**

**Solution notebook**

### 3.4 On-the-Job Search
3.4. ON-THE-JOB SEARCH

Contents

• On-the-Job Search
  – Overview
  – Model
  – Implementation
  – Solving for Policies
  – Exercises
  – Solutions

Overview

In this section we solve a simple on-the-job search model

• based on [LS12], exercise 6.18
• see also [add Jovanovic reference]

Model features

• job-specific human capital accumulation combined with on-the-job search
• infinite horizon dynamic programming with one state variable and two controls

Model

Let

• \( x_t \) denote the time-\( t \) job-specific human capital of a worker employed at a given firm
• \( w_t \) denote current wages

Let \( w_t = x_t(1 - s_t - \phi_t) \), where

• \( \phi_t \) is investment in job-specific human capital for the current role
• \( s_t \) is search effort, devoted to obtaining new offers from other firms.

For as long as the worker remains in the current job, evolution of \( \{x_t\} \) is given by \( x_{t+1} = G(x_t, \phi_t) \)

When search effort at \( t \) is \( s_t \), the worker receives a new job offer with probability \( \pi(s_t) \in [0,1] \)

Value of offer is \( U_{t+1} \), where \( \{U_t\} \) is iid with common distribution \( F \)

Worker has the right to reject the current offer and continue with existing job.

In particular, \( x_{t+1} = U_{t+1} \) if accepts and \( x_{t+1} = G(x_t, \phi_t) \) if rejects

Letting \( b_{t+1} \in \{0,1\} \) be binary with \( b_{t+1} = 1 \) indicating an offer, we can write

\[
x_{t+1} = (1 - b_{t+1})G(x_t, \phi_t) + b_{t+1} \max\{G(x_t, \phi_t), U_{t+1}\}
\] (3.32)
Agent’s objective: maximize expected discounted sum of wages via controls \( \{s_t\} \) and \( \{\phi_t\} \)

Taking the expectation of \( V(x_{t+1}) \) and using (3.32), the Bellman equation for this problem can be written as

\[
V(x) = \max_{s+\phi \leq 1} \left\{ x(1 - s - \phi) + \beta(1 - \pi(s))V[G(x, \phi)] + \beta \pi(s) \int V[G(x, \phi) \vee u]F(du) \right\}. \tag{3.33}
\]

Here nonnegativity of \( s \) and \( \phi \) is understood, while \( a \vee b := \max\{a, b\} \)

**Parameterization**  In the implementation below, we will focus on the parameterization

\[
G(x, \phi) = A(x\phi)^\alpha, \quad \pi(s) = \sqrt{s} \quad \text{and} \quad F = \text{Beta}(2, 2)
\]

with default parameter values

- \( A = 1.4 \)
- \( \alpha = 0.6 \)
- \( \beta = 0.96 \)

The Beta(2,2) distribution is supported on \((0, 1)\). It has a unimodal, symmetric density peaked at 0.5.

**Back-of-the-Envelope Calculations**  Before we solve the model, let’s make some quick calculations that provide intuition on what the solution should look like.

To begin, observe that the worker has two instruments to build capital and hence wages:

1. invest in capital specific to the current job via \( \phi \)
2. search for a new job with better job-specific capital match via \( s \)

Since wages are \( x(1 - s - \phi) \), marginal cost of investment via either \( \phi \) or \( s \) is identical

Our risk neutral worker should focus on whatever instrument has the highest expected return

The relative expected return will depend on \( x \)

For example, suppose first that \( x = 0.05 \)

- If \( s = 1 \) and \( \phi = 0 \), then since \( G(x, \phi) = 0 \), taking expectations of (3.32) gives expected next period capital equal to \( \pi(s)EU = EU = 0.5 \)
- If \( s = 0 \) and \( \phi = 1 \), then next period capital is \( G(x, \phi) = G(0.05, 1) \approx 0.23 \)

Both rates of return are good, but the return from search is better

Next suppose that \( x = 0.4 \)

- If \( s = 1 \) and \( \phi = 0 \), then expected next period capital is again 0.5
- If \( s = 0 \) and \( \phi = 1 \), then \( G(x, \phi) = G(0.4, 1) \approx 0.8 \)

Return from investment via \( \phi \) dominates expected return from search

Combining these observations gives us two informal predictions:
3.4. ON-THE-JOB SEARCH

1. At any given state $x$, the two controls $\phi$ and $s$ will function primarily as substitutes — worker will focus on whichever instrument has the higher expected return.

2. For sufficiently small $x$, search will be preferable to investment in job-specific human capital. For larger $x$, the reverse will be true.

Now let’s turn to implementation, and see if we can match our predictions.

**Implementation**

The QuantEcon package provides some code for solving the DP problem described above. See in particular `jv.jl`, which is repeated here for convenience.

```julia
#+ @author: Spencer Lyon <spencer.lyon@nyu.edu>
@date: 2014-06-27

References
----------

Simple port of the file `quantecon.models.jv`

http://quant-econ.net/jv.html
```
```
3.4. ON-THE-JOB SEARCH

\[ F = \text{Beta}(2, 2) \]

# Set up grid over the state space for DP
# Max of grid is the max of a large quantile value for F and the
# fixed point \( y = G(y, 1) \).
grid_max = \text{max}(A^{1.0 / (1.0 - \alpha)}, \text{quantile}(F, 1 - \epsilon))

# range for linspace(epsilon, grid_max, grid_size). Needed for
# CoordInterpGrid below
x_grid = linspace_range(epsilon, grid_max, grid_size)

JvWorker(A, alpha, bet, x_grid, G, pi_func, F)
end

# make kwarg version
JvWorker(;A=1.4, alpha=0.6, bet=0.96, grid_size=50) = JvWorker(A, alpha, bet,
grid_size)

# TODO: as of 2014-08-13 there is no simple constrained optimizer in Julia
# so, we default to the brute force gridsearch approach for this
# problem

# NOTE: this function is not type stable because it returns either
# Array(Float64, 2) or (Array(Float64, 2), Array(Float64, 2))
# depending on the value of ret_policies. This is probably not a
# huge deal, but it is something to be aware of
function bellman_operator!(jv::JvWorker, V::Vector,
out::Union(Vector, (Vector, Vector));
brute_force=true, ret_policies=false)

# simplify notation

# prepare interpoland of value function
Vf = CoordInterpGrid(jv.x_grid, V, BCnearest, InterpLinear)

# prepare integration bounds
a, b, = \text{quantile}(F, 0.005), \text{quantile}(F, 0.995)

# instantiate variables so they are available outside loop and exist
# within it
if ret_policies
    if !(typeof(out) <: (Vector, Vector))
        msg = "You asked for policies, but only provided one output array"
        msg *= "\the there are two policies so two arrays must be given"
        error(msg)
    end
    s_policy, phi_policy = out[1], out[2]
else
    c1(z) = 1.0 - \text{sum}(z)
    c2(z) = z[1] - \epsilon
    c3(z) = z[2] - \epsilon
    guess = (0.2, 0.2)
constraints = ["type" => "ineq", "fun" => i for i in [c1, c2, c3]]
if typeof(out) <: Tuple
    msg = "Multiple output arrays given. There is only one value"
    msg = " function. Did you mean to pass ret_policies=true?"
    error(msg)
end
new_V = out

# instantiate the linesearch variables if we need to
if brute_force
    max_val = -1.0
    cur_val = 0.0
    max_s = 1.0
    max_phi = 1.0
    search_grid = linspace(epsilon, 1.0, 15)
end
for (i, x) in enumerate(jv.x_grid)
    function w(z)
        s, phi = z
        h(u) = Vf[max(G(x, phi), u)] * pdf(F, u)
        integral, err = quadgk(h, a, b)
        q = pi_func(s) * integral + (1.0 - pi_func(s)) * Vf[G(x, phi)]
        return - x * (1.0 - phi - s) - bet * q
    end
    if brute_force
        for s in search_grid
            for phi in search_grid
                if s + phi <= 1.0
                    cur_val = -w((s, phi))
                else
                    cur_val = -1.0
                end
                if cur_val > max_val
                    max_val, max_s, max_phi = cur_val, s, phi
                end
            end
        end
    else
        max_s, max_phi = minimize(w, guess, constraints=constraints,
            options="disp" => 0,
            method="SLSQP")["x"]
        max_val = -w((max_s, max_phi), x, a, b, Vf, jv)
    end
    if ret_policies
3.4. ON-THE-JOB SEARCH

\[
s_{\text{policy}}[i], \phi_{\text{policy}}[i] = \max_s, \max_{\phi}\ \\
\text{else} \quad \text{new}_V[i] = \max_{\text{val}}\ \\
\text{end} \quad \text{end}
\]

\[
\text{function bellman_operator}(jv\!::\!\text{JvWorker}, V\!::\!\text{Vector}; \text{brute-force}=true, \text{ret-policies}=false) \\
\quad \text{if ret-policies} \\
\quad \quad \text{out} = (\text{similar}(V), \text{similar}(V)) \\
\quad \text{else} \\
\quad \quad \text{out} = \text{similar}(V) \\
\quad \text{end} \\
\quad \text{bellman_operator!}(jv, V, \text{out}, \text{brute-force}=\text{brute-force}, \text{ret-policies}=\text{ret-policies}) \\
\quad \text{return out}
\]

\[
\text{function get_greedy!(jv\!::\!\text{JvWorker}, V\!::\!\text{Vector}, \text{out}\!::\!(\text{Vector}, \text{Vector}); \text{brute-force}=true) \\
\quad \text{bellman_operator!}(jv, V, \text{out}, \text{ret-policies}=true)
\]

\[
\text{function get_greedy}(jv\!::\!\text{JvWorker}, V\!::\!\text{Vector}; \text{brute-force}=true) \\
\quad \text{bellman_operator}(jv, V, \text{ret-policies}=true)
\]

The code is written to be relatively generic—and hence reusable

- For example, we use generic \(G(x, \phi)\) instead of specific \(A(x, \phi)\)

Regarding the imports

- \texttt{fixed_quad} is a simple non-adaptive integration routine
- \texttt{fmin_slsqp} is a minimization routine that permits inequality constraints

Next we build a type called \texttt{JvWorker} that

- packages all the parameters and other basic attributes of a given model
- Implements the method \texttt{bellman_operator} for value function iteration

The \texttt{bellman_operator} method takes a candidate value function \(V\) and updates it to \(TV\) via

\[
TV(x) = \min_{s+\phi \leq 1} w(s, \phi)
\]

where

\[
w(s, \phi) := - \left\{ x(1 - s - \phi) + \beta(1 - \pi(s))V[G(x, \phi)] + \beta\pi(s) \int V[G(x, \phi) \lor u]F(du) \right\} \quad (3.34)
\]

Here we are minimizing instead of maximizing to fit with SciPy’s optimization routines
When we represent $V$, it will be with a Julia array $V$ giving values on grid $x_{\text{grid}}$

But to evaluate the right-hand side of (3.34), we need a function, so we replace the arrays $V$ and $x_{\text{grid}}$ with a function $Vf$ that gives linear interpolation of $V$ on $x_{\text{grid}}$

Hence in the preliminaries of `bellman_operator`

- from the array $V$ we define a linear interpolation $Vf$ of its values
  - $c1$ is used to implement the constraint $s + \phi \leq 1$
  - $c2$ is used to implement $s \geq \epsilon$, a numerically stable
    alternative to the true constraint $s \geq 0$
  - $c3$ does the same for $\phi$

Inside the `for` loop, for each $x$ in the grid over the state space, we set up the function $w(z) = w(s, \phi)$ defined in (3.34).

The function is minimized over all feasible $(s, \phi)$ pairs, either by

- a relatively sophisticated solver from SciPy called `fmin_slsqp`, or
- brute force search over a grid

The former is much faster, but convergence to the global optimum is not guaranteed. Grid search is a simple way to check results

**Solving for Policies**

Let’s plot the optimal policies and see what they look like

The code is in a file `examples/jv_test.jl` from the main repository and looks as follows

```
# === plot policies === #
fig, ax = subplots()
ax[:set_xlim](0, maximum(wp.x_grid))
ax[:set_ylim](-0.1, 1.1)
ax[:plot](wp.x_grid, phi_policy, "b-", label="phi")
ax[:plot](wp.x_grid, s_policy, "g-", label="s")
ax[:set_xlabel]("x")
ax[:legend]()
plt.show()
```

It produces the following figure

The horizontal axis is the state $x$, while the vertical axis gives $s(x)$ and $\phi(x)$

Overall, the policies match well with our predictions from section Back-of-the-Envelope Calculations.

- Worker switches from one investment strategy to the other depending on relative return
- For low values of $x$, the best option is to search for a new job
- Once $x$ is larger, worker does better by investing in human capital specific to the current position
3.4. ON-THE-JOB SEARCH

Figure 3.2: Optimal policies

Exercises

Exercise 1  Let’s look at the dynamics for the state process \( \{x_t\} \) associated with these policies.

The dynamics are given by (3.32) when \( \phi_t \) and \( s_t \) are chosen according to the optimal policies, and \( P\{b_{t+1} = 1\} = \pi(st) \).

Since the dynamics are random, analysis is a bit subtle

One way to do it is to plot, for each \( x \) in a relatively fine grid called `plot_grid`, a large number \( K \) of realizations of \( x_{t+1} \) given \( x_t = x \). Plot this with one dot for each realization, in the form of a 45 degree diagram. Set:

\[
K = 50 \\
\text{plot\_grid\_max, plot\_grid\_size} = 1.2, 100 \\
\text{plot\_grid} = \text{linspace}(0, \text{plot\_grid\_max}, \text{plot\_grid\_size}) \\
\text{fig, ax} = \text{subplots()} \\
\text{ax[:set_xlim]}(0, \text{plot\_grid\_max}) \\
\text{ax[:set_ylim]}(0, \text{plot\_grid\_max})
\]

By examining the plot, argue that under the optimal policies, the state \( x_t \) will converge to a constant value \( \bar{x} \) close to unity.

Argue that at the steady state, \( s_t \approx 0 \) and \( \phi_t \approx 0.6 \).

Exercise 2  In the preceding exercise we found that \( s_t \) converges to zero and \( \phi_t \) converges to about 0.6.

Since these results were calculated at a value of \( \beta \) close to one, let’s compare them to the best choice for an *infinitely* patient worker.

Intuitively, an infinitely patient worker would like to maximize steady state wages, which are a function of steady state capital.
You can take it as given—it’s certainly true—that the infinitely patient worker does not search in the long run (i.e., \( s_t = 0 \) for large \( t \)).

Thus, given \( \phi \), steady state capital is the positive fixed point \( x^*(\phi) \) of the map \( x \mapsto G(x, \phi) \).

Steady state wages can be written as \( w^*(\phi) = x^*(\phi)(1 - \phi) \).

Graph \( w^*(\phi) \) with respect to \( \phi \), and examine the best choice of \( \phi \).

Can you give a rough interpretation for the value that you see?
### Model features

- Infinite horizon dynamic programming with two states and one binary control
- Bayesian updating to learn the unknown distribution

### Model

Let’s first recall the basic McCall model [McC70] and then add the variation we want to consider

#### The Basic McCall Model

Consider an unemployed worker who is presented in each period with a permanent job offer at wage $w_t$

At time $t$, our worker has two choices

1. Accept the offer and work permanently at constant wage $w_t$
2. Reject the offer, receive unemployment compensation $c$, and reconsider next period

The wage sequence $\{w_t\}$ is iid and generated from known density $h$

The worker aims to maximize the expected discounted sum of earnings $E \sum_{t=0}^{\infty} \beta^t y_t$

Trade-off:

- Waiting too long for a good offer is costly, since the future is discounted
- Accepting too early is costly, since better offers will arrive with probability one

Let $V(w)$ denote the maximal expected discounted sum of earnings that can be obtained by an unemployed worker who starts with wage offer $w$ in hand

The function $V$ satisfies the recursion

$$V(w) = \max \left\{ \frac{w}{1-\beta}, c + \beta \int V(w') h(w') dw' \right\} \quad (3.35)$$

where the two terms on the r.h.s. are the respective payoffs from accepting and rejecting the current offer $w$

The optimal policy is a map from states into actions, and hence a binary function of $w$

Not surprisingly, it turns out to have the form $1\{w \geq \bar{w}\}$, where

- $\bar{w}$ is a constant depending on $(\beta, h, c)$ called the reservation wage
- $1\{w \geq \bar{w}\}$ is an indicator function returning 1 if $w \geq \bar{w}$ and 0 otherwise
- 1 indicates “accept” and 0 indicates “reject”

For further details see [LS12], section 6.3
Offer Distribution Unknown  Now let’s extend the model by considering the variation presented in [LS12], section 6.6

The model is as above, apart from the fact that

- the density $h$ is unknown
- the worker learns about $h$ by starting with a prior and updating based on wage offers that he/she observes

The worker knows there are two possible distributions $F$ and $G$ — with densities $f$ and $g$

At the start of time, “nature” selects $h$ to be either $f$ or $g$ — the wage distribution from which the entire sequence $\{w_t\}$ will be drawn

This choice is not observed by the worker, who puts prior probability $\pi_0$ on $f$ being chosen

Update rule: worker’s time $t$ estimate of the distribution is $\pi_t f + (1 - \pi_t)g$, where $\pi_t$ updates via

$$\pi_{t+1} = \frac{\pi_t f(w_{t+1})}{\pi_t f(w_{t+1}) + (1 - \pi_t)g(w_{t+1})}$$  \hspace{1cm} (3.36)

This last expression follows from Bayes’ rule, which tells us that

$$P\{h = f \mid W = w\} = \frac{P\{W = w \mid h = f\}P\{h = f\}}{P\{W = w\}} \quad \text{and} \quad P\{W = w\} = \sum_{\psi \in \{f, g\}} P\{W = w \mid h = \psi\} P\{h = \psi\}$$

The fact that (3.36) is recursive allows us to progress to a recursive solution method

Letting

$$h_\pi(w) := \pi f(w) + (1 - \pi)g(w) \quad \text{and} \quad q(w, \pi) := \frac{\pi f(w)}{\pi f(w) + (1 - \pi)g(w)}$$

we can express the value function for the unemployed worker recursively as follows

$$V(w, \pi) = \max \left\{ \frac{w}{1 - \beta} c + \beta \int V(w', \pi') h_\pi(w') \, dw' \right\} \text{ where } \pi' = q(w', \pi)$$  \hspace{1cm} (3.37)

Notice that the current guess $\pi$ is a state variable, since it affects the worker’s perception of probabilities for future rewards

Parameterization  Following section 6.6 of [LS12], our baseline parameterization will be

- $f = \text{Beta}(1, 1)$ and $g = \text{Beta}(3, 1.2)$
- $\beta = 0.95$ and $c = 0.6$

The densities $f$ and $g$ have the following shape

Looking Forward  What kind of optimal policy might result from (3.37) and the parameterization specified above?

Intuitively, if we accept at $w_a$ and $w_a \leq w_b$, then — all other things being given — we should also accept at $w_b$
This suggests a policy of accepting whenever \( w \) exceeds some threshold value \( \bar{w} \).

But \( \bar{w} \) should depend on \( \pi \) — in fact it should be decreasing in \( \pi \) because

- \( f \) is a less attractive offer distribution than \( g \)
- larger \( \pi \) means more weight on \( f \) and less on \( g \)

Thus larger \( \pi \) depresses the worker’s assessment of her future prospects, and relatively low current offers become more attractive.

**Summary:** We conjecture that the optimal policy is of the form \( 1 \{ w \geq \bar{w}(\pi) \} \) for some decreasing function \( \bar{w} \).

---

**Take 1: Solution by VFI**

Let’s set about solving the model and see how our results match with our intuition.

We begin by solving via value function iteration (VFI), which is natural but ultimately turns out to be second best.

VFI is implemented in the file `odu.jl` contained in the QuantEcon package.

The code is as follows:

```julia
#=
Solves the “Offer Distribution Unknown” Model by value function iteration and a second faster method discussed in the corresponding quantecon lecture.

@author: Spencer Lyon <spencer.lyon@nyu.edu>
```
3.5. SEARCH WITH OFFER DISTRIBUTION UNKNOWN

@date: 2014-08-14

References
---------

http://quant-econ.net/odu.html

=#

type SearchProblem
  bet::Real
  c::Real
  F::Distribution
  G::Distribution
  f::Function
  g::Function
  n_w::Int
  w_max::Real
  w_grid::Union(Vector, Range)
  n_pi::Int
  pi_min::Real
  pi_max::Real
  pi_grid::Union(Vector, Range)
end

function SearchProblem(bet=0.95, c=0.6, F_a=1, F_b=1, G_a=3, G_b=1.2, w_max=2, w_grid_size=40, pi_grid_size=40)
  F = Beta(F_a, F_b)
  G = Beta(G_a, G_b)

  # NOTE: the x./w_max)./w_max in these functions makes our dist match
  #      the scipy one with scale=w_max given
  f(x) = pdf(F, x./w_max)./w_max
  g(x) = pdf(G, x./w_max)./w_max

  pi_min = 1e-3  # avoids instability
  pi_max = 1 - pi_min

  w_grid = linspace_range(0, w_max, w_grid_size)
  pi_grid = linspace_range(pi_min, pi_max, pi_grid_size)

  SearchProblem(bet, c, F, G, f, g,
                w_grid_size, w_max, w_grid,
                pi_grid_size, pi_min, pi_max, pi_grid)
end

# make kwarg version
function SearchProblem(;bet=0.95, c=0.6, F_a=1, F_b=1, G_a=3, G_b=1.2, w_max=2, w_grid_size=40, pi_grid_size=40)
  SearchProblem(bet, c, F_a, F_b, G_a, G_b, w_max, w_grid_size, pi_grid_size)
function q(sp::SearchProblem, w, pi_val)
    new_pi = 1.0 ./ (1 + ((1 - pi_val).*sp.g(w))./(pi_val.*sp.f(w)))
    # Return new_pi when in [pi_min, pi_max] and else end points
    return clamp(new_pi, sp.pi_min, sp.pi_max)
end

function bellman_operator!(sp::SearchProblem, v::Matrix, out::Matrix;
    ret_policy::Bool=false)
    # Simplify names
    f, g, bet, c = sp.f, sp.g, sp.bet, sp.c
    vf = CoordInterpGrid((sp.w_grid, sp.pi_grid), v, BCnan, InterpLinear)
    # set up quadrature nodes/weights
    # q_nodes, q_weights = qnwlege(21, 0.0, sp.w_max)
    for w_i=1:sp.n_w
        w = sp.w_grid[w_i]
        # calculate v1
        v1 = w / (1 - bet)
        for pi_j=1:sp.n_pi
            _pi = sp.pi_grid[pi_j]
            # calculate v2
            integrand(m) = vf[m, q(sp, m, _pi)] * (_pi*f(m) + (1-_pi)*g(m))
            integral, error = quadgk(integrand, 0, sp.w_max)
            # integral = do_quad(integrand, q_nodes, q_weights)
            v2 = c + bet * integral
            # return policy if asked for, otherwise return max of values
            out[w_i, pi_j] = ret_policy ? v1 > v2 : max(v1, v2)
        end
    end
    return out
end

function bellman_operator(sp::SearchProblem, v::Matrix;
    ret_policy::Bool=false)
    out_type = ret_policy ? Bool : Float64
    out = Array(out_type, sp.n_w, sp.n_pi)
    bellman_operator!(sp, v, out, ret_policy=ret_policy)
end

function get_greedy!(sp::SearchProblem, v::Matrix, out::Matrix)
    bellman_operator!(sp, v, out, ret_policy=true)
end
get_greedy(sp::SearchProblem, v::Matrix) = bellman_operator(sp, v, ret_policy=true)

function res_wage_operator!(sp::SearchProblem, phi::Vector, out::Vector)
    # Simplify name
    f, g, bet, c = sp.f, sp.g, sp.bet, sp.c

    # Construct interpolator over pi_grid, given phi
    phi_f = CoordInterpGrid(sp.pi_grid, phi, BCnearest, InterpLinear)

    # set up quadrature nodes/weights
    q_nodes, q_weights = qnwlege(7, 0.0, sp.w_max)

    for (i, _pi) in enumerate(sp.pi_grid)
        integrand(x) = max(x, phi_f[q(sp, x, _pi)]) * (_pi*f(x) + (1-_pi)*g(x))
        integral = do_quad(integrand, q_nodes, q_weights)
        out[i] = (1 - bet)*c + bet*integral
    end
end

function res_wage_operator(sp::SearchProblem, phi::Vector)
    out = similar(phi)
    res_wage_operator!(sp, phi, out)
    return out
end

The type `SearchProblem` is used to store parameters and methods needed to compute optimal actions.

The Bellman operator is implemented as the method `bellman_operator()`, while `get_greedy()` computes an approximate optimal policy from a guess $v$ of the value function.

We will omit a detailed discussion of the code because there is a more efficient solution method.

These ideas are implemented in the `res_wage_operator` method.

Before explaining it let’s look quickly at solutions computed from value function iteration:

Here’s the value function:

The optimal policy:

Code for producing these figures can be found in file `examples/odu_vfi_plots.jl` from the main repository.

The code takes several minutes to run.

The results fit well with our intuition from section *Looking Forward*:

- The black line in the figure above corresponds to the function $ar{w}(\pi)$ introduced there
- decreasing as expected
3.5. SEARCH WITH OFFER DISTRIBUTION UNKNOWN

\begin{figure}
\centering
\includegraphics[width=\textwidth]{figure1.png}
\caption{Graph showing wage distribution and offer acceptance/rejection thresholds.}
\end{figure}
Take 2: A More Efficient Method

Our implementation of VFI can be optimized to some degree,
But instead of pursuing that, let’s consider another method to solve for the optimal policy
Uses iteration with an operator having the same contraction rate as the Bellman operator, but
• one dimensional rather than two dimensional
• no maximization step
As a consequence, the algorithm is orders of magnitude faster than VFI

This section illustrates the point that when it comes to programming, a bit of mathematical analysis goes a long way

Another Functional Equation
To begin, note that when \( w = \bar{w}(\pi) \), the worker is indifferent between accepting and rejecting
Hence the two choices on the right-hand side of (3.37) have equal value:
\[
\frac{\bar{w}(\pi)}{1 - \beta} = c + \beta \int V(w', \pi') h_{\pi}(w') \, dw'
\]
(3.38)
Together, (3.37) and (3.38) give
\[
V(w, \pi) = \max \left\{ \frac{w}{1 - \beta}, \frac{\bar{w}(\pi)}{1 - \beta} \right\}
\]
(3.39)
Combining (3.38) and (3.39), we obtain
\[
\frac{\bar{w}(\pi)}{1 - \beta} = c + \beta \int \max \left\{ \frac{w'}{1 - \beta'}, \frac{\bar{w}(\pi')}{1 - \beta'} \right\} h_{\pi}(w') \, dw'
\]
Multiplying by \( 1 - \beta \), substituting in \( \pi' = q(w', \pi) \) and using \( \circ \) for composition of functions yields
\[
\bar{w}(\pi) = (1 - \beta)c + \beta \int \max \{ w', \bar{w} \circ q(w', \pi) \} h_{\pi}(w') \, dw'
\]
(3.40)
Equation (3.40) can be understood as a functional equation, where \( \bar{w} \) is the unknown function
• Let’s call it the reservation wage functional equation (RWFE)
• The solution \( \bar{w} \) to the RWFE is the object that we wish to compute

Solving the RWFE
To solve the RWFE, we will first show that its solution is the fixed point of a contraction mapping
To this end, let
• \( b[0, 1] \) be the bounded real-valued functions on \([0, 1]\)
• \( \|\psi\| := \sup_{x \in [0,1]} |\psi(x)| \)
Consider the operator $Q$ mapping $\psi \in b[0,1]$ into $Q\psi \in b[0,1]$ via
\[(Q\psi)(\pi) = (1 - \beta)c + \beta \int \max \{ w', \psi \circ q(w', \pi) \} \ h_{\pi}(w') \ dw' \quad (3.41)\]

Comparing (3.40) and (3.41), we see that the set of fixed points of $Q$ exactly coincides with the set of solutions to the RWFE

- If $Q\bar{w} = \bar{w}$ then $\bar{w}$ solves (3.40) and vice versa

Moreover, for any $\psi, \phi \in b[0,1]$, basic algebra and the triangle inequality for integrals tells us that
\[|(Q\psi)(\pi) - (Q\phi)(\pi)| \leq \beta \int \max \{ w', \psi \circ q(w', \pi) \} - \max \{ w', \phi \circ q(w', \pi) \} \ h_{\pi}(w') \ dw' \quad (3.42)\]

Working case by case, it is easy to check that for real numbers $a, b, c$ we always have
\[|\max\{a, b\} - \max\{a, c\}| \leq |b - c| \quad (3.43)\]

Combining (3.42) and (3.43) yields
\[|(Q\psi)(\pi) - (Q\phi)(\pi)| \leq \beta \int |\psi \circ q(w', \pi) - \phi \circ q(w', \pi)| \ h_{\pi}(w') \ dw' \leq \beta \|\psi - \phi\| \quad (3.44)\]

Taking the supremum over $\pi$ now gives us
\[\|Q\psi - Q\phi\| \leq \beta \|\psi - \phi\| \quad (3.45)\]

In other words, $Q$ is a contraction of modulus $\beta$ on the complete metric space $(b[0,1], \| \cdot \|)$

Hence

- A unique solution $\bar{w}$ to the RWFE exists in $b[0,1]$
- $Q^k \psi \to \bar{w}$ uniformly as $k \to \infty$, for any $\psi \in b[0,1]$

**Implementation** These ideas are implemented in the `res_wage_operator` method from `odu.jl` as shown above

The method corresponds to action of the operator $Q$

The following exercise asks you to exploit these facts to compute an approximation to $\bar{w}$

**Exercises**

**Exercise 1** Use the default parameters and the `res_wage_operator` method to compute an optimal policy

Your result should coincide closely with the figure for the optimal policy *shown above*

Try experimenting with different parameters, and confirm that the change in the optimal policy coincides with your intuition
3.6 Optimal Savings

Overview

Next we study the standard optimal savings problem for an infinitely lived consumer—the “common ancestor” described in [LS12], section 1.3

- Also known as the income fluctuation problem
- An important sub-problem for many representative macroeconomic models
  - [Aiy94]
  - [Hug93]
  - etc.
- Useful references include [Dea91], [DH10], [Kuh13], [Rab02], [Rei09] and [SE77]

Our presentation of the model will be relatively brief

- For further details on economic intuition, implication and models, see [LS12]
- Proofs of all mathematical results stated below can be found in this paper

In this lecture we will explore an alternative to value function iteration (VFI) called policy function iteration (PFI)

- Based on the Euler equation, and not to be confused with Howard’s policy iteration algorithm
- Globally convergent under mild assumptions, even when utility is unbounded (both above and below)
- Numerically, turns out to be faster and more efficient than VFI for this model
Model features

- Infinite horizon dynamic programming with two states and one control

The Optimal Savings Problem

Consider a household that chooses a state-contingent consumption plan \( \{c_t\}_{t \geq 0} \) to maximize

\[
E \sum_{t=0}^{\infty} \beta^t u(c_t)
\]

subject to

\[
c_t + a_{t+1} \leq Ra_t + z_t, \quad c_t \geq 0, \quad a_t \geq -b \quad t = 0, 1, \ldots
\]  (3.46)

Here

- \( \beta \in (0, 1) \) is the discount factor
- \( a_t \) is asset holdings at time \( t \), with ad-hoc borrowing constraint \( a_t \geq -b \)
- \( c_t \) is consumption
- \( z_t \) is non-capital income (wages, unemployment compensation, etc.)
- \( R := 1 + r \), where \( r > 0 \) is the interest rate on savings

Assumptions

1. \( \{z_t\} \) is a finite Markov process with Markov matrix \( \Pi \) taking values in \( Z \)
2. \( |Z| < \infty \) and \( Z \subset (0, \infty) \)
3. \( r > 0 \) and \( \beta R < 1 \)
4. \( u \) is smooth, strictly increasing and strictly concave with \( \lim_{c \to 0} u'(c) = \infty \) and \( \lim_{c \to \infty} u'(c) = 0 \)

The asset space is \([-b, \infty)\) and the state is the pair \((a, z) \in S := [-b, \infty) \times Z\)

A feasible consumption path from \((a, z) \in S\) is a consumption sequence \( \{c_t\} \) such that \( \{c_t\} \) and its induced asset path \( \{a_t\} \) satisfy

1. \((a_0, z_0) = (a, z)\)
2. the feasibility constraints in (3.46), and
3. measurability of \( c_t \) w.r.t. the filtration generated by \( \{z_1, \ldots, z_t\} \)

The meaning of the third point is just that consumption at time \( t \) can only be a function of outcomes that have already been observed

The value function \( V : S \to \mathbb{R} \) is defined by

\[
V(a, z) := \sup \mathbb{E} \left\{ \sum_{t=0}^{\infty} \beta^t u(c_t) \right\}
\]  (3.47)

where the supremum is over all feasible consumption paths from \((a, z)\).
An optimal consumption path from \((a, z)\) is a feasible consumption path from \((a, z)\) that attains the supremum in (3.47).

Given our assumptions, it is known that

1. For each \((a, z) \in S\), a unique optimal consumption path from \((a, z)\) exists
2. This path is the unique feasible path from \((a, z)\) satisfying the Euler equality

\[
u'(c_t) = \max \left\{ \beta R \mathbb{E}_t[u'(c_{t+1})], \ u'(Ra_t + z_t + b) \right\}
\] (3.48)

and the transversality condition

\[
\lim_{t \to \infty} \beta^t \mathbb{E} [u'(c_t)a_{t+1}] = 0.
\] (3.49)

Moreover, there exists an optimal consumption function \(c^*: S \to [0, \infty)\) such that the path from \((a, z)\) generated by

\[
(a_0, z_0) = (a, z), \quad z_{t+1} \sim \Pi(z_t, dy), \quad c_t = c^*(a_t, z_t) \quad \text{and} \quad a_{t+1} = Ra_t + z_t - c_t
\]

satisfies both (3.48) and (3.49), and hence is the unique optimal path from \((a, z)\)

In summary, to solve the optimization problem, we need to compute \(c^*\)

**Computation**

There are two standard ways to solve for \(c^*\)

1. Value function iteration (VFI)
2. Policy function iteration (PFI) using the Euler equality

**Policy function iteration**

We can rewrite (3.48) to make it a statement about functions rather than random variables.

In particular, consider the functional equation

\[
u' \circ c (a, z) = \max \left\{ \gamma \int u' \circ c \left\{ Ra + z - c(a, z), \bar{z} \right\} \Pi(z, d\bar{z}), \ u'(Ra + z + b) \right\}
\] (3.50)

where \(\gamma := \beta R\) and \(u' \circ c(s) := u'(c(s))\)

Equation (3.50) is a functional equation in \(c\)

In order to identify a solution, let \(\mathcal{C}\) be the set of candidate consumption functions \(c: S \to \mathbb{R}\) such that

- each \(c \in \mathcal{C}\) is continuous and (weakly) increasing
- \(\min Z \leq c(a, z) \leq Ra + z + b\) for all \((a, z) \in S\)

In addition, let \(K: \mathcal{C} \to \mathcal{C}\) be defined as follows:

For given \(c \in \mathcal{C}\), the value \(Kc(a, z)\) is the unique \(t \in J(a, z)\) that solves

\[
u'(t) = \max \left\{ \gamma \int u' \circ c \left\{ Ra + z - t, \bar{z} \right\} \Pi(z, d\bar{z}), \ u'(Ra + z + b) \right\}
\] (3.51)
3.6. OPTIMAL SAVINGS

where

\[ J(a, z) := \{ t \in \mathbb{R} : \min Z \leq t \leq Ra + z + b \} \]  \hspace{1cm} (3.52)

We refer to \( K \) as Coleman’s policy function operator [Col90].

It is known that

- \( K \) is a contraction mapping on \( \mathcal{C} \) under the metric

\[ \rho(c, d) := \| u' \circ c - u' \circ d \| := \sup_{s \in S} | u'(c(s)) - u'(d(s)) | \quad (c, d \in \mathcal{C}) \]

- The metric \( \rho \) is complete on \( \mathcal{C} \)
- Convergence in \( \rho \) implies uniform convergence on compacts

In consequence, \( K \) has a unique fixed point \( c^* \in \mathcal{C} \) and \( K^n c \to c^* \) as \( n \to \infty \) for any \( c \in \mathcal{C} \).

By the definition of \( K \), the fixed points of \( K \) in \( \mathcal{C} \) coincide with the solutions to (3.50) in \( \mathcal{C} \).

In particular, it can be shown that the path \( \{ c_t \} \) generated from \( (a_0, z_0) \in S \) using policy function \( c^* \) is the unique optimal path from \( (a_0, z_0) \in S \).

**TL;DR** The unique optimal policy can be computed by picking any \( c \in \mathcal{C} \) and iterating with the operator \( K \) defined in (3.51).

**Value function iteration**

The Bellman operator for this problem is given by

\[ T\nu(a, z) = \max_{0 \leq c \leq Ra + z + b} \left\{ u(c) + \beta \int \nu(Ra + z - c, \hat{z}) \Pi(z, d\hat{z}) \right\} \]  \hspace{1cm} (3.53)

We have to be careful with VFI (i.e., iterating with \( T \)) in this setting because \( u \) is not assumed to be bounded

- In fact typically unbounded both above and below — e.g. \( u(c) = \log c \)
- In which case, the standard DP theory does not apply
- \( T^n \nu \) is not guaranteed to converge to the value function for arbitrary continuous bounded \( \nu \)

Nonetheless, we can always try the strategy “iterate and hope”

- In this case we can check the outcome by comparing with PFI
- The latter is known to converge, as described above

**Implementation** The code in `ifp.jl` from QuantEcon provides implementations of both VFI and PFI.

The code is repeated here and a description and clarifications are given below

```
# Tools for solving the standard optimal savings / income fluctuation problem for an infinitely lived consumer facing an exogenous income process that evolves according to a Markov chain.
```
3.6. OPTIMAL SAVINGS

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@date: 2014-08-18

References
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http://quant-econ.net/ifp.html

#= 
# using PyCall
# @pyimport scipy.optimize as opt
# brentq = opt.brentq

type ConsumerProblem
    u::Function
    du::Function
    r::Real
    R::Real
    bet::Real
    b::Real
    Pi::Matrix
    z_vals::Vector
    asset_grid::Union(Vector, Range)
end

default_du{T <: Real}(x::T) = 1.0 / x

function ConsumerProblem(r=0.01, bet=0.96, Pi=[0.6 0.4; 0.05 0.95],
z_vals=[0.5, 1.0], b=0.0, grid_max=16, grid_size=50,
u=log, du=default_du)
    R = 1 + r
    asset_grid = linspace_range(-b, grid_max, grid_size)
    ConsumerProblem(u, du, r, R, bet, b, Pi, z_vals, asset_grid)
end

# make kwarg version
function ConsumerProblem(;r=0.01, beta=0.96, Pi=[0.6 0.4; 0.05 0.95],
z_vals=[0.5, 1.0], b=0.0, grid_max=16, grid_size=50,
u=log, du=x -> 1./x)
    ConsumerProblem(r, beta, Pi, z_vals, b, grid_max, grid_size, u, du)
end

function bellman_operator!(cp::ConsumerProblem, V::Matrix, out::Matrix;
    ret_policy::Bool=false)
    # simplify names, set up arrays
    R, Pi, bet, u, b = cp.R, cp.Pi, cp.bet, cp.u, cp.b
    asset_grid, z_vals = cp.asset_grid, cp.z_vals

    new_V = similar(V)
    new_c = similar(V)

    bellman_operator!(cp::ConsumerProblem, V::Matrix, out::Matrix;
        ret_policy::Bool=false)
z_idx = 1:length(z_vals)

# Linear interpolation of V along the asset grid
vf(a, i_z) = CoordInterpGrid(asset_grid, V[:, i_z], BCnearest, InterpLinear)[a]

# compute lower bound for optimization
opt_lb = minimum(z_vals) - 1e-5

# solve for RHS of Bellman equation
for (i_z, z) in enumerate(z_vals)
    for (i_a, a) in enumerate(asset_grid)
        function obj(c)
            y = sum([vf(R*a+z-c, j) * Pi[i_z, j] for j=z_idx])
            return -u(c) - bet * y
        end
        res = optimize(obj, opt_lb, R.*a.+z.+b)
        c_star = res.minimum
        if ret_policy
            out[i_a, i_z] = c_star
        else
            out[i_a, i_z] = - obj(c_star)
        end
    end
end

function bellman_operator(cp::ConsumerProblem, V::Matrix; ret_policy=false)
    out = similar(V)
    bellman_operator!(cp, V, out, ret_policy=ret_policy)
    return out
end

function get_greedy!(cp::ConsumerProblem, V::Matrix, out::Matrix)
    bellman_operator!(cp, v, out, ret_policy=true)
end

function get_greedy(cp::ConsumerProblem, V::Matrix)
    bellman_operator(cp, V, ret_policy=true)
end

function coleman_operator!(cp::ConsumerProblem, c::Matrix, out::Matrix)
    # simplify names, set up arrays
    R, Pi, bet, du, b = cp.R, cp.Pi, cp.bet, cp.du, cp.b
    asset_grid, z_vals = cp.asset_grid, cp.z_vals
    z_size = length(z_vals)
    gam = R * bet
    vals = Array(Float64, z_size)

    T
# linear interpolation to get consumption function. Updates vals inplace

```javascript
function cf!(a, vals)
    for i=1:z_size
        vals[i] = CoordInterpGrid(asset_grid, c[:, i], BCnearest, InterpLinear)[a]
    end

    nothing
end
```

# compute lower_bound for optimization

```javascript
opt_lb = minimum(z_vals) - 1e-5
for (i_z, z) in enumerate(z_vals)
    for (i_a, a) in enumerate(asset_grid)
        function h(t)
            cf!(R*a+z-t, vals) # update vals
            expectation = dot(du(vals), Pi[i_z, :])
            return abs(du(t) - max(gam * expectation, du(R*a+z+b)))
        end

        res = optimize(h, opt_lb, R*a + z + b, method=:brent)
        out[i_a, i_z] = res.minimum
    end
end

return out
```

```javascript
function coleman_operator(cp::ConsumerProblem, c::Matrix)
    out = similar(c)
    coleman_operator!(cp, c, out)
    return out
end
```

```javascript
function init_values(cp::ConsumerProblem)
    # simplify names, set up arrays
    R, bet, u, b = cp.R, cp.bet, cp.u, cp.b
    asset_grid, z_vals = cp.asset_grid, cp.z_vals
    shape = length(asset_grid), length(z_vals)
    V, c = Array(Float64, shape...), Array(Float64, shape...)

    # Populate V and c
    for (i_z, z) in enumerate(z_vals)
        for (i_a, a) in enumerate(asset_grid)
            c_max = R*a + z + b
            c[i_a, i_z] = c_max
            V[i_a, i_z] = u(c_max) ./ (1 - bet)
        end
    end

    return V, c
end
```

The code contains a type called ConsumerProblem that
3.6. OPTIMAL SAVINGS

- stores all the relevant parameters of a given model
- defines methods
  - `bellman_operator`, which implements the Bellman operator \( T \) specified above
  - `coleman_operator`, which implements the Coleman operator \( K \) specified above
  - `initialize`, which generates suitable initial conditions for iteration

The methods `bellman_operator` and `coleman_operator` both use linear interpolation along the asset grid to approximate the value and consumption functions.

The following exercises walk you through several applications where policy functions are computed.

In exercise 1 you will see that while VFI and PFI produce similar results, the latter is much faster.
- Because we are exploiting analytically derived first order conditions.

Another benefit of working in policy function space rather than value function space is that value functions typically have more curvature.
- Makes them harder to approximate numerically.

**Exercises**

**Exercise 1**  The first exercise is to replicate the following figure, which compares PFI and VFI as solution methods.

The figure shows consumption policies computed by iteration of \( K \) and \( T \) respectively.
- In the case of iteration with \( T \), the final value function is used to compute the observed policy.
Consumption is shown as a function of assets with income $z$ held fixed at its smallest value. The following details are needed to replicate the figure:

- The parameters are the default parameters in the definition of `consumerProblem`.
- The initial conditions are the default ones from `initialize(cp)`.
- Both operators are iterated 80 times.

When you run your code you will observe that iteration with $K$ is faster than iteration with $T$. In the Julia console, a comparison of the operators can be made as follows:

```julia
julia> using QuantEcon

julia> cp = ConsumerProblem();

julia> v, c, = initialize(cp);

julia> @time bellman_operator(cp, v);
elapsed time: 0.095017748 seconds (24212168 bytes allocated, 30.48% gc time)

julia> @time coleman_operator(cp, c);
elapsed time: 0.0696242 seconds (23937576 bytes allocated)
```

**Exercise 2** Next let’s consider how the interest rate affects consumption.

Reproduce the following figure, which shows (approximately) optimal consumption policies for different interest rates:

- Other than $r$, all parameters are at their default values.
• $r$ steps through $\text{linspace}(0, 0.04, 4)$

• Consumption is plotted against assets for income shock fixed at the smallest value

The figure shows that higher interest rates boost savings and hence suppress consumption

**Exercise 3** Now let’s consider the long run asset levels held by households

We’ll take $r = 0.03$ and otherwise use default parameters

The following figure is a 45 degree diagram showing the law of motion for assets when consumption is optimal

The green line and blue line represent the function

$$a' = h(a, z) := Ra + z - c^*(a, z)$$

when income $z$ takes its high and low values respectively

The dashed line is the 45 degree line

We can see from the figure that the dynamics will be stable — assets do not diverge

In fact there is a unique stationary distribution of assets that we can calculate by simulation

• Can be proved via theorem 2 of [HP92]

• Represents the long run dispersion of assets across households when households have idiosyncratic shocks

Ergodicity is valid here, so stationary probabilities can be calculated by averaging over a single long time series
• Hence to approximate the stationary distribution we can simulate a long time series for assets and histogram, as in the following figure

Your task is to replicate the figure
• Parameters are as discussed above
• The histogram in the figure used a single time series \( \{a_t\} \) of length 500,000
• Given the length of this time series, the initial condition \((a_0, z_0)\) will not matter
• You might find it helpful to use the function `mc_sample_path` from `quantecon`

**Exercise 4** Following on from exercises 2 and 3, let’s look at how savings and aggregate asset holdings vary with the interest rate
• Note: [LS12] section 18.6 can be consulted for more background on the topic treated in this exercise

For a given parameterization of the model, the mean of the stationary distribution can be interpreted as aggregate capital in an economy with a unit mass of *ex-ante* identical households facing idiosyncratic shocks

Let’s look at how this measure of aggregate capital varies with the interest rate and borrowing constraint

The next figure plots aggregate capital against the interest rate for \( b \) in \((1, 3)\)

As is traditional, the price (interest rate) is on the vertical axis

The horizontal axis is aggregate capital computed as the mean of the stationary distribution

Exercise 4 is to replicate the figure, making use of code from previous exercises
Try to explain why the measure of aggregate capital is equal to $-b$ when $r = 0$ for both cases shown here.

**Solutions**

**Solution notebook**

### 3.7 Robustness

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**Overview**

This lecture modifies a Bellman equation to express a decision maker’s doubts about transition dynamics.

His specification doubts make the decision maker want a robust decision rule.
Robust means insensitive to misspecification of transition dynamics

The decision maker has a single approximating model

He calls it approximating to acknowledge that he doesn’t completely trust it

He fears that outcomes will actually be determined by another model that he cannot describe explicitly

All that he knows is that the actual data-generating model is in some (uncountable) set of models that surrounds his approximating model

He quantifies the discrepancy between his approximating model and the genuine data-generating model by using a quantity called entropy

(We’ll explain what entropy means below)

He wants a decision rule that will work well enough no matter which of those other models actually governs outcomes

This is what it means for his decision rule to be “robust to misspecification of an approximating model”

This may sound like too much to ask for, but . . .

. . . a secret weapon is available to design robust decision rules

The secret weapon is max-min control theory

A value-maximizing decision maker enlists the aid of an (imaginary) value-minimizing model chooser to construct bounds on the value attained by a given decision rule under different models of the transition dynamics

The original decision maker uses those bounds to construct a decision rule with an assured performance level, no matter which model actually governs outcomes

Note: In reading this lecture, please don’t think that our decision maker is paranoid when he conducts a worst-case analysis. By designing a rule that works well against a worst-case, his intention is to construct a rule that will work well across a set of models.

Sets of Models Imply Sets Of Values  Our “robust” decision maker wants to know how well a given rule will work when he does not know a single transition law . . .

. . . he wants to know sets of values that will be attained by a given decision rule \( F \) under a set of transition laws

Ultimately, he wants to design a decision rule \( F \) that shapes these sets of values in ways that he prefers

With this in mind, consider the following graph, which relates to a particular decision problem to be explained below

The figure shows a value-entropy correspondence for a particular decision rule \( F \)
The shaded set is the graph of the correspondence, which maps entropy to a set of values associated with a set of models that surround the decision maker’s approximating model.

Here

- **Value** refers to a sum of discounted rewards obtained by applying the decision rule $F$ when the state starts at some fixed initial state $x_0$.

- **Entropy** is a nonnegative number that measures the size of a set of models surrounding the decision maker’s approximating model.
  - Entropy is zero when the set includes only the approximating model, indicating that the decision maker completely trusts the approximating model.
  - Entropy is bigger, and the set of surrounding models is bigger, the less the decision maker trusts the approximating model.

The shaded region indicates that for all models having entropy less than or equal to the number on the horizontal axis, the value obtained will be somewhere within the indicated set of values.

Now let’s compare sets of values associated with two different decision rules, $F_r$ and $F_b$.

In the next figure,

- The red set shows the value-entropy correspondence for decision rule $F_r$.
- The blue set shows the value-entropy correspondence for decision rule $F_b$.

The blue correspondence is skinnier than the red correspondence.

This conveys the sense in which the decision rule $F_b$ is *more robust* than the decision rule $F_r$.

- *More robust* means that the set of values is less sensitive to *increasing misspecification* as measured by entropy.
Notice that the less robust rule $F_r$ promises higher values for small misspecifications (small entropy).

(But it is more fragile in the sense that it is more sensitive to perturbations of the approximating model.)

Below we’ll explain in detail how to construct these sets of values for a given $F$, but for now . . .

Here is a hint about the secret weapons we’ll use to construct these sets:

- We’ll use some min problems to construct the lower bounds
- We’ll use some max problems to construct the upper bounds

We will also describe how to choose $F$ to shape the sets of values.

This will involve crafting a skinnier set at the cost of a lower level (at least for low values of entropy)

**Inspiring Video** If you want to understand more about why one serious quantitative researcher is interested in this approach, we recommend [Lars Peter Hansen’s Nobel lecture](#).

**Other References** Our discussion in this lecture is based on

- [HS00]
- [HS08]
The Model

For simplicity, we present ideas in the context of a class of problems with linear transition laws and quadratic objective functions.

To fit in with our earlier lecture on LQ control, we will treat loss minimization rather than value maximization.

To begin, recall the infinite horizon LQ problem, where an agent chooses a sequence of controls \( \{u_t\} \) to minimize

\[
\sum_{t=0}^\infty \beta^t \{x_t^\prime R x_t + u_t^\prime Q u_t\}
\]  

(3.54)

subject to the linear law of motion

\[
x_{t+1} = Ax_t + Bu_t + Cw_{t+1}, \quad t = 0, 1, 2, \ldots
\]  

(3.55)

As before,

- \( x_t \) is \( n \times 1 \), \( A \) is \( n \times n \)
- \( u_t \) is \( k \times 1 \), \( B \) is \( n \times k \)
- \( w_t \) is \( j \times 1 \), \( C \) is \( n \times j \)
- \( R \) is \( n \times n \) and \( Q \) is \( k \times k \)

Here \( x_t \) is the state, \( u_t \) is the control, and \( w_t \) is a shock vector.

For now we take \( \{w_t\} := \{w_t\}_{t=1}^\infty \) to be deterministic — a single fixed sequence.

We also allow for model uncertainty on the part of the agent solving this optimization problem.

In particular, the agent takes \( w_t = 0 \) for all \( t \geq 0 \) as a benchmark model, but admits the possibility that this model might be wrong.

As a consequence, she also considers a set of alternative models expressed in terms of sequences \( \{w_t\} \) that are “close” to the zero sequence.

She seeks a policy that will do well enough for a set of alternative models whose members are pinned down by sequences \( \{w_t\} \).

Soon we’ll quantify the quality of a model specification in terms of the maximal size of the expression \( \sum_{t=0}^\infty \beta^{t+1} w_{t+1}^\prime w_{t+1} \).

Constructing More Robust Policies

If our agent takes \( \{w_t\} \) as a given deterministic sequence, then, drawing on intuition from earlier lectures on dynamic programming, we can anticipate Bellman equations such as

\[
J_{t-1}(x) = \min_u \{x^\prime R x + u^\prime Q u + \beta J_t(Ax + Bu + Cw_t)\}
\]

(Here \( J \) depends on \( t \) because the sequence \( \{w_t\} \) is not recursive).
Our tool for studying robustness is to construct a rule that works well even if an adverse sequence \{w_t\} occurs.

In our framework, “adverse” means “loss increasing.”

As we’ll see, this will eventually lead us to construct the Bellman equation

\[
J(x) = \min_u \max_w \{ x'Rx + u'Qu + \beta [ J(Ax + Bu + Cw) - \theta w'w ] \} \tag{3.56}
\]

Notice that we’ve added the penalty term \(-\theta w'w\)

Since \(w'w = \|w\|^2\), this term becomes influential when \(w\) moves away from the origin.

The penalty parameter \(\theta\) controls how much we penalize the maximizing agent for “harming” the minimizing agent.

By raising \(\theta\) more and more, we more and more limit the ability of maximizing agent to distort outcomes relative to the approximating model.

So bigger \(\theta\) is implicitly associated with smaller distortion sequences \{w_t\}

**Analyzing the Bellman equation**  
So what does \(J\) in (3.56) look like?

As with the ordinary LQ control model, \(J\) takes the form \(J(x) = x'Px\) for some symmetric positive definite matrix \(P\).

One of our main tasks will be to analyze and compute the matrix \(P\).

Related tasks will be to study associated feedback rules for \(u_t\) and \(w_{t+1}\).

First, using matrix calculus, you will be able to verify that

\[
\max_w \{(Ax + Bu + Cw)'P(Ax + Bu + Cw) - \theta w'w\} = (Ax + Bu)'D(P)(Ax + Bu) \tag{3.57}
\]

where

\[
D(P) := P + PC(\theta I - C'PC)^{-1}C'P \tag{3.58}
\]

and \(I\) is a \(j \times j\) identity matrix. Substituting this expression for the maximum into (3.56) yields

\[
x'Px = \min_u \{ x'Rx + u'Qu + \beta (Ax + Bu)'D(P)(Ax + Bu) \} \tag{3.59}
\]

Using similar mathematics, the solution to this minimization problem is \(u = -Fx\) where \(F := (Q + \beta B'D(P)B)^{-1}\beta B'D(P)A\).

Substituting this minimizer back into (3.59) and working through the algebra gives \(x'Px = x'B(D(P))x\) for all \(x\), or, equivalently,

\[
P = B(D(P))
\]

where \(D\) is the operator defined in (3.58) and

\[
B(P) := R - \beta^2 A'PB(Q + \beta B'PB)^{-1}B'PA + \beta A'PA
\]
The operator $B$ is the standard (i.e., non-robust) LQ Bellman operator, and $P = B(P)$ is the standard matrix Riccati equation coming from the Bellman equation—see this discussion. Under some regularity conditions (see [HS08]), the operator $B \circ D$ has a unique positive definite fixed point, which we denote below by $\hat{P}$.

A robust policy, indexed by $\theta$, is $u = -\hat{F}x$ where

$$\hat{F} := (Q + \beta B' D(\hat{P}) B)^{-1} \beta B' D(\hat{P}) A$$

(3.60)

We also define

$$\hat{K} := (\theta I - C' \hat{P} C)^{-1} C' \hat{P} (A - B \hat{F})$$

(3.61)

The interpretation of $\hat{K}$ is that $w_{t+1} = \hat{K}x_t$ on the worst-case path of $\{x_t\}$, in the sense that this vector is the maximizer of (3.57) evaluated at the fixed rule $u = -\hat{F}x$.

Note that $\hat{P}$, $\hat{F}$, $\hat{K}$ are all determined by the primitives and $\theta$.

Note also that if $\theta$ is very large, then $D$ is approximately equal to the identity mapping. Hence, when $\theta$ is large, $\hat{P}$ and $\hat{F}$ are approximately equal to their standard LQ values.

Furthermore, when $\theta$ is large, $\hat{K}$ is approximately equal to zero. Conversely, smaller $\theta$ is associated with greater fear of model misspecification, and greater concern for robustness.

**Robustness as Outcome of a Two-Person Zero-Sum Game**

What we have done above can be interpreted in terms of a two-person zero-sum game in which $\hat{F}$, $\hat{K}$ are Nash equilibrium objects.

Agent 1 is our original agent, who seeks to minimize loss in the LQ program while admitting the possibility of misspecification.

Agent 2 is an imaginary malevolent player.

Agent 2’s malevolence helps the original agent to compute bounds on his value function across a set of models.

We begin with agent 2’s problem.

**Agent 2’s Problem** Agent 2

1. knows a fixed policy $F$ specifying the behavior of agent 1, in the sense that $u_t = -Fx_t$ for all $t$

2. responds by choosing a shock sequence $\{w_t\}$ from a set of paths sufficiently close to the benchmark sequence $\{0, 0, 0, \ldots\}$.

A natural way to say “sufficiently close to the zero sequence” is to restrict the summed inner product $\sum_{t=1}^{\infty} \bar{w}_t' w_t$ to be small.
However, to obtain a time-invariant recursive formulation, it turns out to be convenient to restrict
a discounted inner product
\[ \sum_{t=1}^{\infty} \beta^t w_t w_t' \leq \eta \] (3.62)
Now let \( F \) be a fixed policy, and let \( J_F(x_0, w) \) be the present-value cost of that policy given sequence
\( w := \{w_t\} \) and initial condition \( x_0 \in \mathbb{R}^n \)
Substituting \(-Fx_t\) for \( u_t\) in (3.54), this value can be written as
\[ J_F(x_0, w) := \sum_{t=0}^{\infty} \beta^t x_t' (R + F'QF)x_t \] (3.63)
where
\[ x_{t+1} = (A - BF)x_t + Cw_{t+1} \] (3.64)
and the initial condition \( x_0 \) is as specified in the left side of (3.63)
Agent 2 chooses \( w \) to maximize agent 1’s loss \( J_F(x_0, w) \) subject to (3.62)
Using a Lagrangian formulation, we can express this problem as
\[ \max_w \sum_{t=0}^{\infty} \beta^t \left\{ x_t' (R + F'QF)x_t - \beta \theta (w_{t+1}'w_{t+1} - \eta) \right\} \]
where \( \{x_t\} \) satisfied (3.64) and \( \theta \) is a Lagrange multiplier on constraint (3.62)
For the moment, let’s take \( \theta \) as fixed, allowing us to drop the constant \( \beta \theta \eta \) term in the objective
function, and hence write the problem as
\[ \max_w \sum_{t=0}^{\infty} \beta^t \left\{ x_t' (R + F'QF)x_t - \beta \theta w_{t+1}'w_{t+1} \right\} \]

or, equivalently,
\[ \min_w \sum_{t=0}^{\infty} \beta^t \left\{ -x_t' (R + F'QF)x_t + \beta \theta w_{t+1}'w_{t+1} \right\} \] (3.65)
subject to (3.64)

What’s striking about this optimization problem is that it is once again an LQ discounted dynamic
programming problem, with \( w = \{w_t\} \) as the sequence of controls
The expression for the optimal policy can be found by applying the usual LQ formula (see here)
We denote it by \( K(F, \theta) \), with the interpretation \( w_{t+1} = K(F, \theta)x_t \)
The remaining step for agent 2’s problem is to set \( \theta \) to enforce the constraint (3.62), which can be
done by choosing \( \theta = \theta_\eta \) such that
\[ \beta \sum_{t=0}^{\infty} \beta^t x_t K(F, \theta_\eta)'K(F, \theta_\eta)x_t = \eta \] (3.66)
Here \( x_t \) is given by (3.64) — which in this case becomes \( x_{t+1} = (A - BF + CK(F, \theta))x_t \)
Using Agent 2’s Problem to Construct Bounds on the Value Sets

The Lower Bound Define the minimized object on the right side of problem (3.65) as \( R_\theta(x_0, F) \).
Because “minimizers minimize” we have
\[
R_\theta(x_0, F) \leq \sum_{t=0}^{\infty} \beta^t \left\{ -x_t'(R + F'QF)x_t \right\} + \beta \sum_{t=0}^{\infty} \beta^t w_{t+1}^{t+1},
\]
where \( x_{t+1} = (A - BF + CK(F, \theta))x_t \) and \( x_0 \) is a given initial condition.
This inequality in turn implies the inequality
\[
R_\theta(x_0, F) - \theta \text{ ent} \leq \sum_{t=0}^{\infty} \beta^t \left\{ -x_t'(R + F'QF)x_t \right\}
\]
where
\[
\text{ent} := \beta \sum_{t=0}^{\infty} \beta^t w_{t+1}^{t+1}
\]
The left side of inequality (3.67) is a straight line with slope \(-\theta\)
Technically, it is a “separating hyperplane”
At a particular value of entropy, the line is tangent to the lower bound of values as a function of entropy
In particular, the lower bound on the left side of (3.67) is attained when
\[
\text{ent} = \beta \sum_{t=0}^{\infty} \beta^t x_t'K(F, \theta)'K(F, \theta)x_t
\]
To construct the lower bound on the set of values associated with all perturbations \( w \) satisfying the entropy constraint (3.62) at a given entropy level, we proceed as follows:

- For a given \( \theta \), solve the minimization problem (3.65)
- Compute the minimizer \( R_\theta(x_0, F) \) and the associated entropy using (3.68)
- Compute the lower bound on the value function \( R_\theta(x_0, F) - \theta \text{ ent} \) and plot it against ent
- Repeat the preceding three steps for a range of values of \( \theta \) to trace out the lower bound

Note: This procedure sweeps out a set of separating hyperplanes indexed by different values for the Lagrange multiplier \( \theta \)

The Upper Bound To construct an upper bound we use a very similar procedure
We simply replace the minimization problem (3.65) with the maximization problem
\[
V_\theta(x_0, F) = \max_w \sum_{t=0}^{\infty} \beta^t \left\{ -x_t'(R + F'QF)x_t - \beta \theta w_{t+1}^{t+1} \right\}
\]
where now \( \hat{\theta} > 0 \) penalizes the choice of \( w \) with larger entropy.

(Notice that \( \hat{\theta} = -\theta \) in problem (3.65))

Because “maximizers maximize” we have

\[
V_{\hat{\theta}}(x_0, F) \geq \sum_{t=0}^{\infty} \beta^t \left\{ -x'_t(R + F'QF)x_t \right\} - \hat{\theta} \sum_{t=0}^{\infty} \beta^t w'_{t+1} w_{t+1}
\]

which in turn implies the inequality

\[
V_{\hat{\theta}}(x_0, F) + \hat{\theta} \text{ent} \geq \sum_{t=0}^{\infty} \beta^t \left\{ -x'_t(R + F'QF)x_t \right\}
\]

(3.70)

where

\[
\text{ent} \equiv \beta \sum_{t=0}^{\infty} \beta^t w'_{t+1} w_{t+1}
\]

The left side of inequality (3.70) is a straight line with slope \( \hat{\theta} \)

The upper bound on the left side of (3.70) is attained when

\[
\text{ent} = \beta \sum_{t=0}^{\infty} \beta^t x'_t K(F, \hat{\theta})' K(F, \hat{\theta}) x_t
\]

(3.71)

To construct the upper bound on the set of values associated all perturbations \( w \) with a given entropy we proceed much as we did for the lower bound

- For a given \( \hat{\theta} \), solve the maximization problem (3.69)
- Compute the maximizer \( V_{\hat{\theta}}(x_0, F) \) and the associated entropy using (3.71)
- Compute the upper bound on the value function \( V_{\hat{\theta}}(x_0, F) + \hat{\theta} \text{ent} \) and plot it against ent
- Repeat the preceding three steps for a range of values of \( \hat{\theta} \) to trace out the upper bound

**Reshaping the set of values**  Now in the interest of *reshaping* these sets of values by choosing \( F \), we turn to agent 1’s problem

**Agent 1’s Problem**  Now we turn to agent 1, who solves

\[
\min_{\{u_t\}} \sum_{t=0}^{\infty} \beta^t \left\{ x'_t R x_t + u'_t Q u_t - \beta \theta w'_{t+1} w_{t+1} \right\}
\]

(3.72)

where \( \{w_{t+1}\} \) satisfies \( w_{t+1} = Kx_t \)

In other words, agent 1 minimizes

\[
\sum_{t=0}^{\infty} \beta^t \left\{ x'_t (R - \beta \theta K'K)x_t + u'_t Q u_t \right\}
\]

(3.73)

subject to

\[
x_{t+1} = (A + CK)x_t + Bu_t
\]

(3.74)

Once again, the expression for the optimal policy can be found here — we denote it by \( \hat{F} \)
Nash Equilibrium Clearly the \( \tilde{F} \) we have obtained depends on \( K \), which, in agent 2’s problem, depended on an initial policy \( F \). Holding all other parameters fixed, we can represent this relationship as a mapping \( \Phi \), where

\[
\tilde{F} = \Phi(K(F, \theta))
\]

The map \( F \mapsto \Phi(K(F, \theta)) \) corresponds to a situation in which

1. agent 1 uses an arbitrary initial policy \( F \)
2. agent 2 best responds to agent 1 by choosing \( K(F, \theta) \)
3. agent 1 best responds to agent 2 by choosing \( \tilde{F} = \Phi(K(F, \theta)) \)

As you may have already guessed, the robust policy \( \hat{F} \) defined in (3.60) is a fixed point of the mapping \( \Phi \).

In particular, for any given \( \theta \),

1. \( K(\hat{F}, \theta) = \hat{K} \), where \( \hat{K} \) is as given in (3.61)
2. \( \Phi(\hat{K}) = \hat{F} \)

A sketch of the proof is given in the appendix.

The Stochastic Case

Now we turn to the stochastic case, where the sequence \( \{w_t\} \) is treated as an iid sequence of random vectors.

In this setting, we suppose that our agent is uncertain about the conditional probability distribution of \( w_{t+1} \).

The agent takes the standard normal distribution \( N(0, I) \) as the baseline conditional distribution, while admitting the possibility that other “nearby” distributions prevail.

These alternative conditional distributions of \( w_{t+1} \) might depend nonlinearly on the history \( x_s, s \leq t \).

To implement this idea, we need a notion of what it means for one distribution to be near another one.

Here we adopt a very useful measure of closeness for distributions known as the relative entropy, or Kullback-Leibler divergence.

For densities \( p, q \), the Kullback-Leibler divergence of \( q \) from \( p \) is defined as

\[
D_{KL}(p, q) := \int \ln \left[ \frac{p(x)}{q(x)} \right] p(x) \, dx
\]

Using this notation, we replace (3.56) with the stochastic analogue

\[
J(x) = \min_u \max_{\psi \in \mathcal{P}} \left\{ x'Rx + u'Qu + \beta \left[ \int J(Ax + Bu + C \psi) \, d\psi - \theta D_{KL}(\psi, \phi) \right] \right\}
\]  (3.75)
Here \( \mathcal{P} \) represents the set of all densities on \( \mathbb{R}^n \) and \( \phi \) is the benchmark distribution \( N(0, I) \).

The distribution \( \phi \) is chosen as the least desirable conditional distribution in terms of next period outcomes, while taking into account the penalty term \( \theta D_{KL}(\psi, \phi) \).

This penalty term plays a role analogous to the one played by the deterministic penalty \( \theta w'w \) in (3.56), since it discourages large deviations from the benchmark.

**Solving the Model**
The maximization problem in (3.75) appears highly nontrivial — after all, we are maximizing over an infinite dimensional space consisting of the entire set of densities. However, it turns out that the solution is tractable, and in fact also falls within the class of normal distributions.

First, we note that \( J \) has the form \( J(x) = x'Px + d \) for some positive definite matrix \( P \) and constant real number \( d \).

Moreover, it turns out that if \( (I - \theta^{-1}C'PC)^{-1} \) is nonsingular, then

\[
\max_{\psi \in \mathcal{P}} \left\{ \int (Ax + Bu + Cw)'P(Ax + Bu + Cw) \psi(dw) - \theta D_{KL}(\psi, \phi) \right\} = (Ax + Bu)'D(P)(Ax + Bu) + \kappa(\theta, P) \tag{3.76}
\]

where 
\[
\kappa(\theta, P) := \theta \ln[\det(I - \theta^{-1}C'PC)^{-1}]
\]

and the maximizer is the Gaussian distribution

\[
\psi = N \left( (\theta I - C'PC)^{-1}C'P(Ax + Bu), (I - \theta^{-1}C'PC)^{-1} \right) \tag{3.77}
\]

Substituting the expression for the maximum into Bellman equation (3.75) and using \( J(x) = x'Px + d \) gives

\[
x'Px + d = \min_u \{ x'Rx + u'Qu + \beta (Ax + Bu)'D(P)(Ax + Bu) + \beta [d + \kappa(\theta, P)] \} \tag{3.78}
\]

Since constant terms do not affect minimizers, the solution is the same as (3.59), leading to 

\[
x'Px + d = x'B(D(P))x + \beta [d + \kappa(\theta, P)]
\]

To solve this Bellman equation, we take \( \hat{P} \) to be the positive definite fixed point of \( B \circ D \).

In addition, we take \( \hat{d} \) as the real number solving \( d = \beta [d + \kappa(\theta, P)] \), which is 

\[
\hat{d} := \frac{\beta}{1 - \beta} \kappa(\theta, P) \tag{3.79}
\]

The robust policy in this stochastic case is the minimizer in (3.78), which is once again \( u = -\hat{F}x \) for \( \hat{F} \) given by (3.60).

Substituting the robust policy into (3.77) we obtain the worst case shock distribution:

\[
w_{t+1} \sim N(\hat{K}x_t, (I - \theta^{-1}C'\hat{P}C)^{-1})
\]

where \( \hat{K} \) is given by (3.61).

Note that the mean of the worst-case shock distribution is equal to the same worst-case \( w_{t+1} \) as in the earlier deterministic setting.
Computing Other Quantities  Before turning to implementation, we briefly outline how to compute several other quantities of interest

Worst-Case Value of a Policy  One thing we will be interested in doing is holding a policy fixed and computing the discounted loss associated with that policy

So let $F$ be a given policy and let $J_F(x)$ be the associated loss, which, by analogy with (3.75), satisfies

$$J_F(x) = \max_{\psi \in \mathcal{P}} \left\{ x' (R + F'QF)x + \beta \left[ \int J_F((A - BF)x + Cw) \psi(dw) - \theta D_{KL}(\psi, \phi) \right] \right\}$$

Writing $J_F(x) = x'P_Fx + d_F$ and applying the same argument used to derive (3.76) we get

$$x'P_Fx + d_F = x'(R + F'QF)x + \beta \left[ x'(A - BF)'D(P_F)(A - BF)x + d_F + \kappa(\theta, P_F) \right]$$

To solve this we take $P_F$ to be the fixed point

$$P_F = R + F'QF + \beta(A - BF)'D(P_F)(A - BF)$$

and

$$d_F := \frac{\beta}{1 - \beta} \kappa(\theta, P_F) = \frac{\beta}{1 - \beta} \theta \ln[\det(I - \theta^{-1}C'P_F C)^{-1}] \quad (3.80)$$

If you skip ahead to the appendix, you will be able to verify that $-P_F$ is the solution to the Bellman equation in agent 2’s problem discussed above — we use this in our computations

Implementation

The QuantEcon package provides a type called RBLQ for implementation of robust LQ optimal control

Here’s the relevant code, from file robustlq.jl

```
#=
Provides a type called RBLQ for solving robust linear quadratic control problems.

@author : Spencer Lyon <spencer.lyon@nyu.edu>

@date : 2014-08-19

References
-----

Simple port of the file quantecon.robstulq

http://quant-econ.net/robustness.html
#

type RBLQ
   A::Matrix
```

---

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B::Matrix
C::Matrix
Q::Matrix
R::Matrix
k::Int
n::Int
j::Int
bet::Real
theta::Real

end

function RBLQ(Q::ScalarOrArray, R::ScalarOrArray, A::ScalarOrArray,
               B::ScalarOrArray, C::ScalarOrArray, bet::Real, theta::Real)
    k = size(Q, 1)
n = size(R, 1)
j = size(C, 2)
    # coerce sizes
    A = reshape([A], n, n)
    B = reshape([B], n, k)
    C = reshape([C], n, j)
    R = reshape([R], n, n)
    Q = reshape([Q], k, k)
    RBLQ(A, B, C, Q, R, k, n, j, bet, theta)
end

function d_operator(rlq::RBLQ, P::Matrix)
    C, theta, I = rlq.C, rlq.theta, eye(rlq.j)
    P + P*C*((theta.*I - C'*P*C) \ (C'*P))
end

function b_operator(rlq::RBLQ, P::Matrix)
    F = (Q+bet.*B'*P*B) \ (bet.*B'*P*A)
    bP = R - bet.*A'*P*B * F + bet.*A'*P*A
    F, bP
end

function robust_rule(rlq::RBLQ)
    bet, theta, k, j = rlq.bet, rlq.theta, rlq.k, rlq.j
    I = eye(j)
    Z = zeros(k, j)
    Ba = [B C]
    Qa = [Q Z
           Z' -bet.*I.*theta]
    lq = LQ(Qa, R, A, Ba, bet=bet)
    # Solve and convert back to robust problem
3.7. ROBUSTNESS

P, f, d = stationary_values(lq)
F = f[1:k, :]
K = -f[k+1:end, :]

return F, K, P
end

function robust_rule_simple(rlq::RBLQ,
P::Matrix=zeros(Float64, rlq.n, rlq.n);
max_iter=80,
tol=1e-8)
    # Simplify notation
    bet, theta, k, j = rlq.bet, rlq.theta, rlq.k, rlq.j
    iterate, e = 0, tol + 1.0
    F = similar(P)  # instantiate so available after loop
    while iterate <= max_iter && e > tol
        F, new_P = b_operator(rlq, d_operator(rlq, P))
        e = sqrt(sum((new_P - P).^2))
        iterate += 1
        P = new_P
    end
    if iterate >= max_iter
        warn("Maximum iterations in robust_rul_simple")
    end
    I = eye(j)
    K = (theta.*I - C'*P*C)\(C'*P)*(A - B*F)
    return F, K, P
end

function F_to_K(rlq::RBLQ, F::Matrix)
    # simplify notation
    bet, theta = rlq.bet, rlq.theta

    # set up lq
    Q2 = bet * theta
    R2 = - R - F'*Q*F
    A2 = A - B*F
    B2 = C
    lq = LQ(Q2, R2, A2, B2, bet=bet)

    neg_P, neg_K, d = stationary_values(lq)
    return -neg_K, -neg_P
end
```plaintext
function K_to_F(rlq::RBLQ, K::Matrix)
    bet, theta = rlq.bet, rlq.theta

    A1, B1, Q1, R1 = A+C*K, B, Q, R-bet*theta.*K
    lq = LQ(Q1, R1, A1, B1, bet=bet)

    P, F, d = stationary_values(lq)

    return F, P
end

function compute_deterministic_entropy(rlq::RBLQ, F, K, x0)
    B, C, bet = rlq.B, rlq.C, rlq.bet
    H0 = K'*K
    C0 = zeros(Float64, rlq.n, 1)
    A0 = A - B*F + C*K

    return var_quadratic_sum(A0, C0, H0, bet, x0)
end

function evaluate_F(rlq::RBLQ, F::Matrix)
    bet, theta, j = rlq.bet, rlq.theta, rlq.j

    # Solve for policies and costs using agent 2's problem
    K_F, P_F = F_to_K(rlq, F)
    I = eye(j)
    H = inv(I - C'*P_F*C./theta)
    d_F = log(det(H))

    # compute O_F and o_F
    sig = -1.0 / theta
    A0 = sqrt(bet) .* (A - B*F + C*K_F)
    O_F = solve_discrete_lyapunov(A0', bet*K_F'*K_F)
    ho = (trace(H - 1) - d_F) / 2.0
    tr = trace(O_F*C*H*C')
    o_F = (ho + bet*tr) / (1 - bet)

    return K_F, P_F, d_F, O_F, o_F
end
```

Here is a brief description of the methods of the type

- `d_operator()` and `b_operator()` implement $D$ and $B$ respectively
- `robust_rule()` and `robust_rule_simple()` both solve for the triple $\hat{F}, \hat{K}, \hat{P}$, as described in equations (3.60) – (3.61) and the surrounding discussion
  - `robust_rule()` is more efficient
  - `robust_rule_simple()` is more transparent and easier to follow
• \texttt{K_to_F()} and \texttt{F_to_K()} solve the decision problems of agent 1 and agent 2 respectively
• \texttt{compute_deterministic_entropy()} computes the left-hand side of (3.66)
• \texttt{evaluate_F()} computes the loss and entropy associated with a given policy — see this discussion

**Application**

Let us consider a monopolist similar to this one, but now facing model uncertainty

The inverse demand function is

\[ p_t = a_0 - a_1 y_t + d_t \]

where

\[ d_{t+1} = \rho d_t + \sigma_d w_{t+1}, \quad \{w_t\} \overset{iid}{\sim} N(0,1) \]

and all parameters are strictly positive

The period return function for the monopolist is

\[ r_t = p_t y_t - \gamma \frac{(y_{t+1} - y_t)^2}{2} - c y_t \]

Its objective is to maximize expected discounted profits, or, equivalently, to minimize

\[ \mathbb{E} \sum_{t=0}^{\infty} \beta^t (-r_t) \]

To form a linear regulator problem, we take the state and control to be

\[ x_t = \begin{bmatrix} 1 \\ y_t \\ d_t \end{bmatrix} \quad \text{and} \quad u_t = y_{t+1} - y_t \]

Setting \( b := (a_0 - c)/2 \) we define

\[ R = - \begin{bmatrix} 0 & b & 0 \\ b & -a_1 & 1/2 \\ 0 & 1/2 & 0 \end{bmatrix} \quad \text{and} \quad Q = \gamma / 2 \]

For the transition matrices we set

\[ A = \begin{bmatrix} 1 & 0 & 0 \\ 0 & 1 & 0 \\ 0 & 0 & \rho \end{bmatrix}, \quad B = \begin{bmatrix} 0 \\ 1 \\ 0 \end{bmatrix}, \quad C = \begin{bmatrix} 0 \\ 0 \\ \sigma_d \end{bmatrix} \]

Our aim is to compute the value-entropy correspondences shown above

The parameters are

\[ a_0 = 100, a_1 = 0.5, \rho = 0.9, \sigma_d = 0.05, \beta = 0.95, c = 2, \gamma = 50.0 \]

The standard normal distribution for \( w_t \) is understood as the agent’s baseline, with uncertainty parameterized by \( \theta \)

We compute value-entropy correspondences for two policies
3.7. ROBUSTNESS

1. The no concern for robustness policy $F_0$, which is the ordinary LQ loss minimizer

2. A “moderate” concern for robustness policy $F_\theta$, with $\theta = 0.02$

The code for producing the graph shown above, with blue being for the robust policy, is given in examples/robust_monopolist.jl

We repeat it here for convenience

```julia
# The robust control problem for a monopolist with adjustment costs. The
# inverse demand curve is:
#
# $p_t = a_0 - a_1 y_t + d_t$
#
# where $d_{t+1} = \rho d_t + \sigma_d w_{t+1}$ for $w_t \sim N(0,1)$ and iid.
# The period return function for the monopolist is
#
# $r_t = p_t y_t - \lambda (y_{t+1} - y_t)^2 / 2 - \gamma y_t$
#
# The objective of the firm is $E_t \sum_{t=0}^\infty \beta^t r_t$
#
# For the linear regulator, we take the state and control to be
#
# $x_t = (1, y_t, d_t)$ and $u_t = y_{t+1} - y_t$
#
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@Date: 2014-07-05

References
---------

Simple port of the file examples/robust_monopolist.py

http://quant-econ.net/robustness.html#application

#=
using QuantEcon
using PyPlot
using Grid

# model parameters
a_0 = 100
a_1 = 0.5
rho = 0.9
sigma_d = 0.05
bet = 0.95
c = 2
gam = 50.0
theta = 0.002
ac = (a_0 - c) / 2.0

# Define LQ matrices
```
R = \begin{bmatrix} 0 & ac & 0 \\ ac & -a_1 & 0.5 \\ 0 & 0.5 & 0 \end{bmatrix}
R = -R \quad \# \text{For minimization}
Q = \begin{bmatrix} \gamma / 2.0 \end{bmatrix}
A = \begin{bmatrix} 1. & 0. & 0. \\ 0. & 1. & 0. \\ 0. & 0. & \rho \end{bmatrix}
B = \begin{bmatrix} 0. & 1. & 0. \end{bmatrix}
C = \begin{bmatrix} 0. & 0. & \sigma_d \end{bmatrix}

## Functions

```plaintext
function evaluate_policy(theta, F)
    rlq = RBLQ(Q, R, A, B, C, bet, theta)
    K_F, P_F, d_F, O_F, o_F = evaluate_F(rlq, F)
    x0 = \begin{bmatrix} 1.0 & 0.0 & 0.0 \end{bmatrix}
    value = -x0'*P_F*x0 - d_F
    entropy = x0'*O_F*x0 + o_F
    return value[1], entropy[1] \quad \# \text{return scalars}
end
```

```plaintext
function value_and_entropy(emax, F, bw, grid_size=1000)
    if lowercase(bw) == "worst"
        thetas = 1 ./ linspace(1e-8, 1000, grid_size)
    else
        thetas = -1 ./ linspace(1e-8, 1000, grid_size)
    end

data = Array(Float64, grid_size, 2)
for (i, theta) in enumerate(thetas)
    data[i, :] = collect(evaluate_policy(theta, F))
    if data[i, 2] >= emax \quad \# \text{stop at this entropy level}
        data = data[1:i, :]
        break
    end
end
return data
end
```

## Main

```plaintext
# compute optimal rule
optimal_lq = LQ(Q, R, A, B, C, bet)
P0, F0, D0 = stationary_values(optimal_lq)

# compute robust rule for our theta
baseline_robust = RBLQ(Q, R, A, B, C, bet, theta)
Fb, Kb, Pb = robust_rule(baseline_robust)

# Check the positive definiteness of worst-case covariance matrix to
```

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# ensure that theta exceeds the breakdown point

```python
test_matrix = eye(size(Pb, 1)) - (C' * Pb * C ./ theta)[1]
eigenvals, eigenvecs = eig(test_matrix)
assert all(eigenvals .>= 0)
```

e_max = 1.6e6

# compute values and entropies

```python
optimal_best_case = value_and_entropy(e_max, F_0, "best")
robust_best_case = value_and_entropy(e_max, F_b, "best")
optimal_worst_case = value_and_entropy(e_max, F_0, "worst")
robust_worst_case = value_and_entropy(e_max, F_b, "worst")
```

# plot results

```python
fig, ax = subplots()
ax[:set_xlim](0, e_max)
ax[:set_ylabel]("Value")
ax[:set_xlabel]("Entropy")
ax[:grid]()

for axis in ["x", "y"]
    plt.ticklabel_format(style="sci", axis=axis, scilimits=(0,0))
end

plot_args = {:lw => 2, :alpha => 0.7}
colors = ("r", "b")

# we reverse order of "worst_case"s so values are ascending

data_pairs = ((optimal_best_case, optimal_worst_case),
              (robust_best_case, robust_worst_case))

egrid = linspace(0, e_max, 100)
egrid_data = Array{Float64}[]
for (c, data_pair) in zip(colors, data_pairs)
    for data in data_pair
        x, y = data[:, 2], data[:, 1]
curve(z) = InterpIrregular(x, y, BCnearest, InterpLinear)[z]
ax[:plot](egrid, curve(egrid), color=c; plot_args...)
push!(egrid_data, curve(egrid))
end
ax[:fill_between](egrid, egrid_data[1], egrid_data[2],
                  color=colors[1], alpha=0.1)
ax[:fill_between](egrid, egrid_data[3], egrid_data[4],
                  color=colors[2], alpha=0.1)
end
plt.show()
```

Here’s another such figure, with \( \theta = 0.002 \) instead of 0.02

Can you explain the different shape of the value-entropy correspondence for the robust policy?
We sketch the proof only of the first claim in this section, which is that, for any given $\theta$, $K(\hat{F}, \theta) = \hat{K}$, where $\hat{K}$ is as given in (3.61).

This is the content of the next lemma

**Lemma.** If $\hat{P}$ is the fixed point of the map $B \circ D$ and $\hat{F}$ is the robust policy as given in (3.60), then

$$K(\hat{F}, \theta) = (\theta I - C'\hat{P}C)^{-1}C'\hat{P}(A - B\hat{F})$$

(3.81)

**Proof:** As a first step, observe that when $F = \hat{F}$, the Bellman equation associated with the LQ problem (3.64) – (3.65) is

$$\hat{P} = -R - \hat{F}'Q\hat{F} - \beta^2(A - B\hat{F})'\hat{P}(\beta\theta I + \beta C'\hat{P}C)^{-1}C'\hat{P}(A - B\hat{F}) + \beta(A - B\hat{F})'\hat{P}(A - B\hat{F})$$

(3.82)

(revisit this discussion if you don’t know where (3.82) comes from) and the optimal policy is

$$w_{t+1} = -\beta(\beta\theta I + \beta C'\hat{P}C)^{-1}C'\hat{P}(A - B\hat{F})x_t$$

Suppose for a moment that $-\hat{P}$ solves the Bellman equation (3.82)

In this case the policy becomes

$$w_{t+1} = (\theta I - C'\hat{P}C)^{-1}C'\hat{P}(A - B\hat{F})x_t$$

which is exactly the claim in (3.81)

Hence it remains only to show that $-\hat{P}$ solves (3.82), or, in other words,

$$\hat{P} = R + \hat{F}'Q\hat{F} + \beta(A - B\hat{F})'\hat{P}(\theta I + \beta C'\hat{P}C)^{-1}C'\hat{P}(A - B\hat{F}) + \beta(A - B\hat{F})'\hat{P}(A - B\hat{F})$$

Using the definition of $D$, we can rewrite the right-hand side more simply as

$$R + \hat{F}'Q\hat{F} + \beta(A - B\hat{F})'D(\hat{P})(A - B\hat{F})$$

Although it involves a substantial amount of algebra, it can be shown that the latter is just $\hat{P}$

(Hint: Use the fact that $\hat{P} = B(D(\hat{P}))$)
3.8 Covariance Stationary Processes

Contents

• Covariance Stationary Processes
  – Overview
  – Introduction
  – Spectral Analysis
  – Implementation

Overview

In this lecture we study covariance stationary linear stochastic processes, a class of models routinely used to study economic and financial time series. This class has the advantage of being

1. simple enough to be described by an elegant and comprehensive theory
2. relatively broad in terms of the kinds of dynamics it can represent

We consider these models in both the time and frequency domain

ARMA Processes We will focus much of our attention on linear covariance stationary models with a finite number of parameters.

In particular, we will study stationary ARMA processes, which form a cornerstone of the standard theory of time series analysis.

It’s well known that every ARMA processes can be represented in linear state space form.

However, ARMA have some important structure that makes it valuable to study them separately

Spectral Analysis Analysis in the frequency domain is also called spectral analysis.

In essence, spectral analysis provides an alternative representation of the autocovariance of a covariance stationary process.

Having a second representation of this important object

• shines new light on the dynamics of the process in question
• allows for a simpler, more tractable representation in certain important cases

The famous Fourier transform and its inverse are used to map between the two representations

Other Reading For supplementary reading, see

• [LS12], chapter 2
3.8. COVARIANCE STATIONARY PROCESSES

- [Sar87], chapter 11
- John Cochrane’s notes on time series analysis, chapter 8
- [Shi95], chapter 6
- [CC08], all

Introduction

Consider a sequence of random variables \{X_t\} indexed by \(t \in \mathbb{Z}\) and taking values in \(\mathbb{R}\)

Thus, \{X_t\} begins in the infinite past and extends to the infinite future — a convenient and standard assumption

As in other fields, successful economic modeling typically requires identifying some deep structure in this process that is relatively constant over time

If such structure can be found, then each new observation \(X_t, X_{t+1}, \ldots\) provides additional information about it — which is how we learn from data

For this reason, we will focus in what follows on processes that are \textit{stationary} — or become so after some transformation (differencing, cointegration, etc.)

Definitions

A real-valued stochastic process \(\{X_t\}\) is called \textit{covariance stationary} if

1. Its mean \(\mu := \mathbb{E}X_t\) does not depend on \(t\)
2. For all \(k\) in \(\mathbb{Z}\), the \(k\)-th autocovariance \(\gamma(k) := \mathbb{E}(X_t - \mu)(X_{t+k} - \mu)\) is finite and depends only on \(k\)

The function \(\gamma: \mathbb{Z} \rightarrow \mathbb{R}\) is called the \textit{autocovariance function} of the process

Throughout this lecture, we will work exclusively with zero-mean (i.e., \(\mu = 0\)) covariance stationary processes

The zero-mean assumption costs nothing in terms of generality, since working with non-zero-mean processes involves no more than adding a constant

Example 1: White Noise

Perhaps the simplest class of covariance stationary processes is the white noise processes

A process \(\{\epsilon_t\}\) is called a \textit{white noise process} if

1. \(\mathbb{E}\epsilon_t = 0\)
2. \(\gamma(k) = \sigma^2 \mathbf{1}\{k = 0\}\) for some \(\sigma > 0\)

(Here \(\mathbf{1}\{k = 0\}\) is defined to be 1 if \(k = 0\) and zero otherwise)
Example 2: General Linear Processes  From the simple building block provided by white noise, we can construct a very flexible family of covariance stationary processes — the general linear processes

$$X_t = \sum_{j=0}^{\infty} \psi_j \epsilon_{t-j}, \quad t \in \mathbb{Z}$$

(3.83)

where

- \(\{\epsilon_t\}\) is white noise
- \(\{\psi_t\}\) is a square summable sequence in \(\mathbb{R}\) (that is, \(\sum_{t=0}^{\infty} \psi_t^2 < \infty\))

The sequence \(\{\psi_t\}\) is often called a linear filter

With some manipulations it is possible to confirm that the autocovariance function for (3.83) is

$$\gamma(k) = \sigma^2 \sum_{j=0}^{\infty} \psi_j \psi_{j+k}$$

(3.84)

By the Cauchy-Schwartz inequality one can show that the last expression is finite. Clearly it does not depend on \(t\)

Wold’s Decomposition  Remarkably, the class of general linear processes goes a long way towards describing the entire class of zero-mean covariance stationary processes

In particular, Wold’s theorem states that every zero-mean covariance stationary process \(\{X_t\}\) can be written as

$$X_t = \sum_{j=0}^{\infty} \psi_j \epsilon_{t-j} + \eta_t$$

where

- \(\{\epsilon_t\}\) is white noise
- \(\{\psi_t\}\) is square summable
- \(\eta_t\) can be expressed as a linear function of \(X_{t-1}, X_{t-2}, \ldots\) and is perfectly predictable over arbitrarily long horizons

For intuition and further discussion, see [Sar87], p. 286

AR and MA  General linear processes are a very broad class of processes, and it often pays to specialize to those for which there exists a representation having only finitely many parameters

(In fact, experience shows that models with a relatively small number of parameters typically perform better than larger models, especially for forecasting)

One very simple example of such a model is the AR(1) process

$$X_t = \phi X_{t-1} + \epsilon_t \quad \text{where} \quad |\phi| < 1 \quad \text{and} \quad \{\epsilon_t\} \text{ is white noise}$$

(3.85)

By direct substitution, it is easy to verify that \(X_t = \sum_{j=0}^{\infty} \phi^j \epsilon_{t-j}\)

Hence \(\{X_t\}\) is a general linear process
Applying (3.84) to the previous expression for $X_t$, we get the AR(1) autocovariance function

$$\gamma(k) = \phi^k \frac{\sigma^2}{1 - \phi^2}, \quad k = 0, 1, \ldots$$  (3.86)

The next figure plots this function for $\phi = 0.8$ and $\phi = -0.8$ with $\sigma = 1$.

Another very simple process is the MA(1) process

$$X_t = \epsilon_t + \theta \epsilon_{t-1}$$

You will be able to verify that

$$\gamma(0) = \sigma^2(1 + \theta^2), \quad \gamma(1) = \sigma^2 \theta, \quad \text{and} \quad \gamma(k) = 0 \quad \forall \ k > 1$$

The AR(1) can be generalized to an AR($p$) and likewise for the MA(1).

Putting all of this together, we get the

**ARMA Processes**  A stochastic process $\{X_t\}$ is called an *autoregressive moving average process*, or ARMA($p, q$), if it can be written as

$$X_t = \phi_1 X_{t-1} + \cdots + \phi_p X_{t-p} + \epsilon_t + \theta_1 \epsilon_{t-1} + \cdots + \theta_q \epsilon_{t-q}$$  (3.87)

where $\{\epsilon_t\}$ is white noise.

There is an alternative notation for ARMA processes in common use, based around the *lag operator* $L$.

**Def.** Given arbitrary variable $Y_t$, let $L^k Y_t := Y_{t-k}$.

It turns out that

- lag operators can lead to very succinct expressions for linear stochastic processes.
• algebraic manipulations treating the lag operator as an ordinary scalar often are legitimate. Using $L$, we can rewrite (3.87) as

$$L^0 X_t - \phi_1 L^1 X_t - \cdots - \phi_p L^p X_t = L^0 \epsilon_t + \theta_1 L^1 \epsilon_t + \cdots + \theta_q L^q \epsilon_t$$  

(3.88)

If we let $\phi(z)$ and $\theta(z)$ be the polynomials

$$\phi(z) := 1 - \phi_1 z - \cdots - \phi_p z^p \quad \text{and} \quad \theta(z) := 1 + \theta_1 z + \cdots + \theta_q z^q$$

(3.89)

then (3.88) simplifies further to

$$\phi(L) X_t = \theta(L) \epsilon_t$$  

(3.90)

In what follows we always assume that the roots of the polynomial $\phi(z)$ lie outside the unit circle in the complex plane. This condition is sufficient to guarantee that the ARMA($p, q$) process is covariance stationary. In fact it implies that the process falls within the class of general linear processes described above. That is, given an ARMA($p, q$) process \{X_t\} satisfying the unit circle condition, there exists a square summable sequence \{ψ_t\} with $X_t = \sum_{j=0}^{\infty} \psi_t \epsilon_{t-j}$ for all $t$.

The sequence \{ψ_t\} can be obtained by a recursive procedure outlined on page 79 of [CC08]. In this context, the function $t \mapsto \psi_t$ is often called the impulse response function.

**Spectral Analysis**

Autocovariance functions provide a great deal of information about covariance stationary processes. In fact, for zero-mean Gaussian processes, the autocovariance function characterizes the entire joint distribution. Even for non-Gaussian processes, it provides a significant amount of information. It turns out that there is an alternative representation of the autocovariance function of a covariance stationary process, called the spectral density. At times, the spectral density is easier to derive, easier to manipulate and provides additional intuition.

**Complex Numbers** Before discussing the spectral density, we invite you to recall the main properties of complex numbers (or skip to the next section).

It can be helpful to remember that, in a formal sense, complex numbers are just points $(x, y) \in \mathbb{R}^2$ endowed with a specific notion of multiplication. When $(x, y)$ is regarded as a complex number, $x$ is called the real part and $y$ is called the imaginary part.

The modulus or absolute value of a complex number $z = (x, y)$ is just its Euclidean norm in $\mathbb{R}^2$, but is usually written as $|z|$ instead of $\|z\|$.
The product of two complex numbers \((x, y)\) and \((u, v)\) is defined to be \((xu - vy, xv + yu)\), while addition is standard pointwise vector addition.

When endowed with these notions of multiplication and addition, the set of complex numbers forms a field — addition and multiplication play well together, just as they do in \(\mathbb{R}\).

The complex number \((x, y)\) is often written as \(x + iy\), where \(i\) is called the imaginary unit, and is understood to obey \(i^2 = -1\).

The \(x + iy\) notation can be thought of as an easy way to remember the definition of multiplication given above, because, proceeding naively,

\[(x + iy)(u + iv) = xu - yv + i(xv + yu)\]

Converted back to our first notation, this becomes \((xu - vy, xv + yu)\), which is the same as the product of \((x, y)\) and \((u, v)\) from our previous definition.

Complex numbers are also sometimes expressed in their polar form \(re^{i\omega}\), which should be interpreted as \(re^{i\omega} = r(\cos(\omega) + i \sin(\omega))\).

**Spectral Densities** Let \(\{X_t\}\) be a covariance stationary process with autocovariance function \(\gamma\) satisfying \(\sum_k \gamma(k)^2 < \infty\).

The spectral density \(f\) of \(\{X_t\}\) is defined as the discrete time Fourier transform of its autocovariance function \(\gamma\)

\[f(\omega) := \sum_{k \in \mathbb{Z}} \gamma(k)e^{-i\omega k}, \quad \omega \in \mathbb{R}\]

(Some authors normalize the expression on the right by constants such as \(1/\pi\) — the chosen convention makes little difference provided you are consistent.)

Using the fact that \(\gamma\) is even, in the sense that \(\gamma(t) = \gamma(-t)\) for all \(t\), you should be able to show that

\[f(\omega) = \gamma(0) + 2\sum_{k \geq 1} \gamma(k) \cos(\omega k)\] (3.91)

It is not difficult to confirm that \(f\) is

- real-valued
- even \((f(\omega) = f(-\omega)\), and
- \(2\pi\)-periodic, in the sense that \(f(2\pi + \omega) = f(\omega)\) for all \(\omega\).

It follows that the values of \(f\) on \([0, \pi]\) determine the values of \(f\) on all of \(\mathbb{R}\) — the proof is an exercise.

For this reason it is standard to plot the spectral density only on the interval \([0, \pi]\).

**Example 1: White Noise** Consider a white noise process \(\{\epsilon_t\}\) with standard deviation \(\sigma\).

It is simple to check that in this case we have \(f(\omega) = \sigma^2\). In particular, \(f\) is a constant function.

As we will see, this can be interpreted as meaning that “all frequencies are equally present”.

---

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(White light has this property when frequency refers to the visible spectrum, a connection that provides the origins of the term “white noise”)

**Example 2: AR and MA and ARMA** It is an exercise to show that the MA(1) process $X_t = \theta \epsilon_{t-1} + \epsilon_t$ has spectral density

$$f(\omega) = \sigma^2 (1 + 2\theta \cos(\omega) + \theta^2)$$  \hspace{1cm} (3.92)

With a bit more effort, it’s possible to show (see, e.g., p. 261 of [Sar87]) that the spectral density of the AR(1) process $X_t = \phi X_{t-1} + \epsilon_t$ is

$$f(\omega) = \frac{\sigma^2}{1 - 2\phi \cos(\omega) + \phi^2}$$  \hspace{1cm} (3.93)

More generally, it can be shown that the spectral density of the ARMA process (3.87) is

$$f(\omega) = \left| \frac{\theta(e^{i\omega})}{\phi(e^{i\omega})} \right|^2 \sigma^2$$  \hspace{1cm} (3.94)

where

- $\sigma$ is the standard deviation of the white noise process $\{\epsilon_t\}$
- the polynomials $\phi(\cdot)$ and $\theta(\cdot)$ are as defined in (3.89)

The derivation of (3.94) uses the fact that convolutions become products under Fourier transformations.

The proof is elegant and can be found in many places — see, for example, [Sar87], chapter 11, section 4.

It’s a nice exercise to verify that (3.92) and (3.93) are indeed special cases of (3.94).

**Interpreting the Spectral Density** Plotting (3.93) reveals the shape of the spectral density for the AR(1) model when $\phi$ takes the values 0.8 and -0.8 respectively.

These spectral densities correspond to the autocovariance functions for the AR(1) process shown above.

Informally, we think of the spectral density as being large at those $\omega \in [0, \pi]$ such that the autocovariance function exhibits significant cycles at this “frequency.”

To see the idea, let’s consider why, in the lower panel of the preceding figure, the spectral density for the case $\phi = -0.8$ is large at $\omega = \pi$.

Recall that the spectral density can be expressed as

$$f(\omega) = \gamma(0) + 2 \sum_{k \geq 1} \gamma(k) \cos(\omega k) = \gamma(0) + 2 \sum_{k \geq 1} (-0.8)^k \cos(\omega k)$$  \hspace{1cm} (3.95)

When we evaluate this at $\omega = \pi$, we get a large number because $\cos(\pi k)$ is large and positive when $(-0.8)^k$ is positive, and large in absolute value and negative when $(-0.8)^k$ is negative.
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Hence the product is always large and positive, and hence the sum of the products on the right-hand side of (3.95) is large.

These ideas are illustrated in the next figure, which has $k$ on the horizontal axis (click to enlarge).

On the other hand, if we evaluate $f(\omega)$ at $\omega = \pi/3$, then the cycles are not matched, the sequence $\gamma(k) \cos(\omega k)$ contains both positive and negative terms, and hence the sum of these terms is much smaller.

In summary, the spectral density is large at frequencies $\omega$ where the autocovariance function exhibits cycles.

**Inverting the Transformation** We have just seen that the spectral density is useful in the sense that it provides a frequency-based perspective on the autocovariance structure of a covariance stationary process.

Another reason that the spectral density is useful is that it can be “inverted” to recover the autocovariance function via the inverse Fourier transform.

In particular, for all $k \in \mathbb{Z}$, we have

$$\gamma(k) = \frac{1}{2\pi} \int_{-\pi}^{\pi} f(\omega) e^{i\omega k} d\omega \quad (3.96)$$

This is convenient in situations where the spectral density is easier to calculate and manipulate than the autocovariance function.

(For example, the expression (3.94) for the ARMA spectral density is much easier to work with than the expression for the ARMA autocovariance).

**Mathematical Theory** This section is loosely based on [Sar87], p. 249-253, and included for those who...
• would like a bit more insight into spectral densities
• and have at least some background in Hilbert space theory

Others should feel free to skip to the next section — none of this material is necessary to progress to computation

Recall that every separable Hilbert space \( H \) has a countable orthonormal basis \( \{ h_k \} \)

The nice thing about such a basis is that every \( f \in H \) satisfies

\[
f = \sum_k a_k h_k \quad \text{where} \quad a_k := \langle f, h_k \rangle \tag{3.97}
\]

where \( \langle \cdot, \cdot \rangle \) denotes the inner product in \( H \)

Thus, \( f \) can be represented to any degree of precision by linearly combining basis vectors.

The scalar sequence \( a = \{ a_k \} \) is called the Fourier coefficients of \( f \), and satisfies \( \sum_k |a_k|^2 < \infty \)

In other words, \( a \) is in \( \ell_2 \), the set of square summable sequences.

Consider an operator \( T \) that maps \( a \in \ell_2 \) into its expansion \( \sum_k a_k h_k \in H \)

The Fourier coefficients of \( T a \) are just \( a = \{ a_k \} \), as you can verify by confirming that \( \langle T a, h_k \rangle = a_k \)

Using elementary results from Hilbert space theory, it can be shown that

• \( T \) is one-to-one — if \( a \) and \( \beta \) are distinct in \( \ell_2 \), then so are their expansions in \( H \)
• \( T \) is onto — if \( f \in H \) then its preimage in \( \ell_2 \) is the sequence \( a \) given by \( a_k = \langle f, h_k \rangle \)
• \( T \) is a linear isometry — in particular \( \langle a, \beta \rangle = \langle T a, T \beta \rangle \)

Summarizing these results, we say that any separable Hilbert space is isometrically isomorphic to \( \ell_2 \)

In essence, this says that each separable Hilbert space we consider is just a different way of looking at the fundamental space \( \ell_2 \)

With this in mind, let’s specialize to a setting where

• \( \gamma \in \ell_2 \) is the autocovariance function of a covariance stationary process, and \( f \) is the spectral density
• \( H = L_2 \), where \( L_2 \) is the set of square summable functions on the interval \( [-\pi, \pi] \), with inner product \( \langle g, h \rangle = \int_{-\pi}^{\pi} g(\omega)h(\omega)d\omega \)
• \( \{ h_k \} = \) the orthonormal basis for \( L_2 \) given by the set of trigonometric functions

\[
h_k(\omega) = \frac{e^{ik\omega}}{\sqrt{2\pi}}, \quad k \in \mathbb{Z}, \quad \omega \in [-\pi, \pi]
\]

Using the definition of \( T \) from above and the fact that \( f \) is even, we now have

\[
T\gamma = \sum_{k \in \mathbb{Z}} \gamma(k) \frac{e^{ik\omega}}{\sqrt{2\pi}} = \frac{1}{\sqrt{2\pi}} f(\omega)
\tag{3.98}
\]

In other words, apart from a scalar multiple, the spectral density is just an transformation of \( \gamma \in \ell_2 \) under a certain linear isometry — a different way to view \( \gamma \)
In particular, it is an expansion of the autocovariance function with respect to the trigonometric basis functions in $L_2$.

As discussed above, the Fourier coefficients of $T\gamma$ are given by the sequence $\gamma$, and, in particular, $\gamma(k) = \langle T\gamma, h_k \rangle$.

Transforming this inner product into its integral expression and using (3.98) gives (3.96), justifying our earlier expression for the inverse transform.

**Implementation**

Most code for working with covariance stationary models deals with ARMA models.

Julia code for studying ARMA models can be found in the DSP.jl package.

Since this code doesn’t quite cover our needs — particularly vis-a-vis spectral analysis — we’ve put together the module arma.jl, which is part of QuantEcon package.

The module provides functions for mapping ARMA($p, q$) models into their

1. impulse response function
2. simulated time series
3. autocovariance function
4. spectral density

In addition to individual plots of these entities, we provide functionality to generate 2x2 plots containing all this information.

In other words, we want to replicate the plots on pages 68–69 of [LS12].

Here’s an example corresponding to the model $X_t = 0.5X_{t-1} + \epsilon_t - 0.8\epsilon_{t-2}$.

---

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**March 21, 2015**
Code  For interest’s sake, “arma.jl” is printed below

#= 

@authors: John Stachurski
Date: Thu Aug 21 11:09:30 EST 2014

Provides functions for working with and visualizing scalar ARMA processes. Ported from Python module quantecon.arma, which was written by Doc-Jin Jang, Jerry Choi, Thomas Sargent and John Stachurski

References
-------

http://quant-econ.net/arma.html

An example of usage is

using QuantEcon
phi = 0.5
theta = [0.0, -0.8]
sigma = 1.0
lp = ARMA(phi, theta, sigma)
require(joinpath(Pkg.dir("QuantEcon"), "examples", "arma_plots.jl"))
quad_plot(lp)

=#

type ARMA
  phi::Vector  # AR parameters phi_1, ..., phi_p
  theta::Vector  # MA parameters theta_1, ..., theta_q
  p::Integer  # Number of AR coefficients
  q::Integer  # Number of MA coefficients
  sigma::Real  # Variance of white noise
  ma_poly::Vector  # MA polynomial --- filtering representation
  ar_poly::Vector  # AR polynomial --- filtering representation
end

# constructors to coerce phi/theta to vectors
ARMA(phi::Real, theta::Real=0.0, sigma::Real=1.0) = ARMA([phi], [theta], sigma)
ARMA(phi::Real, theta::Vector=[0.0], sigma::Real=1.0) = ARMA([phi], theta, sigma)
ARMA(phi::Vector, theta::Real=0.0, sigma::Real=1.0) = ARMA(phi, theta, sigma)

function ARMA(phi::Vector, theta::Vector=[0.0], sigma::Real=1.0)
  p = length(phi)
  q = length(theta)
  ma_poly = [1.0, theta]
  ar_poly = [1.0, -phi]
  return ARMA(phi, theta, p, q, sigma, ma_poly, ar_poly)
end
function spectral_density(arma::ARMA; res=1200, two_pi=true)
    # Compute the spectral density associated with ARMA process arma
    wmax = two_pi ? 2pi : pi
    w = linspace(0, wmax, res)
    tf = TFFilter(reverse(arma.ma_poly), reverse(arma.ar_poly))
    h = freqz(tf, w)
    spect = arma.sigma^2 * abs(h).^2
    return w, spect
end

function autocovariance(arma::ARMA; num_autocov=16)
    # Compute the autocovariance function associated with ARMA process arma
    # Computation is via the spectral density and inverse FFT
    (w, spect) = spectral_density(arma)
    acov = real(Base.ifft(spect))
    # num_autocov should be <= len(acov) / 2
    return acov[1:num_autocov]
end

function impulse_response(arma::ARMA; impulse_length=30)
    # Compute the impulse response function associated with ARMA process arma
    err_msg = "Impulse length must be greater than number of AR coefficients"
    @assert impulse_length >= arma.p err_msg
    # == Pad theta with zeros at the end ==#
    theta = [arma.theta, zeros(impulse_length - arma.q)]
    psi_zero = 1.0
    psi = Array(Float64, impulse_length)
    for j = 1:impulse_length
        psi[j] = theta[j]
        for i = 1:min(j, arma.p)
        end
    end
    return [psi_zero, psi[1:end-1]]
end

function simulation(arma::ARMA; ts_length=90, impulse_length=30)
    # Simulate the ARMA process arma assuming Gaussian shocks
    J = impulse_length
    T = ts_length
    psi = impulse_response(arma, impulse_length=impulse_length)
    epsilon = arma.sigma * randn(T + J)
    X = Array(Float64, T)
    for t=1:T
        X[t] = dot(epsilon[t:J+t-1], psi)
    end
    return X
end

Here’s an example of usage

julia> using QuantEcon
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```julia
julia> using QuantEcon
julia> phi = 0.5;
julia> theta = [0, -0.8];
julia> lp = ARMA(phi, theta);
julia> QuantEcon.quad_plot(lp)
```

**Explanation** The call

```julia
lp = ARMA(phi, theta, sigma)
```

creates an instance `lp` that represents the ARMA$(p, q)$ model

$$X_t = \phi_1 X_{t-1} + \ldots + \phi_p X_{t-p} + \epsilon_t + \theta_1 \epsilon_{t-1} + \ldots + \theta_q \epsilon_{t-q}$$

If `phi` and `theta` are arrays or sequences, then the interpretation will be

- `phi` holds the vector of parameters $(\phi_1, \phi_2, \ldots, \phi_p)$
- `theta` holds the vector of parameters $(\theta_1, \theta_2, \ldots, \theta_q)$

The parameter `sigma` is always a scalar, the standard deviation of the white noise.

We also permit `phi` and `theta` to be scalars, in which case the model will be interpreted as

$$X_t = \phi X_{t-1} + \epsilon_t + \theta \epsilon_{t-1}$$

The two numerical packages most useful for working with ARMA models are DSP.jl and the `fft` routine in Julia.

**Computing the Autocovariance Function** As discussed above, for ARMA processes the spectral density has a *simple representation* that is relatively easy to calculate.

Given this fact, the easiest way to obtain the autocovariance function is to recover it from the spectral density via the inverse Fourier transform.

Here we use Julia’s Fourier transform routine `fft`, which wraps a standard C-based package called FFTW.

A look at the `fft` documentation shows that the inverse transform `ifft` takes a given sequence $A_0, A_1, \ldots, A_{n-1}$ and returns the sequence $a_0, a_1, \ldots, a_{n-1}$ defined by

$$a_k = \frac{1}{n} \sum_{t=0}^{n-1} A_t e^{ik2\pi t/n}$$

Thus, if we set $A_t = f(\omega_t)$, where $f$ is the spectral density and $\omega_t := 2\pi t/n$, then

$$a_k = \frac{1}{n} \sum_{t=0}^{n-1} f(\omega_t) e^{i\omega_t k} = \frac{2\pi}{n} \sum_{t=0}^{n-1} f(\omega_t) e^{i\omega_t k}, \quad \omega_t := 2\pi t/n$$
For $n$ sufficiently large, we then have

$$a_k \approx \frac{1}{2\pi} \int_{0}^{2\pi} f(\omega)e^{i\omega k} d\omega = \frac{1}{2\pi} \int_{-\pi}^{\pi} f(\omega)e^{i\omega k} d\omega$$

(You can check the last equality)

In view of (3.96) we have now shown that, for $n$ sufficiently large, $a_k \approx \gamma(k)$ — which is exactly what we want to compute

### 3.9 Estimation of Spectra

#### Overview

In a previous lecture we covered some fundamental properties of covariance stationary linear stochastic processes

One objective for that lecture was to introduce spectral densities — a standard and very useful technique for analyzing such processes

In this lecture we turn to the problem of estimating spectral densities and other related quantities from data

Estimates of the spectral density are computed using what is known as a periodogram — which in turn is computed via the famous fast Fourier transform

Once the basic technique has been explained, we will apply it to the analysis of several key macroeconomic time series

For supplementary reading, see [Sar87] or [CC08].

#### Periodograms

*Recall that* the spectral density $f$ of a covariance stationary process with autocorrelation function $\gamma$ can be written as

$$f(\omega) = \gamma(0) + 2 \sum_{k \geq 1} \gamma(k) \cos(\omega k), \quad \omega \in \mathbb{R}$$
Now consider the problem of estimating the spectral density of a given time series, when \( \gamma \) is unknown.

In particular, let \( X_0, \ldots, X_{n-1} \) be \( n \) consecutive observations of a single time series that is assumed to be covariance stationary.

The most common estimator of the spectral density of this process is the *periodogram* of \( X_0, \ldots, X_{n-1} \), which is defined as

\[
I(\omega) := \frac{1}{n} \left| \sum_{t=0}^{n-1} X_t e^{it\omega} \right|^2, \quad \omega \in \mathbb{R}
\]

(Recall that \( |z| \) denotes the modulus of complex number \( z \)).

Alternatively, \( I(\omega) \) can be expressed as

\[
I(\omega) = \frac{1}{n} \left\{ \left[ \sum_{t=0}^{n-1} X_t \cos(\omega t) \right]^2 + \left[ \sum_{t=0}^{n-1} X_t \sin(\omega t) \right]^2 \right\}
\]

It is straightforward to show that the function \( I \) is even and 2\( \pi \)-periodic (i.e., \( I(\omega) = I(-\omega) \) and \( I(\omega + 2\pi) = I(\omega) \) for all \( \omega \in \mathbb{R} \)).

From these two results, you will be able to verify that the values of \( I \) on \([0, \pi]\) determine the values of \( I \) on all of \( \mathbb{R} \).

The next section helps to explain the connection between the periodogram and the spectral density.

**Interpretation** To interpret the periodogram, it is convenient to focus on its values at the *Fourier frequencies*

\[
\omega_j := \frac{2\pi j}{n}, \quad j = 0, \ldots, n-1
\]

In what sense is \( I(\omega_j) \) an estimate of \( f(\omega_j) \)?

The answer is straightforward, although it does involve some algebra.

With a bit of effort one can show that, for any integer \( j > 0 \),

\[
\sum_{t=0}^{n-1} e^{it\omega_j} = \sum_{t=0}^{n-1} \exp \left\{ i2\pi j \frac{t}{n} \right\} = 0
\]

Letting \( \bar{X} \) denote the sample mean \( n^{-1} \sum_{t=0}^{n-1} X_t \), we then have

\[
nI(\omega_j) = \left[ \sum_{t=0}^{n-1} (X_t - \bar{X}) e^{it\omega_j} \right]^2 = \sum_{t=0}^{n-1} (X_t - \bar{X}) e^{it\omega_j} \sum_{r=0}^{n-1} (X_r - \bar{X}) e^{-ir\omega_j}
\]

By carefully working through the sums, one can transform this to

\[
nI(\omega_j) = \sum_{t=0}^{n-1} (X_t - \bar{X})^2 + 2 \sum_{k=1}^{n-1} \sum_{t=k}^{n-1} (X_t - \bar{X})(X_{t-k} - \bar{X}) \cos(\omega_j k)
\]
Now let
\[ \hat{\gamma}(k) := \frac{1}{n} \sum_{t=k}^{n-1} (X_t - \bar{X})(X_{t-k} - \bar{X}), \quad k = 0, 1, \ldots, n - 1 \]

This is the sample autocovariance function, the natural “plug-in estimator” of the autocovariance function \( \gamma \)

(‘Plug-in estimator” is an informal term for an estimator found by replacing expectations with sample means)

With this notation, we can now write
\[ I(\omega_j) = \hat{\gamma}(0) + 2 \sum_{k=1}^{n-1} \hat{\gamma}(k) \cos(\omega_j k) \]

Recalling our expression for \( f \) given above, we see that \( I(\omega_j) \) is just a sample analog of \( f(\omega_j) \)

**Calculation** Let’s now consider how to compute the periodogram as defined in (3.99)

There are already functions available that will do this for us — an example is `periodogram` in the DSP.jl package

However, it is very simple to replicate their results, and this will give us a platform to make useful extensions

The most common way to calculate the periodogram is via the discrete Fourier transform, which in turn is implemented through the fast Fourier transform algorithm

In general, given a sequence \( a_0, \ldots, a_{n-1} \), the discrete Fourier transform computes the sequence
\[ A_j := \frac{1}{n} \sum_{t=0}^{n-1} a_t \exp \left\{ i \frac{2\pi}{n} t j \right\}, \quad j = 0, \ldots, n - 1 \]

With \( a_0, \ldots, a_{n-1} \) stored in Julia array \( a \), the function call `fft(a)` returns the values \( A_0, \ldots, A_{n-1} \) as a Julia array

It follows that, when the data \( X_0, \ldots, X_{n-1} \) is stored in array \( X \), the values \( I(\omega_j) \) at the Fourier frequencies, which are given by
\[ \frac{1}{n} \left| \sum_{t=0}^{n-1} X_t \exp \left\{ i \frac{2\pi}{n} t j \right\} \right|^2, \quad j = 0, \ldots, n - 1 \]

can be computed by `abs(fft(X)).^2 ./ length(X)`

Note: The Julia function `abs` acts elementwise, and correctly handles complex numbers (by computing their modulus, which is exactly what we need)

Here’s a function that puts all this together

```julia
function periodogram(x::Array):
    n = length(x)
    I_w = abs(fft(x)).^2 / n
    w = 2pi * [0:n-1] ./ n # Fourier frequencies
    return I_w
end
```
3.9. ESTIMATION OF SPECTRA

```python
w, I_w = w[1:int(n/2)], I_w[1:int(n/2)]  # Truncate to interval [0, pi]
return w, I_w
```

Let’s generate some data for this function using the ARMA type from QuantEcon

(See the lecture on linear processes for details on this class)

Here’s a code snippet that, once the preceding code has been run, generates data from the process

\[ X_t = 0.5X_{t-1} + \epsilon_t - 0.8\epsilon_{t-2} \]  \hspace{1cm} (3.100)

where \( \{\epsilon_t\} \) is white noise with unit variance, and compares the periodogram to the actual spectral density

```python
import PyPlot: plt
import QuantEcon: ARMA

n = 40  # Data size
phi, theta = 0.5, [0, -0.8]  # AR and MA parameters
lp = ARMA(phi, theta)
X = simulation(lp, ts_length=n)

fig, ax = plt.subplots()
x, y = periodogram(X)
ax[0].plot(x, y, "b-", lw=2, alpha=0.5, label="periodogram")
x_sd, y_sd = spectral_density(lp, two_pi=False, resolution=120)
ax[0].plot(x_sd, y_sd, "r-", lw=2, alpha=0.8, label="spectral density")
ax[0].legend()
plt.show()
```

Running this should produce a figure similar to this one

![Figure showing periodogram and spectral density comparison](image)

This estimate looks rather disappointing, but the data size is only 40, so perhaps it’s not surprising that the estimate is poor.
However, if we try again with $n = 1200$ the outcome is not much better.

The periodogram is far too irregular relative to the underlying spectral density. This brings us to our next topic:

**Smoothing**

There are two related issues here:

One is that, given the way the fast Fourier transform is implemented, the number of points $\omega$ at which $I(\omega)$ is estimated increases in line with the amount of data. In other words, although we have more data, we are also using it to estimate more values. A second issue is that densities of all types are fundamentally hard to estimate without parameteric assumptions. Typically, nonparametric estimation of densities requires some degree of smoothing. The standard way that smoothing is applied to periodograms is by taking local averages. In other words, the value $I(\omega_j)$ is replaced with a weighted average of the adjacent values

$$I(\omega_{j-p}), I(\omega_{j-p+1}), \ldots, I(\omega_{j}), \ldots, I(\omega_{j+p})$$

This weighted average can be written as

$$I_S(\omega_j) := \sum_{\ell = -p}^{p} w(\ell)I(\omega_{j+\ell}) \quad (3.101)$$

where the weights $w(-p), \ldots, w(p)$ are a sequence of $2p + 1$ nonnegative values summing to one. In generally, larger values of $p$ indicate more smoothing — more on this below.
The next figure shows the kind of sequence typically used

Note the smaller weights towards the edges and larger weights in the center, so that more distant values from $I(\omega_j)$ have less weight than closer ones in the sum (3.101)

**Estimation with Smoothing**  Our next step is to provide code that will not only estimate the periodogram but also provide smoothing as required

Such functions have been written in `estspec.jl` and are available via QuantEcon

The file `estspec.jl` are printed below

```julia
#=
Functions for working with periodograms of scalar data.

@author : Spencer Lyon <spencer.lyon@nyu.edu>

@date : 2014-08-21

References
--------

Simple port of the file quantecon.estspec

http://quant-econ.net/estspec.html

=#
import DSP

function smooth(x::Array, window_len::Int=7, window::String="hanning")
    if length(x) < window_len
        throw(ArgumentError("Input vector length must be >= window length"))
    end

    if window_len < 3
        throw(ArgumentError("Window length must be at least 3."))
    end

```
if iseven(window_len)
    window_len += 1
    println("Window length must be odd, reset to \$window_len")
end

windows = {
    "hanning" => DSP.hanning,
    "hamming" => DSP.hamming,
    "bartlett" => DSP.bartlett,
    "blackman" => DSP.blackman,
    "flat" => DSP.rect  # moving average
}

# Reflect x around x[0] and x[-1] prior to convolution
k = int(window_len / 2)
xb = x[1:k]  # First k elements
xt = x[end-k+1:end]  # Last k elements
s = [reverse(xb), x, reverse(xt)]

# === Select window values === #
if !haskey(windows, window)
    msg = "Unrecognized window type \'$window'"
    print(msg * " Defaulting to hanning")
    window = "hanning"
end

w = windows[window](window_len)
return conv(w ./ sum(w), s)[window_len+1:end-window_len]
end

function smooth(x::Array; window_len::Int=7, window::String="hanning")
    smooth(x, window_len, window)
end

function periodogram(x::Vector)
    n = length(x)
    I_w = abs(fft(x)).^2 ./ n
    w = 2pi * [0:n-1] ./ n  # Fourier frequencies

    # int rounds to nearest integer. We want to round up or take 1/2 + 1 to
    # make sure we get the whole interval from [0, pi]
    ind = iseven(n) ? int(n / 2 + 1) : int(n / 2)
    w, I_w = w[1:ind], I_w[1:ind]
    return w, I_w
end

function periodogram(x::Vector, window::String, window_len::Int=7)
    w, I_w = periodogram(x)
The listing displays three functions, `smooth()`, `periodogram()`, `ar_periodogram()`. We will discuss the first two here and the third one below.

The `periodogram()` function returns a periodogram, optionally smoothed via the `smooth()` function.

Regarding the `smooth()` function, since smoothing adds a nontrivial amount of computation, we have applied a fairly terse array-centric method based around `conv`.

Readers are left to either explore or simply use this code according to their interests.

The next three figures each show smoothed and unsmoothed periodograms, as well as the true spectral density.

(The model is the same as before — see equation (3.100) — and there are 400 observations.)

From top figure to bottom, the window length is varied from small to large.

In looking at the figure, we can see that for this model and data size, the window length chosen in the middle figure provides the best fit.

Relative to this value, the first window length provides insufficient smoothing, while the third gives too much smoothing.

Of course in real estimation problems the true spectral density is not visible and the choice of appropriate smoothing will have to be made based on judgement/priors or some other theory.

Pre-Filtering and Smoothing In the code listing above we showed three functions from the file `estspec.jl`
3.9. ESTIMATION OF SPECTRA

![Graphs showing periodogram, spectral density, and smoothed periodogram for different window lengths.](image-url)
3.9. ESTIMATION OF SPECTRA

The third function in the file (ar_periodogram()) adds a pre-processing step to periodogram smoothing.

First we describe the basic idea, and after that we give the code.

The essential idea is to

1. Transform the data in order to make estimation of the spectral density more efficient.
2. Compute the periodogram associated with the transformed data.
3. Reverse the effect of the transformation on the periodogram, so that it now estimates the spectral density of the original process.

Step 1 is called pre-filtering or pre-whitening, while step 3 is called recoloring.

The first step is called pre-whitening because the transformation is usually designed to turn the data into something closer to white noise.

Why would this be desirable in terms of spectral density estimation?

The reason is that we are smoothing our estimated periodogram based on estimated values at nearby points — recall (3.101).

The underlying assumption that makes this a good idea is that the true spectral density is relatively regular — the value of \( I(\omega) \) is close to that of \( I(\omega') \) when \( \omega \) is close to \( \omega' \).

This will not be true in all cases, but it is certainly true for white noise.

For white noise, \( I \) is as regular as possible — it is a constant function.

In this case, values of \( I(\omega') \) at points \( \omega' \) near to \( \omega \) provided the maximum possible amount of information about the value \( I(\omega) \).

Another way to put this is that if \( I \) is relatively constant, then we can use a large amount of smoothing without introducing too much bias.

**The AR(1) Setting**  Let’s examine this idea more carefully in a particular setting — where the data is assumed to be AR(1).

(More general ARMA settings can be handled using similar techniques to those described below.)

Suppose in particular that \( \{X_t\} \) is covariance stationary and AR(1), with

\[
X_{t+1} = \mu + \phi X_t + \epsilon_{t+1}
\]

(3.102)

where \( \mu \) and \( \phi \in (-1, 1) \) are unknown parameters and \( \{\epsilon_t\} \) is white noise.

It follows that if we regress \( X_{t+1} \) on \( X_t \) and an intercept, the residuals will approximate white noise.

Let

- \( g \) be the spectral density of \( \{\epsilon_t\} \) — a constant function, as discussed above.
- \( I_0 \) be the periodogram estimated from the residuals — an estimate of \( g \).
- \( f \) be the spectral density of \( \{X_t\} \) — the object we are trying to estimate.
In view of an earlier result we obtained while discussing ARMA processes, \( f \) and \( g \) are related by

\[
f(\omega) = \left| \frac{1}{1 - \phi e^{i\omega}} \right|^2 g(\omega)
\]

(3.103)

This suggests that the recoloring step, which constructs an estimate \( I \) of \( f \) from \( I_0 \), should set

\[
I(\omega) = \left| \frac{1}{1 - \hat{\phi} e^{i\omega}} \right|^2 I_0(\omega)
\]

where \( \hat{\phi} \) is the OLS estimate of \( \phi \)

The code for `ar_periodogram()` — the third function in `estspec.jl` — does exactly this. (See the code [here](#))

The next figure shows realizations of the two kinds of smoothed periodograms

1. “standard smoothed periodogram”, the ordinary smoothed periodogram, and
2. “AR smoothed periodogram”, the pre-whitened and recolored one generated by `ar_periodogram()`

The periodograms are calculated from time series drawn from (3.102) with \( \mu = 0 \) and \( \phi = -0.9 \)

Each time series is of length 150

The difference between the three subfigures is just randomness — each one uses a different draw of the time series

In all cases, periodograms are fit with the “hamming” window and window length of 65

Overall, the fit of the AR smoothed periodogram is much better, in the sense of being closer to the true spectral density

**Exercises**

**Exercise 1** Replicate this figure (modulo randomness)

The model is as in equation (3.100) and there are 400 observations

For the smoothed periodogram, the window type is “hamming”

**Exercise 2** Replicate this figure (modulo randomness)

The model is as in equation (3.102), with \( \mu = 0, \phi = -0.9 \) and 150 observations in each time series

All periodograms are fit with the “hamming” window and window length of 65

**Exercise 3** To be written. The exercise will be to use the code from this lecture to download FRED data and generate periodograms for different kinds of macroeconomic data.
3.9. ESTIMATION OF SPECTRA

- spectral density
- standard smoothed periodogram
- AR smoothed periodogram

March 21, 2015
Overview

In this lecture we study optimal fiscal policy in a linear quadratic setting.

We slightly modify a well-known model of Robert Lucas and Nancy Stokey [LS83] so that convenient formulas for solving linear-quadratic models can be applied to simplify the calculations.

The economy consists of a representative household and a benevolent government.

The government finances an exogenous stream of government purchases with state-contingent loans and a linear tax on labor income.

A linear tax is sometimes called a flat-rate tax.

The household maximizes utility by choosing paths for consumption and labor, taking prices and the government’s tax rate and borrowing plans as given.

Maximum attainable utility for the household depends on the government’s tax and borrowing plans.

The Ramsey problem [Ram27] is to choose tax and borrowing plans that maximize the household’s welfare, taking the household’s optimizing behavior as given.

There is a large number of competitive equilibria indexed by different government fiscal policies. The Ramsey planner chooses the best competitive equilibrium.

We want to study the dynamics of tax rates, tax revenues, government debt under a Ramsey plan.

Because the Lucas and Stokey model features state-contingent government debt, the government debt dynamics differ substantially from those in a model of Robert Barro [Bar79].
The treatment given here closely follows this manuscript, prepared by Thomas J. Sargent and Francois R. Velde

We cover only the key features of the problem in this lecture, leaving you to refer to that source for additional results and intuition

**Model Features**

- Linear quadratic (LQ) model
- Representative household
- Stochastic dynamic programming over an infinite horizon
- Distortionary taxation

**The Ramsey Problem**

We begin by outlining the key assumptions regarding technology, households and the government sector

**Technology** Labor can be converted one-for-one into a single, non-storable consumption good

In the usual spirit of the LQ model, the amount of labor supplied in each period is unrestricted

This is unrealistic, but helpful when it comes to solving the model

Realistic labor supply can be induced by suitable parameter values

**Households** Consider a representative household who chooses a path \( \{ \ell_t, c_t \} \) for labor and consumption to maximize

\[
- \mathbb{E} \frac{1}{2} \sum_{t=0}^{\infty} \beta^t \left[ (c_t - b_t)^2 + \ell_t^2 \right]
\]

subject to the budget constraint

\[
\mathbb{E} \sum_{t=0}^{\infty} \beta^t p_t^0 \left[ d_t + (1 - \tau_t) \ell_t + s_t - c_t \right] = 0
\]

Here

- \( \beta \) is a discount factor in \((0, 1)\)
- \( p_t^0 \) is state price at time \( t \)
- \( b_t \) is a stochastic preference parameter
- \( d_t \) is an endowment process
- \( \tau_t \) is a flat tax rate on labor income
- \( s_t \) is a promised time-\( t \) coupon payment on debt issued by the government
The budget constraint requires that the present value of consumption be restricted to equal the present value of endowments, labor income and coupon payments on bond holdings.

**Government** The government imposes a linear tax on labor income, fully committing to a stochastic path of tax rates at time zero. The government also issues state-contingent debt. Given government tax and borrowing plans, we can construct a competitive equilibrium with distorting government taxes. Among all such competitive equilibria, the Ramsey plan is the one that maximizes the welfare of the representative consumer.

**Exogenous Variables** Endowments, government expenditure, the preference parameter $b_t$ and promised coupon payments on initial government debt $s_t$ are all exogenous, and given by

\[
\begin{align*}
  d_t &= S_d x_t \\
  g_t &= S_g x_t \\
  b_t &= S_b x_t \\
  s_t &= S_s x_t
\end{align*}
\]

The matrices $S_d, S_g, S_b, S_s$ are primitives and $\{x_t\}$ is an exogenous stochastic process taking values in $\mathbb{R}^k$.

We consider two specifications for $\{x_t\}$

1. Discrete case: $\{x_t\}$ is a discrete state Markov chain with transition matrix $P$
2. VAR case: $\{x_t\}$ obeys $x_{t+1} = Ax_t + Cw_{t+1}$ where $\{w_t\}$ is independent zero mean Gaussian with identify covariance matrix

**Feasibility** The period-by-period feasibility restriction for this economy is

\[
c_t + g_t = d_t + \ell_t
\]  

(3.106)

A labor-consumption process $\{\ell_t, c_t\}$ is called feasible if (3.106) holds for all $t$.

**Government budget constraint** Where $p_0^t$ is a scaled Arrow-Debreu price, the time zero government budget constraint is

\[
\mathbb{E} \sum_{t=0}^{\infty} \beta^t p_0^t (s_t + g_t - \tau_t \ell_t) = 0
\]  

(3.107)

**Equilibrium** An equilibrium is a feasible allocation $\{\ell_t, c_t\}$, a sequence of prices $\{p_t\}$, and a tax system $\{\tau_t\}$ such that

1. The allocation $\{\ell_t, c_t\}$ is optimal for the household given $\{p_t\}$ and $\{\tau_t\}$

---

THOMAS SARGENT AND JOHN STACHURSKI  
March 21, 2015
2. The government’s budget constraint (3.107) is satisfied

The Ramsey problem is to choose the equilibrium \( \{ \ell_t, c_t, \tau_t, p_t \} \) that maximizes the household’s welfare.

If \( \{ \ell_t, c_t, \tau_t, p_t \} \) is a solution to the Ramsey problem, then \( \{ \tau_t \} \) is called the Ramsey plan.

The solution procedure we adopt is

1. Use the first order conditions from the household problem to pin down prices and allocations given \( \{ \tau_t \} \)
2. Use these expressions to rewrite the government budget constraint (3.107) in terms of exogenous variables and allocations
3. Maximize the household’s objective function (3.104) subject to the constraint constructed in step 2 and the feasibility constraint (3.106)

The solution to this maximization problem pins down all quantities of interest.

**Solution** Step one is to obtain the first order conditions for the household’s problem, taking taxes and prices as given.

Letting \( \mu \) be the Lagrange multiplier on (3.105), the first order conditions are

\[
p_t = \frac{b_t - c_t}{b_0 - c_0} \quad \text{and} \quad \tau_t = 1 - \frac{\ell_t}{b_t - c_t}
\]

Rearranging and normalizing at \( \mu = b_0 - c_0 \), we can write these conditions as

\[
p_t = \frac{b_t - c_t}{b_0 - c_0} \quad \text{and} \quad \tau_t = 1 - \frac{\ell_t}{b_t - c_t}
\]

Substituting (3.108) into the government’s budget constraint (3.107) yields

\[
E \sum_{t=0}^{\infty} \beta^t \left[ (b_t - c_t)(s_t + g_t - \ell_t) + \ell_t^2 \right] = 0
\]

The Ramsey problem now amounts to maximizing (3.104) subject to (3.109) and (3.106)

The associated Lagrangian is

\[
\mathcal{L} = E \sum_{t=0}^{\infty} \beta^t \left\{ -\frac{1}{2} [(c_t - b_t)^2 + \ell_t^2] + \lambda \left[ (b_t - c_t)(\ell_t - s_t - g_t) - \ell_t^2 \right] + \mu_t [d_t + \ell_t - c_t - g_t] \right\}
\]

The first order conditions associated with \( c_t \) and \( \ell_t \) are

\[
-(c_t - b_t) + \lambda[-\ell_t + (g_t + s_t)] = \mu_t
\]

and

\[
\ell_t - \lambda[(b_t - c_t) - 2\ell_t] = \mu_t
\]

Combining these last two equalities with (3.106) and working through the algebra, one can show that

\[
\ell_t = \tilde{\ell}_t - \nu \mu_t \quad \text{and} \quad c_t = \tilde{c}_t - \nu m_t
\]
3.10. OPTIMAL TAXATION

- $\nu := \lambda / (1 + 2\lambda)$
- $\bar{\ell}_t := (b_t - d_t + g_t)/2$
- $\bar{c}_t := (b_t + d_t - g_t)/2$
- $m_t := (b_t - d_t - s_t)/2$

Apart from $\nu$, all of these quantities are expressed in terms of exogenous variables.

To solve for $\nu$, we can use the government’s budget constraint again.

The term inside the brackets in (3.109) is $(b_t - c_t)(s_t + g_t) - (b_t - c_t)\ell_t + \ell_t^2$

Using (3.111), the definitions above and the fact that $\bar{\ell} = b - \bar{c}$, this term can be rewritten as

$$(b_t - \bar{c}_t)(g_t + s_t) + 2m_t^2(\nu^2 - \nu)$$

Reinserting into (3.109), we get

$$\mathbb{E} \left\{ \sum_{t=0}^{\infty} \beta^t (b_t - \bar{c}_t)(g_t + s_t) \right\} + (\nu^2 - \nu)\mathbb{E} \left\{ \sum_{t=0}^{\infty} \beta^t 2m_t^2 \right\} = 0 \quad (3.112)$$

Although it might not be clear yet, we are nearly there:

- The two expectations terms in (3.112) can be solved for in terms of model primitives
- This in turn allows us to solve for the Lagrange multiplier $\nu$
- With $\nu$ in hand, we can go back and solve for the allocations via (3.111)
- Once we have the allocations, prices and the tax system can be derived from (3.108)

Solving the Quadratic Term  Let’s consider how to obtain the term $\nu$ in (3.112)

If we can solve the two expected geometric sums

$$b_0 := \mathbb{E} \left\{ \sum_{t=0}^{\infty} \beta^t (b_t - \bar{c}_t)(g_t + s_t) \right\} \quad \text{and} \quad a_0 := \mathbb{E} \left\{ \sum_{t=0}^{\infty} \beta^t 2m_t^2 \right\} \quad (3.113)$$

then the problem reduces to solving

$$b_0 + a_0(\nu^2 - \nu) = 0$$

for $\nu$

Provided that $4b_0 < a_0$, there is a unique solution $\nu \in (0, 1/2)$, and a unique corresponding $\lambda > 0$

Let’s work out how to solve the expectations terms in (3.113)

For the first one, the random variable $(b_t - \bar{c}_t)(g_t + s_t)$ inside the summation can be expressed as

$$\frac{1}{2} x_t' (S_b - S_d + S_g)' (S_g + S_s) x_t$$

For the second expectation in (3.113), the random variable $2m_t^2$ can be written as

$$\frac{1}{2} x_t' (S_b - S_d - S_s)' (S_b - S_d - S_s) x_t$$
3.10. OPTIMAL TAXATION

It follows that both of these expectations terms are special cases of the expression

\[ q(x_0) = \mathbb{E} \sum_{t=0}^{\infty} \beta^t x'_t H x_t \]  

(3.114)

where \( H \) is a conformable matrix, and \( x'_t \) is the transpose of column vector \( x_t \).

Suppose first that \( \{x_t\} \) is the Gaussian VAR described above. In this case, the formula for computing \( q(x_0) \) is known to be

\[ q(x_0) = x'_0 Q x_0 + v, \]

where \( Q \) is the solution to

\[ Q = H + \beta A' Q A, \]

and

\[ v = \text{trace} \left( C' Q C \right) \beta / (1 - \beta) \]

The first equation is known as a discrete Lyapunov equation, and can be solved using this function.

Next suppose that \( \{x_t\} \) is the discrete Markov process described above. Suppose further that each \( x_t \) takes values in the state space \( \{x_1, \ldots, x_N\} \subset \mathbb{R}^k \).

Let \( h: \mathbb{R}^k \to \mathbb{R} \) be a given function, and suppose that we wish to evaluate

\[ q(x_0) = \mathbb{E} \sum_{t=0}^{\infty} \beta^t h(x_t) \text{ given } x_0 = x^j \]

For example, in the discussion above, \( h(x_t) = x'_t H x_t \)

It is legitimate to pass the expectation through the sum, leading to

\[ q(x_0) = \sum_{t=0}^{\infty} \beta^t (P^t h)[j] \]  

(3.115)

Here

- \( P^t \) is the \( t \)-th power of the transition matrix \( P \)
- \( h \) is, with some abuse of notation, the vector \( (h(x^1), \ldots, h(x^N)) \)
- \( (P^t h)[j] \) indicates the \( j \)-th element of \( P^t h \)

It can be shown that (3.115) is in fact equal to the \( j \)-th element of the vector \( (I - \beta P)^{-1} h \)

This last fact is applied in the calculations below.

Other Variables We are interested in tracking several other variables besides the ones described above.

One is the present value of government obligations outstanding at time \( t \), which can be expressed as

\[ B_t := \mathbb{E}_t \sum_{j=0}^{\infty} \beta^j p_{t+j}^t \left( r_{t+j} - g_{t+j} \right) \]  

(3.116)

Using our expression for prices and the Ramsey plan, we can also write \( B_t \) as

\[ B_t = \mathbb{E}_t \sum_{j=0}^{\infty} \beta^j \frac{(b_{t+j} - c_{t+j})(\ell_{t+j} - g_{t+j}) - \ell^2_{t+j}}{b_t - c_t} \]
This variation is more convenient for computation
Yet another way to write $B_t$ is

$$B_t = \sum_{j=0}^{\infty} R_{tj}^{-1}(\tau_{t+j}\ell_{t+j} - g_{t+j})$$

where

$$R_{tj}^{-1} := E_t^p t^j$$

Here $R_{tj}$ can be thought of as the gross $j$-period risk-free rate on holding government debt between $t$ and $j$

Furthermore, letting $R_t$ be the one-period risk-free rate, we define

$$\pi_{t+1} := B_{t+1} - R_t[B_t - (\tau_t\ell_t - g_t)]$$

and

$$\Pi_t := \sum_{s=0}^{t} \pi_t$$

The term $\pi_{t+1}$ is the payout on the public's portfolio of government debt

As shown in the original manuscript, if we distort one-step-ahead transition probabilities by the adjustment factor

$$\xi_t := \frac{p_{t+1}^t}{E_t p_{t+1}^t}$$

then $\Pi_t$ is a martingale under the distorted probabilities

See the treatment in the manuscript for more discussion and intuition

For now we will concern ourselves with computation

**Implementation**

The following code provides functions for

1. Solving for the Ramsey plan given a specification of the economy
2. Simulating the dynamics of the major variables

The file is `examples/lqramsey.jl` from the main repository

Description and clarifications are given below

This module provides code to compute Ramsey equilibria in a LQ economy with distortionary taxation. The program computes allocations (consumption, leisure), tax rates, revenues, the net present value of the debt and other related quantities.

Functions for plotting the results are also provided below.

@author: Spencer Lyon <spencer.lyon@nyu.edu>
3.10. OPTIMAL TAXATION

@date: 2014-08-21

References
-------

Simple port of the file examples/lqramsey.py

http://quant-econ.net/lqramsey.html

#= using QuantEcon
  using PyPlot

abstract AbstractStochProcess

  type ContStochProcess <: AbstractStochProcess
    A::Matrix
    C::Matrix
  end

  type DiscreteStochProcess <: AbstractStochProcess
    P::Matrix
    x_vals::Array
  end

  type Economy{SP <: AbstractStochProcess}
    bet::Real
    Sg::Matrix
    Sd::Matrix
    Sb::Matrix
    Ss::Matrix
    is_discrete::Bool
    proc::SP
  end

  type Path
    g
db
sc
lp
tau
rvn
B
R
pi
Pi
function compute_exog_sequences(econ::Economy, x)
    # Compute exogenous variable sequences
    Sg, Sd, Sb, Ss = econ.Sg, econ.Sd, econ.Sb, econ.Ss
    g, d, b, s = [squeeze(S * x, 1) for S in (Sg, Sd, Sb, Ss)]

    #= Solve for Lagrange multiplier in the govt budget constraint
    In fact we solve for \( \nu = \lambda / (1 + 2\lambda) \). Here \( \nu \) is the
    solution to a quadratic equation \( a(\nu^2 - \nu) + b = 0 \) where
    \( a \) and \( b \) are expected discounted sums of quadratic forms of the state. =#
    Sm = Sb - Sd - Ss

    return g, d, b, s, Sm
end

function compute_allocation(econ::Economy, Sm, nu, x, b)
    Sg, Sd, Sb, Ss = econ.Sg, econ.Sd, econ.Sb, econ.Ss

    # Solve for the allocation given \( \nu \) and \( x \)
    Sc = 0.5 .* (Sb + Sd - Sg - nu .* Sm)
    Sl = 0.5 .* (Sb - Sd + Sg - nu .* Sm)
    c = squeeze(Sc * x, 1)
    l = squeeze(Sl * x, 1)
    p = squeeze((Sb - Sc) * x, 1) # Price without normalization
    tau = 1 .- l ./ (b .- c)
    rvn = l .* tau

    return Sc, Sl, c, l, p, tau, rvn
end

function compute_nu(a0, b0)
    disc = a0^2 - 4a0*b0

    if disc >= 0
        nu = 0.5 *(a0 - sqrt(disc)) / a0
    else
        println("There is no Ramsey equilibrium for these parameters.")
        error("Government spending (economy.g) too low")
    end

    # Test that the Lagrange multiplier has the right sign
    if nu * (0.5 - nu) < 0
        println("Negative multiplier on the government budget constraint.")
        error("Government spending (economy.g) too low")
    end

    return nu
end
function compute_Pi(B, R, rvn, g, xi)
    Pi = cumsum(pi .* xi)
    return pi, Pi
end

function compute_paths(econ::Economy{DiscreteStochProcess}, T)
    # simplify notation
    bet, Sg, Sd, Sb, Ss = econ.bet, econ.Sg, econ.Sd, econ.Sb, econ.Ss
    P, x_vals = econ.proc.P, econ.proc.x_vals

    state = mc_sample_path(P, 1, T)
    x = x_vals[:, state]

    # Compute exogenous sequence
    g, d, b, s, Sm = compute_exog_sequences(econ, x)

    # compute a0, b0
    ns = size(P, 1)
    F = eye(ns) - bet.*P
    a0 = (F \ ((Sm * x_vals)' * 2))[1] ./ 2
    H = ((Sb - Sd + Sg) * x_vals) .* ((Sg - Ss)*x_vals)
    b0 = (F \ H')[1] ./ 2

    # compute lagrange multiplier
    nu = compute_nu(a0, b0)

    # Solve for the allocation given nu and x
    Sc, Sl, c, l, p, tau, rvn = compute_allocation(econ, Sm, nu, x, b)

    # compute remaining variables
    H = ((Sb - Sc)*x_vals) .* ((Sl - Sg)*x_vals) - (Sl*x_vals).^2
    temp = squeeze(F*H', 2)
    B = temp[state] ./ p
    H = squeeze(P[state, :] * ((Sb - Sc)*x_vals)', 2)
    R = p ./ (bet .* H)
    temp = squeeze(P[state, :] *((Sb - Sc) * x_vals)', 2)
    xi = p[2:end] ./ temp[1:end-1]

    # compute pi
    pi, Pi = compute_Pi(B, R, rvn, g, xi)

    Path(g, d, b, s, c, l, p, tau, rvn, B, R, pi, Pi, xi)
end

function compute_paths(econ::Economy{ContStochProcess}, T)
    # simplify notation
    bet, Sg, Sd, Sb, Ss = econ.bet, econ.Sg, econ.Sd, econ.Sb, econ.Ss
# Generate an initial condition x0 satisfying x0 = A x0
mx, nx = size(A)
x0 = null((eye(mx) - A))
x0 = x0[end] < 0 ? -x0 : x0
x0 = x0 ./ x0[end]
x0 = squeeze(x0, 2)

# Generate a time series x of length T starting from x0
mx, nw = size(C)
x = zeros(nx, T)
w = randn(nw, T)
x[:, 1] = x0
for t=2:T
    x[:, t] = A * x[:, t-1] + C * w[:, t]
end

# Compute exogenous sequence
g, d, b, s, Sm = compute_exog_sequences(econ, x)

# compute a0 and b0
H = Sm'

a0 = 0.5 * var_quadratic_sum(A, C, H, bet, x0)
H = (Sb - Sd + Sg)'*(Sg + Ss)
b0 = 0.5 * var_quadratic_sum(A, C, H, bet, x0)

# compute lagrange multiplier
nu = compute_nu(a0, b0)

# Solve for the allocation given nu and x
Sc, Sl, c, l, p, tau, rvn = compute_allocation(econ, Sm, nu, x, b)

# compute remaining variables
H = Sl'Sl - (Sb - Sc) '* (Sl - Sg)
L = Array(Float64, T)
for t=1:T
    L[t] = var_quadratic_sum(A, C, H, bet, x[:, t])
end
B = L ./ p
Rinv = squeeze(bet .* (Sb- Sc)*A*x[:, 1] ./ p
R = 1 ./ Rinv
AF1 = (Sb - Sc) * x[:, 2:end]
AF2 = (Sb - Sc) * A * x[:, 1:end-1]
xi = AF1 ./ AF2
xi = squeeze(xi, 1)

# compute pi
pi, Pi = compute_Pi(B, R, rvn, g, xi)

Path(g, d, b, s, c, l, p, tau, rvn, B, R, pi, Pi, xi)
end

function gen_fig_1(path::Path)
T = length(path.c)

num_rows, num_cols = 2, 2
fig, axes = subplots(num_rows, num_cols, figsize=(14, 10))
plt.subplots_adjust(hspace=0.4)
for i=1:num_rows
    for j=1:num_cols
        axes[i, j][[:grid]]()
        axes[i, j][[:set_xlabel]]("Time")
    end
end
bbox = (0., 1.02, 1., .102)
legend_args = {:bbox_to_anchor => bbox, :loc => 3, :mode => :expand}
p_args = {:lw => 2, :alpha => 0.7}

# Plot consumption, govt expenditure and revenue
ax = axes[1, 1]
ax[[:plot]](path.rvn, label="$\tau_t \ell_t$"; p_args...)
ax[[:plot]](path.g, label="$g_t$"; p_args...)
ax[[:plot]](path.c, label="$c_t$"; p_args...)
ax[[:legend]](ncol=3; legend_args...)

# Plot govt expenditure and debt
ax = axes[1, 2]
ax[[:plot]](1:T, path.rvn, label="$\tau_t \ell_t$"; p_args...)
ax[[:plot]](1:T, path.g, label="$g_t$"; p_args...)
ax[[:plot]](1:T-1, path.B[2:end], label="$B_{t+1}$"; p_args...)
ax[[:legend]](ncol=3; legend_args...)

# Plot risk free return
ax = axes[2, 1]
ax[[:plot]](1:T, path.R - 1, label="$R_{t - 1}$"; p_args...)
ax[[:legend]](ncol=1; legend_args...)

# Plot revenue, expenditure and risk free rate
ax = axes[2, 2]
ax[[:plot]](1:T, path.rvn, label="$\tau_t \ell_t$"; p_args...)
ax[[:plot]](1:T, path.g, label="$g_t$"; p_args...)
ax[[:plot]](1:T-1, path.pi, label="$\pi_{t+1}$"; p_args...)
ax[[:legend]](ncol=3; legend_args...)

plt.show()

end

function gen_fig_2(path::Path)
    T = length(path.c)

    # Prepare axes
    num_rows, num_cols = 2, 1
fig, axes = subplots(num_rows, num_cols, figsize=(10, 10))
plt.subplots_adjust(hspace=0.5)
bbox = (0., 1.02, 1., .102)
legend_args = {:bbox_to_anchor => bbox, :loc => 3, :mode => :expand}
p_args = {:lw => 2, :alpha => 0.7}

# Plot adjustment factor
ax = axes[1]
ax[:plot](2:T, path.xi, label=L"\xi_t"; p_args...)
ax[:grid]()
ax[:set_xlabel]("Time")
ax[:legend](ncol=1; legend_args...)

# Plot adjusted cumulative return
ax = axes[2]
ax[:plot](2:T, path.Pi, label=L"\Pi_t"; p_args...)
ax[:grid]()
ax[:set_xlabel]("Time")
ax[:legend](ncol=1; legend_args...)

plt.show()

end

Comments on the Code The function var_quadratic_sum From QuantEcon.jl is for computing the value of (3.114) when the exogenous process \{x_t\} is of the VAR type described above. This code defines two Types: Economy and Path. The first is used to collect all the parameters and primitives of a given LQ economy, while the second collects output of the computations.

Examples

Let’s look at two examples of usage.

The Continuous Case Our first example adopts the VAR specification described above. Regarding the primitives, we set

- \( \beta = 1/1.05 \)
- \( b_t = 2.135 \) and \( s_t = d_t = 0 \) for all \( t \)

Government spending evolves according to

\[ g_{t+1} - \mu_g = \rho (g_t - \mu_g) + C_g w_{g,t+1} \]

with \( \rho = 0.7, \mu_g = 0.35 \) and \( C_g = \mu_g \sqrt{1 - \rho^2}/10 \)

Here’s the code, from file examples/lqramsey_ar1.jl.
Example 1: Govt spending is AR(1) and state is (g, i).

@author: Spencer Lyon <spencer.lyon@nyu.edu>

@date: 2014-08-21

References

Simple port of the file examples/lqramsey_ar1.py

http://quant-econ.net/lqramsey.html

---

include("lqramsey.jl")

# == Parameters == #

bet = 1 / 1.05
rho, mg = .7, .35
A = eye(2)
A = [rho mg*(1 - rho); 0.0 1.0]
C = [sqrt(1 - rho^2)*mg/10 0.0]
Sg = [1.0 0.0]
Sd = [0.0 0.0]
Sb = [0 2.135]
Ss = [0.0 0.0]
discrete = false
proc = ContStochProcess(A, C)

econ = Economy(bet, Sg, Sd, Sb, Ss, discrete, proc)
T = 50

path = compute_paths(econ, T)

Running the program produces the figure

The legends on the figures indicate the variables being tracked

Most obvious from the figure is tax smoothing in the sense that tax revenue is much less variable than government expenditure

After running the code above, if you then execute gen_fig_2(path) from your Julia console you will produce the figure

See the original manuscript for comments and interpretation

The Discrete Case  Our second example adopts a discrete Markov specification for the exogenous process

Here’s the code, from file examples/lqramsey_discrete.jl
Example 2: LQ Ramsey model with discrete exogenous process.

@author: Spencer Lyon <spencer.lyon@nyu.edu>

@date: 2014-08-21

References
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Simple port of the file examples/lqramsey_discrete.py

http://quant-econ.net/lqramsey.html

include("lqramsey.jl")

# Parameters
bet = 1 / 1.05
P = [0.8 0.2 0.0
     0.0 0.5 0.5
     0.0 0.0 1.0]

# Possible states of the world
# Each column is a state of the world. The rows are [g d b s t]
x_vals = [0.5 0.5 0.25
          0.0 0.0 0.0
          2.2 2.2 2.2
          0.0 0.0 0.0
          1.0 1.0 1.0]

Sg = [1.0 0.0 0.0 0.0 0.0]
Sd = [0.0 1.0 0.0 0.0 0.0]
Sb = [0.0 0.0 1.0 0.0 0.0]
Ss = [0.0 0.0 0.0 1.0 0.0]

discrete = true
proc = DiscreteStochProcess(P, x_vals)

econ = Economy(bet, Sg, Sd, Sb, Ss, discrete, proc)
T = 15

path = compute_paths(econ, T)

The call gen_fig_1(path) generates the figure
while gen_fig_2(path) generates
See the original manuscript for comments and interpretation
Exercises

Exercise 1  Modify the VAR example given above, setting

\[ g_{t+1} - \mu_g = \rho (g_{t-3} - \mu_g) + C_g w_{g,t+1} \]

with \( \rho = 0.95 \) and \( C_g = 0.7 \sqrt{1 - \rho^2} \)

Produce the corresponding figures

Solutions

Solution notebook

3.11 History Dependent Public Policies

Contents

- History Dependent Public Policies
  - Overview
  - Two Sources of History Dependence
  - Competitive equilibrium
  - Ramsey Problem
  - Time Inconsistency
  - Concluding remarks

Overview

This lecture describes history-dependent public policies and some of their representations

History dependent policies are decision rules that depend on the entire past history of the state variables

History dependent policies naturally emerge in Ramsey problems

A Ramsey planner (typically interpreted as a government) devises a plan of actions at time \( t = 0 \) to follow at all future dates and for all contingencies

In order to make a plan, he takes as given Euler equations expressing private agents’ first-order necessary conditions

He also takes into account that his future actions affect earlier decisions by private agents, an avenue opened up by the maintained assumption of rational expectations

Another setting in which history dependent policies naturally emerge is where instead of a Ramsey planner there is a sequence of government administrators whose time \( t \) member takes as given the policies used by its successors
We study these ideas in the context of a model in which a benevolent tax authority is forced
- to raise a prescribed present value of revenues
- to do so by imposing a distorting flat rate tax on the output of a competitive representative
  firm
The firm faces costs of adjustment and lives within a competitive equilibrium, which in turn im-
poses restrictions on the tax authority \(^1\)

**References**  The presentation below is based on a recent paper by Evans and Sargent [ES13]
Regarding techniques, we will make use of the methods described in
1. the linear regulator lecture
2. the upcoming lecture on solving linear quadratic Stackelberg models

**Two Sources of History Dependence**

We compare two timing protocols
1. An infinitely lived benevolent tax authority solves a Ramsey problem
2. There is a sequence of tax authorities, each choosing only a time \(t\) tax rate
Under both timing protocols, optimal tax policies are *history-dependent*
But history dependence captures different economic forces across the two timing protocols
In the first timing protocol, history dependence expresses the *time-inconsistency of the Ramsey plan*
In the second timing protocol, history dependence reflects the unfolding of constraints that assure
that a time \(t\) government wants to confirm the representative firm’s expectations about govern-
ment actions
We describe recursive representations of history-dependent tax policies under both timing proto-
ocols

**Ramsey Timing Protocol**  The first timing protocol models a policy maker who can be said to
‘commit’, choosing a sequence of tax rates once-and-for-all at time 0
To obtain a *recursive* representation of a Ramsey policy, we compare two methods
We first apply a method proposed by Kydland and Prescott [KP80] that uses a promised marginal
utility to augment natural state variables
We then apply a closely related method due to [MS85], [PCL86], and [BD86]
This method uses a ‘co-state on a co-state’ variable to augment the authentic state variables
After applying both methods, we describe links between them and confirm that they recover the
same Ramsey plan

\(^1\) We could also call a competitive equilibrium a rational expectations equilibrium.
Sequence of Governments Timing Protocol  For the second timing protocol we use the notion of a sustainable plan proposed in [CK90], also referred to as a credible public policy in [Sto89]

A key idea here is that history-dependent policies can be arranged so that, when regarded as a representative firm’s forecasting functions, they confront policy makers with incentives to confirm them

We follow Chang [Cha98] in expressing such history-dependent plans recursively

Credibility considerations contribute an additional auxiliary state variable in the form of a promised value to the planner

It expresses how decisions must unfold to give the government the incentive to confirm private sector expectations when the government chooses sequentially

Note: We occasionally hear confusion about the consequences of recursive representations of government policies under our two timing protocols. It is incorrect to regard a recursive representation of the Ramsey plan as in any way ‘solving a time-inconsistency problem’. On the contrary, the evolution of the auxiliary state variable that augments the authentic ones under our first timing protocol ought to be viewed as expressing the time-inconsistency of a Ramsey plan. Despite that, in literatures about practical monetary policy one sometimes hears interpretations that sell Ramsey plans in settings where our sequential timing protocol is the one that more accurately characterizes decision making. Please beware of discussions that toss around claims about credibility if you don’t also see recursive representations of policies with the complete list of state variables appearing in our [Cha98] -like analysis that we present below.

Competitive equilibrium

A representative competitive firm sells output $q_t$ at price $p_t$ when market-wide output is $Q_t$

The market as a whole faces a downward sloping inverse demand function

$$ p_t = A_0 - A_1 Q_t, \quad A_0 > 0, A_1 > 0 $$

(3.117)

The representative firm

• has given initial condition $q_0$

• endures quadratic adjustment costs $\frac{d}{2}(q_{t+1} - q_t)^2$

• pays a flat rate tax $\tau_t$ per unit of output

• treats $\{p_t, \tau_t\}_{t=0}^{\infty}$ as exogenous

• chooses $\{q_{t+1}\}_{t=0}^{\infty}$ to maximize

$$ \sum_{t=0}^{\infty} \beta^t \{ p_t q_t - \frac{d}{2} (q_{t+1} - q_t)^2 - \tau_t q_t \} $$

(3.118)

Let $u_t := q_{t+1} - q_t$ be the firm’s ‘control variable’ at time $t$

First-order conditions for the representative firm’s problem are

$$ u_t = \frac{\beta}{d} p_{t+1} + \beta u_{t+1} - \frac{\beta}{d} \tau_{t+1}, \quad t = 0, 1, \ldots $$

(3.119)
To compute a competitive equilibrium, it is appropriate to take (3.119), eliminate $p_t$ in favor of $Q_t$ by using (3.117), and then set $q_t = Q_t$.

This last step makes the representative firm be representative $^2$

We arrive at

$$u_t = \frac{\beta}{d} (A_0 - A_1 Q_{t+1}) + \beta u_{t+1} - \frac{\beta}{d} \tau_{t+1}$$  \hspace{1cm} (3.120)

$$Q_{t+1} = Q_t + u_t$$  \hspace{1cm} (3.121)

**Notation:** For any scalar $x_t$, let $\vec{x} = \{x_t\}_{t=0}^{\infty}$

Given a tax sequence $\{\tau_{t+1}\}_{t=0}^{\infty}$, a competitive equilibrium is a price sequence $\vec{p}$ and an output sequence $\vec{Q}$ that satisfy (3.117), (3.120), and (3.121).

For any sequence $\vec{x} = \{x_t\}_{t=0}^{\infty}$, the sequence $\vec{x}_1 := \{x_t\}_{t=1}^{\infty}$ is called the continuation sequence or simply the continuation.

Note that a competitive equilibrium consists of a first period value $u_0 = Q_1 - Q_0$ and a continuation competitive equilibrium with initial condition $Q_1$.

Also, a continuation of a competitive equilibrium is a competitive equilibrium.

Following the lead of [Cha98], we shall make extensive use of the following property:

- A continuation $\vec{\tau}_1 = \{\tau_{t+1}\}_{t=1}^{\infty}$ of a tax policy $\vec{\tau}$ influences $u_0$ via (3.120) entirely through its impact on $u_1$.

A continuation competitive equilibrium can be indexed by a $u_1$ that satisfies (3.120).

With some abuse of language, in the spirit of [KP80], we shall use $u_{t+1}$ to describe what we shall call a promised marginal value that a competitive equilibrium offers to a representative firm $^3$

Define $\vec{Q} := [Q_0, \ldots, Q_t]$.

A history-dependent tax policy is a sequence of functions $\{\sigma_t\}_{t=0}^{\infty}$ with $\sigma_t$ mapping $\vec{Q}$ into a choice of $\tau_{t+1}$.

Below, we shall

- Study history-dependent tax policies that either solve a Ramsey plan or are credible.
- Describe recursive representations of both types of history-dependent policies.

**Ramsey Problem**

The planner’s objective is cast in terms of consumer surplus net of the firm’s adjustment costs.

$^2$ It is important not to set $q_t = Q_t$ prematurely. To make the firm a price taker, this equality should be imposed after and not before solving the firm’s optimization problem.

$^3$ We could instead, perhaps with more accuracy, define a promised marginal value as $\beta(A_0 - A_1 Q_{t+1}) - \beta \tau_{t+1} + u_{t+1}/\beta$, since this is the object to which the firm’s first-order condition instructs it to equate to the marginal cost $du_t$ of $u_t = q_{t+1} - q_t$. This choice would align better with how Chang [Cha98] chose to express his competitive equilibrium recursively. But given $(u_t, Q_t)$, the representative firm knows $(Q_{t+1}, \tau_{t+1})$, so it is adequate to take $u_{t+1}$ as the intermediate variable that summarizes how $\tau_{t+1}$ affects the firm’s choice of $u_t$. 

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**THOMAS SARGENT AND JOHN STACHURSKI**  
March 21, 2015
Consumer surplus is
\[
\int_0^Q (A_0 - A_1 x) dx = A_0 Q - \frac{A_1}{2} Q^2
\]
Hence the planner’s one-period return function is
\[
A_0 Q_t - \frac{A_1}{2} Q_t^2 - \frac{d}{2} u_t^2
\]
At time \( t = 0 \), a Ramsey planner faces the intertemporal budget constraint
\[
\sum_{t=1}^{\infty} \beta^t \tau_t Q_t = G_0
\]
Note that (3.123) precludes taxation of initial output \( Q_0 \)

The **Ramsey problem** is to choose a tax sequence \( \tau \) and a competitive equilibrium outcome \( (\bar{Q}, \bar{u}) \) that maximize
\[
\sum_{t=0}^{\infty} \beta^t \left\{ A_0 Q_t - \frac{A_1}{2} Q_t^2 - \frac{d}{2} u_t^2 \right\}
\]
subject to (3.123)

Thus, the Ramsey timing protocol is:

1. At time 0, knowing \( (Q_0, G_0) \), the Ramsey planner chooses \( \{\tau_{t+1}\}_{t=0}^{\infty} \)
2. Given \( (Q_0, \{\tau_{t+1}\}_{t=0}^{\infty}) \), a competitive equilibrium outcome \( \{u_t, Q_{t+1}\}_{t=0}^{\infty} \) emerges

**Note:** In bringing out the timing protocol associated with a Ramsey plan, we run head on into a set of issues analyzed by Bassetti [Bas05]. This is because our definition of the Ramsey timing protocol doesn’t completely describe all conceivable actions by the government and firms as time unfolds. For example, the definition is silent about how the government would respond if firms, for some unspecified reason, were to choose to deviate from the competitive equilibrium associated with the Ramsey plan, possibly prompting violation of government budget balance. This is an example of the issues raised by [Bas05], who identifies a class of government policy problems whose proper formulation requires supplying a complete and coherent description of all actors’ behavior across all possible histories. Implicitly, we are assuming that a more complete description of a government strategy could be specified that (a) agrees with ours along the Ramsey outcome, and (b) suffices uniquely to implement the Ramsey plan by deterring firms from taking actions that deviate from the Ramsey outcome path.

**Computing a Ramsey Plan** The planner chooses \( \{u_t\}_{t=0}^{\infty}, \{\tau_t\}_{t=1}^{\infty} \) to maximize (3.124) subject to (3.120), (3.121), and (3.123)

To formulate this problem as a Lagrangian, attach a Lagrange multiplier \( \mu \) to the budget constraint (3.123)

Then the planner chooses \( \{u_t\}_{t=0}^{\infty}, \{\tau_t\}_{t=1}^{\infty} \) to maximize and the Lagrange multiplier \( \mu \) to minimize
\[
\sum_{t=0}^{\infty} \beta^t (A_0 Q_t - \frac{A_1}{2} Q_t^2 - \frac{d}{2} u_t^2) + \mu \left[ \sum_{t=0}^{\infty} \beta^t \tau_t Q_t - G_0 - \tau_0 Q_0 \right]
\]
subject to and (3.120) and (3.121)
**Implementability Multiplier Approach**  
The Ramsey problem is a special case of the linear quadratic dynamic Stackelberg problem analyzed in the Stackelberg lecture.

The idea is to construct a recursive representation of a Ramsey plan by including among the state variables Lagrange multipliers on implementability constraints.

These multipliers require the Ramsey planner to choose among competitive equilibrium allocations.

Their motions through time become components of a recursive representation of a history-dependent plan for taxes.

For us, the key implementability conditions are (3.120) for $t \geq 0$.

Holding fixed $\mu$ and $G_0$, the Lagrangian for the planning problem can be abbreviated as

$$
\max_{\{u_t, \tau_{t+1}\}} \sum_{t=0}^{\infty} \beta^t \left\{ A_0 Q_t - \frac{A_1}{2} Q^2_t - \frac{d}{2} u^2_t + \mu \tau_t Q_t \right\}
$$

Define

$$
z_t := \begin{bmatrix} 1 \\ Q_t \\ \tau_t \end{bmatrix} \quad \text{and} \quad y_t := \begin{bmatrix} z_t \\ u_t \end{bmatrix} = \begin{bmatrix} 1 \\ Q_t \\ \tau_t \\ u_t \end{bmatrix}
$$

Here the elements of $z_t$ are genuine state variables and $u_t$ is a jump variable.

We include $\tau_t$ as a state variable for bookkeeping purposes: it helps to map the problem into a linear regulator problem with no cross products between states and controls.

However, it will be a redundant state variable in the sense that the optimal tax $\tau_{t+1}$ will not depend on $\tau_t$.

The government chooses $\tau_{t+1}$ at time $t$ as a function of the time $t$ state.

Thus, we can rewrite the Ramsey problem as

$$
\max_{\{y_t, \tau_{t+1}\}} -\sum_{t=0}^{\infty} \beta^t y_t^T R y_t
$$

subject to $z_0$ given and the law of motion

$$
y_{t+1} = Ay_t + B \tau_{t+1}
$$

where

$$
R = \begin{bmatrix} 0 & -\frac{A_0}{2} & 0 & 0 \\ -\frac{A_0}{2} & \frac{A_1}{2} & \frac{-\mu}{2} & 0 \\ 0 & \frac{-\mu}{2} & 0 & 0 \\ 0 & 0 & 0 & \frac{d}{2} \end{bmatrix}, \quad A = \begin{bmatrix} 1 & 0 & 0 & 0 \\ 0 & 1 & 0 & 1 \\ 0 & 0 & 0 & 0 \\ -\frac{A_0}{d} & \frac{A_1}{d} & 0 & \frac{A_1}{d} + \frac{1}{p} \end{bmatrix}, \quad B = \begin{bmatrix} 0 \\ 0 \\ 1 \\ \frac{1}{p} \end{bmatrix}
$$

Because this problem falls within the framework, we can proceed as follows.

Letting $\lambda_t$ be a vector of Lagrangian multipliers on the transition laws summarized in (3.127), it follows that

- $\lambda_t = Py_t$, where $P$ solves the *Riccati equation* $P = R + \beta A'PA - \beta A'PB(B'PB)^{-1}B'PA$.
• \( \tau_{t+1} = -Fy_t \), where \( F \) is the associated policy matrix \( F = (B'PB)^{-1}B'PA' \)

We can rewrite \( \lambda_t = Py_t \) as

\[
\begin{bmatrix}
\lambda_{zt} \\
\lambda_{ut}
\end{bmatrix} =
\begin{bmatrix}
P_{11} & P_{12} \\
P_{21} & P_{22}
\end{bmatrix}
\begin{bmatrix}
z_t \\
u_t
\end{bmatrix}
\]

Solve for \( u_t \) to get

\[ u_t = -P_{22}^{-1}P_{21}z_t + P_{22}^{-1}\lambda_{ut} \]

Now the multiplier \( \lambda_{ut} \) becomes our authentic state variable — one that measures the cost to the government of confirming the representative firm’s prior expectations about time \( t \) government actions

The complete state at time \( t \) becomes \( [z_t \lambda_{ut}] \), and

\[ y_t = \begin{bmatrix} z_t \\ u_t \end{bmatrix} = \begin{bmatrix} I & 0 \\ -P_{22}^{-1}P_{21} & P_{22}^{-1} \end{bmatrix} \begin{bmatrix} z_t \\ \lambda_{ut} \end{bmatrix} \]

so

\[ \tau_{t+1} = -F \begin{bmatrix} I & 0 \\ -P_{22}^{-1}P_{21} & P_{22}^{-1} \end{bmatrix} \begin{bmatrix} z_t \\ \lambda_{ut} \end{bmatrix} \]

The evolution of the state is

\[
\begin{bmatrix}
z_{t+1} \\
\lambda_{ut+1}
\end{bmatrix} =
\begin{bmatrix}
I & 0 \\
P_{21} & P_{22}
\end{bmatrix}
\begin{bmatrix}
A - BF
\end{bmatrix}
\begin{bmatrix}
I & 0 \\
-P_{22}^{-1}P_{21} & P_{22}^{-1}
\end{bmatrix}
\begin{bmatrix}
z_t \\
\lambda_{ut}
\end{bmatrix}
\]

with initial state

\[
\begin{bmatrix}
z_0 \\
\lambda_{u0}
\end{bmatrix} =
\begin{bmatrix}
1 \\
Q_0 \\
\tau_0 \\
0
\end{bmatrix}
\tag{3.128}
\]

Equation (3.128) incorporates the finding that the Ramsey planner finds it optimal to set \( \lambda_{u0} \) to zero

**Kydland-Prescott Approach**  Kydland and Prescott [KP80] or Chang [Cha98] would formulate our Ramsey problem in terms of the Bellman equation

\[ v(Q_t, \tau_t, u_t) = \max_{\tau_{t+1}} \left\{ A_0 Q_t - \frac{A_1}{2} Q_t^2 - \frac{d}{2} u_t^2 + \mu \tau_t Q_t + \beta v(Q_{t+1}, \tau_{t+1}, u_{t+1}) \right\} \]

where the maximization is subject to the constraints

\[ Q_{t+1} = Q_t + u_t \]

and

\[ u_{t+1} = -\frac{A_0}{d} + \frac{A_1}{d} Q_t + \frac{A_1}{d} + \frac{1}{\beta} u_t + \frac{1}{d} \tau_{t+1} \]

We now regard \( u_t \) as a state

It plays the role of a promised marginal utility in the Kydland-Prescott framework
Define the state vector to be

\[ y_t = \begin{bmatrix} 1 \\ Q_t \\ \tau_t \\ u_t \end{bmatrix} = \begin{bmatrix} z_t \\ u_t \end{bmatrix}, \]

where \( z_t = [1 \; Q_t \; \tau_t]' \) are authentic state variables and \( u_t \) is a variable whose time 0 value is a 'jump' variable but whose values for dates \( t \geq 1 \) will become state variables that encode history dependence in the Ramsey plan.

Write a dynamic programming problem in the style of [KP80] as

\[ v(y_t) = \max_{\tau_{t+1}} \{-y_t' R y_t + \beta v(y_{t+1})\} \tag{3.129} \]

where the maximization is subject to the constraint

\[ y_{t+1} = A y_t + B \tau_{t+1} \]

and where

\[
R = \begin{bmatrix}
0 & -\frac{\mu}{2} & 0 & 0 \\
-\frac{\mu}{2} & -\frac{\mu}{2} & 0 & 0 \\
0 & 0 & 0 & \frac{\beta}{2} \\
0 & 0 & 0 & \frac{\beta}{2}
\end{bmatrix},
\quad
A = \begin{bmatrix}
1 & 0 & 0 & 0 \\
0 & 1 & 0 & 1 \\
0 & 0 & 0 & 0 \\
-\frac{\mu}{\beta} & \frac{\mu}{\beta} & 0 & \frac{\mu}{\beta} + \frac{1}{\beta}
\end{bmatrix},
\quad
B = \begin{bmatrix} 0 \\ 0 \end{bmatrix}.
\]

Functional equation (3.129) has solution

\[ v(y_t) = -y_t' P y_t \]

where

- \( P \) solves the algebraic matrix Riccati equation \( P = R + \beta A' PA - \beta A' PB (B' PB)^{-1} B'PA \)
- the optimal policy function is given by \( \tau_{t+1} = -F y_t \) for \( F = (B' PB)^{-1} B'PA \)

Since the formulas for \( A, B, \) and \( R \) are identical, it follows that \( F \) and \( P \) are the same as in the Lagrangian multiplier approach of the preceding section.

The optimal choice of \( u_0 \) satisfies \( \frac{\partial v}{\partial u_0} = 0 \)

If we partition \( P \) as

\[ P = \begin{bmatrix} P_{11} & P_{12} \\ P_{21} & P_{22} \end{bmatrix} \]

then we have

\[ 0 = \frac{\partial}{\partial u_0} (z_0' P_{11} z_0 + z_0' P_{12} u_0 + u_0' P_{21} z_0 + u_0' P_{22} u_0) = P_{11}' z_0 + P_{21} u_0 + 2P_{22} u_0 \]

which implies

\[ u_0 = -P_{22}^{-1} P_{21} z_0 \tag{3.130} \]

Thus, the Ramsey plan is

\[ \tau_{t+1} = -F \begin{bmatrix} z_t \\ u_t \end{bmatrix} \quad \text{and} \quad \begin{bmatrix} z_{t+1} \\ u_{t+1} \end{bmatrix} = (A - BF) \begin{bmatrix} z_t \\ u_t \end{bmatrix} \]

with initial state \( \begin{bmatrix} z_0 & -P_{22}^{-1} P_{21} z_0 \end{bmatrix}' \).
Comparison  We can compare the outcome from the Kydland-Prescott approach to the outcome of the Lagrangian multiplier on the implementability constraint approach of the preceding section.

Using the formula
$$
\begin{bmatrix}
z_t \\
u_t
\end{bmatrix} =
\begin{bmatrix}
I & 0 \\
-P_{21}^{-1}P_{22} & P_{22}^{-1}
\end{bmatrix}
\begin{bmatrix}
z_t \\
\lambda_{ut}
\end{bmatrix}
$$
and applying it to the evolution of the state
$$
\begin{bmatrix}
z_{t+1} \\
\lambda_{ut+1}
\end{bmatrix} =
\begin{bmatrix}
I & 0 \\
P_{21} & P_{22}
\end{bmatrix}
(A - BF)
\begin{bmatrix}
z_t \\
\lambda_{ut}
\end{bmatrix}
$$
we get
$$
\begin{bmatrix}
z_{t+1} \\
u_{t+1}
\end{bmatrix} =
(A - BF)
\begin{bmatrix}
z_t \\
u_t
\end{bmatrix}
$$
(3.131)
or $y_{t+1} = AFy_t$ where $AF := A - BF$.

Then using the initial state value $\lambda_{u,0} = 0$, we obtain
$$
\begin{bmatrix}
z_0 \\
u_0
\end{bmatrix} =
\begin{bmatrix}
z_0 \\
-P_{22}^{-1}P_{22}z_0
\end{bmatrix}
$$
(3.132)
This is identical to the initial state delivered by the Kydland-Prescott approach.

Recursive Representation  An outcome of the preceding results is that the Ramsey plan can be represented recursively as the choice of an initial marginal utility (or rate of growth of output) according to a function
$$
u_0 = v(Q_0|\mu)
$$
(3.133)
that obeys (3.132) and the following updating equations for $t \geq 0$:
$$
\tau_{t+1} = \tau(Q_t, u_t|\mu)
$$
(3.134)
$$
Q_{t+1} = Q_t + u_t
$$
(3.135)
$$
u_{t+1} = u(Q_t, u_t|\mu)
$$
(3.136)
We have conditioned the functions $\nu$, $\tau$, and $u$ by $\mu$ to emphasize how the dependence of $F$ on $G_0$ appears indirectly through the Lagrange multiplier $\mu$.

An Example Calculation  We’ll discuss how to compute $\mu$ below but first consider the following numerical example.

We take the parameter set $[A_0, A_1, d, \beta, Q_0] = [100, .05, .2, .95, 100]$ and compute the Ramsey plan with the following piece of code.
In the following, `\( \hat{u} \)` and `\( \hat{\tau} \)` are what the planner would choose if he could reset at time \( t \), `\( \hat{u}_{\text{dif}} \)` and `\( \hat{\tau}_{\text{dif}} \)` are the difference between those and what the planner is constrained to choose. The variable `\( \mu \)` is the Lagrange multiplier associated with the constraint at time \( t \).

For more complete description of inputs and outputs see the website.

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@date: 2014-08-21

References
--------

Simple port of the file examples/evans_sargent.py

http://quant-econ.net/hist_dep_policies.html

```python
#
using QuantEcon
using Optim
using PyPlot

type HistDepRamsey
   # These are the parameters of the economy
   A0::Real
   A1::Real
   d::Real
   Q0::Real
   tau0::Real
   mu0::Real
   bet::Real

   # These are the LQ fields and stationary values
   R::Matrix
   A::Matrix
   B::Matrix
   Q::Real
   P::Matrix
   F::Matrix
   lq::LQ
end

type RamseyPath
   y::Matrix
   uhat::Vector
   uhatdif::Vector
   tauhat::Vector
   tauhatdif::Vector
```
function HistDepRamsey(A0, A1, d, Q0, tau0, mu, bet)
    # Create Matrices for solving Ramsey problem
    R = [0.0, -A0/2, 0.0, 0.0, -A0/2, A1/2, -mu/2, 0.0, 0.0, -mu/2, 0.0, 0.0, 0.0, 0.0, d/2]
    A = [1.0, 0.0, 0.0, 0.0, 0.0, 1.0, 0.0, 0.0, 0.0, 0.0, 0.0, 0.0, 0.0, A1/d, A1/d, A1/d+1.0/bet]
    B = [0.0, 0.0, 1.0, 1.0/d]
    Q = 0.0
    # Use LQ to solve the Ramsey Problem.
    lq = LQ(Q, -R, A, B, bet=bet)
    P, F, _d = stationary_values(lq)
    HistDepRamsey(A0, A1, d, Q0, tau0, mu0, bet, R, A, B, Q, P, F, lq)
end

function compute_G(hdr::HistDepRamsey, mu)
    # simplify notation
    bet = hdr.bet
    # Need y_0 to compute government tax revenue.
    u0 = compute_u0(hdr, P)
    y0 = vcat([1.0 Q0 tau0]', u0)
    # Define A_F and S matrices
    AF = A - B * F
    S = [0.0, 1.0, 0.0, 0] * [0.0, 0.0, 1.0, 0]
    # Solves equation (25)
    Omega = solve_discrete_lyapunov(sqrt(bet) .* AF', bet .* AF' * S * AF)
    T0 = y0' * Omega * y0
    return T0[1], A, B, F, P
end

function compute_u0(hdr::HistDepRamsey, P::Matrix)

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# simplify notation
Q0, tau0 = hdr.Q0, hdr.tau0

P21 = P[4, 1:3]
P22 = P[4, 4]
z0 = [1.0 Q0 tau0]'
u0 = -P22^(-1) .* P21*(z0)

return u0[1]
end

function init_path(hdr::HistDepRamsey, mu0, T::Int=20)
    # Construct starting values for the path of the Ramsey economy
    G0, A, B, F, P = compute_G(hdr, mu0)
    # Compute the optimal u0
    u0 = compute_u0(hdr, P)
    # Initialize vectors
    y = Array(Float64, 4, T)
uhat = Array(Float64, T)
uhatdif = Array(Float64, T)
tauhat = Array(Float64, T)
tauhatdif = Array(Float64, T-1)
mu = Array(Float64, T)
G = Array(Float64, T)
GPay = Array(Float64, T)

    # Initial conditions
    G[1] = G0
    mu[1] = mu0
    uhatdif[1] = 0.0
    uhat[1] = u0
    y[:, 1] = vcat([1.0 hdr.Q0 hdr.tau0]', u0)

    return RamseyPath(y, uhat, uhatdif, tauhat, tauhatdif, mu, G, GPay)
end

function compute_ramsey_path!(hdr::HistDepRamsey, rp::RamseyPath)
    # simplify notation
    y, uhat, uhatdif, tauhat, = rp.y, rp.uhat, rp.uhatdif, rp.tauhat
tauhatdif, mu, G, GPay = rp.tauhatdif, rp.mu, rp.G, rp.GPay
    bet = hdr.bet

    G0, A, B, F, P = compute_G(hdr, mu[1])

    for t=2:T
        # iterate government policy
        y[:, t] = (A - B * F) * y[:, t-1]
    end

    return rp
end
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```python
# update G
G[t] = (G[t-1] - bet*y[2, t]*y[3, t])/bet
GPay[t] = bet.*y[2, t]*y[3, t]

# Compute the mu if the government were able to reset its plan
if is the tax revenues the government would receive if they reset the
plan with Lagrange multiplier mu minus current G
ff(mu) = abs(compute_G(hdr, mu)[1]-G[t])

# find ff = 0
mu[t] = optimize(ff, mu[t-1]-1e4, mu[t-1]+1e4).minimum

# Compute alternative decisions
P21temp = Ptemp[4, 1:3]
P22temp = P[4, 4]
what[t] = (-P22temp^(-1) .* P21temp * y[1:3, t])[1]

yhat = (Atemp-Btemp * Ftemp) * [y[1:3, t-1], what[t-1]]
tauhat[t] = yhat[3]
tauhatdif[t-1] = tauhat[t] - y[3, t]
whatdif[t] = what[t] - y[3, t]

end

return rp
end

function plot1(rp::RamseyPath)
    tt = 1:length(rp.mu)  # it is used to make the plot time index correct.
y = rp.y

    n_rows = 3
    fig, axes = subplots(n_rows, 1, figsize=(10, 12))
    subplots_adjust(hspace=0.5)
    for ax in axes
        ax[:grid]()
        ax[:set_xlim](0, 15)
    end

    bbox = (0., 1.02, 1., .102)
    legend_args = {:bbox_to_anchor => bbox, :loc => 3, :mode => "expand"}
    p_args = {:lw => 2, :alpha => 0.7}

    ax = axes[1]
    ax[:plot](tt, squeeze(y[2, :], 1), "b-", label="output"; p_args...)
    ax[:set_ylabel]([$Q$], fontsize=16)
    ax[:legend](ncol=1; legend_args...)

    ax = axes[2]
```

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```python
ax[:plot](tt, squeeze(y[3, :], 1), "b-", label="tax rate"; p_args...)
ax[:set_ylabel](L"$\tau$", fontsize=16)
ax[:set_yticks]((0.0, 0.2, 0.4, 0.6, 0.8))
ax[:legend](ncol=1; legend_args...)

ax = axes[3]
ax[:plot](tt, squeeze(y[4, :], 1), "b-", label="first difference in output";
p_args...)
ax[:set_ylabel](L"$u$", fontsize=16)
ax[:set_yticks]((0, 100, 200, 300, 400))
ax[:legend](ncol=1; legend_args...)
ax[:set_xlabel](L"time", fontsize=16)
plt.show()
end

function plot2(rp::RamseyPath)
y, uhatdif, tauhatdif, mu = rp.y, rp.uhatdif, rp.tauhatdif, rp.mu
G, GPay = rp.G, rp.GPay
T = length(rp.mu)

tt = 1:T  # it is used to make the plot time index correct.
tt2 = 1:T-1

n_rows = 4
fig, axes = subplots(n_rows, 1, figsize=(10, 16))
plt.subplots_adjust(hspace=0.5)
for ax in axes
    ax[:grid](alpha=.5)
    ax[:set_xlim](-0.5, 15)
end
bbox = (0., 1.02, 1., .102)
legend_args = {:bbox_to_anchor => bbox, :loc => 3, :mode => "expand"}
p_args = { :lw => 2, :alpha => 0.7 }

ax = axes[1]
ax[:plot](tt2, tauhatdif,
        label="time inconsistency differential for tax rate"; p_args...)
ax[:set_ylabel](L"$\Delta \tau$", fontsize=16)
ax[:set_yticks]((0.0, 0.4, 0.8, 1.2))
ax[:legend](ncol=1; legend_args...)

ax = axes[2]
ax[:plot](tt, uhatdif,
        label="time inconsistency differential for $u$"; p_args...)
ax[:set_ylabel](L"$\Delta u$", fontsize=16)
ax[:set_yticks]((-3.0, -2.0, -1.0, 0.0))
ax[:legend](ncol=1; legend_args...)

ax = axes[3]
ax[:plot](tt, mu,
        label="Lagrange multiplier"; p_args...)
ax[:set_ylabel](L"$\mu$", fontsize=16)
```

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The program can also be found in the QuantEcon GitHub repository.

It computes a number of sequences besides the Ramsey plan, some of which have already been discussed, while others will be described below.

The next figure uses the program to compute and show the Ramsey plan for \( \tau \) and the Ramsey outcome for \((Q_t, u_t)\). From top to bottom, the panels show \( Q_t \), \( \tau_t \) and \( u_t := Q_{t+1} - Q_t \) over \( t = 0, \ldots, 15 \).

The optimal decision rule is

\[
\tau_{t+1} = -248.0624 - 0.1242Q_t - 0.3347u_t \tag{3.137}
\]

Notice how the Ramsey plan calls for a high tax at \( t = 1 \) followed by a perpetual stream of lower taxes.

Taxing heavily at first, less later expresses time-inconsistency of the optimal plan for \( \{\tau_t\}_{t=0}^\infty \).

We’ll characterize this formally after first discussing how to compute \( \mu \).

\[
\text{As promised, } \tau_t \text{ does not appear in the Ramsey planner’s decision rule for } \tau_{t+1}.\]
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Computing $\mu$  Define the selector vectors $e_\tau = [0 \ 0 \ 1 \ 0]'$ and $e_Q = [0 \ 1 \ 0 \ 0]'$ and express $\tau_t = e'_\tau y_t$ and $Q_t = e'_Q y_t$ 

Evidently $Q_t \tau_t = y'_t e_Q e'_\tau y_t = y'_t S y_t$ where $S := e_Q e'_\tau$

We want to compute $T_0 = \sum_{t=1}^{\infty} \beta^t \tau_t Q_t = \tau_1 Q_1 + \beta T_1$

where $T_1 = \sum_{t=2}^{\infty} \beta^{t-1} Q_t \tau_t$

The present values $T_0$ and $T_1$ are connected by

$T_0 = \beta y'_0 A'_F S A_F y_0 + \beta T_1$

Guess a solution that takes the form $T_t = y'_t \Omega y_t$, then find an $\Omega$ that satisfies

$\Omega = \beta A'_F S A_F + \beta A'_F \Omega A_F$  \hspace{1cm} (3.138)

Equation (3.138) is a discrete Lyapunov equation that can be solved for $\Omega$ using QuantEcon’s solve_discrete_lyapunov function

The matrix $F$ and therefore the matrix $A_F = A - BF$ depend on $\mu$

To find a $\mu$ that guarantees that $T_0 = G_0$ we proceed as follows:

1. Guess an initial $\mu$, compute a tentative Ramsey plan and the implied $T_0 = y'_0 \Omega(\mu) y_0$
2. If $T_0 > G_0$, lower $\mu$; if $T_0 < \mu$, raise $\mu$
3. Continue iterating on step 3 until $T_0 = G_0$

Time Inconsistency

Recall that the Ramsey planner chooses $\{u_t\}_{t=0}^{\infty}$, $\{\tau_t\}_{t=1}^{\infty}$ to maximize

$\sum_{t=0}^{\infty} \beta^t \left\{ A_0 Q_t - \frac{A_1}{2} Q^2_t - \frac{d}{2} u^2_t \right\}$

subject to (3.120), (3.121), and (3.123)

We express the outcome that a Ramsey plan is time-inconsistent the following way

**Proposition.** A continuation of a Ramsey plan is not a Ramsey plan

Let

$w(Q_0, u_0 | \mu_0) = \sum_{t=0}^{\infty} \beta^t \left\{ A_0 Q_t - \frac{A_1}{2} Q^2_t - \frac{d}{2} u^2_t \right\}$  \hspace{1cm} (3.139)

where

- $\{Q_t, u_t\}_{t=0}^{\infty}$ are evaluated under the Ramsey plan whose recursive representation is given by (3.134), (3.135), (3.136)
- $\mu_0$ is the value of the Lagrange multiplier that assures budget balance, computed as described above
Evidently, these continuation values satisfy the recursion

\[ w(Q_t, u_t | \mu_0) = A_0 Q_t - \frac{A_1}{2} Q_t^2 - \frac{d}{2} u_t^2 + \beta w(Q_{t+1}, u_{t+1} | \mu_0) \]  

(3.140)

for all \( t \geq 0 \), where \( Q_{t+1} = Q_t + u_t \).

Under the timing protocol affiliated with the Ramsey plan, the planner is committed to the outcome of iterations on \((3.134), (3.135), (3.136)\). In particular, when time \( t \) comes, the Ramsey planner is committed to the value of \( u_t \) implied by the Ramsey plan and receives continuation value \( w(Q_t, u_t, \mu_0) \).

That the Ramsey plan is time-inconsistent can be seen by subjecting it to the following ‘revolutionary’ test

First, define continuation revenues \( G_t \) that the government raises along the original Ramsey outcome by

\[ G_t = \beta^{-t} (G_0 - \sum_{s=1}^{t} \beta^s \tau_s Q_s) \]  

(3.141)

where \( \{\tau_t, Q_t\}_{t=0}^{\infty} \) is the original Ramsey outcome \(^5\).

Then at time \( t \geq 1 \),

1. take \((Q_t, G_t)\) inherited from the original Ramsey plan as initial conditions
2. invite a brand new Ramsey planner to compute a new Ramsey plan, solving for a new \( u_t \), to be called \( \hat{u}_t \), and for a new \( \mu \), to be called \( \hat{\mu}_t \).

The revised Lagrange multiplier \( \hat{\mu}_t \) is chosen so that, under the new Ramsey plan, the government is able to raise enough continuation revenues \( G_t \) given by \((3.141)\).

Would this new Ramsey plan be a continuation of the original plan? The answer is no because along a Ramsey plan, for \( t \geq 1 \), in general it is true that

\[ w(Q_t, v(Q_t | \hat{\mu}) | \hat{\mu}) > w(Q_t, u_t | \mu_0) \]  

(3.142)

Inequality \((3.142)\) expresses a continuation Ramsey planner’s incentive to deviate from a time 0 Ramsey plan by

1. resetting \( u_t \) according to \((3.133)\)
2. adjusting the Lagrange multiplier on the continuation appropriately to account for tax revenues already collected \(^6\).

Inequality \((3.142)\) expresses the time-inconsistency of a Ramsey plan

---

\(^5\) The continuation revenues \( G_t \) are the time \( t \) present value of revenues that must be raised to satisfy the original time 0 government intertemporal budget constraint, taking into account the revenues already raised from \( s = 1, \ldots, t \) under the original Ramsey plan.

\(^6\) For example, let the Ramsey plan yield time 1 revenues \( Q_1 \tau_1 \). Then at time 1, a continuation Ramsey planner would want to raise continuation revenues, expressed in units of time 1 goods, of \( \check{G}_1 := \frac{G - \beta Q_1 \tau_1}{\beta} \). To finance the remainder revenues, the continuation Ramsey planner would find a continuation Lagrange multiplier \( \check{\mu} \) by applying the three-step procedure from the previous section to revenue requirements \( \check{G}_1 \).
A Simulation To bring out the time inconsistency of the Ramsey plan, we compare

- the time $t$ values of $\tau_{t+1}$ under the original Ramsey plan with
- the value $\tilde{\tau}_{t+1}$ associated with a new Ramsey plan begun at time $t$ with initial conditions $(Q_t, G_t)$ generated by following the original Ramsey plan

Here again $G_t := \beta^{-1}(G_0 - \sum_{s=1}^{t} \beta^{s} \tau_s Q_s)$

The difference $\Delta \tau_t := \tilde{\tau}_t - \tau_t$ is shown in the top panel of the following figure

In the second panel we compare the time $t$ outcome for $u_t$ under the original Ramsey plan with the time $t$ value of this new Ramsey problem starting from $(Q_t, G_t)$

To compute $u_t$ under the new Ramsey plan, we use the following version of formula (3.130):

$$\tilde{u}_t = -P_{22}^{-1}(\tilde{\mu}_t)P_{21}(\tilde{\mu}_t)z_t$$

Here $z_t$ is evaluated along the Ramsey outcome path, where we have included $\tilde{\mu}_t$ to emphasize the dependence of $P$ on the Lagrange multiplier $\mu_0$.

To compute $u_t$ along the Ramsey path, we just iterate the recursion starting (3.131) from the initial $Q_0$ with $u_0$ being given by formula (3.130)

Thus the second panel indicates how far the reinitialized value $\tilde{u}_t$ value departs from the time $t$ outcome along the Ramsey plan

Note that the restarted plan raises the time $t+1$ tax and consequently lowers the time $t$ value of $u_t$

Associated with the new Ramsey plan at $t$ is a value of the Lagrange multiplier on the continuation government budget constraint

This is the third panel of the figure

The fourth panel plots the required continuation revenues $G_t$ implied by the original Ramsey plan

These figures help us understand the time inconsistency of the Ramsey plan

Further Intuition One feature to note is the large difference between $\tilde{\tau}_{t+1}$ and $\tau_{t+1}$ in the top panel of the figure

If the government is able to reset to a new Ramsey plan at time $t$, it chooses a significantly higher tax rate than if it were required to maintain the original Ramsey plan

The intuition here is that the government is required to finance a given present value of expenditures with distorting taxes $\tau$

The quadratic adjustment costs prevent firms from reacting strongly to variations in the tax rate for next period, which tilts a time $t$ Ramsey planner toward using time $t+1$ taxes

As was noted before, this is evident in the first figure, where the government taxes the next period heavily and then falls back to a constant tax from then on

---

7 It can be verified that this formula puts non-zero weight only on the components 1 and $Q_t$ of $z_t$. 

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This can also been seen in the third panel of the second figure, where the government pays off a significant portion of the debt using the first period tax rate. The similarities between the graphs in the last two panels of the second figure reveals that there is a one-to-one mapping between $G$ and $\mu$.

The Ramsey plan can then only be time consistent if $G_t$ remains constant over time, which will not be true in general.

Credible Policy  We express the theme of this section in the following: In general, a continuation of a Ramsey plan is not a Ramsey plan. This is sometimes summarized by saying that a Ramsey plan is not credible.

On the other hand, a continuation of a credible plan is a credible plan.

The literature on a credible public policy ([CK90] and [Sto89]) arranges strategies and incentives so that public policies can be implemented by a sequence of government decision makers instead of a single Ramsey planner who chooses an entire sequence of history-dependent actions once and for all at time $t = 0$.

Here we confine ourselves to sketching how recursive methods can be used to characterize credible policies in our model.

A key reference on these topics is [Cha98].

A credibility problem arises because we assume that the timing of decisions differs from those for a Ramsey problem.

A sequential timing protocol is a protocol such that

1. At each $t \geq 0$, given $Q_t$ and expectations about a continuation tax policy $\{\tau_{s+1}\}_{s=t}^\infty$ and a continuation price sequence $\{p_{s+1}\}_{s=t}^\infty$, the representative firm chooses $u_t$.

2. At each $t$, given $(Q_t, u_t)$, a government chooses $\tau_{t+1}$.

Item (2) captures that taxes are now set sequentially, the time $t + 1$ tax being set after the government has observed $u_t$.

Of course, the representative firm sets $u_t$ in light of its expectations of how the government will ultimately choose to set future taxes.

A credible tax plan $\{\tau_{s+1}\}_{s=t}^\infty$

- is anticipated by the representative firm, and
- is one that a time $t$ government chooses to confirm.

We use the following recursion, closely related to but different from (3.140), to define the continuation value function for the government:

$$J_t = A_0 Q_t - A_1 Q_t^2 - \frac{d}{2} u_t^2 + \beta J_{t+1}(\tau_{t+1}, G_{t+1})$$ (3.143)

This differs from (3.140) because

- continuation values are now allowed to depend explicitly on values of the choice $\tau_{t+1}$, and
3.11. HISTORY DEPENDENT PUBLIC POLICIES

- continuation government revenue to be raised $G_{t+1}$ need not be ones called for by the prevailing government policy

Thus, deviations from that policy are allowed, an alteration that recognizes that $\tau_i$ is chosen sequentially

Express the government budget constraint as requiring that $G_0$ solves the difference equation

$$G_t = \beta \tau_{t+1} Q_{t+1} + \beta G_{t+1}, \quad t \geq 0 \quad (3.144)$$

subject to the terminal condition $\lim_{t \to +\infty} \beta^t G_t = 0$

Because the government is choosing sequentially, it is convenient to

- take $G_t$ as a state variable at $t$
- to regard the time $t$ government as choosing $(\tau_{t+1}, G_{t+1})$ subject to constraint (3.144)

To express the notion of a credible government plan concisely, we expand the strategy space by also adding $J_t$ itself as a state variable and allowing policies to take the following recursive forms

8

Regard $J_0$ as an a discounted present value promised to the Ramsey planner and take it as an initial condition.

Then after choosing $u_0$ according to

$$u_0 = v(Q_0, G_0, J_0), \quad (3.145)$$

choose subsequent taxes, outputs, and continuation values according to recursions that can be represented as

$$\hat{\tau}_{t+1} = \tau(Q_t, u_t, G_t, J_t) \quad (3.146)$$

$$u_{t+1} = \hat{\xi}(Q_t, u_t, G_t, J_t, \tau_{t+1}) \quad (3.147)$$

$$G_{t+1} = \beta^{-1} G_t - \tau_{t+1} Q_{t+1} \quad (3.148)$$

$$J_{t+1}(\tau_{t+1}, G_{t+1}) = v(Q_t, u_t, G_{t+1}, J_t, \tau_{t+1}) \quad (3.149)$$

Here

- $\hat{\tau}_{t+1}$ is the time $t + 1$ government action called for by the plan, while
- $\tau_{t+1}$ is possibly some one-time deviation that the time $t + 1$ government contemplates and
- $G_{t+1}$ is the associated continuation tax collections

The plan is said to be credible if, for each $t$ and each state $(Q_t, u_t, G_t, J_t)$, the plan satisfies the incentive constraint

$$J_t = A_0 Q_t - \frac{A_1}{2} Q^2_t - \frac{d}{2} u^2_t + \beta J_{t+1}(\hat{\tau}_{t+1}, \hat{G}_{t+1}) \quad (3.150)$$

$$\geq A_0 Q_t - \frac{A_1}{2} Q^2_t - \frac{d}{2} u^2_t + \beta J_{t+1}(\tau_{t+1}, G_{t+1}) \quad (3.151)$$

8 This choice is the key to what [LS12] call ‘dynamic programming squared’.

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for all tax rates $\tau_{t+1} \in \mathbb{R}$ available to the government

Here $\hat{G}_{t+1} = \frac{G_t - \hat{\tau}_{t+1} Q_{t+1}}{\beta}$

- Inequality expresses that continuation values adjust to deviations in ways that discourage the government from deviating from the prescribed $\hat{\tau}_{t+1}$

- Inequality (3.150) indicates that two continuation values $J_{t+1}$ contribute to sustaining time $t$ promised value $J_t$
  - $J_{t+1}(\hat{\tau}_{t+1}, \hat{G}_{t+1})$ is the continuation value when the government chooses to confirm the private sector’s expectation, formed according to the decision rule (3.146) \(^9\)
  - $J_{t+1}(\tau_{t+1}, G_{t+1})$ tells the continuation consequences should the government disappoint the private sector’s expectations

The internal structure of a credible plan deters deviations from it

That (3.150) maps two continuation values $J_{t+1}(\tau_{t+1}, G_{t+1})$ and $J_{t+1}(\hat{\tau}_{t+1}, \hat{G}_{t+1})$ into one promised value $J_t$ reflects how a credible plan arranges a system of private sector expectations that induces the government to choose to confirm them

Chang [Cha98] builds on how inequality (3.150) maps two continuation values into one

Remark Let $J$ be the set of values associated with credible plans

Every value $J \in J$ can be attained by a credible plan that has a recursive representation of form (3.146), (3.147), (3.148)

The set of values can be computed as the largest fixed point of an operator that maps sets of candidate values into sets of values

Given a value within this set, it is possible to construct a government strategy of the recursive form (3.146), (3.147), (3.148) that attains that value

In many cases, there is a set of values and associated credible plans

In those cases where the Ramsey outcome is credible, a multiplicity of credible plans is a key part of the story because, as we have seen earlier, a continuation of a Ramsey plan is not a Ramsey plan

For it to be credible, a Ramsey outcome must be supported by a worse outcome associated with another plan, the prospect of reversion to which sustains the Ramsey outcome

Concluding remarks

The term ‘optimal policy’, which pervades an important applied monetary economics literature, means different things under different timing protocols

Under the ‘static’ Ramsey timing protocol (i.e., choose a sequence once-and-for-all), we obtain a unique plan

Here the phrase ‘optimal policy’ seems to fit well, since the Ramsey planner optimally reaps early benefits from influencing the private sector’s beliefs about the government’s later actions

\(^9\) Note the double role played by (3.146): as decision rule for the government and as the private sector’s rule for forecasting government actions.
When we adopt the sequential timing protocol associated with credible public policies, ‘optimal policy’ is a more ambiguous description.

There is a multiplicity of credible plans.

True, the theory explains how it is optimal for the government to confirm the private sector’s expectations about its actions along a credible plan.

But some credible plans have very bad outcomes.

These bad outcomes are central to the theory because it is the presence of bad credible plans that makes possible better ones by sustaining the low continuation values that appear in the second line of incentive constraint (3.150).

Recently, many have taken for granted that ‘optimal policy’ means ‘follow the Ramsey plan’.

In pursuit of more attractive ways to describe a Ramsey plan when policy making is in practice done sequentially, some writers have repackaged a Ramsey plan in the following way:

- Take a Ramsey outcome - a sequence of endogenous variables under a Ramsey plan - and reinterpret it (or perhaps only a subset of its variables) as a target path of relationships among outcome variables to be assigned to a sequence of policy makers.

- If appropriate (infinite dimensional) invertibility conditions are satisfied, it can happen that following the Ramsey plan is the only way to hit the target path.

- The spirit of this work is to say, “in a democracy we are obliged to live with the sequential timing protocol, so let’s constrain policy makers’ objectives in ways that will force them to follow a Ramsey plan in spite of their benevolence.”

- By this slight of hand, we acquire a theory of an optimal outcome target path.

This ‘invertibility’ argument leaves open two important loose ends:

1. implementation, and
2. time consistency

As for (1), repackaging a Ramsey plan (or the tail of a Ramsey plan) as a target outcome sequence does not confront the delicate issue of how that target path is to be implemented.

As for (2), it is an interesting question whether the ‘invertibility’ logic can repackage and conceal a Ramsey plan well enough to make policy makers forget or ignore the benevolent intentions that give rise to the time inconsistency of a Ramsey plan in the first place.

To attain such an optimal output path, policy makers must forget their benevolent intentions because there will inevitably occur temptations to deviate from that target path, and the implied relationship among variables like inflation, output, and interest rates along it.

**Remark** The continuation of such an optimal target path is not an optimal target path.

---

10 It is possible to read [Woo03] and [GW10] as making some carefully qualified statements of this type. Some of the qualifications can be interpreted as advice ‘eventually’ to follow a tail of a Ramsey plan.

11 In our model, the Ramsey outcome would be a path \((\vec{\beta}, \vec{Q})\).

12 See [GW10].

13 Sometimes the analysis is framed in terms of following the Ramsey plan only from some future date \(T\) onwards.

14 See [Bas05] and [ACK10].
REFERENCES


REFERENCES


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