Programming for Quantitative Economics

This book presents a set of lectures on python programming for quantitative economics, designed and written by

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Introduction to Python

This first part of the course provides a relatively fast-paced introduction to the Python programming language.

About Python

"Python has gotten sufficiently weapons grade that we don't descend into R anymore. Sorry, R people. I used to be one of you but we no longer descend into R." – Chris Wiggins

Overview

In this lecture we will

- outline what Python is
- · showcase some of its abilities
- compare it to some other languages.

At this stage, it's **not** our intention that you try to replicate all you see.

We will work through what follows at a slow pace later in the lecture series.

Our only objective for this lecture is to give you some feel of what Python is, and what it can do.

What's Python?

<u>Python</u> is a general-purpose programming language conceived in 1989 by Dutch programmer <u>Guido van Rossum</u>.

Python is free and open source, with development coordinated through the <u>Python Software Foundation</u>.

Python has experienced rapid adoption in the last decade and is now one of the most popular programming languages.

Common Uses

Python is a general-purpose language used in almost all application domains such as

- communications
- web development
- CGI and graphical user interfaces
- game development
- multimedia, data processing, security, etc., etc., etc.

Used extensively by Internet services and high tech companies including

- Google
- <u>Dropbox</u>
- Reddit

- YouTube
- Walt Disney Animation.

Python is very beginner-friendly and is often used to <u>teach computer science and programming</u>.

For reasons we will discuss, Python is particularly popular within the scientific community with users including NASA, CERN and practically all branches of academia.

It is also <u>replacing familiar tools like Excel</u> in the fields of finance and banking.

Relative Popularity

The following chart, produced using Stack Overflow Trends, shows one measure of the relative popularity of Python

The figure indicates not only that Python is widely used but also that adoption of Python has accelerated significantly since 2012.

We suspect this is driven at least in part by uptake in the scientific domain, particularly in rapidly growing fields like data science.

For example, the popularity of <u>pandas</u>, a library for data analysis with Python has exploded, as seen here.

(The corresponding time path for MATLAB is shown for comparison)

Note that pandas takes off in 2012, which is the same year that we see Python's popularity begin to spike in the first figure.

Overall, it's clear that

- Python is one of the most popular programming languages worldwide.
- Python is a major tool for scientific computing, accounting for a rapidly rising share of scientific work around the globe.

Features

Python is a <u>high-level language</u> suitable for rapid development.

It has a relatively small core language supported by many libraries.

Other features of Python:

- multiple programming styles are supported (procedural, object-oriented, functional, etc.)
- it is interpreted rather than compiled.

Syntax and Design

One nice feature of Python is its elegant syntax — we'll see many examples later on.

Elegant code might sound superfluous but in fact it's highly beneficial because it makes the syntax easy to read and easy to remember.

Remembering how to read from files, sort dictionaries and other such routine tasks means that you don't need to break your flow in order to hunt down correct syntax.

Closely related to elegant syntax is an elegant design.

Features like iterators, generators, decorators and list comprehensions make Python highly expressive, allowing you to get more done with less code.

Namespaces improve productivity by cutting down on bugs and syntax errors.

Scientific Programming

Python has become one of the core languages of scientific computing.

It's either the dominant player or a major player in

- machine learning and data science
- astronomy
- artificial intelligence
- <u>chemistry</u>
- computational biology
- meteorology

Its popularity in economics is also beginning to rise.

This section briefly showcases some examples of Python for scientific programming.

• All of these topics will be covered in detail later on.

Numerical Programming

Fundamental matrix and array processing capabilities are provided by the excellent <u>NumPy</u> library.

NumPy provides the basic array data type plus some simple processing operations.

For example, let's build some arrays

```
import numpy as np
a = np.linspace(-np.pi, np.pi, 100)
b = np.cos(a)
c = np.sin(a)
# Load the library
# Create even grid from -π to π # Apply cosine to each element of a # Apply sin to each element of a
```

Now let's take the inner product

```
b @ c
```

3.3306690738754696e-16

The number you see here might vary slightly but it's essentially zero.

(For older versions of Python and NumPy you need to use the <u>np.dot</u> function)

The SciPy library is built on top of NumPy and provides additional functionality.

For example, let's calculate $\int_{-2}^2 \phi(z) dz$ where ϕ is the standard normal density.

```
from scipy.stats import norm from scipy.integrate import quad  \phi = \text{norm()}  value, error = quad(\phi.pdf, -2, 2) # Integrate using Gaussian quadrature value
```

0.9544997361036417

SciPy includes many of the standard routines used in

- linear algebra
- integration
- interpolation
- optimization
- <u>distributions and random number generation</u>
- signal processing

See them all $\underline{\text{here}}$.

Graphics

The most popular and comprehensive Python library for creating figures and graphs is <u>Matplotlib</u>, with functionality including

- plots, histograms, contour images, 3D graphs, bar charts etc.
- output in many formats (PDF, PNG, EPS, etc.)
- LaTeX integration

Example 2D plot with embedded LaTeX annotations

Example contour plot

Example 3D plot

More examples can be found in the Matplotlib thumbnail gallery.

Other graphics libraries include

- Plotly
- Bokeh
- <u>VPython</u> 3D graphics and animations

Symbolic Algebra

It's useful to be able to manipulate symbolic expressions, as in Mathematica or Maple.

The **SymPy** library provides this functionality from within the Python shell.

```
from sympy import Symbol x, y = Symbol('x'), Symbol('y') # Treat 'x' and 'y' as algebraic symbols <math>x + x + x + y
```

$$3x + y$$

We can manipulate expressions

```
expression = (x + y)**2
expression.expand()
```

$$x^2 + 2xy + y^2$$

solve polynomials

```
from sympy import solve
solve(x**2 + x + 2)
```

```
[-1/2 - sqrt(7)*I/2, -1/2 + sqrt(7)*I/2]
```

and calculate limits, derivatives and integrals

```
from sympy import limit, sin, diff
limit(1 / x, x, 0)
```

 ∞

```
limit(sin(x) / x, x, 0)
```

```
diff(sin(x), x)
```

 $\cos(x)$

The beauty of importing this functionality into Python is that we are working within a fully fledged programming language.

We can easily create tables of derivatives, generate LaTeX output, add that output to figures and so on.

Statistics

Python's data manipulation and statistics libraries have improved rapidly over the last few years.

Pandas

One of the most popular libraries for working with data is pandas.

Pandas is fast, efficient, flexible and well designed.

Here's a simple example, using some dummy data generated with Numpy's excellent random functionality.

```
df.mean()
```

price 0.411768 weight -0.699135 dtype: float64

Other Useful Statistics Libraries

- <u>statsmodels</u> various statistical routines
- scikit-learn machine learning in Python (sponsored by Google, among others)
- pyMC for Bayesian data analysis
- pystan Bayesian analysis based on stan

Networks and Graphs

Python has many libraries for studying graphs.

One well-known example is <u>NetworkX</u>. Its features include, among many other things:

- standard graph algorithms for analyzing networks
- plotting routines

Here's some example code that generates and plots a random graph, with node color determined by shortest path length from a central node.

```
import networkx as nx
import matplotlib.pyplot as plt
%matplotlib inline
np.random.seed(1234)
g = nx.random_geometric_graph(200, 0.12, pos=p)
pos = nx.get_node_attributes(g, 'pos')
# Find node nearest the center point (0.5, 0.5) dists = [(x - 0.5)**2 + (y - 0.5)**2 for x, y in list(pos.values())] ncenter = np.argmin(dists)
# Plot graph, coloring by path length from central node
p = nx.single_source_shortest_path_length(g, ncenter)
plt.figure()
nx.draw_networkx_edges(g, pos, alpha=0.4)
nx.draw_networkx_nodes(g,
                         nodelist=list(p.keys()),
                         node_size=120, alpha=0.5,
node_color=list(p.values()),
cmap=plt.cm.jet_r)
plt.show()
```

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Cloud Computing

Running your Python code on massive servers in the cloud is becoming easier and easier.

A nice example is Anaconda Enterprise.

See also

- Amazon Elastic Compute Cloud
- The Google App Engine (Python, Java, PHP or Go)
- Pythonanywhere
- Sagemath Cloud

Parallel Processing

Apart from the cloud computing options listed above, you might like to consider

- · Parallel computing through IPython clusters.
- The <u>Starcluster</u> interface to Amazon's EC2.
- GPU programming through PyCuda, PyOpenCL, Theano or similar.

Other Developments

There are many other interesting developments with scientific programming in Python.

Some representative examples include

- <u>Jupyter</u> Python in your browser with interactive code cells, embedded images and other useful features.
- Numba Make Python run at the same speed as native machine code!
- Blaze a generalization of NumPy.
- PyTables manage large data sets.
- CVXPY convex optimization in Python.

Learn More

- Browse some Python projects on GitHub.
- Read more about <u>Python's history and rise in popularity</u>.
- Have a look at <u>some of the Jupyter notebooks</u> people have shared on various scientific topics.
- Visit the Python Package Index.
- View some of the questions people are asking about Python on Stackoverflow.

 Keep up to date on what's happening in the Python community with the <u>Python</u> subreddit.

Setting up Your Python Environment

Overview

In this lecture, you will learn how to

- 1. get a Python environment up and running
- 2. execute simple Python commands
- 3. run a sample program
- 4. install the code libraries that underpin these lectures

Anaconda

The <u>core Python package</u> is easy to install but *not* what you should choose for these lectures.

These lectures require the entire scientific programming ecosystem, which

- the core installation doesn't provide
- is painful to install one piece at a time.

Hence the best approach for our purposes is to install a Python distribution that contains

- 1. the core Python language and
- 2. compatible versions of the most popular scientific libraries.

The best such distribution is **Anaconda**.

Anaconda is

- · very popular
- cross-platform
- comprehensive
- completely unrelated to the Nicki Minaj song of the same name

Note

Anaconda also comes with a great package management system to organize your code libraries.

All of what follows assumes that you adopt this recommendation!

Installing Anaconda

To install Anaconda, $\underline{\text{download}}$ the binary and follow the instructions.

Important points:

- · Install the latest version!
- If you are asked during the installation process whether you'd like to make Anaconda your default Python installation, say yes.

Updating Anaconda

Anaconda supplies a tool called conda to manage and upgrade your Anaconda packages.

One conda command you should execute regularly is the one that updates the whole Anaconda distribution.

As a practice run, please execute the following

- 1. Open up a terminal
- 2. Type conda update anaconda

For more information on conda, type conda help in a terminal.

Jupyter Notebooks

<u>Jupyter</u> notebooks are one of the many possible ways to interact with Python and the scientific libraries.

They use a browser-based interface to Python with

- The ability to write and execute Python commands.
- Formatted output in the browser, including tables, figures, animation, etc.
- The option to mix in formatted text and mathematical expressions.

Because of these features, Jupyter is now a major player in the scientific computing ecosystem.

Here's an image showing execution of some code (borrowed from here) in a Jupyter notebook

While Jupyter isn't the only way to code in Python, it's great for when you wish to

- start coding in Python
- test new ideas or interact with small pieces of code
- share or collaborate scientific ideas with students or colleagues

These lectures are designed for executing in Jupyter notebooks.

Starting the Jupyter Notebook

Once you have installed Anaconda, you can start the Jupyter notebook.

Either

- · search for Jupyter in your applications menu, or
- open up a terminal and type jupyter notebook
 - Windows users should substitute "Anaconda command prompt" for "terminal" in the previous line.

If you use the second option, you will see something like this

The output tells us the notebook is running at http://localhost:8888/

- localhost is the name of the local machine
- 8888 refers to port number 8888 on your computer

Thus, the Jupyter kernel is listening for Python commands on port 8888 of our local machine.

Hopefully, your default browser has also opened up with a web page that looks something like this

What you see here is called the Jupyter dashboard.

If you look at the URL at the top, it should be localhost: 8888 or similar, matching the message above.

Assuming all this has worked OK, you can now click on New at the top right and select Python 3 or similar.

Here's what shows up on our machine:

The notebook displays an $\it active cell$, into which you can type Python commands.

Notebook Basics

Let's start with how to edit code and run simple programs.

Running Cells

Notice that, in the previous figure, the cell is surrounded by a green border.

This means that the cell is in edit mode.

In this mode, whatever you type will appear in the cell with the flashing cursor.

When you're ready to execute the code in a cell, hit Shift-Enter instead of the usual Enter.

(Note: There are also menu and button options for running code in a cell that you can find by exploring)

Modal Editing

The next thing to understand about the Jupyter 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
 - o Indicated by a green border around one cell, plus a blinking cursor
 - Whatever you type appears as is in that cell
- 2. Command mode
 - o The green border is replaced by a grey (or grey and blue) border
 - Keystrokes are interpreted as commands for example, typing b adds a new cell below the current one

To switch to

- command mode from edit mode, hit the Esc key or Ctrl-M
- edit mode from command mode, hit Enter or click in a cell

The modal behavior of the Jupyter notebook is very efficient when you get used to it.

Inserting Unicode (e.g., Greek Letters)

Python supports <u>unicode</u>, allowing the use of characters such as α and β as names in your code.

In a code cell, try typing \alpha and then hitting the tab key on your keyboard.

A Test Program

Let's run a test program.

Here's an arbitrary program we can use:

http://matplotlib.org/3.1.1/gallery/pie and polar charts/polar bar.html.

On that page, you'll see the following code

```
import numpy as np
import matplotlib.pyplot as plt
%matplotlib inline

# Fixing random state for reproducibility
np.random.seed(19680801)

# Compute pie slices
N = 20
0 = np.linspace(0.0, 2 * np.pi, N, endpoint=False)
radii = 10 * np.random.rand(N)
width = np.pi / 4 * np.random.rand(N)
colors = plt.cm.viridis(radii / 10.)

ax = plt.subplot(111, projection='polar')
ax.bar(0, radii, width=width, bottom=0.0, color=colors, alpha=0.5)
plt.show()
```

...../_images/getting_started_0_0.png

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 it into a cell in the notebook.

Hopefully you will get a similar plot.

Working with the Notebook

Here are a few more tips on working with Jupyter notebooks.

Tab Completion

In the previous program, we executed the line import numpy as np

• NumPy is a numerical library we'll work with in depth.

After this import command, functions in NumPy can be accessed with np.function_name type syntax.

• For example, try np.random.randn(3).

We can explore these attributes of np using the Tab key.

For example, here we type np. ran and hit Tab

Jupyter offers up the two possible completions, random and rank.

In this way, the Tab key helps remind you of what's available and also saves you typing.

On-Line Help

To get help on np. rank, say, we can execute np. rank?.

Documentation appears in a split window of the browser, like so

Clicking on the top right of the lower split closes the on-line help.

Other Content

In addition to executing code, the Jupyter 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 Esc to enter command mode and then type m to indicate that we are writing Markdown, a mark-up language similar to (but simpler than) LaTeX.

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

Now we Shift+Enter to produce this

Sharing Notebooks

Notebook files are just text files structured in $\underline{\sf JSON}$ and typically ending with . ipynb.

The notebooks you see on that site are static html representations.

To run one, download it as an ipynb file by clicking on the download icon at the top right.

Save it somewhere, navigate to it from the Jupyter dashboard and then run as discussed above.

QuantEcon Notes

 $\label{eq:QuantEcon} QuantEcon\ has its own site for sharing Jupyter notebooks\ related\ to\ economics\ -\ \underline{QuantEcon}\ \underline{Notes}.$

Notebooks submitted to QuantEcon Notes can be shared with a link, and are open to comments and votes by the community.

Installing Libraries

Most of the libraries we need come in Anaconda.

Other libraries can be installed with pip.

One library we'll be using is QuantEcon.py.

You can install **QuantEcon.py** by starting Jupyter and typing

```
!pip install ——upgrade quantecon
```

into a cell.

Alternatively, you can type the following into a terminal

```
pip install quantecon
```

More instructions can be found on the library page.

To upgrade to the latest version, which you should do regularly, use

```
pip install --upgrade quantecon
```

Another library we will be using is interpolation.py.

This can be installed by typing in Jupyter

```
!pip install interpolation
```

Working with Python Files

So far we've focused on executing Python code entered into a Jupyter notebook cell.

Traditionally most Python code has been run in a different way.

Code is first saved in a text file on a local machine

By convention, these text files have a .py extension.

We can create an example of such a file as follows:

```
%%file foo.py
print("foobar")
```

Overwriting foo.py

This writes the line print ("foobar") into a file called foo.py in the local directory.

Here %file is an example of a cell magic.

Editing and Execution

If you come across code saved in a *.py file, you'll need to consider the following questions:

- 1. how should you execute it?
- 2. How should you modify or edit it?

Option 1: JupyterLab

JupyterLab is an integrated development environment built on top of Jupyter notebooks.

With JupyterLab you can edit and run *.py files as well as Jupyter notebooks.

To start JupyterLab, search for it in the applications menu or type j upyter-lab in a terminal.

Now you should be able to open, edit and run the file foo. py created above by opening it in JupyterLab.

Read the docs or search for a recent YouTube video to find more information.

Option 2: Using a Text Editor

One can also edit files using a text editor and then run them from within Jupyter notebooks.

A text editor is an application that is specifically designed to work with text files — such as Python programs.

Nothing beats the power and efficiency of a good text editor for working with program text.

A good text editor will provide

- efficient text editing commands (e.g., copy, paste, search and replace)
- syntax highlighting, etc.

Right now, an extremely popular text editor for coding is VS Code.

VS Code is easy to use out of the box and has many high quality extensions.

Alternatively, 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 <u>Vim</u>.

Exercises

Exercise 1

If Jupyter is still running, quit by using Ctrl-C at the terminal where you started it.

Now launch again, but this time using jupyter notebook --no-browser.

This should start the kernel without launching the browser.

Note also the startup message: It should give you a URL such as http://localhost:8888 where the notebook is running.

Now

- 1. Start your browser or open a new tab if it's already running.
- 2. Enter the URL from above (e.g. http://localhost:8888) in the address bar at the top. You should now be able to run a standard Jupyter notebook session.

This is an alternative way to start the notebook that can also be handy.

Exercise 2

This exercise will familiarize you with git and GitHub.

<u>Git</u> 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 \underline{GitHub} .

GitHub is a wonderland of collaborative coding projects.

For example, it hosts many of the scientific libraries we'll be using later on, such as this one.

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 Git version
- 2. the various point-and-click GUI versions
 - See, for example, the GitHub version

As the 1st task, try

- 1. Installing Git.
- 2. Getting a copy of QuantEcon.py using Git.

For example, if you've installed the command line version, open up a terminal and enter.

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

(This is just git clone in front of the URL for the repository)

As the 2nd task,

- 1. Sign up to GitHub.
- 2. Look into 'forking' GitHub repositories (forking means making your own copy of a GitHub repository, stored on GitHub).
- 3. Fork QuantEcon.py.
- 4. Clone your fork to some local directory, make edits, commit them, and push them back up to your forked GitHub repo.
- 5. If you made a valuable improvement, send us a pull request!

For reading on these and other topics, try

- The official Git documentation.
- Reading through the docs on GitHub.
- Pro Git Book by Scott Chacon and Ben Straub.
- One of the thousands of Git tutorials on the Net.

An Introductory Example

Overview

We're now ready to start learning the Python language itself.

In this lecture, we will write and then pick apart small Python programs.

The objective is to introduce you to basic Python syntax and data structures.

Deeper concepts will be covered in later lectures.

You should have read the <u>lecture</u> on getting started with Python before beginning this one.

The Task: Plotting a White Noise Process

Suppose we want to simulate and plot the white noise process $\epsilon_0, \epsilon_1, \dots, \epsilon_T$, where each draw ϵ_t is independent standard normal.

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

(Here t is on the horizontal axis and ϵ_t is on the vertical axis.)

 $We'll \ do \ this \ in \ several \ different \ ways, each \ time \ learning \ something \ more \ about \ Python.$

We run the following command first, which helps ensure that plots appear in the notebook if you run it on your own machine.

```
%matplotlib inline
```

Version 1

Here are a few lines of code that perform the task we set

```
import numpy as np
import matplotlib.pyplot as plt

e_values = np.random.randn(100)
plt.plot(∈_values)
plt.show()

././_images/python_by_example_1_0.png
```

Let's break this program down and see how it works.

Imports

The first two lines of the program import functionality from external code libraries.

The first line imports NumPy, a favorite Python package for tasks like

- working with arrays (vectors and matrices)
- common mathematical functions like cos and sqrt
- generating random numbers
- linear algebra, etc.

After import numpy as np we have access to these attributes via the syntax np.attribute.

Here's two more examples

```
np.sqrt(4)

2.0

np.log(4)

1.3862943611198906
```

We could also use the following syntax:

```
import numpy
numpy.sqrt(4)
```

2.0

But the former method (using the short name np) is convenient and more standard.

Why So Many Imports?

Python programs typically require several import statements.

The reason is that the core language is deliberately kept small, so that it's easy to learn and maintain.

When you want to do something interesting with Python, you almost always need to import additional functionality.

Packages

As stated above, NumPy is a Python package.

Packages are used by developers to organize code they wish to share.

In fact, a package is just a directory containing

- 1. files with Python code called **modules** in Python speak
- possibly some compiled code that can be accessed by Python (e.g., functions compiled from C or FORTRAN code)
- a file called __init__.py that specifies what will be executed when we type import package_name

In fact, you can find and explore the directory for NumPy on your computer easily enough if you look around.

On this machine, it's located in

```
anaconda3/lib/python3.7/site-packages/numpy
```

Subpackages

Consider the line ϵ _values = np.random.randn(100).

Here np refers to the package NumPy, while random is a subpackage of NumPy.

Subpackages are just packages that are subdirectories of another package.

Importing Names Directly

Recall this code that we saw above

```
import numpy as np
np.sqrt(4)
```

2.0

Here's another way to access NumPy's square root function

```
from numpy import sqrt
sqrt(4)
```

2.0

This is also fine.

The advantage is less typing if we use sqrt often in our code.

The disadvantage is that, in a long program, these two lines might be separated by many other lines.

Then it's harder for readers to know where sqrt came from, should they wish to.

Random Draws

Returning to our program that plots white noise, the remaining three lines after the import statements are

```
e_values = np.random.randn(100)
plt.plot(e_values)
plt.show()

.../_images/python_by_example_7_0.png
```

The first line generates 100 (quasi) independent standard normals and stores them in ϵ_{values} .

The next two lines genererate the plot.

We can and will look at various ways to configure and improve this plot below.

Alternative Implementations

Let's try writing some alternative versions of <u>our first program</u>, which plotted IID draws from the normal distribution.

The programs below are less efficient than the original one, and hence somewhat artificial.

But they do help us illustrate some important Python syntax and semantics in a familiar setting.

A Version with a For Loop

Here's a version that illustrates for loops and Python lists.

```
ts_length = 100
e_values = [] # empty list

for i in range(ts_length):
    e = np.random.randn()
    e_values.append(e)

plt.plot(e_values)
plt.show()
```

../../_images/python_by_example_8_0.png

In brief,

- The first line sets the desired length of the time series.
- The next line creates an empty list called ϵ_{-} values that will store the ϵ_{t} values as we generate them.
- The statement # empty list is a *comment*, and is ignored by Python's interpreter.
- The next three lines are the for loop, which repeatedly draws a new random number e₁ and appends it to the end of the list e_values.
- The last two lines generate the plot and display it to the user.

Let's study some parts of this program in more detail.

Lists

Consider the statement $\epsilon_{\text{values}} = []$, which creates an empty list.

Lists are a *native Python data structure* used to group a collection of objects.

For example, try

```
x = [10, 'foo', False]
type(x)
```

list

The first element of x is an <u>integer</u>, the next is a <u>string</u>, and the third is a <u>Boolean value</u>.

When adding a value to a list, we can use the syntax list_name.append(some_value)

```
x
[10, 'foo', False]

x.append(2.5)
x
[10, 'foo', False, 2.5]
```

Here append() is what's called a *method*, which is a function "attached to" an object—in this case, the list x.

We'll learn all about methods later on, but just to give you some idea,

- Python objects such as lists, strings, etc. all have methods that are used to manipulate
 the data contained in the object.
- String objects have string methods, list objects have list methods, etc.

Another useful list method is pop()

```
x
[10, 'foo', False, 2.5]

x.pop()

2.5

x
[10, 'foo', False]
```

Lists in Python are zero-based (as in C, Java or Go), so the first element is referenced by x [0]

```
x[0] # first element of x

10

x[1] # second element of x

'foo'
```

The For Loop

Now let's consider the for loop from the program above, which was

Python executes the two indented lines ts_length times before moving on.

These two lines are called a code block, since they comprise the "block" of code that we are looping over.

 $\label{thm:code} \mbox{Unlike most other languages, Python knows the extent of the code block } \emph{only from indentation.}$

In our program, indentation decreases after line $\epsilon_$ values. append (e), telling Python that this line marks the lower limit of the code block.

More on indentation below—for now, let's look at another example of a for loop

```
animals = ['dog', 'cat', 'bird']
for animal in animals:
    print("The plural of " + animal + " is " + animal + "s")

The plural of dog is dogs
The plural of cat is cats
The plural of bird is birds
```

This example helps to clarify how the for loop works: When we execute a loop of the form

The Python interpreter performs the following:

• For each element of the sequence, it "binds" the name variable_name to that element and then executes the code block.

The sequence object can in fact be a very general object, as we'll see soon enough.

A Comment on Indentation

In discussing the for loop, we explained that the code blocks being looped over are delimited by indentation.

In fact, in Python, **all** code blocks (i.e., those occurring inside loops, if clauses, function definitions, etc.) are delimited by indentation.

Thus, unlike most other languages, whitespace in Python code affects the output of the program.

Once you get used to it, this is a good thing: It

- · forces clean, consistent indentation, improving readability
- removes clutter, such as the brackets or end statements used in other languages

On the other hand, it takes a bit of care to get right, so please remember:

• The line before the start of a code block always ends in a colon

```
for i in range(10):if x > y:while x < 100:</li>etc., etc.
```

- All lines in a code block must have the same amount of indentation.
- The Python standard is 4 spaces, and that's what you should use.

While Loops

The for loop is the most common technique for iteration in Python.

But, for the purpose of illustration, let's modify <u>the program above</u> to use a while loop instead.

```
ts_length = 100

e_values = []

i = 0

while i < ts_length:

e = np.random.randn()

e_values.append(e)

i = i + 1

plt.plot(e_values)

plt.show()
```

...../_images/python_by_example_19_0.png

Note that

- the code block for the while loop is again delimited only by indentation
- the statement i = i + 1 can be replaced by i += 1

Another Application

Let's do one more application before we turn to exercises.

In this application, we plot the balance of a bank account over time.

There are no withdraws over the time period, the last date of which is denoted by T.

The initial balance is b_0 and the interest rate is \emph{r} .

The balance updates from period t to t + 1 according to $b_{t+1} = (1 + r)b_t$.

In the code below, we generate and plot the sequence b_0, b_1, \ldots, b_T .

Instead of using a Python list to store this sequence, we will use a NumPy array.

```
r = 0.025  # interest rate
T = 50  # end date
b = np.empty(T+1) # an empty NumPy array, to store all b_t
b[0] = 10  # initial balance

for t in range(T):
    b[t+1] = (1 + r) * b[t]

plt.plot(b, label='bank balance')
plt.legend()
plt.show()
```

../../_images/python_by_example_20_0.png

The statement b = np.empty(T+1) allocates storage in memory for T+1 (floating point) numbers.

These numbers are filled in by the for loop.

Allocating memory at the start is more efficient than using a Python list and append, since the latter must repeatedly ask for storage space from the operating system.

Notice that we added a legend to the plot - a feature you will be asked to use in the exercises.

Exercises

Now we turn to exercises. It is important that you complete them before continuing, since they present new concepts we will need.

Exercise 1

Your first task is to simulate and plot the correlated time series

```
x_{t+1} = \alpha x_t + \epsilon_{t+1} where x_0 = 0 and t = 0, \dots, T
```

The sequence of shocks $\{e_t\}$ is assumed to be IID and standard normal.

In your solution, restrict your import statements to

```
import numpy as np
import matplotlib.pyplot as plt
```

Set T=200 and $\alpha=0.9$.

Exercise 2

Starting with your solution to exercise 2, plot three simulated time series, one for each of the cases $\alpha=0, \alpha=0.8$ and $\alpha=0.98$.

Use a for loop to step through the α values.

If you can, add a legend, to help distinguish between the three time series.

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.
- For the legend, noted that the expression 'foo' + str(42) evaluates to 'foo42'.

Exercise 3

Similar to the previous exercises, plot the time series $% \left(x\right) =\left(x\right) +\left(x\right)$

$$x_{t+1} = \alpha |x_t| + \epsilon_{t+1}$$
 where $x_0 = 0$ and $t = 0, \dots, T$

Use $T=200, \alpha=0.9$ and $\{\epsilon_t\}$ as before.

Search online for a function that can be used to compute the absolute value $|x_t|$.

Exercise 4

One important aspect of essentially all programming languages is branching and conditions.

In Python, conditions are usually implemented with if-else syntax.

Here's an example, that prints -1 for each negative number in an array and 1 for each nonnegative number

```
numbers = [-9, 2.3, -11, 0]

for x in numbers:
    if x < 0:
        print(-1)
    else:
        print(1)</pre>
```

-1 1 -1 1

Now, write a new solution to Exercise 3 that does not use an existing function to compute the absolute value.

Replace this existing function with an if-else condition.

Exercise 5

Here's a harder exercise, that takes some thought and planning.

The task is to compute an approximation to π using Monte Carlo.

Use no imports besides

```
import numpy as np
```

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, $area = \pi * radius^2$.

Solutions

Exercise 1

Here's one solution.

```
α = 0.9
T = 200
x = np.empty(T+1)
x[0] = 0

for t in range(T):
    x[t+1] = α * x[t] + np.random.randn()
plt.plot(x)
plt.show()
```

.../../_images/python_by_example_25_0.png

Exercise 2

```
a_values = [0.0, 0.8, 0.98]
T = 200
x = np.empty(T+1)

for α in α_values:
    x[0] = 0
    for t in range(T):
        x[t+1] = α * x[t] + np.random.randn()
    plt.plot(x, label=f'$\\alpha = {α}$$')

plt.legend()
plt.show()
..../_images/python_by_example_26_0.png
```

Exercise 3

Here's one solution:

```
α = 0.9
T = 200
x = np.empty(T+1)
x[0] = 0

for t in range(T):
    x[t+1] = α * np.abs(x[t]) + np.random.randn()

plt.plot(x)
plt.show()
```

.../../_images/python_by_example_27_0.png

Exercise 4

Here's one way:

```
α = 0.9
T = 200
x = np.empty(T+1)
x[0] = 0

for t in range(T):
    if x[t] < 0:
        abs_x = - x[t]
    else:
        abs_x = x[t]
    x[t+1] = α * abs_x + np.random.randn()

plt.plot(x)
plt.show()</pre>
./../_images/python_by_example_28_0.png
```

Here's a shorter way to write the same thing:

```
α = 0.9
T = 200
x = np.empty(T+1)
x[0] = 0

for t in range(T):
    abs_x = - x[t] if x[t] < 0 else x[t]
    x[t+1] = α * abs_x + np.random.randn()

plt.plot(x)
plt.show()</pre>
```

../../_images/python_by_example_29_0.png

Exercise 5

Consider the circle of diameter 1 embedded in the unit square.

Let A be its area and let r = 1/2 be its radius.

If we know π then we can compute A via $A = \pi r^2$.

But here the point is to compute π , which we can do by $\pi = A/r^2$.

Summary: If we can estimate the area of a circle with diameter 1, then dividing by $r^2=(1/2)^2=1/4$ gives an estimate of π .

We estimate the area by sampling bivariate uniforms and looking at the fraction that falls into the circle.

```
n = 100000
count = 0
for i in range(n):
    u, v = np.random.uniform(), np.random.uniform()
    d = np.sqrt((u - 0.5)**2 + (v - 0.5)**2)
    if d < 0.5:
        count += 1
area_estimate = count / n
print(area_estimate * 4)  # dividing by radius**2</pre>
```

3.14416

Functions

Overview

One construct that's extremely useful and provided by almost all programming languages is **functions**.

We have already met several functions, such as

- the sqrt() function from NumPy and
- the built-in print() function

In this lecture we'll treat functions systematically and begin to learn just how useful and important they are.

One of the things we will learn to do is build our own user-defined functions

We will use the following imports.

```
import numpy as np
import matplotlib.pyplot as plt
%matplotlib inline
```

Function Basics

A function is a named section of a program that implements a specific task.

Many functions exist already and we can use them off the shelf.

First we review these functions and then discuss how we can build our own.

Built-In Functions

Python has a number of built-in functions that are available without import.

We have already met some

```
max(19, 20)

20

print('foobar')

foobar

str(22)
```

```
type(22)
```

int

Two more useful built-in functions are any() and all()

```
bools = False, True, True
all(bools) # True if all are True and False otherwise
```

False

```
any(bools) # False if all are False and True otherwise
```

True

The full list of Python built-ins is here.

Third Party Functions

If the built-in functions don't cover what we need, we either need to import functions or create our own.

Examples of importing and using functions were given in the <u>previous lecture</u>

Here's another one, which tests whether a given year is a leap year:

```
import calendar
calendar.isleap(2020)
```

True

Defining Functions

In many instances, it is useful to be able to define our own functions.

This will become clearer as you see more examples.

Let's start by discussing how it's done.

Syntax

Here's a very simple Python function, that implements the mathematical function f(x)=2x+1

```
def f(x):
    return 2 * x + 1
```

Now that we've defined this function, let's call it and check whether it does what we expect:

```
f(1)
```

3

```
f(10)
```

21

Here's a longer function, that computes the absolute value of a given number.

(Such a function already exists as a built-in, but let's write our own for the exercise.)

```
def new_abs_function(x):
    if x < 0:
        abs_value = -x
    else:
        abs_value = x
    return abs_value</pre>
```

Let's review the syntax here.

- def is a Python keyword used to start function definitions.
- def new_abs_function(x): indicates that the function is called new_abs_function
 and that it has a single argument x.
- The indented code is a code block called the *function body*.
- The return keyword indicates that abs_value is the object that should be returned to the calling code.

This whole function definition is read by the Python interpreter and stored in memory.

Let's call it to check that it works:

```
print(new_abs_function(3))
print(new_abs_function(-3))
3
3
```

Why Write 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 almost always a bad idea)

We will say more about this $\underline{\mathsf{later}}$.

Applications

Random Draws

Consider again this code from the <u>previous lecture</u>

```
ts_length = 100
e_values = [] # empty list

for i in range(ts_length):
    e = np.random.randn()
    e_values.append(e)

plt.plot(e_values)
plt.show()

.../_images/functions_13_0.png
```

We will break this program into two parts:

- 1. A user-defined function that generates a list of random variables.
- 2. The main part of the program that
 - 1. calls this function to get data
 - 2. plots the data

This is accomplished in the next program

When the interpreter gets to the expression $generate_data(100)$, it executes the function body with n set equal to 100.

The net result is that the name data is bound to the list ϵ _values returned by the function.

Adding Conditions

Our function generate_data() is rather limited.

Let's make it slightly more useful by giving it the ability to return either standard normals or uniform random variables on (0,1) as required.

This is achieved in the next piece of code.

....._images/functions_15_0.png

Hopefully, the syntax of the if/else clause is self-explanatory, with indentation again delimiting the extent of the code blocks.

Note

Notes

- We are passing the argument U as a string, which is why we write it as 'U'.
- Notice that equality is tested with the == syntax, not =.
 - \circ For example, the statement a = 10 assigns the name a to the value 10.
 - The expression a == 10 evaluates to either True or False, depending on the value of a.

Now, there are several ways that we can simplify the code above.

For example, we can get rid of the conditionals all together by just passing the desired generator type as a function.

To understand this, consider the following version.

.../../_images/functions_16_0.png

Now, when we call the function $generate_data()$, we pass np.random.uniform as the second argument.

This object is a function.

When the function call generate_data(100, np.random.uniform) is executed, Python runs the function code block with n equal to 100 and the name generator_type "bound" to the function np.random.uniform.

 While these lines are executed, the names generator_type and np.random.uniform are "synonyms", and can be used in identical ways.

This principle works more generally—for example, consider the following piece of code

```
max(7, 2, 4) # max() is a built-in Python function
```

7

Here we created another name for the built-in function \max (), which could then be used in identical ways.

In the context of our program, the ability to bind new names to functions means that there is no problem *passing a function as an argument to another function*—as we did above.

Exercises

Exercise 1

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

There are functions to compute this in various modules, but let's write our own version as an exercise.

In particular, write a function factorial such that factorial(n) returns n! for any positive integer n.

Exercise 2

The <u>binomial random variable</u> $Y \sim Bin(n, p)$ represents the number of successes in n binary trials, where each trial succeeds with probability p.

Without any import besides from numpy. random import uniform, 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 $\mathbf{U}<\mathbf{p}$ evaluates to True with probability p.

Exercise 3

First, write a function that returns one realization of the following random device

- 1. Flip an unbiased coin 10 times.
- If a head occurs k or more times consecutively within this sequence at least once, pay one dollar.
- 3. If not, pay nothing.

Second, write another function that does the same task except that the second rule of the above random device becomes

• If a head occurs k or more times within this sequence, pay one dollar.

Use no import besides from numpy.random import uniform.

Solutions

Exercise 1

Here's one solution.

```
def factorial(n):
    k = 1
    for i in range(n):
        k = k * (i + 1)
    return k
factorial(4)
```

24

Exercise 2

```
from numpy.random import uniform

def binomial_rv(n, p):
    count = 0
    for i in range(n):
        U = uniform()
        if U < p:
            count = count + 1 # Or count += 1
    return count

binomial_rv(10, 0.5)</pre>
```

7

Exercise 3

Here's a function for the first random device.

```
from numpy.random import uniform

def draw(k): # pays if k consecutive successes in a sequence

payoff = 0
    count = 0

for i in range(10):
    U = uniform()
    count = count + 1 if U < 0.5 else 0
    print(count) # print counts for clarity
    if count == k:
        payoff = 1

return payoff

draw(3)</pre>
```

1

Here's another function for the second random device.

```
def draw_new(k): # pays if k successes in a sequence
  payoff = 0
  count = 0

for i in range(10):
    U = uniform()
    count = count + ( 1 if U < 0.5 else 0 )
    print(count)
    if count == k:
        payoff = 1

  return payoff

draw_new(3)</pre>
```

0

Python Essentials

Overview

We have covered a lot of material quite quickly, with a focus on examples.

Now let's cover some core features of Python in a more systematic way.

This approach is less exciting but helps clear up some details.

Data Types

Computer programs typically keep track of a range of data types.

For example, 1.5 is a floating point number, while 1 is an integer.

Programs need to distinguish between these two types for various reasons.

One is that they are stored in memory differently.

Another is that arithmetic operations are different

• For example, floating point arithmetic is implemented on most machines by a specialized Floating Point Unit (FPU).

In general, floats are more informative but arithmetic operations on integers are faster and more accurate.

Python provides numerous other built-in Python data types, some of which we've already met

• strings, lists, etc.

Let's learn a bit more about them.

Primitive Data Types

One simple data type is Boolean values, which can be either True or False

```
x = True
x
```

True

We can check the type of any object in memory using the type () function.

```
type(x)
bool
```

In the next line of code, the interpreter evaluates the expression on the right of = and binds y to this value

```
y = 100 < 10
y
```

False

```
type(y)
```

bool

In arithmetic expressions, True is converted to 1 and False is converted 0.

This is called **Boolean arithmetic** and is often useful in programming.

Here are some examples

```
x + y
```

1

```
x * y
```

0

```
True + True
```

2

```
bools = [True, True, False, True] # List of Boolean values
sum(bools)
```

3

Complex numbers are another primitive data type in Python

```
x = complex(1, 2)
y = complex(2, 1)
print(x * y)

type(x)
```

5j

 ${\tt complex}$

Containers

 $Python\ has\ several\ basic\ types\ for\ storing\ collections\ of\ (possibly\ heterogeneous)\ data.$

We've already discussed lists.

A related data type is **tuples**, which are "immutable" lists

```
x = ('a', 'b') # Parentheses instead of the square brackets
x = 'a', 'b' # Or no brackets --- the meaning is identical
x
```

```
('a', 'b')
```

```
type(x)
```

tuple

In Python, an object is called **immutable** if, once created, the object cannot be changed.

Conversely, an object is **mutable** if it can still be altered after creation.

Python lists are mutable

```
x = [1, 2]
x[0] = 10
x
```

[10, 2]

But tuples are not

```
x = (1, 2)
x[0] = 10
```

TypeError: 'tuple' object does not support item assignment

We'll say more about the role of mutable and immutable data a bit later.

Tuples (and lists) can be "unpacked" as follows

```
integers = (10, 20, 30)
x, y, z = integers
x
```

10

```
у
```

20

You've actually $\underline{\text{seen an example of this}}$ already.

Tuple unpacking is convenient and we'll use it often.

Slice Notation

To access multiple elements of a list or tuple, you can use Python's slice notation.

For example,

```
a = [2, 4, 6, 8]
a[1:]
```

[4, 6, 8]

```
a[1:3]
```

[4, 6]

The general rule is that a[m:n] returns n - m elements, starting at a[m].

Negative numbers are also permissible

```
a[-2:] # Last two elements of the list
```

[6, 8]

The same slice notation works on tuples and strings

```
s = 'foobar'
s[-3:] # Select the last three elements
```

'bar'

Sets and Dictionaries

Two other container types we should mention before moving on are <u>sets</u> and <u>dictionaries</u>.

Dictionaries are much like lists, except that the items are named instead of numbered

```
d = {'name': 'Frodo', 'age': 33}
type(d)

dict

d['age']
```

33

The names 'name' and 'age' are called the keys.

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

Sets are unordered collections without duplicates, and set methods provide the usual set-theoretic operations $\frac{1}{2} \left(\frac{1}{2} \right) = \frac{1}{2} \left(\frac{1}{2} \right) \left(\frac{1}{2} \right$

```
s1 = {'a', 'b'}
type(s1)
set
```

```
s2 = {'b', 'c'}
s1.issubset(s2)
```

False

```
s1.intersection(s2)
{'b'}
```

The $\operatorname{\mathsf{set}}$ () function creates sets from sequences

```
s3 = set(('foo', 'bar', 'foo'))
s3
{'bar', 'foo'}
```

Input and Output

Let's briefly review reading and writing to text files, starting with writing

```
f = open('newfile.txt', 'w')  # Open 'newfile.txt' for writing
f.write('Testing\n')  # Here '\n' means new line
f.write('Testing again')
f.close()
```

Here

- The built-in function open () creates a file object for writing to.
- Both write() and close() are methods of file objects.

Where is this file that we've created?

Recall that Python maintains a concept of the present working directory (pwd) that can be located from with Jupyter or IPython via

```
%pwd
```

'/Users/junaluzi/Documents/ExecutableBookProject/quantecon-example'

If a path is not specified, then this is where Python writes to.

We can also use Python to read the contents of newline.txt as follows

```
f = open('newfile.txt', 'r')
out = f.read()
out
```

'Testing\nTesting again'

```
print(out)
```

Testing Testing again

Paths

Note that if newfile.txt is not in the present working directory then this call to open() fails.

In this case, you can shift the file to the pwd or specify the <u>full path</u> to the file

```
f = open('insert_full_path_to_file/newfile.txt', 'r')
```

Iterating

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

One of Python's strengths is its simple, flexible interface to this kind of iteration via the for loop.

Looping over Different Objects

Many Python objects are "iterable", in the sense that they can be looped over.

To give an example, let's write the file $us_cities.txt$, which lists US cities and their population, to the present working directory.

```
%%file us_cities.txt
new york: 8244910
los angeles: 3819702
chicago: 2707120
houston: 2145146
philadelphia: 1536471
phoenix: 1469471
san antonio: 1359758
san diego: 1326179
dallas: 1223229
```

Overwriting us_cities.txt

Here %file is an IPython cell magic.

Suppose that we want to make the information more readable, by capitalizing names and adding commas to mark thousands.

The program below reads the data in and makes the conversion:

```
data_file = open('us_cities.txt', 'r')
for line in data_file:
     city, population = line.split(':')
                                                              # Tuple unpacking
# Capitalize city names
# Add commas to numbers
     city = city.title()
population = f'{int(population):,}'
     print(city.ljust(15) + population)
data_file.close()
New York
                    8,244,910
                    3,819,702
Los Angeles
                    2,707,120
Chicago
Houston
                    2,145,146
Philadelphia
                    1,536,471
Phoenix
                    1,469,471
San Antonio
San Diego
                    1,359,758
1,326,179
Dallas
                    1,223,229
```

Here format() is a string method used for inserting variables into strings.

The reformatting of each line is the result of three different string methods, the details of which can be left till later.

The interesting part of this program for us is line 2, which shows that

- 1. The file object data_file is iterable, in the sense that it can be placed to the right of in within a for loop.
- 2. Iteration steps through each line in the file.

This leads to the clean, convenient syntax shown in our program.

Many other kinds of objects are iterable, and we'll discuss some of them later on.

Looping without Indices

One thing you might have noticed is that Python tends to favor looping without explicit indexing.

For example,

```
x_values = [1, 2, 3] # Some iterable x
for x in x_values:
    print(x * x)
1
4
9
```

is preferred to

```
for i in range(len(x_values)):
    print(x_values[i] * x_values[i])

1
4
9
```

When you compare these two alternatives, you can see why the first one is preferred.

 $Python\ provides\ some\ facilities\ to\ simplify\ looping\ without\ indices.$

One is ${\tt zip}(\tt),$ which is used for stepping through pairs from two sequences.

For example, try running the following code

```
countries = ('Japan', 'Korea', 'China')
cities = ('Tokyo', 'Seoul', 'Beijing')
for country, city in zip(countries, cities):
    print(f'The capital of {country} is {city}')
```

```
The capital of Japan is Tokyo
The capital of Korea is Seoul
The capital of China is Beijing
```

The zip() function is also useful for creating dictionaries — for example

```
names = ['Tom', 'John']
marks = ['E', 'F']
dict(zip(names, marks))

{'Tom': 'E', 'John': 'F'}
```

If we actually need the index from a list, one option is to use enumerate().

To understand what ${\tt enumerate}$ () does, consider the following example

```
letter_list = ['a', 'b', 'c']
for index, letter in enumerate(letter_list):
    print(f"letter_list[{index}] = '{letter}'")

letter_list[0] = 'a'
letter_list[1] = 'b'
letter_list[2] = 'c'
```

List Comprehensions

We can also simplify the code for generating the list of random draws considerably by using something called a *list comprehension*.

<u>List comprehensions</u> are an elegant Python tool for creating lists.

Consider the following example, where the list comprehension is on the right-hand side of the second line

```
animals = ['dog', 'cat', 'bird']
plurals = [animal + 's' for animal in animals]
plurals
['dogs', 'cats', 'birds']
```

Here's another example

```
range(8)

range(0, 8)

doubles = [2 * x for x in range(8)]

doubles

[0, 2, 4, 6, 8, 10, 12, 14]
```

Comparisons and Logical Operators

Comparisons

Many different kinds of expressions evaluate to one of the Boolean values (i.e., True or False).

A common type is comparisons, such as

```
x, y = 1, 2
x < y
```

True

```
x > y
```

False

One of the nice features of Python is that we can *chain* inequalities

```
1 < 2 < 3
```

True

```
1 <= 2 <= 3
```

True

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

```
x = 1  # Assignment
x == 2  # Comparison
```

False

For "not equal" use !=

```
1 != 2
```

True

Note that when testing conditions, we can use any valid Python expression

```
x = 'yes' if 42 else 'no'
x
```

'yes'

```
x = 'yes' if [] else 'no'
x
```

'no'

What's going on here?

The rule is:

- Expressions that evaluate to zero, empty sequences or containers (strings, lists, etc.) and None are all equivalent to False.
 - $\circ~$ for example, [] and () are equivalent to False in an if clause
- All other values are equivalent to True.
 - $\circ~$ for example, 42 is equivalent to True in an if clause

Combining Expressions

We can combine expressions using and, or and not.

These are the standard logical connectives (conjunction, disjunction and denial)

```
1 < 2 and 'f' in 'foo'
```

True

```
1 < 2 and 'g' in 'foo'
```

False

```
1 < 2 or 'g' in 'foo'
```

True

```
not True
```

False

```
not not True
```

True

Remember

- P and Q is True if both are True, else False
- P or Q is False if both are False, else True

More Functions

Let's talk a bit more about functions, which are all important for good programming style.

The Flexibility of Python Functions

As we discussed in the previous lecture, Python functions are very flexible.

In particular

- Any number of functions can be defined in a given file.
- Functions can be (and often are) defined inside other functions.
- Any object can be passed to a function as an argument, including other functions.
- A function can return any kind of object, including functions.

We already gave an example of how straightforward it is to pass a function to a function.

Note that a function can have arbitrarily many return statements (including zero).

Execution of the function terminates when the first return is hit, allowing code like the following example

```
def f(x):
    if x < 0:
        return 'negative'
    return 'nonnegative'</pre>
```

Functions without a return statement automatically return the special Python object None.

Docstrings

 $Python\ has\ a\ system\ for\ adding\ comments\ to\ functions,\ modules,\ etc.\ called\ \emph{docstrings}.$

The nice thing about docstrings is that they are available at run-time.

Try running this

```
def f(x):
    """
    This function squares its argument
    return x**2
```

After running this code, the docstring is available

```
f?
```

```
Type: function
String Form:<function f at 0x2223320>
File: /home/john/temp/temp.py
Definition: f(x)
Docstring: This function squares its argument
```

```
f??
```

```
Type: function
String Form:<function f at 0x2223320>
File: /home/john/temp/temp.py
Definition: f(x)
Source:
def f(x):
    This function squares its argument
    """
    return x**2
```

With one question mark we bring up the docstring, and with two we get the source code as well.

One-Line Functions: lambda

The lambda keyword is used to create simple functions on one line.

For example, the definitions

```
def f(x):
    return x**3
```

and

```
f = lambda x: x**3
```

are entirely equivalent.

To see why lambda is useful, suppose that we want to calculate $\int_0^2 x^3 dx$ (and have forgotten our high-school calculus).

The SciPy library has a function called quad that will do this calculation for us.

The syntax of the quad function is quad (f, a, b) where f is a function and a and b are numbers.

To create the function $f(x) = x^3$ we can use lambda as follows

```
from scipy.integrate import quad
quad(lambda x: x**3, 0, 2)
```

```
(4.0, 4.440892098500626e-14)
```

Here the function created by lambda is said to be *anonymous* because it was never given a name.

Keyword Arguments

In a previous lecture, you came across the statement

```
plt.plot(x, 'b-', label="white noise")
```

In this call to Matplotlib's plot function, notice that the last argument is passed in name=argument syntax.

This is called a *keyword argument*, with label being the keyword.

Non-keyword arguments are called *positional arguments*, since their meaning is determined by order

plot(x, 'b-', label="white noise") is different from plot('b-', x, label="white noise")

Keyword arguments are particularly useful when a function has a lot of arguments, in which case it's hard to remember the right order.

You can adopt keyword arguments in user-defined functions with no difficulty.

The next example illustrates the syntax

```
def f(x, a=1, b=1):
    return a + b * x
```

The keyword argument values we supplied in the definition of f become the default values

```
f(2)
```

3

They can be modified as follows

```
f(2, a=4, b=5)
```

14

Coding Style and PEP8

To learn more about the Python programming philosophy type import this at the prompt.

Among other things, Python strongly favors consistency in programming style.

We've all heard the saying about consistency and little minds.

In programming, as in mathematics, the opposite is true

• A mathematical paper where the symbols ∪ and ∩ were reversed would be very hard to read, even if the author told you so on the first page.

In Python, the standard style is set out in PEP8.

(Occasionally we'll deviate from PEP8 in these lectures to better match mathematical notation)

Exercises

Solve the following exercises.

(For some, the built-in function $\operatorname{\mathsf{sum}}($) comes in handy).

Exercise 1

Part 1: Given two numeric lists or tuples x_{vals} and y_{vals} of equal length, compute their inner product using zip().

Part 2: In one line, count the number of even numbers in 0,...,99.

• Hint: x % 2 returns 0 if x is even, 1 otherwise.

Part 3: Given pairs = ((2, 5), (4, 2), (9, 8), (12, 10)), count the number of pairs (a, b) such that both a and b are even.

Exercise 2

Consider the polynomial

$$p(x) = a_0 + a_1 x + a_2 x^2 + \dots + a_n x^n = \sum_{i=0}^n a_i x^i$$
 (1)

Write a function p such that p(x, coeff) that computes the value in (1) given a point x and a list of coefficients coeff.

Try to use enumerate() in your loop.

Exercise 3

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

```
Hint: 'foo'.upper() returns 'F00'.
```

Exercise 4

Write a function that takes two sequences seq_a and seq_b as arguments and returns True if every element in seq_a is also an element of seq_b, else False.

- By "sequence" we mean a list, a tuple or a string.
- Do the exercise without using sets and set methods.

Exercise 5

When we cover the numerical libraries, we will see they include many alternatives for interpolation and function approximation.

 $Nevertheless, let's \ write \ our \ own \ function \ approximation \ routine \ as \ an \ exercise.$

In particular, without using any imports, write a function 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 <= x <= b.

and returns the <u>piecewise linear interpolation</u> of f at x, based on n evenly spaced grid points a = point[0] < point[1] < ... < point[n-1] = b.

Aim for clarity, not efficiency.

Exercise 6

Using list comprehension syntax, we can simplify the loop in the following code.

```
import numpy as np

n = 100

e_values = []
for i in range(n):
    e = np.random.randn()
    e_values.append(e)
```

Solutions

Exercise 1

Part 1 Solution:

Here's one possible solution

```
x_vals = [1, 2, 3]
y_vals = [1, 1, 1]
sum([x * y for x, y in zip(x_vals, y_vals)])
```

6

```
sum(x * y for x, y in zip(x_vals, y_vals))
```

6

Part 2 Solution:

One solution is

```
sum([x % 2 == 0 for x in range(100)])
```

50

This also works:

```
sum(x % 2 == 0 for x in range(100))
```

50

Some less natural alternatives that nonetheless help to illustrate the flexibility of list comprehensions are

```
len([x for x in range(100) if x % 2 == 0])
```

50

and

```
sum([1 for x in range(100) if x % 2 == 0])
```

50

Part 3 Solution

Here's one possibility

```
pairs = ((2, 5), (4, 2), (9, 8), (12, 10))
sum([x % 2 == 0 and y % 2 == 0 for x, y in pairs])
```

2

Exercise 2

```
def p(x, coeff):
    return sum(a * x**i for i, a in enumerate(coeff))
```

```
p(1, (2, 4))
```

6

Exercise 3

Here's one solution:

```
def f(string):
    count = 0
    for letter in string:
        if letter == letter.upper() and letter.isalpha():
            count += 1
    return count

f('The Rain in Spain')
```

An alternative, more pythonic solution:

```
def count_uppercase_chars(s):
    return sum([c.isupper() for c in s])
count_uppercase_chars('The Rain in Spain')
```

3

Exercise 4

Here's a solution:

```
def f(seq_a, seq_b):
    is_subset = True
    for a in seq_a:
        if a not in seq_b:
        is_subset = False
    return is_subset

# == test == #

print(f([1, 2], [1, 2, 3]))
print(f([1, 2, 3], [1, 2]))
```

True False

Of course, if we use the sets data type then the solution is easier

```
def f(seq_a, seq_b):
    return set(seq_a).issubset(set(seq_b))
```

Exercise 5

```
def linapprox(f, a, b, n, x):
    Evaluates the piecewise linear interpolant of f at x on the interval
    [a, b], with n evenly spaced grid points.
    Parameters
        f : function
              The function to approximate
        x, a, b : scalars (floats or integers)
Evaluation point and endpoints, with a <= x <= b
        n : integer
             Number of grid points
    Returns
        A float. The interpolant evaluated at x
    length\_of\_interval = b - a
    \begin{array}{l} num\_subintervals = n-1 \\ step = length\_of\_interval \ / \ num\_subintervals \end{array}
    \# === find first grid point larger than x === #
    point = a
    while point <= x:</pre>
        point += step
    # === x must lie between the gridpoints (point - step) and point ===
    u, v = point - step, point
    return f(u) + (x - u) * (f(v) - f(u)) / (v - u)
```

Exercise 6

Here's one solution.

```
\begin{bmatrix} n = 100 \\ \epsilon\_values = [np.random.randn() for i in range(n)] \end{bmatrix}
```

OOP I: Introduction to Object Oriented Programming

Overview

OOP is one of the major paradigms in programming.

The traditional programming paradigm (think Fortran, C, MATLAB, etc.) is called *procedural*.

It works as follows

- The program has a state corresponding to the values of its variables.
- Functions are called to act on these data.
- Data are passed back and forth via function calls.

In contrast, in the OOP paradigm

• data and functions are "bundled together" into "objects"

(Functions in this context are referred to as methods)

Python and OOP

Python is a pragmatic language that blends object-oriented and procedural styles, rather than taking a purist approach.

However, at a foundational level, Python is object-oriented.

In particular, in Python, everything is an object.

In this lecture, we explain what that statement means and why it matters.

Objects

In Python, an $\it object$ is a collection of data and instructions held in computer memory that consists of

- 1. a type
- 2. a unique identity
- 3. data (i.e., content)
- 4. methods

These concepts are defined and discussed sequentially below.

Type

Python provides for different types of objects, to accommodate different categories of data.

For example

```
s = 'This is a string'
type(s)
str
```

```
x = 42  # Now let's create an integer
type(x)
```

int

The type of an object matters for many expressions.

For example, the addition operator between two strings means concatenation

```
'300' + 'cc'
```

On the other hand, between two numbers it means ordinary addition

```
300 + 400
```

700

Consider the following expression

Here we are mixing types, and it's unclear to Python whether the user wants to

- convert '300' to an integer and then add it to 400, or
- convert 400 to string and then concatenate it with '300'

Some languages might try to guess but Python is strongly typed

- Type is important, and implicit type conversion is rare.
- Python will respond instead by raising a TypeError.

To avoid the error, you need to clarify by changing the relevant type.

For example,

```
int('300') + 400  # To add as numbers, change the string to an integer
700
```

Identity

In Python, each object has a unique identifier, which helps Python (and us) keep track of the object.

The identity of an object can be obtained via the ${\tt id}$ () function

```
y = 2.5
z = 2.5
id(y)
```

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```
id(z)
```

4469478800

In this example, y and z happen to have the same value (i.e., 2.5), but they are not the same object.

The identity of an object is in fact just the address of the object in memory.

Object Content: Data and Attributes

If we set x = 42 then we create an object of type int that contains the data 42.

In fact, it contains more, as the following example shows

```
\begin{array}{rcl}
x &= & 42 \\
x
\end{array}
```

42

```
x.imag
```

0

```
x.__class__
```

int

When Python creates this integer object, it stores with it various auxiliary information, such as the imaginary part, and the type.

Any name following a dot is called an *attribute* of the object to the left of the dot.

• e.g.,imag and __class__ are attributes of x.

We see from this example that objects have attributes that contain auxiliary information.

They also have attributes that act like functions, called methods.

These attributes are important, so let's discuss them in-depth.

Methods

Methods are functions that are bundled with objects.

Formally, methods are attributes of objects that are callable (i.e., can be called as functions)

```
x = ['foo', 'bar']
callable(x.append)

True

callable(x.__doc__)
```

False

Methods typically act on the data contained in the object they belong to, or combine that data with other data ${\bf x}$

```
x = ['a', 'b']
x.append('c')
s = 'This is a string'
s.upper()
```

'THIS IS A STRING'

```
s.lower()
```

'this is a string'

```
s.replace('This', 'That')
```

'That is a string'

A great deal of Python functionality is organized around method calls.

For example, consider the following piece of code

```
x = ['a', 'b']
x[0] = 'aa' # Item assignment using square bracket notation
x
```

```
['aa', 'b']
```

It doesn't look like there are any methods used here, but in fact the square bracket assignment notation is just a convenient interface to a method call.

What actually happens is that Python calls the __setitem__ method, as follows

```
x = ['a', 'b']
x.__setitem__(0, 'aa') # Equivalent to x[0] = 'aa'
x
['aa', 'b']
```

(If you wanted to you could modify the <u>__setitem__</u> method, so that square bracket assignment does something totally different)

Summary

In Python, everything in memory is treated as an object.

This includes not just lists, strings, etc., but also less obvious things, such as

- functions (once they have been read into memory)
- · modules (ditto)
- files opened for reading or writing
- integers, etc.

Consider, for example, functions.

When Python reads a function definition, it creates a function object and stores it in memory.

The following code illustrates

```
def f(x): return x**2
f

<function __main__.f(x)>

type(f)

function

id(f)

4470432960

f.__name__
```

We can see that f has type, identity, attributes and so on—just like any other object.

It also has methods.

One example is the <u>__call__</u> method, which just evaluates the function

```
f.__call__(3)
```

Another is the $__dir__$ method, which returns a list of attributes.

Modules loaded into memory are also treated as objects

```
import math
id(math)
```

4433854528

This uniform treatment of data in Python (everything is an object) helps keep the language simple and consistent.

OOP II: Building Classes

Overview

In an <u>earlier lecture</u>, we learned some foundations of object-oriented programming.

The objectives of this lecture are

- · cover OOP in more depth
- · learn how to build our own objects, specialized to our needs

For example, you already know how to

- · create lists, strings and other Python objects
- use their methods to modify their contents

So imagine now you want to write a program with consumers, who can

- hold and spend cash
- · consume goods
- · work and earn cash

A natural solution in Python would be to create consumers as objects with

- data, such as cash on hand
- methods, such as buy or work that affect this data

Python makes it easy to do this, by providing you with class definitions.

 ${\it Classes are blueprints that help you build objects according to your own specifications.}$

It takes a little while to get used to the syntax so we'll provide plenty of examples.

We'll use the following imports:

```
import numpy as np
import matplotlib.pyplot as plt
%matplotlib inline
```

OOP Review

OOP is supported in many languages:

- JAVA and Ruby are relatively pure OOP.
- Python supports both procedural and object-oriented programming.
- Fortran and MATLAB are mainly procedural, some OOP recently tacked on.
- C is a procedural language, while C++ is C with OOP added on top.

Let's cover general OOP concepts before we specialize to Python.

Key Concepts

As discussed an <u>earlier lecture</u>, in the OOP paradigm, data and functions are **bundled together** into "objects".

An example is a Python list, which not only stores data but also knows how to sort itself, etc.

```
x = [1, 5, 4]
x.sort()
x
```

As we now know, sort is a function that is "part of" the list object — and hence called a *method*.

If we want to make our own types of objects we need to use class definitions.

A *class definition* is a blueprint for a particular class of objects (e.g., lists, strings or complex numbers).

It describes

- What kind of data the class stores
- What methods it has for acting on these data

An object or instance is a realization of the class, created from the blueprint

- Each instance has its own unique data.
- Methods set out in the class definition act on this (and other) data.

In Python, the data and methods of an object are collectively referred to as attributes.

Attributes are accessed via "dotted attribute notation"

- object_name.data
- object_name.method_name()

In the example

```
x = [1, 5, 4]
x.sort()
x.__class__
```

list

- x is an object or instance, created from the definition for Python lists, but with its own particular data.
- x.sort() and x.__class__ are two attributes of x.
- dir(x) can be used to view all the attributes of x.

Why is OOP Useful?

OOP is useful for the same reason that abstraction is useful: for recognizing and exploiting the common structure.

For example,

- a Markov chain consists of a set of states and a collection of transition probabilities for moving across states
- a general equilibrium theory consists of a commodity space, preferences, technologies, and an equilibrium definition
- a game consists of a list of players, lists of actions available to each player, player payoffs as functions of all players' actions, and a timing protocol

These are all abstractions that collect together "objects" of the same "type".

Recognizing common structure allows us to employ common tools.

In economic theory, this might be a proposition that applies to all games of a certain type.

In Python, this might be a method that's useful for all Markov chains (e.g., simulate).

When we use OOP, the simulate method is conveniently bundled together with the Markov chain object.

Defining Your Own Classes

Let's build some simple classes to start off.

Example: A Consumer Class

First, we'll build a Consumer class with

- a wealth attribute that stores the consumer's wealth (data)
- an earn method, where earn(y) increments the consumer's wealth by y
- a spend method, where spend(x) either decreases wealth by x or returns an error if
 insufficient funds exist

Admittedly a little contrived, this example of a class helps us internalize some new syntax.

Here's one implementation

```
class Consumer:
    def __init__(self, w):
        "Initialize consumer with w dollars of wealth"
        self.wealth = w

    def earn(self, y):
        "The consumer earns y dollars"
        self.wealth += y

    def spend(self, x):
        "The consumer spends x dollars if feasible"
        new_wealth = self.wealth - x
        if new_wealth < 0:
            print("Insufficent funds")
        else:
        self.wealth = new_wealth</pre>
```

There's some special syntax here so let's step through carefully

The class keyword indicates that we are building a class.

This class defines instance data wealth and three methods: __init__, earn and spend

 wealth is instance data because each consumer we create (each instance of the Consumer class) will have its own separate wealth data.

The ideas behind the earn and spend methods were discussed above.

Both of these act on the instance data wealth.

The __init__ method is a constructor method.

Whenever we create an instance of the class, this method will be called automatically.

 ${\sf Calling_init_sets}\ up\ a\ ``namespace''\ to\ hold\ the\ instance\ data-more\ on\ this\ soon.$

We'll also discuss the role of self just below.

Usage

Here's an example of usage

```
c1 = Consumer(10) # Create instance with initial wealth 10
c1.spend(5)
c1.wealth
```

5

```
c1.earn(15)
c1.spend(100)
```

Insufficent funds

We can of course create multiple instances each with its own data

```
c1 = Consumer(10)
c2 = Consumer(12)
c2.spend(4)
c2.wealth
```

8

```
c1.wealth
```

In fact, each instance stores its data in a separate namespace dictionary

```
c1.__dict__
{'wealth': 10}

c2.__dict__
{'wealth': 8}
```

When we access or set attributes we're actually just modifying the dictionary maintained by the instance.

Self

If you look at the Consumer class definition again you'll see the word self throughout the code.

The rules with self are that

- · Any instance data should be prepended with self
 - o e.g., the earn method references self.wealth rather than just wealth
- Any method defined within the class should have self as its first argument
 - e.g., def earn(self, y) rather than just def earn(y)
- Any method referenced within the class should be called as self.method_name

There are no examples of the last rule in the preceding code but we will see some shortly.

Details

In this section, we look at some more formal details related to classes and self

- You might wish to skip to the next section <oop_solow_growth> on first pass of this lecture.
- You can return to these details after you've familiarized yourself with more examples.

 $\label{thm:methods} \mbox{Methods actually live inside a class object formed when the interpreter reads the class definition$

```
print(Consumer.__dict__) # Show __dict__ attribute of class object

{'__module__': '__main__', '__init__': <function Consumer.__init__ at
0x11da83ca0>, 'earn': <function Consumer.earn at 0x11da83d30>, 'spend':
<function Consumer.spend at 0x11da83dc0>, '__dict__': <attribute
'__dict__' of 'Consumer' objects>, '__weakref__': <attribute
'__weakref__' of 'Consumer' objects>, '__doc__': None}
```

Note how the three methods __init__, earn and spend are stored in the class object.

Consider the following code

```
c1 = Consumer(10)
c1.earn(10)
c1.wealth
```

20

When you call earn via c1.earn(10) the interpreter passes the instance c1 and the argument 10 to Consumer.earn.

In fact, the following are equivalent

- c1.earn(10)
- Consumer.earn(c1, 10)

In the function call Consumer.earn(c1, 10) note that c1 is the first argument.

Recall that in the definition of the earn method, self is the first parameter

```
def earn(self, y):
    "The consumer earns y dollars"
    self.wealth += y
```

The end result is that self is bound to the instance c1 inside the function call.

That's why the statement self.wealth += y inside earn ends up modifying c1.wealth.

Example: The Solow Growth Model

For our next example, let's write a simple class to implement the Solow growth model.

The Solow growth model is a neoclassical growth model where the amount of capital stock per capita k_t evolves according to the rule

$$k_{t+1} = \frac{szk_t^{\alpha} + (1 - \delta)k_t}{1 + n}$$
 (2)

Here

- s is an exogenously given savings rate
- z is a productivity parameter
- α is capital's share of income
- \bullet *n* is the population growth rate
- ullet δ is the depreciation rate

The **steady state** of the model is the k that solves (2) when $k_{t+1} = k_t = k$.

Here's a class that implements this model.

Some points of interest in the code are

- An instance maintains a record of its current capital stock in the variable self.k.
- The h method implements the right-hand side of (2).
- The update method uses h to update capital as per (2).
 - Notice how inside update the reference to the local method h is self.h.

The methods $steady_state$ and $generate_sequence$ are fairly self-explanatory

```
class Solow:
    Implements the Solow growth model with the update rule
         k_{t+1} = [(s \ z \ k^{\alpha}_t) + (1 - \delta)k_t] / (1 + n)
    0.00
    def __init__(self, n=0.05, # population growth rate
                            s=0.25, # savings rate
                            δ=0.1,
                                       # depreciation rate
                            \alpha=0.3,
                                      # share of labor
                            z=2.0, # productivity
k=1.0): # current capital stock
         self.n, self.s, self.\delta, self.\alpha, self.z = n, s, \delta, \alpha, z
         self_k = k
    def h(self):
           'Evaluate the h function"
         # Unpack parameters (get rid of self to simplify notation)
         \alpha, s, \beta, \alpha, z= self.\alpha, self.\alpha, self.\alpha, self.\alpha
         return (s * z * self.k**\alpha + (1 - \delta) * self.k) / (1 + n)
    def update(self):
           Update the current state (i.e., the capital stock)."
          self.k = self.h()
    def steady_state(self):
         "Compute the steady state value of capital."

# Unpack parameters (get rid of self to simplify notation)
         # Compute and return steady state 3 suff.\alpha, self.\alpha, self.\alpha
         return ((s * z) / (n + \delta))**(1 / (1 - \alpha))
    def generate_sequence(self, t):
         "Generate and return a time series of length t" path = []
          for i in range(t):
              path.append(self.k)
              self.update()
         \textcolor{return}{\textbf{return}} \ \texttt{path}
```

Here's a little program that uses the class to compute time series from two different initial conditions

The common steady state is also plotted for comparison

Example: A Market

Next, let's write a class for a simple one good market where agents are price takers.

The market consists of the following objects:

- A linear demand curve $Q = a_d b_d p$
- A linear supply curve $Q = a_z + b_z(p t)$

Here

- p is price paid by the consumer, Q is quantity and t is a per-unit tax.
- Other symbols are demand and supply parameters.

The class provides methods to compute various values of interest, including competitive equilibrium price and quantity, tax revenue raised, consumer surplus and producer surplus.

Here's our implementation.

(It uses a function from SciPy called quad for numerical integration—a topic we will say more about later on.)

```
from scipy.integrate import quad
class Market:
     def __init__(self, ad, bd, az, bz, tax):
           Set up market parameters. All parameters are scalars. See
           https://lectures.quantecon.org/py/python_oop.html for
interpretation.
           self.ad, self.bd, self.az, self.bz, self.tax = ad, bd, az, bz,
tax
           if ad < az:</pre>
                raise ValueError('Insufficient demand.')
     def price(self):
           "Return equilibrium price"
           return (self.ad - self.az + self.bz * self.tax) / (self.bd +
self.bz)
     def quantity(self):
            'Compute equilibrium quantity"
           return self.ad - self.bd * self.price()
     def consumer_surp(self):
           "Compute consumer surplus"

# == Compute area under inverse demand function == #
           integrand = lambda x: (self.ad / self.bd) - (1 / self.bd) * x
area, error = quad(integrand, 0, self.quantity())
return area - self.price() * self.quantity()
     def producer_surp(self):
            'Compute producer surplus"
          # == Compute area above inverse supply curve, excluding tax == #
integrand = lambda x: -(self.az / self.bz) + (1 / self.bz) * x
area, error = quad(integrand, 0, self.quantity())
return (self.price() - self.tax) * self.quantity() - area
     def taxrev(self):
           "Compute tax revenue"
           return self.tax * self.quantity()
     def inverse_demand(self, x):
    "Compute inverse demand"
    return self.ad / self.bd - (1 / self.bd)* x
     def inverse_supply(self, x):
           "Compute inverse supply curve"
return -(self.az / self.bz) + (1 / self.bz) * x + self.tax
     def inverse_supply_no_tax(self, x):
           "Compute inverse supply curve without tax" return -(self.az / self.bz) + (1 / self.bz) * x
```

Here's a sample of usage

```
baseline_params = 15, .5, -2, .5, 3
m = Market(*baseline_params)
print("equilibrium price = ", m.price())

equilibrium price = 18.5

print("consumer surplus = ", m.consumer_surp())

consumer surplus = 33.0625
```

Here's a short program that uses this class to plot an inverse demand curve together with inverse supply curves with and without taxes

```
# Baseline ad, bd, az, bz, tax
baseline_params = 15, .5, -2, .5, 3

m = Market(*baseline_params)

q_max = m.quantity() * 2
q_grid = np.linspace(0.0, q_max, 100)
pd = m.inverse_demand(q_grid)
ps = m.inverse_supply(q_grid)
psno = m.inverse_supply_no_tax(q_grid)

fig, ax = plt.subplots()
ax.plot(q_grid, pd, lw=2, alpha=0.6, label='demand')
ax.plot(q_grid, ps, lw=2, alpha=0.6, label='supply')
ax.plot(q_grid, psno, '--k', lw=2, alpha=0.6, label='supply without tax')
ax.set_xlabel('quantity', fontsize=14)
ax.set_ylabel('price', fontsize=14)
ax.set_ylabel('price', fontsize=14)
ax.legend(loc='lower right', frameon=False, fontsize=14)
plt.show()

..../_images/python_oop_18_0.png
```

The next program provides a function that

- takes an instance of Market as a parameter
- computes dead weight loss from the imposition of the tax

```
def deadw(m):
    "Computes deadweight loss for market m."
    # == Create analogous market with no tax == #
    m_no_tax = Market(m.ad, m.bd, m.az, m.bz, 0)
# == Compare surplus, return difference == #
    surp1 = m_no_tax.consumer_surp() + m_no_tax.producer_surp()
    surp2 = m.consumer_surp() + m.producer_surp() + m.taxrev()
    return surp1 - surp2
```

Here's an example of usage

```
baseline_params = 15, .5, -2, .5, 3
m = Market(*baseline_params)
deadw(m) # Show deadweight loss
```

1.125

Example: Chaos

Let's look at one more example, related to chaotic dynamics in nonlinear systems.

One simple transition rule that can generate complex dynamics is the logistic map

$$x_{t+1} = rx_t(1 - x_t), \quad x_0 \in [0, 1], \quad r \in [0, 4]$$
(3)

Let's write a class for generating time series from this model.

Here's one implementation

```
class Chaos:
    """
    Models the dynamical system with :math:`x_{t+1} = r x_t (1 - x_t)`
    """
    def __init__(self, x0, r):
        Initialize with state x0 and parameter r
        self.x, self.r = x0, r

def update(self):
        "Apply the map to update state."
        self.x = self.r * self.x *(1 - self.x)

def generate_sequence(self, n):
        "Generate and return a sequence of length n."
    path = []
    for i in range(n):
        path.append(self.x)
        self.update()
    return path
```

Here's an example of usage

```
ch = Chaos(0.1, 4.0) \# \times \emptyset = 0.1 and r = 0.4 ch.generate_sequence(5) \# First \ 5 iterates
```

```
[0.1, 0.3600000000000004, 0.9216, 0.2890137600000006, 0.8219392261226498]
```

This piece of code plots a longer trajectory

```
ch = Chaos(0.1, 4.0)
ts_length = 250

fig, ax = plt.subplots()
ax.set_xlabel('$t$', fontsize=14)
ax.set_ylabel('$x_t$', fontsize=14)
x = ch.generate_sequence(ts_length)
ax.plot(range(ts_length), x, 'bo-', alpha=0.5, lw=2, label='$x_t$')
plt.show()

..../_images/python_oop_23_0.png
```

The next piece of code provides a bifurcation diagram

```
fig, ax = plt.subplots()
ch = Chaos(0.1, 4)
r = 2.5
while r < 4:
    ch.r = r
    t = ch.generate_sequence(1000)[950:]
    ax.plot([r] * len(t), t, 'b.', ms=0.6)
    r = r + 0.005

ax.set_xlabel('$r$', fontsize=16)
ax.set_ylabel('$x_t$', fontsize=16)
plt.show()</pre>
```

==..,..,_.mages/python_oop_2 1_o.png

On the horizontal axis is the parameter r in (3).

The vertical axis is the state space [0, 1].

For each r we compute a long time series and then plot the tail (the last 50 points).

The tail of the sequence shows us where the trajectory concentrates after settling down to some kind of steady state, if a steady state exists.

Whether it settles down, and the character of the steady state to which it does settle down, depend on the value of r.

For r between about 2.5 and 3, the time series settles into a single fixed point plotted on the vertical axis.

For r between about 3 and 3.45, the time series settles down to oscillating between the two values plotted on the vertical axis.

For r a little bit higher than 3.45, the time series settles down to oscillating among the four values plotted on the vertical axis.

Notice that there is no value of r that leads to a steady state oscillating among three values.

Special Methods

Python provides special methods with which some neat tricks can be performed.

For example, recall that lists and tuples have a notion of length and that this length can be queried via the len function

```
x = (10, 20)
len(x)
```

2

If you want to provide a return value for the len function when applied to your user-defined object, use the __len__ special method

```
class Foo:
    def __len__(self):
        return 42
```

Now we get

```
f = Foo()
len(f)
```

42

A special method we will use regularly is the <u>__call__</u> method.

This method can be used to make your instances callable, just like functions

```
class Foo:
    def __call__(self, x):
        return x + 42
```

After running we get

```
f = Foo()
f(8) # Exactly equivalent to f.__call__(8)
```

50

Exercise 1 provides a more useful example.

Exercises

Exercise 1

The <u>empirical cumulative distribution function (ecdf)</u> corresponding to a sample $\{X_i\}_{i=1}^n$ is defined as

$$F_n(x) := \frac{1}{n} \sum_{i=1}^n \mathbf{1} \{ X_i \le x \} \qquad (x \in \mathbb{R})$$
 (4)

Here $\mathbf{1}\{X_i \leq x\}$ is an indicator function (one if $X_i \leq x$ and zero otherwise) and hence $F_n(x)$ is the fraction of the sample that falls below x.

The Glivenko-Cantelli Theorem states that, provided that the sample is IID, the ecdf F_n converges to the true distribution function F.

Implement F_n as a class called ECDF, where

- A given sample $\{X_i\}_{i=1}^n$ are the instance data, stored as self.observations.
- The class implements a __call__ method that returns $F_n(x)$ for any x.

Your code should work as follows (modulo randomness)

```
from random import uniform

samples = [uniform(0, 1) for i in range(10)]
F = ECDF(samples)
F(0.5) # Evaluate ecdf at x = 0.5
```

```
F.observations = [uniform(0, 1) for i in range(1000)]
F(0.5)
```

Aim for clarity, not efficiency.

Exercise 2

In an <u>earlier exercise</u>, you wrote a function for evaluating polynomials.

This exercise is an extension, where the task is to build a simple class called Polynomial for representing and manipulating polynomial functions such as

$$p(x) = a_0 + a_1 x + a_2 x^2 + \dots + a_N x^N = \sum_{n=0}^{N} a_n x^n \qquad (x \in \mathbb{R})$$
 (5)

The instance data for the class Polynomial will be the coefficients (in the case of (5), the numbers a_0, \ldots, a_N).

Provide methods that

- 1. Evaluate the polynomial (5), returning p(x) for any x.
- 2. Differentiate the polynomial, replacing the original coefficients with those of its derivative p'.

Avoid using any import statements.

Solutions

Exercise 1

```
# == test == #
from random import uniform
samples = [uniform(0, 1) for i in range(10)]
F = ECDF(samples)
print(F(0.5)) # Evaluate ecdf at x = 0.5
F.observations = [uniform(0, 1) for i in range(1000)]
print(F(0.5))
```

0.9 0.516

Exercise 2

```
class Polynomial:
    def __init__(self, coefficients):
        Creates an instance of the Polynomial class representing
             p(x) = a_0 x^0 + ... + a_N x^N,
        where a_i = coefficients[i].
        self.coefficients = coefficients
    def __call__(self, x):
   "Evaluate the polynomial at x."
        for i, a in enumerate(self.coefficients):
            y += a * x**i
        return y
    def differentiate(self):
         "Reset self.coefficients to those of p' instead of p."
         new_coefficients = []
        for i, a in enumerate(self.coefficients):
             new\_coefficients.append(i * a)
        # Remove the first element, which is zero
        del new_coefficients[0]
        # And reset coefficients data to new values
self.coefficients = new_coefficients
         return new_coefficients
```

The Scientific Libraries

Next we cover the third party libraries most useful for scientific work in Python

Python for Scientific Computing

In addition to what's in Anaconda, this lecture will need the following libraries:

!pip install --upgrade quantecon

Overview

Python is extremely popular for scientific computing, due to such factors as

- the accessible and flexible nature of the language itself,
- the huge range of high quality scientific libraries now available,
- the fact that the language and libraries are open source,
- the popular Anaconda Python distribution, which simplifies installation and management of those libraries, and
- the recent surge of interest in using Python for machine learning and artificial intelligence.

In this lecture we give a short overview of scientific computing in Python, addressing the following questions:

- What are the relative strengths and weaknesses of Python for these tasks?
- What are the main elements of the scientific Python ecosystem?
- How is the situation changing over time?

Scientific Libraries

Let's briefly review Python's scientific libraries, starting with why we need them.

The Role of Scientific Libraries

One obvious reason we use scientific libraries is because they implement routines we want to use.

For example, it's almost always better to use an existing routine for root finding than to write a new one from scratch.

(For standard algorithms, efficiency is maximized if the community can coordinate on a common set of implementations, written by experts and tuned by users to be as fast and robust as possible.)

But this is not the only reason that we use Python's scientific libraries.

Another is that pure Python, while flexible and elegant, is not fast.

So we need libraries that are designed to accelerate execution of Python code.

As we'll see below, there are now Python libraries that can do this extremely well.

Python's Scientific Ecosystem

In terms of popularity, the big four in the world of scientific Python libraries are

- NumPy
- SciPy
- Matplotlib
- Pandas

For us, there's another (relatively new) library that will also be essential for numerical computing:

Numba

Over the next few lectures we'll see how to use these libraries.

But first, let's quickly review how they fit together.

- NumPy forms the foundations by providing a basic array data type (think of vectors and matrices) and functions for acting on these arrays (e.g., matrix multiplication).
- SciPy builds on NumPy by adding the kinds of numerical methods that are routinely
 used in science (interpolation, optimization, root finding, etc.).
- Matplotlib is used to generate figures, with a focus on plotting data stored in NumPy arrays.
- Pandas provides types and functions for empirical work (e.g., manipulating data).
- Numba accelerates execution via JIT compilation we'll learn about this soon.

The Need for Speed

Now let's discuss execution speed.

Higher-level languages like Python are optimized for humans.

This means that the programmer can leave many details to the runtime environment

- specifying variable types
- memory allocation/deallocation, etc.

The upside is that, compared to low-level languages, Python is typically faster to write, less error-prone and easier to debug.

The downside is that Python is harder to optimize — that is, turn into fast machine code — than languages like C or Fortran.

Indeed, the standard implementation of Python (called CPython) cannot match the speed of compiled languages such as C or Fortran.

Does that mean that we should just switch to C or Fortran for everything?

The answer is: No, no and one hundred times no!

(This is what you should say to the senior professor insisting that the model needs to be rewritten in Fortran or C++.)

There are two reasons why:

First, for any given program, relatively few lines are ever going to be time-critical.

Hence it is far more efficient to write most of our code in a high productivity language like Python.

Second, even for those lines of code that *are* time-critical, we can now achieve the same speed as C or Fortran using Python's scientific libraries.

Where are the Bottlenecks?

Before we learn how to do this, let's try to understand why plain vanilla Python is slower than C or Fortran.

This will, in turn, help us figure out how to speed things up.

Dynamic Typing

Consider this Python operation

```
a, b = 10, 10
a + b
```

Even for this simple operation, the Python interpreter has a fair bit of work to do.

For example, in the statement a + b, the interpreter has to know which operation to invoke.

If a and b are strings, then a + b requires string concatenation

```
a, b = 'foo', 'bar'
a + b
```

'foobar'

If a and b are lists, then a + b requires list concatenation

```
a, b = ['foo'], ['bar']
a + b

['foo', 'bar']
```

(We say that the operator + is *overloaded* — its action depends on the type of the objects on which it acts)

As a result, Python must check the type of the objects and then call the correct operation.

This involves substantial overheads.

Static Types

Compiled languages avoid these overheads with explicit, static types.

For example, consider the following C code, which sums the integers from 1 to 10

```
#include <stdio.h>
int main(void) {
    int i;
    int sum = 0;
    for (i = 1; i <= 10; i++) {
        sum = sum + i;
    }
    printf("sum = %d\n", sum);
    return 0;
}</pre>
```

The variables i and sum are explicitly declared to be integers.

Hence, the meaning of addition here is completely unambiguous.

Data Access

Another drag on speed for high-level languages is data access.

To illustrate, let's consider the problem of summing some data — say, a collection of integers.

Summing with Compiled Code

In C or Fortran, these integers would typically be stored in an array, which is a simple data structure for storing homogeneous data.

Such an array is stored in a single contiguous block of memory

- In modern computers, memory addresses are allocated to each byte (one byte = 8 bits).
- For example, a 64 bit integer is stored in 8 bytes of memory.
- An array of n such integers occupies 8n consecutive memory slots.

Moreover, the compiler is made aware of the data type by the programmer.

• In this case 64 bit integers

Hence, each successive data point can be accessed by shifting forward in memory space by a known and fixed amount.

• In this case 8 bytes

Summing in Pure Python

Python tries to replicate these ideas to some degree.

For example, in the standard Python implementation (CPython), list elements are placed in memory locations that are in a sense contiguous.

However, these list elements are more like pointers to data rather than actual data.

Hence, there is still overhead involved in accessing the data values themselves.

This is a considerable drag on speed.

In fact, it's generally true that memory traffic is a major culprit when it comes to slow execution.

Let's look at some ways around these problems.

Vectorization

There is a clever method called **vectorization** that can be used to speed up high level languages in numerical applications.

The key idea is to send array processing operations in batch to pre-compiled and efficient native machine code.

The machine code itself is typically compiled from carefully optimized C or Fortran.

For example, when working in a high level language, the operation of inverting a large matrix can be subcontracted to efficient machine code that is pre-compiled for this purpose and supplied to users as part of a package.

This clever idea dates back to MATLAB, which uses vectorization extensively.

Vectorization can greatly accelerate many numerical computations (but not all, as we shall see).

Let's see how vectorization works in Python, using NumPy.

Operations on Arrays

First, let's run some imports

```
import random
import numpy as np
import quantecon as qe
```

Next let's try some non-vectorized code, which uses a native Python loop to generate, square and then sum a large number of random variables:

```
n = 1_000_000

%%time
y = 0  # Will accumulate and store sum
for i in range(n):
    x = random.uniform(0, 1)
    y += x**2
```

```
CPU times: user 353 ms, sys: 777 \mus, total: 354 ms Wall time: 354 ms
```

The following vectorized code achieves the same thing.

```
%%time
x = np.random.uniform(0, 1, n)
y = np.sum(x**2)
```

CPU times: user 11.2 ms, sys: 3.15 ms, total: 14.4 ms Wall time: 14.2 ms $\,$

As you can see, the second code block runs much faster. Why?

The second code block breaks the loop down into three basic operations

- 1. draw n uniforms
- 2. square them
- 3. sum them

These are sent as batch operators to optimized machine code.

Apart from minor overheads associated with sending data back and forth, the result is ${\sf C}$ or Fortran-like speed.

When we run batch operations on arrays like this, we say that the code is vectorized.

Vectorized code is typically fast and efficient.

It is also surprisingly flexible, in the sense that many operations can be vectorized.

The next section illustrates this point.

Universal Functions

Many functions provided by NumPy are so-called universal functions — also called ufuncs.

This means that they

- map scalars into scalars, as expected
- map arrays into arrays, acting element-wise

For example, np. cos is a ufunc:

```
np.cos(1.0)
```

0.5403023058681398

```
np.cos(np.linspace(0, 1, 3))
```

array([1. , 0.87758256, 0.54030231])

By exploiting ufuncs, many operations can be vectorized.

For example, consider the problem of maximizing a function f of two variables (x, y) over the square $[-a, a] \times [-a, a]$.

For f and a let's choose

$$f(x, y) = \frac{\cos(x^2 + y^2)}{1 + x^2 + y^2}$$
 and $a = 3$

Here's a plot of f

```
import matplotlib.pyplot as plt
%matplotlib inline
from mpl_toolkits.mplot3d.axes3d import Axes3D
from matplotlib import cm
def f(x, y):
     return np.cos(x**2 + y**2) / (1 + x**2 + y**2)
xgrid = np.linspace(-3, 3, 50)
ygrid = xgrid
x, y = np.meshgrid(xgrid, ygrid)
fig = plt.figure(figsize=(8, 6))
ax = fig.add_subplot(111, projection='3d')
ax.plot_surface(x,
                   y,
f(x, y),
rstride=2, cstride=2,
cmap=cm.jet,
alpha=0.7,
                   linewidth=0.25)
ax.set_zlim(-0.5, 1.0)
ax.set_xlabel('$x$', fontsize=14)
ax.set_ylabel('$y$', fontsize=14)
plt.show()
```

.../../_images/need_for_speed_10_0.png

To maximize it, we're going to use a naive grid search:

- 1. Evaluate f for all (x, y) in a grid on the square.
- 2. Return the maximum of observed values.

The grid will be

```
grid = np.linspace(-3, 3, 1000)
```

Here's a non-vectorized version that uses Python loops.

```
%%time
m = -np.inf
for x in grid:
    for y in grid:
z = f(x, y)
         if z > m:
              m = z
```

```
CPU times: user 3.51 s, sys: 2.43 ms, total: 3.51 s
Wall time: 3.16 s
```

And here's a vectorized version

```
%%time
x, y = np.meshgrid(grid, grid)
np.max(f(x, y))
CPU times: user 20.6 ms, sys: 5.46 ms, total: 26.1 ms
Wall time: 25.8 ms
0.9999819641085747
```

In the vectorized version, all the looping takes place in compiled code.

As you can see, the second version is **much** faster.

(We'll make it even faster again later on, using more scientific programming tricks.)

Beyond Vectorization

At its best, vectorization yields fast, simple code.

However, it's not without disadvantages.

One issue is that it can be highly memory-intensive.

For example, the vectorized maximization routine above is far more memory intensive than the non-vectorized version that preceded it.

This is because vectorization tends to create many intermediate arrays before producing the final calculation.

Another issue is that not all algorithms can be vectorized.

In these kinds of settings, we need to go back to loops.

Fortunately, there are alternative ways to speed up Python loops that work in almost any setting.

For example, in the last few years, a new Python library called [Numba] (http://numba.pydata.org/) has appeared that solves the main problems with vectorization listed above.

It does so through something called **just in time (JIT) compilation**, which can generate extremely fast and efficient code.

We'll learn how to use Numba soon.

NumPy

"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

NumPy is a first-rate library for numerical programming

- Widely used in academia, finance and industry.
- Mature, fast, stable and under continuous development.

We have already seen some code involving NumPy in the preceding lectures.

In this lecture, we will start a more systematic discussion of both

- NumPy arrays and
- $\bullet \;\;$ the fundamental array processing operations provided by NumPy.

References

• The official NumPy documentation.

NumPy Arrays

The essential problem that NumPy solves is fast array processing.

The most important structure that NumPy defines is an array data type formally called a <u>numpy.ndarray</u>.

NumPy arrays power a large proportion of the scientific Python ecosystem.

Let's first import the library.

```
import numpy as np
```

To create a NumPy array containing only zeros we use <u>np.zeros</u>

```
a = np.zeros(3)
a
```

```
array([0., 0., 0.])
```

```
type(a)
```

numpy.ndarray

NumPy arrays are somewhat like native Python lists, except that

- Data must be homogeneous (all elements of the same type).
- These types must be one of the <u>data types</u> (dtypes) provided by NumPy.

The most important of these dtypes are:

- float64: 64 bit floating-point number
- int64: 64 bit integer
- bool: 8 bit True or False

There are also dtypes to represent complex numbers, unsigned integers, etc.

On modern machines, the default dtype for arrays is float64

```
a = np.zeros(3)
type(a[0])
```

numpy.float64

If we want to use integers we can specify as follows:

```
a = np.zeros(3, dtype=int)
type(a[0])
```

numpy.int64

Shape and Dimension

Consider the following assignment

```
z = np.zeros(10)
```

Here ${\bf z}$ is a ${\it flat}$ array with no dimension — neither row nor column vector.

The dimension is recorded in the shape attribute, which is a tuple

```
z.shape (10,)
```

Here the shape tuple has only one element, which is the length of the array (tuples with one element end with a comma).

To give it dimension, we can change the $\ensuremath{\mathsf{shape}}$ attribute

```
z = np.zeros(4)
z.shape = (2, 2)
z
```

```
array([[0., 0.],
[0., 0.]])
```

In the last case, to make the 2 by 2 array, we could also pass a tuple to the zeros() function, as in z = np.zeros((2, 2)).

Creating Arrays

As we've seen, the np. zeros function creates an array of zeros.

You can probably guess what np.ones creates.

Related is np.empty, which creates arrays in memory that can later be populated with data

```
z = np.empty(3)
z
array([0., 0., 0.])
```

The numbers you see here are garbage values.

(Python allocates 3 contiguous 64 bit pieces of memory, and the existing contents of those memory slots are interpreted as float64 values)

To set up a grid of evenly spaced numbers use np.linspace

```
z = np.linspace(2, 4, 5) # From 2 to 4, with 5 elements
```

To create an identity matrix use either np.identity or np.eye

In addition, NumPy arrays can be created from Python lists, tuples, etc. using np.array

```
type(z)
```

numpy.ndarray

```
z = np.array((10, 20), dtype=float) # Here 'float' is equivalent to 'np.float64' z
```

```
array([10., 20.])
```

```
z = np.array([[1, 2], [3, 4]])  # 2D array from a list of lists
z
```

```
array([[1, 2],
[3, 4]])
```

See also np. asarray, which performs a similar function, but does not make a distinct copy of data already in a NumPy array.

```
na = np.linspace(10, 20, 2)
na is np.asarray(na) # Does not copy NumPy arrays
```

True

```
na is np.array(na) # Does make a new copy --- perhaps unnecessarily
```

False

To read in the array data from a text file containing numeric data use np.loadtxt or np.genfromtxt—see the documentation for details.

Array Indexing

For a flat array, indexing is the same as Python sequences:

```
z = np.linspace(1, 2, 5)
z
array([1. , 1.25, 1.5 , 1.75, 2. ])

z[0]
1.0

z[0:2] # Two elements, starting at element 0
array([1. , 1.25])

z[-1]
2.0
```

For 2D arrays the index syntax is as follows:

And so on.

Note that indices are still zero-based, to maintain compatibility with Python sequences.

Columns and rows can be extracted as follows

```
z[0, :]
array([1, 2])

z[:, 1]
array([2, 4])
```

 $\label{eq:num-py} \mbox{NumPy arrays of integers can also be used to extract elements}$

```
z = np.linspace(2, 4, 5)
z
```

```
array([2. , 2.5, 3. , 3.5, 4. ])
```

```
indices = np.array((0, 2, 3))
z[indices]
```

```
array([2. , 3. , 3.5])
```

Finally, an array of dtype bool can be used to extract elements

```
z
array([2. , 2.5, 3. , 3.5, 4. ])

d = np.array([0, 1, 1, 0, 0], dtype=bool)
d
array([False, True, True, False, False])

z[d]
array([2.5, 3. ])
```

We'll see why this is useful below.

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

```
z = np.empty(3)
z
array([2. , 3. , 3.5])

z[:] = 42
z
array([42., 42., 42.])
```

Array Methods

Arrays have useful methods, all of which are carefully optimized

Max

```
a = np.array((4, 3, 2, 1))

array([4, 3, 2, 1])

a.sort()  # Sorts a in place
array([1, 2, 3, 4])

a.sum()  # Sum

10

a.mean()  # Mean

2.5
```

a.max()

```
a.argmax()
                      # Returns the index of the maximal element
3
a.cumsum()
                      # Cumulative sum of the elements of a
array([ 1, 3, 6, 10])
a.cumprod()
                      # Cumulative product of the elements of a
array([ 1, 2, 6, 24])
a.var()
                      # Variance
1.25
                      # Standard deviation
a.std()
1.118033988749895
a.shape = (2, 2)
                      # Equivalent to a.transpose()
array([[1, 3],
[2, 4]])
```

Another method worth knowing is searchsorted().

If z is a nondecreasing array, then z \cdot searchsorted(a) returns the index of the first element of z that is >= a

```
z = np.linspace(2, 4, 5)
z
array([2. , 2.5, 3. , 3.5, 4. ])
z.searchsorted(2.2)
```

Many of the methods discussed above have equivalent functions in the NumPy namespace

```
a = np.array((4, 3, 2, 1))
np.sum(a)

10

np.mean(a)
```

2.5

Operations on Arrays

Arithmetic Operations

The operators +, -, *, / and ** all act *elementwise* on arrays

```
a = np.array([1, 2, 3, 4])
b = np.array([5, 6, 7, 8])
a + b
```

```
array([ 6, 8, 10, 12])
```

```
a * b
array([ 5, 12, 21, 32])
```

We can add a scalar to each element as follows

```
a + 10
array([11, 12, 13, 14])
```

Scalar multiplication is similar

```
a * 10
array([10, 20, 30, 40])
```

The two-dimensional arrays follow the same general rules

In particular, A * B is *not* the matrix product, it is an element-wise product.

Matrix Multiplication

With Anaconda's scientific Python package based around Python 3.5 and above, one can use the @symbol for matrix multiplication, as follows:

```
A = np.ones((2, 2))
B = np.ones((2, 2))
A @ B

array([[2., 2.],
[2., 2.]])
```

(For older versions of Python and NumPy you need to use the <u>np.dot</u> function)

We can also use $\ensuremath{\text{@}}$ to take the inner product of two flat arrays

```
A = np.array((1, 2))
B = np.array((10, 20))
A @ B
```

50

In fact, we can use @ when one element is a Python list or tuple

```
A = np.array(((1, 2), (3, 4)))
A
```

```
array([[1, 2], [3, 4]])
```

```
A @ (0, 1)
array([2, 4])
```

Since we are post-multiplying, the tuple is treated as a column vector.

Mutability and Copying Arrays

NumPy arrays are mutable data types, like Python lists.

In other words, their contents can be altered (mutated) in memory after initialization.

We already saw examples above.

Here's another example:

```
a = np.array([42, 44])
array([42, 44])

a[-1] = 0 # Change last element to θ
array([42, 0])
```

 $\label{thm:matching} \mbox{Mutability leads to the following behavior (which can be shocking to MATLAB programmers...)}$

```
a = np.random.randn(3)
a
array([0.9779746 , 0.6735592 , 0.90282662])

b = a
b[0] = 0.0
a
```

What's happened is that we have changed a by changing b.

The name b is bound to a and becomes just another reference to the array (the Python assignment model is described in more detail <u>later in the course</u>).

, 0.6735592 , 0.90282662])

Hence, it has equal rights to make changes to that array.

This is in fact the most sensible default behavior!

It means that we pass around only pointers to data, rather than making copies.

Making copies is expensive in terms of both speed and memory.

Making Copies

array([0.

It is of course possible to make b an independent copy of a when required.

This can be done using $\mathsf{np}_{\:\raisebox{1pt}{\text{\circle*{1.5}}}}\mathsf{copy}$

```
a = np.random.randn(3)
a
```

```
array([0.47334726, 0.85466086, 1.27929065])
```

```
b = np.copy(a)
b
```

```
array([0.47334726, 0.85466086, 1.27929065])
```

Now b is an independent copy (called a deep copy)

```
b[:] = 1
b
array([1., 1., 1.])

a
array([0.47334726, 0.85466086, 1.27929065])
```

Note that the change to b has not affected a.

Additional Functionality

Let's look at some other useful things we can do with NumPy.

Vectorized Functions

NumPy provides versions of the standard functions \log , \exp , \sin , etc. that act *element-wise* on arrays

```
z = np.array([1, 2, 3])
np.sin(z)
array([0.84147098, 0.90929743, 0.14112001])
```

This eliminates the need for explicit element-by-element loops such as

```
n = len(z)
y = np.empty(n)
for i in range(n):
    y[i] = np.sin(z[i])
```

Because they act element-wise on arrays, these functions are called *vectorized functions*.

In NumPy-speak, they are also called *ufuncs*, which stands for "universal functions".

As we saw above, the usual arithmetic operations (+, *, etc.) also work element-wise, and combining these with the utuncs gives a very large set of fast element-wise functions.

```
z

array([1, 2, 3])

(1 / np.sqrt(2 * np.pi)) * np.exp(- 0.5 * z**2)

array([0.24197072, 0.05399097, 0.00443185])
```

Not all user-defined functions will act element-wise.

For example, passing the function f defined below a NumPy array causes a ValueError

```
def f(x):
    return 1 if x > 0 else 0
```

The NumPy function np.where provides a vectorized alternative:

```
x = np.random.randn(4)
x
```

```
array([-0.22108355, -0.00622957, 0.10792989, -0.08008869])
```

You can also use np. vectorize to vectorize a given function

```
array([0, 0, 1, 0])
```

However, this approach doesn't always obtain the same speed as a more carefully crafted vectorized function.

Comparisons

As a rule, comparisons on arrays are done element-wise

```
z = np.array([2, 3])
y = np.array([2, 3])
z == y
```

```
array([ True, True])
```

```
y[0] = 5

z == y
```

array([False, True])

```
z != y
```

```
array([ True, False])
```

The situation is similar for >, <, >= and <=.

We can also do comparisons against scalars

```
z = np.linspace(0, 10, 5)
z
```

```
array([ 0. , 2.5, 5. , 7.5, 10. ])
```

```
z > 3
```

```
array([False, False, True, True, True])
```

This is particularly useful for conditional extraction

```
b = z > 3
b
```

```
array([False, False, True, True, True])
```

```
z[b]
```

```
array([ 5. , 7.5, 10. ])
```

Of course we can—and frequently do—perform this in one step

```
z[z > 3]
```

```
array([ 5. , 7.5, 10. ])
```

Sub-packages

 $\label{lem:numPy} NumPy\ provides\ some\ additional\ functional ity\ related\ to\ scientific\ programming\ through\ its\ sub-packages.$

We've already seen how we can generate random variables using np. random

```
z = np.random.randn(10000)  # Generate standard normals
y = np.random.binomial(10, 0.5, size=1000)  # 1,000 draws from Bin(10, 0.5)
y.mean()
```

5.014

Another commonly used subpackage is np.linalg

```
A = np.array([[1, 2], [3, 4]])
np.linalg.det(A) # Compute the determinant
```

-2.000000000000000004

Much of this functionality is also available in <u>SciPy</u>, a collection of modules that are built on top of NumPy.

We'll cover the SciPy versions in more detail soon.

For a comprehensive list of what's available in NumPy see this documentation.

Exercises

Exercise 1

Consider the polynomial expression

$$p(x) = a_0 + a_1 x + a_2 x^2 + \dots + a_N x^N = \sum_{n=0}^{N} a_n x^n$$
 (6)

<u>Earlier</u>, you wrote a simple function p(x, coeff) to evaluate $(\underline{6})$ without considering efficiency.

Now write a new function that does the same job, but uses NumPy arrays and array operations for its computations, rather than any form of Python loop.

(Such functionality is already implemented as np.poly1d, but for the sake of the exercise don't use this class)

• Hint: Use np.cumprod()

Exercise 2

Let q be a NumPy array of length n with q.sum() == 1.

Suppose that q represents a <u>probability mass function</u>.

We wish to generate a discrete random variable x such that $\mathbb{P}\{x=i\}=q_i$.

In other words, x takes values in range (len(g)) and x = i with probability g[i].

The standard (inverse transform) algorithm is as follows:

- Divide the unit interval [0,1] into n subintervals I_0,I_1,\ldots,I_{n-1} such that the length of I_i is a_i
- Draw a uniform random variable U on [0,1] and return the i such that $U \in I_i$.

The probability of drawing i is the length of I_i , which is equal to q_i .

We can implement the algorithm as follows

```
from random import uniform

def sample(q):
    a = 0.0
    U = uniform(0, 1)
    for i in range(len(q)):
        if a < U <= a + q[i]:
            return i
        a = a + q[i]</pre>
```

If you can't see how this works, try thinking through the flow for a simple example, such as q = [0.25, 0.75] It helps to sketch the intervals on paper.

Your exercise is to speed it up using NumPy, avoiding explicit loops

• Hint: Use np. searchsorted and np. cumsum

If you can, implement the functionality as a class called ${\tt DiscreteRV}$, where

- the data for an instance of the class is the vector of probabilities q
- the class has a draw() method, which returns one draw according to the algorithm described above

If you can, write the method so that draw(k) returns k draws from q.

Exercise 3

Recall our <u>earlier discussion</u> of the empirical cumulative distribution function.

Your task is to

- 1. Make the __call__ method more efficient using NumPy.
- 2. Add a method that plots the ECDF over [a, b], where a and b are method parameters.

Solutions

```
import matplotlib.pyplot as plt
%matplotlib inline
```

Exercise 1

This code does the job

```
def p(x, coef):
    X = np.ones_like(coef)
    X[1:] = x
    y = np.cumprod(X)  # y = [1, x, x**2,...]
    return coef @ y
```

Let's test it

```
x = 2
coef = np.linspace(2, 4, 3)
print(coef)
print(p(x, coef))
# For comparison
q = np.poly1d(np.flip(coef))
print(q(x))
```

```
[2. 3. 4.]
24.0
24.0
```

Exercise 2

Here's our first pass at a solution:

```
from numpy import cumsum
from numpy.random import uniform

class DiscreteRV:
    Generates an array of draws from a discrete random variable with
vector of
    probabilities given by q.

def __init__(self, q):
    The argument q is a NumPy array, or array like, nonnegative and
sums

to 1
    self.q = q
    self.Q = cumsum(q)

def draw(self, k=1):
    Returns k draws from q. For each such draw, the value i is
returned
    with probability q[i].
    return self.Q.searchsorted(uniform(0, 1, size=k))
```

The logic is not obvious, but if you take your time and read it slowly, you will understand.

There is a problem here, however.

Suppose that g is altered after an instance of discreteRV is created, for example by

```
q = (0.1, 0.9)
d = DiscreteRV(q)
d.q = (0.5, 0.5)
```

The problem is that Q does not change accordingly, and Q is the data used in the draw method.

To deal with this, one option is to compute Q every time the draw method is called.

But this is inefficient relative to computing **Q** once-off.

A better option is to use descriptors.

A solution from the $\underline{\text{quantecon library}}$ using descriptors that behaves as we desire can be found $\underline{\text{here}}$.

Exercise 3

An example solution is given below.

In essence, we've just taken this code from QuantEcon and added in a plot method

```
0.00
Modifies ecdf.py from QuantEcon to add in a plot method
class ECDF:
    One-dimensional empirical distribution function given a vector of
    Parameters
    observations : array_like
         An array of observations
    Attributes
    observations : array_like
An array of observations
         __init__(self, observations):
self.observations = np.asarray(observations)
    def _
    def __call__(self, x):
         Evaluates the ecdf at x
         Parameters
         x : scalar(float)
             The x at which the ecdf is evaluated
         Returns
         scalar(float)
              Fraction of the sample less than x
         return np.mean(self.observations <= x)</pre>
    def plot(self, ax, a=None, b=None):
         Plot the ecdf on the interval [a, b].
         Parameters
         a : scalar(float), optional(default=None)
Lower endpoint of the plot interval
         b: scalar(float), optional(default=None)
Upper endpoint of the plot interval
         .....
         # === choose reasonable interval if [a, b] not specified === #
         if a is None:
              a = self.observations.min() - self.observations.std()
         if b is None:
             b = self.observations.max() + self.observations.std()
         # === generate plot === #
         x_vals = np.linspace(a, b, num=100)
f = np.vectorize(self.__call__)
         ax.plot(x_vals, f(x_vals))
plt.show()
```

Here's an example of usage

```
fig, ax = plt.subplots()
X = np.random.randn(1000)
F = ECDF(X)
F.plot(ax)

.../_images/numpy_95_0.png
```

Matplotlib

Overview

We've already generated quite a few figures in these lectures using Matplotlib.

 $Matplot lib is an outstanding \ graphics \ library, \ designed \ for \ scientific \ computing, \ with$

- high-quality 2D and 3D plots
- output in all the usual formats (PDF, PNG, etc.)
- LaTeX integration

- fine-grained control over all aspects of presentation
- animation, etc.

Matplotlib's Split Personality

Matplotlib is unusual in that it offers two different interfaces to plotting.

One is a simple MATLAB-style API (Application Programming Interface) that was written to help MATLAB refugees find a ready home.

The other is a more "Pythonic" object-oriented API.

For reasons described below, we recommend that you use the second API.

But first, let's discuss the difference.

The APIs

The MATLAB-style API

Here's the kind of easy example you might find in introductory treatments

```
import matplotlib.pyplot as plt
%matplotlib inline
import numpy as np

x = np.linspace(0, 10, 200)
y = np.sin(x)

plt.plot(x, y, 'b-', linewidth=2)
plt.show()
```

....._images/matplotlib_0_0.png

This is simple and convenient, but also somewhat limited and un-Pythonic.

For example, in the function calls, a lot of objects get created and passed around without making themselves known to the programmer.

Python programmers tend to prefer a more explicit style of programming (run import this in a code block and look at the second line).

This leads us to the alternative, object-oriented Matplotlib API.

The Object-Oriented API

Here's the code corresponding to the preceding figure using the object-oriented API

```
fig, ax = plt.subplots()
ax.plot(x, y, 'b-', linewidth=2)
plt.show()

.../_images/matplotlib_1_0.png
```

Here the call fig, ax = plt.subplots() returns a pair, where

- fig is a Figure instance—like a blank canvas.
- ax is an AxesSubplot instance—think of a frame for plotting in.

The plot() function is actually a method of ax.

While there's a bit more typing, the more explicit use of objects gives us better control.

This will become more clear as we go along.

Tweaks

Here we've changed the line to red and added a legend

```
fig, ax = plt.subplots()
ax.plot(x, y, 'r-', linewidth=2, label='sine function', alpha=0.6)
ax.legend()
plt.show()
.../_images/matplotlib_2_0.png
```

We've also used alpha to make the line slightly transparent—which makes it look smoother.

The location of the legend can be changed by replacing ax.legend() with ax.legend(loc='upper center').

```
fig, ax = plt.subplots()
ax.plot(x, y, 'r-', linewidth=2, label='sine function', alpha=0.6)
ax.legend(loc='upper center')
plt.show()

.../_images/matplotlib_3_0.png
```

If everything is properly configured, then adding LaTeX is trivial

```
fig, ax = plt.subplots()
ax.plot(x, y, 'r-', linewidth=2, label='$y=\sin(x)$', alpha=0.6)
ax.legend(loc='upper center')
plt.show()

.../_images/matplotlib_4_0.png
```

Controlling the ticks, adding titles and so on is also straightforward $% \left(1\right) =\left(1\right) \left(1\right) \left($

```
fig, ax = plt.subplots()
ax.plot(x, y, 'r-', linewidth=2, label='$y=\sin(x)$', alpha=0.6)
ax.legend(loc='upper center')
ax.set_yticks([-1, 0, 1])
ax.set_title('Test plot')
plt.show()
```

More Features

Matplotlib has a huge array of functions and features, which you can discover over time as you have need for them.

We mention just a few.

Multiple Plots on One Axis

It's straightforward to generate multiple plots on the same axes.

Here's an example that randomly generates three normal densities and adds a label with their mean

```
from scipy.stats import norm
from random import uniform

fig, ax = plt.subplots()
x = np.linspace(-4, 4, 150)
for i in range(3):
    m, s = uniform(-1, 1), uniform(1, 2)
    y = norm.pdf(x, loc=m, scale=s)
    current_label = f'$\mu = {m:.2}$'
    ax.plot(x, y, linewidth=2, alpha=0.6, label=current_label)
ax.legend()
plt.show()
```

.../../_images/matplotlib_6_0.png

Multiple Subplots

Sometimes we want multiple subplots in one figure.

Here's an example that generates 6 histograms $\,$

```
num_rows, num_cols = 3, 2
fig, axes = plt.subplots(num_rows, num_cols, figsize=(10, 12))
for i in range(num_rows):
    for j in range(num_cols):
        m, s = uniform(-1, 1), uniform(1, 2)
        x = norm.rvs(loc=m, scale=s, size=100)
        axes[i, j].hist(x, alpha=0.6, bins=20)
        t = f'$\mu = {m:.2}, \quad \sigma = {s:.2}$'
        axes[i, j].set(title=t, xticks=[-4, 0, 4], yticks=[])
plt.show()

..../_images/matplotlib_7_0.png
```

., ., _....ages, ...acpret...s_, _e

3D Plots

Matplotlib does a nice job of 3D plots — here is one example

.../../_images/matplotlib_8_0.png

A Customizing Function

Perhaps you will find a set of customizations that you regularly use.

Suppose we usually prefer our axes to go through the origin, and to have a grid.

Here's a nice example from <u>Matthew Doty</u> of how the object-oriented API can be used to build a custom subplots function that implements these changes.

Read carefully through the code and see if you can follow what's going on

```
def subplots():
    "Custom subplots with axes through the origin"
    fig, ax = plt.subplots()

# Set the axes through the origin
    for spine in ['left', 'bottom']:
        ax.spines[spine].set_position('zero')
    for spine in ['right', 'top']:
        ax.spines[spine].set_color('none')

    ax.grid()
    return fig, ax

fig, ax = subplots() # Call the local version, not plt.subplots()
    x = np.linspace(-2, 10, 200)
    y = np.sin(x)
    ax.plot(x, y, 'r-', linewidth=2, label='sine function', alpha=0.6)
    ax.legend(loc='lower right')
    plt.show()

...J._images/matplotlib_9_0.png
```

The custom subplots function

- 1. calls the standard plt.subplots function internally to generate the fig, ax pair,
- 2. makes the desired customizations to ax, and
- 3. passes the fig, ax pair back to the calling code.

Further Reading

- The Matplotlib gallery provides many examples.
- A nice Matplotlib tutorial by Nicolas Rougier, Mike Muller and Gael Varoquaux.
- mpltools allows easy switching between plot styles.
- Seaborn facilitates common statistics plots in Matplotlib.

Exercises

Exercise 1

Plot the function

```
f(x) = \cos(\pi\theta x) \exp(-x)
```

over the interval [0,5] for each θ in np.linspace(0, 2, 10).

Place all the curves in the same figure.

The output should look like this

Solutions

Exercise 1

Here's one solution

```
def f(x, θ):
    return np.cos(np.pi * θ * x ) * np.exp(- x)

θ_vals = np.linspace(0, 2, 10)
x = np.linspace(0, 5, 200)
fig, ax = plt.subplots()

for θ in θ_vals:
    ax.plot(x, f(x, θ))
plt.show()

plt.show()
```

SciPy

Overview

SciPy builds on top of NumPy to provide common tools for scientific programming such as

- <u>linear algebra</u>
- numerical integration
- interpolation
- optimization
- <u>distributions and random number generation</u>
- signal processing
- etc., etc

Like NumPy, SciPy is stable, mature and widely used.

Many SciPy routines are thin wrappers around industry-standard Fortran libraries such as <u>LAPACK</u>, <u>BLAS</u>, etc.

It's not really necessary to "learn" SciPy as a whole.

A more common approach is to get some idea of what's in the library and then look up documentation as required.

In this lecture, we aim only to highlight some useful parts of the package.

SciPy versus NumPy

SciPy is a package that contains various tools that are built on top of NumPy, using its array data type and related functionality.

In fact, when we import SciPy we also get NumPy, as can be seen from this excerpt the SciPy initialization file:

```
# Import numpy symbols to scipy namespace
from numpy import *
from numpy.random import rand, randn
from numpy.fft import fft, ifft
from numpy.lib.scimath import *
```

However, it's more common and better practice to use NumPy functionality explicitly

```
import numpy as np
a = np.identity(3)
```

What is useful in SciPy is the functionality in its sub-packages

• scipy.optimize, scipy.integrate, scipy.stats, etc.

Let's explore some of the major sub-packages.

Statistics

The scipy.stats subpackage supplies

- numerous random variable objects (densities, cumulative distributions, random sampling, etc.)
- some estimation procedures
- · some statistical tests

Random Variables and Distributions

Recall that numpy. random provides functions for generating random variables

```
np.random.beta(5, 5, size=3)
array([0.65925144, 0.74668457, 0.48906918])
```

This generates a draw from the distribution with the density function below when a, b = 5,

$$f(x; a, b) = \frac{x^{(a-1)}(1-x)^{(b-1)}}{\int_0^1 u^{(a-1)}(1-u)^{(b-1)} du} \qquad (0 \le x \le 1)$$

Sometimes we need access to the density itself, or the cdf, the quantiles, etc.

For this, we can use scipy. stats, which provides all of this functionality as well as random number generation in a single consistent interface.

Here's an example of usage

```
from scipy.stats import beta
import matplotlib.pyplot as plt
%matplotlib inline

q = beta(5, 5)  # Beta(a, b), with a = b = 5
obs = q.rvs(2000)  # 2000 observations
grid = np.linspace(0.01, 0.99, 100)

fig, ax = plt.subplots()
ax.hist(obs, bins=40, density=True)
ax.plot(grid, q.pdf(grid), 'k-', linewidth=2)
plt.show()
```

The object q that represents the distribution has additional useful methods, including

```
q.cdf(0.4) # Cumulative distribution function
```

0.26656768000000003

```
q.ppf(0.8) # Quantile (inverse cdf) function
```

0.6339134834642708

```
q.mean()
```

0.5

The general syntax for creating these objects that represent distributions (of type rv_frozen) is

```
name = scipy.stats.distribution_name(shape_parameters, loc=c,
scale=d)
```

Here distribution_name is one of the distribution names in scipy.stats.

The loc and scale parameters transform the original random variable X into Y=c+dX.

Alternative Syntax

There is an alternative way of calling the methods described above.

For example, the code that generates the figure above can be replaced by

```
obs = beta.rvs(5, 5, size=2000)
grid = np.linspace(0.01, 0.99, 100)

fig, ax = plt.subplots()
ax.hist(obs, bins=40, density=True)
ax.plot(grid, beta.pdf(grid, 5, 5), 'k-', linewidth=2)
plt.show()

..../_images/scipy_7_0.png
```

Other Goodies in scipy.stats

There are a variety of statistical functions in scipy.stats.

 $For example, \verb|scipy.stats.linregress| implements simple linear regression$

```
from scipy.stats import linregress

x = np.random.randn(200)
y = 2 * x + 0.1 * np.random.randn(200)
gradient, intercept, r_value, p_value, std_err = linregress(x, y)
gradient, intercept

(1.9987406665842973, -0.0017072286417713356)
```

To see the full list, consult the documentation.

Roots and Fixed Points

A **root** or **zero** 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

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

(7)

```
f = lambda x: np.sin(4 * (x - 1/4)) + x + x**20 - 1
x = np.linspace(0, 1, 100)

fig, ax = plt.subplots()
ax.plot(x, f(x))
ax.axhline(ls='--', c='k', label='$f(x)$')
ax.set_xlabel('$x$', fontsize=12)
ax.set_ylabel('$f(x)$', fontsize=12)
ax.legend(fontsize=12)
plt.show()
```

The unique root is approximately 0.408.

Let's consider some numerical techniques for finding roots.

Bisection

One of the most common algorithms for numerical root-finding is bisection.

To understand the idea, recall the well-known game where

- Player A thinks of a secret number between 1 and 100
- Player B asks if it's less than 50

```
If yes, B asks if it's less than 25If no, B asks if it's less than 75
```

And so on.

This is bisection.

Here's a simplistic implementation of the algorithm in Python.

It works for all sufficiently well behaved increasing continuous functions with f(a) < 0 < f(b)

Let's test it using the function f defined in (7)

```
bisect(f, 0, 1)
```

0.408294677734375

Not surprisingly, SciPy provides its own bisection function.

Let's test it using the same function f defined in (7)

```
from scipy.optimize import bisect
bisect(f, 0, 1)
```

0.4082935042806639

The Newton-Raphson Method

Another very common root-finding algorithm is the Newton-Raphson method.

In SciPy this algorithm is implemented by scipy.optimize.newton.

Unlike bisection, the Newton-Raphson method uses local slope information in an attempt to increase the speed of convergence.

Let's investigate this using the same function f defined above.

With a suitable initial condition for the search we get convergence:

```
from scipy.optimize import newton newton(f, 0.2) # Start the search at initial condition x = 0.2
```

0.40829350427935673

But other initial conditions lead to failure of convergence:

```
newton(f, 0.7) # Start the search at x = 0.7 instead
```

0.70017000000000279

Hybrid Methods

A general principle of numerical methods is as follows:

- If you have specific knowledge about a given problem, you might be able to exploit it to generate efficiency.
- If not, then the choice of algorithm involves a trade-off between speed and robustness. In practice, most default algorithms for root-finding, optimization and fixed points use *hybrid* methods.

These methods typically combine a fast method with a robust method in the following manner:

- 1. Attempt to use a fast method
- 2. Check diagnostics
- 3. If diagnostics are bad, then switch to a more robust algorithm

In scipy.optimize, the function brentq is such a hybrid method and a good default

```
from scipy.optimize import brentq
brentq(f, 0, 1)
```

0.40829350427936706

Here the correct solution is found and the speed is better than bisection:

```
%timeit brentq(f, 0, 1)

16.7 µs ± 309 ns per loop (mean ± std. dev. of 7 runs, 100000 loops each)

%timeit bisect(f, 0, 1)

61.1 µs ± 1.51 µs per loop (mean ± std. dev. of 7 runs, 10000 loops each)
```

Multivariate Root-Finding

Use scipy.optimize.fsolve, a wrapper for a hybrid method in MINPACK.

See the <u>documentation</u> for details.

A **fixed point** of a real function f on [a, b] is an $x \in [a, b]$ such that f(x) = x.

SciPy has a function for finding (scalar) fixed points too

```
from scipy.optimize import fixed_point
fixed_point(lambda x: x**2, 10.0) # 10.0 is an initial guess
array(1.)
```

If you don't get good results, you can always switch back to the brentq root finder, since the fixed point of a function f is the root of g(x) := x - f(x).

Optimization

Most numerical packages provide only functions for minimization.

Maximization can be performed by recalling that the maximizer of a function f on domain D is the minimizer of -f on D.

Minimization is closely related to root-finding: For smooth functions, interior optima correspond to roots of the first derivative.

The speed/robustness trade-off described above is present with numerical optimization too.

Unless you have some prior information you can exploit, it's usually best to use hybrid methods.

For constrained, univariate (i.e., scalar) minimization, a good hybrid option is fminbound

```
from scipy.optimize import fminbound
fminbound(lambda x: x**2, -1, 2) # Search in [-1, 2]
0.0
```

Multivariate Optimization

 $\label{lem:multivariate} \textbf{Multivariate local optimizers include minimize, fmin, fmin_powell, fmin_cg, fmin_bfgs, and fmin_ncg.}$

Constrained multivariate local optimizers include fmin_l_bfgs_b, fmin_tnc, fmin_cobyla.

See the documentation for details.

Integration

Most numerical integration methods work by computing the integral of an approximating polynomial.

The resulting error depends on how well the polynomial fits the integrand, which in turn depends on how "regular" the integrand is.

In SciPy, the relevant module for numerical integration is scipy. integrate.

A good default for univariate integration is quad

```
from scipy.integrate import quad
integral, error = quad(lambda x: x**2, 0, 1)
integral
```

```
0.3333333333333333
```

In fact, quad is an interface to a very standard numerical integration routine in the Fortran library QUADPACK.

It uses <u>Clenshaw-Curtis quadrature</u>, based on expansion in terms of Chebychev polynomials.

There are other options for univariate integration—a useful one is fixed_quad, which is fast and hence works well inside for loops.

There are also functions for multivariate integration.

See the documentation for more details.

Linear Algebra

We saw that NumPy provides a module for linear algebra called linalg.

SciPy also provides a module for linear algebra with the same name.

The latter is not an exact superset of the former, but overall it has more functionality.

We leave you to investigate the set of available routines.

Exercises

Exercise 1

Previously we discussed the concept of recursive function calls.

Try to write a recursive implementation of homemade bisection function described above.

Test it on the function (7).

Solutions

Exercise 1

Here's a reasonable solution:

We can test it as follows

```
f = lambda x: np.sin(4 * (x - 0.25)) + x + x**20 - 1
bisect(f, 0, 1)
Current mid point = 0.5
Current mid point = 0.25
Current mid point = 0.375
Current mid point = 0.4375
Current mid point = 0.40625
Current mid point = 0.421875
Current mid point = 0.4140625
Current mid point = 0.41015625
Current mid point = 0.408203125
Current mid point = 0.4091796875
Current mid point = 0.40869140625
Current mid point = 0.408447265625
Current mid point = 0.4083251953125
Current mid point = 0.40826416015625
```

Numba

In addition to what's in Anaconda, this lecture will need the following libraries:

```
!pip install --upgrade quantecon
```

Please also make sure that you have the latest version of Anaconda, since old versions are a <u>common source of errors</u>.

Let's start with some imports:

```
import numpy as np
import quantecon as qe
import matplotlib.pyplot as plt
%matplotlib inline
```

Overview

In an <u>earlier lecture</u> we learned about vectorization, which is one method to improve speed and efficiency in numerical work.

Vectorization involves sending array processing operations in batch to efficient low-level code.

However, as <u>discussed previously</u>, vectorization has several weaknesses.

One is that it is highly memory-intensive when working with large amounts of data.

Another is that the set of algorithms that can be entirely vectorized is not universal.

In fact, for some algorithms, vectorization is ineffective.

Fortunately, a new Python library called Numba solves many of these problems.

It does so through something called $\it just$ in time (JIT) compilation.

The key idea is to compile functions to native machine code instructions on the fly.

When it succeeds, the compiled code is extremely fast.

Numba is specifically designed for numerical work and can also do other tricks such as multithreading.

Numba will be a key part of our lectures — especially those lectures involving dynamic programming.

This lecture introduces the main ideas.

Compiling Functions

As stated above, Numba's primary use is compiling functions to fast native machine code during runtime.

An Example

Let's consider a problem that is difficult to vectorize: generating the trajectory of a difference equation given an initial condition.

We will take the difference equation to be the quadratic map

$$x_{t+1} = \alpha x_t (1 - x_t)$$

In what follows we set

Here's the plot of a typical trajectory, starting from $x_0 = 0.1$, with t on the x-axis

```
def qm(x0, n):
    x = np.empty(n+1)
    x[0] = x0
    for t in range(n):
        x[t+1] = α * x[t] * (1 - x[t])
    return x

x = qm(0.1, 250)
fig, ax = plt.subplots()
ax.plot(x, 'b-', lw-2, alpha=0.8)
ax.set_xlabel('$t$', fontsize=12)
ax.set_ylabel('$t$', fontsize = 12)
plt.show()
```

Notice the "chaotic" nature of the trajectory — an excellent discussion of these kinds of systems can be found in [$\underline{LM13}$].

To speed the function qm up using Numba, our first step is

```
from numba import jit
qm_numba = jit(qm)
```

The function qm_numba is a version of qm that is "targeted" for JIT-compilation.

We will explain what this means momentarily.

Let's time and compare identical function calls across these two versions, starting with the original function qm:

```
n = 10_000_000

qe.tic()
qm(0.1, int(n))
time1 = qe.toc()
```

TOC: Elapsed: 0:00:7.39

Now let's try qm_numba

```
qe.tic()
qm_numba(0.1, int(n))
time2 = qe.toc()
```

TOC: Elapsed: 0:00:0.09

This is already a massive speed gain.

In fact, the next time and all subsequent times it runs even faster as the function has been compiled and is in memory:

```
qe.tic()
qm_numba(0.1, int(n))
time3 = qe.toc()
```

TOC: Elapsed: 0:00:0.02

```
time1 / time3 # Calculate speed gain
```

286.82018659097

This kind of speed gain is huge relative to how simple and clear the implementation is.

How and When it Works

Numba attempts to generate fast machine code using the infrastructure provided by the LLVM Project.

It does this by inferring type information on the fly.

(See our earlier lecture on scientific computing for a discussion of types.)

The basic idea is this:

- Python is very flexible and hence we could call the function [qm]{.title-ref} with many types.
 - o e.g., x0 could be a NumPy array or a list, n could be an integer or a float, etc.
- This makes it hard to *pre*-compile the function.
- However, when we do actually call the function, say by executing qm(0.5, 10), the types of x0 and n become clear.
- Moreover, the types of other variables in qm can be inferred once the input is known.
- So the strategy of Numba and other JIT compilers is to wait until this moment, and then
 compile the function.

That's why it is called "just-in-time" compilation.

Note that, if you make the call qm(0.5, 10) and then follow it with qm(0.9, 20), compilation only takes place on the first call.

The compiled code is then cached and recycled as required.

Decorators and "nopython" Mode

In the code above we created a JIT compiled version of qm via the call

```
qm_numba = jit(qm)
```

In practice this would typically be done using an alternative *decorator* syntax.

(We will explain all about decorators in a later lecture but you can skip the details at this stage.)

Let's see how this is done.

Decorator Notation

To target a function for JIT compilation we can put @jit before the function definition.

Here's what this looks like for qm

```
@jit
def qm(x0, n):
    x = np.empty(n+1)
    x[0] = x0
    for t in range(n):
        x[t+1] = α * x[t] * (1 - x[t])
    return x
```

This is equivalent to qm = jit(qm).

The following now uses the jitted version:

Type Inference and "nopython" Mode

Clearly type inference is a key part of JIT compilation.

As you can imagine, inferring types is easier for simple Python objects (e.g., simple scalar data types such as floats and integers).

Numba also plays well with NumPy arrays.

In an ideal setting, Numba can infer all necessary type information.

This allows it to generate native machine code, without having to call the Python runtime environment.

In such a setting, Numba will be on par with machine code from low-level languages.

When Numba cannot infer all type information, some Python objects are given generic object status and execution falls back to the Python runtime.

When this happens, Numba provides only minor speed gains or none at all.

We generally prefer to force an error when this occurs, so we know effective compilation is failing.

This is done by using either @jit(nopython=True) or, equivalently, @njit instead of @jit.

For example,

```
from numba import njit

@njit
def qm(x0, n):
    x = np.empty(n+1)
    x[0] = x0
    for t in range(n):
        x[t+1] = 4 * x[t] * (1 - x[t])
    return x
```

Compiling Classes

As mentioned above, at present Numba can only compile a subset of Python.

However, that subset is ever expanding.

For example, Numba is now quite effective at compiling classes.

If a class is successfully compiled, then its methods act as JIT-compiled functions.

To give one example, let's consider the class for analyzing the Solow growth model we created in <u>this lecture</u>.

To compile this class we use the @jitclass decorator:

```
from numba import jitclass, float64
```

Notice that we also imported something called float64.

This is a data type representing standard floating point numbers.

We are importing it here because Numba needs a bit of extra help with types when it trys to deal with classes.

Here's our code:

```
solow_data = [
    ('n', float64),
    ('s', float64),
    ('ô', float64),
    ('z', float64),
    ('z', float64),
    ('k', float64)
       ('k', float64)
@jitclass(solow_data)
class Solow:
       Implements the Solow growth model with the update rule
              k_{t+1} = [(s \ z \ k^{\alpha}_{t}) + (1 - \delta)k_{t}] / (1 + n)
       .....
       def __init__(self, n=0.05, # population growth rate
                                       s=0.25, # savings rate
                                                      # depreciation rate
                                        δ=0.1,
                                       α=0.3,
                                                      # share of labor
                                        z=2.0
                                                       # productivity
                                        k=1.0): # current capital stock
             self.n, self.s, self.0, self.\alpha, self.z = n, s, \delta, \alpha, z self.k = k
       def h(self):
               "Evaluate the h function"
             # Unpack parameters (get rid of self to simplify notation) n, s, \delta, \alpha, z = self.n, self.s, self.\delta, self.\alpha, self.z # Apply the update rule return (s * z * self.k**\alpha + (1 - \delta) * self.k) / (1 + n)
       def update(self):
              "Update the current state (i.e., the capital stock)." self.k = self.h()
       def steady_state(self):
             "Compute the steady state value of capital."

# Unpack parameters (get rid of self to simplify notation)

n, s, \delta, \alpha, z = self.n, self.s, self.\delta, self.\alpha, self.z

# Compute and return steady state

return ((s * z) / (n + \delta))**(1 / (1 - \alpha))
       def generate_sequence(self, t):
              "Generate and return a time series of length t" path = []
              for i in range(t):
                    path.append(self.k)
                     self_update()
              return path
```

First we specified the types of the instance data for the class in solow_data.

After that, targeting the class for JIT compilation only requires adding @jitclass(solow_data) before the class definition.

When we call the methods in the class, the methods are compiled just like functions.

```
s1 = Solow()
s2 = Solow(k=8.0)

T = 60
fig, ax = plt.subplots()

# Plot the common steady state value of capital
ax.plot([s1.steady_state()]*T, 'k-', label='steady state')

# Plot time series for each economy
for s in s1, s2:
    lb = f'capital series from initial state {s.k}'
    ax.plot(s.generate_sequence(T), 'o-', lw=2, alpha=0.6, label=lb)
ax.set_ylabel('$k_{f}}*, fontsize=12)
ax.set_xlabel('$t$*', fontsize=12)
ax.legend()
plt.show()
```

Alternatives to Numba

There are additional options for accelerating Python loops.

Here we quickly review them.

However, we do so only for interest and completeness.

If you prefer, you can safely skip this section.

Cython

Like <u>Numba</u>, <u>Cython</u> provides an approach to generating fast compiled code that can be used from Python.

As was the case with Numba, a key problem is the fact that Python is dynamically typed.

As you'll recall, Numba solves this problem (where possible) by inferring type.

 $\label{eq:cython} \mbox{Cython's approach is different-programmers add type definitions directly to their "Python" code. \\$

As such, the Cython language can be thought of as Python with type definitions.

In addition to a language specification, Cython is also a language translator, transforming Cython code into optimized C and C++ code.

Cython also takes care of building language extensions — the wrapper code that interfaces between the resulting compiled code and Python.

While Cython has certain advantages, we generally find it both slower and more cumbersome than Numba.

Interfacing with Fortran via F2Py

If you are comfortable writing Fortran you will find it very easy to create extension modules from Fortran code using $\underline{F2Py}$.

F2Py is a Fortran-to-Python interface generator that is particularly simple to use.

Robert Johansson provides a <u>nice introduction</u> to F2Py, among other things.

Recently, <u>a Jupyter cell magic for Fortran</u> has been developed — you might want to give it a try.

Summary and Comments

Let's review the above and add some cautionary notes.

Limitations

As we've seen, Numba needs to infer type information on all variables to generate fast machine-level instructions.

For simple routines, Numba infers types very well.

For larger ones, or for routines using external libraries, it can easily fail.

Hence, it's prudent when using Numba to focus on speeding up small, time-critical snippets of code.

This will give you much better performance than blanketing your Python programs with @jit statements.

A Gotcha: Global Variables

Here's another thing to be careful about when using Numba.

Consider the following example

```
a = 1
@jit
def add_a(x):
    return a + x
print(add_a(10))
```

11

```
a = 2
print(add_a(10))
```

11

Notice that changing the global had no effect on the value returned by the function.

When Numba compiles machine code for functions, it treats global variables as constants to ensure type stability.

Exercises

Exercise 1

<u>Previously</u> we considered how to approximate π by Monte Carlo.

Use the same idea here, but make the code efficient using Numba.

Compare speed with and without Numba when the sample size is large.

Exercise 2

In the <u>Introduction to Quantitative Economics with Python</u> lecture series you can learn all about finite-state Markov chains.

For now, let's just concentrate on simulating a very simple example of such a chain.

Suppose that the volatility of returns on an asset can be in one of two regimes - high or low.

The transition probabilities across states are as follows

For example, let the period length be one day, and suppose the current state is high.

We see from the graph that the state tomorrow will be

- high with probability 0.8
- low with probability 0.2

Your task is to simulate a sequence of daily volatility states according to this rule.

Set the length of the sequence to n $\,=\,$ 1_000_000 and start in the high state.

Implement a pure Python version and a Numba version, and compare speeds.

To test your code, evaluate the fraction of time that the chain spends in the low state.

If your code is correct, it should be about 2/3.

Hints:

- Represent the low state as 0 and the high state as 1.
- If you want to store integers in a NumPy array and then apply JIT compilation, use x = np.empty(n, dtype=np.int_).

Exercise 1

Here is one solution:

```
from random import uniform
@njit
def calculate_pi(n=1_000_000):
    count = 0
    for i in range(n):
        u, v = uniform(0, 1), uniform(0, 1)
        d = np.sqrt((u - 0.5)**2 + (v - 0.5)**2)
        if d < 0.5:
            count += 1

    area_estimate = count / n
    return area_estimate * 4 # dividing by radius**2</pre>
```

Now let's see how fast it runs:

```
%time calculate_pi()

CPU times: user 136 ms, sys: 1.34 ms, total: 137 ms
Wall time: 138 ms

3.142872

%time calculate_pi()

CPU times: user 12.5 ms, sys: 61 µs, total: 12.6 ms
Wall time: 12.6 ms

3.144484
```

If we switch of JIT compilation by removing @njit, the code takes around 150 times as long on our machine.

So we get a speed gain of 2 orders of magnitude–which is huge–by adding four characters.

Exercise 2

We let

- 0 represent "low"
- 1 represent "high"

```
p, q = 0.1, 0.2 # Prob of leaving low and high state respectively
```

Here's a pure Python version of the function

```
def compute_series(n):
    x = np.empty(n, dtype=np.int_)
    x[0] = 1  # Start in state 1
    U = np.random.uniform(0, 1, size=n)
    for t in range(1, n):
        current_x = x[t-1]
        if current_x == 0:
              x[t] = U[t]  q
    return x
```

Let's run this code and check that the fraction of time spent in the low state is about 0.666

```
n = 1_000_000
x = compute_series(n)
print(np.mean(x == 0)) # Fraction of time x is in state 0
```

0.66609

This is (approximately) the right output.

Now let's time it:

```
qe.tic()
compute_series(n)
qe.toc()
```

TOC: Elapsed: 0:00:0.70

0.7064418792724609

Next let's implement a Numba version, which is easy

```
from numba import jit
compute_series_numba = jit(compute_series)
```

Let's check we still get the right numbers

```
x = compute_series_numba(n)
print(np.mean(x == 0))
```

0.666374

Let's see the time

```
qe.tic()
compute_series_numba(n)
qe.toc()
```

TOC: Elapsed: 0:00:0.00

0.008621931076049805

This is a nice speed improvement for one line of code!

LM13

Andrzej Lasota and Michael C Mackey. *Chaos, fractals, and noise: stochastic aspects of dynamics.* Volume 97. Springer Science & Business Media, 2013.

Parallelization

In addition to what's in Anaconda, this lecture will need the following libraries:

```
!pip install --upgrade quantecon
```

Overview

The growth of CPU clock speed (i.e., the speed at which a single chain of logic can be run) has slowed dramatically in recent years.

This is unlikely to change in the near future, due to inherent physical limitations on the construction of chips and circuit boards.

Chip designers and computer programmers have responded to the slowdown by seeking a different path to fast execution: parallelization.

Hardware makers have increased the number of cores (physical CPUs) embedded in each machine.

For programmers, the challenge has been to exploit these multiple CPUs by running many processes in parallel (i.e., simultaneously).

This is particularly important in scientific programming, which requires handling

- large amounts of data and
- CPU intensive simulations and other calculations.

In this lecture we discuss parallelization for scientific computing, with a focus on

- 1. the best tools for parallelization in Python and
- 2. how these tools can be applied to quantitative economic problems.

Let's start with some imports:

```
import numpy as np
import quantecon as qe
import matplotlib.pyplot as plt
%matplotlib inline
```

Types of Parallelization

Large textbooks have been written on different approaches to parallelization but we will keep a tight focus on what's most useful to us.

We will briefly review the two main kinds of parallelization commonly used in scientific computing and discuss their pros and cons.

Multiprocessing

 $\label{thm:multiple} \mbox{Multiprocessing means concurrent execution of multiple processes using more than one processor.}$

In this context, a process is a chain of instructions (i.e., a program).

Multiprocessing can be carried out on one machine with multiple CPUs or on a collection of machines connected by a network.

In the latter case, the collection of machines is usually called a **cluster**.

With multiprocessing, each process has its own memory space, although the physical memory chip might be shared.

Multithreading

Multithreading is similar to multiprocessing, except that, during execution, the threads all share the same memory space.

Native Python struggles to implement multithreading due to some <u>legacy design features</u>.

But this is not a restriction for scientific libraries like NumPy and Numba.

Functions imported from these libraries and JIT-compiled code run in low level execution environments where Python's legacy restrictions don't apply.

Advantages and Disadvantages

Multithreading is more lightweight because most system and memory resources are shared by the threads.

In addition, the fact that multiple threads all access a shared pool of memory is extremely convenient for numerical programming.

On the other hand, multiprocessing is more flexible and can be distributed across clusters.

For the great majority of what we do in these lectures, multithreading will suffice.

Implicit Multithreading in NumPy

Actually, you have already been using multithreading in your Python code, although you might not have realized it.

(We are, as usual, assuming that you are running the latest version of Anaconda Python.)

This is because NumPy cleverly implements multithreading in a lot of its compiled code.

Let's look at some examples to see this in action.

A Matrix Operation

The next piece of code computes the eigenvalues of a large number of randomly generated matrices.

It takes a few seconds to run.

```
n = 20
m = 1000
for i in range(n):
    X = np.random.randn(m, m)
    λ = np.linalg.eigvals(X)
```

Now, let's look at the output of the htop system monitor on our machine while this code is running:

We can see that 4 of the 8 CPUs are running at full speed.

This is because NumPy's eigvals routine neatly splits up the tasks and distributes them to different threads.

A Multithreaded Ufunc

Over the last few years, NumPy has managed to push this kind of multithreading out to more and more operations.

For example, let's return to a maximization problem discussed previously:

```
def f(x, y):
    return np.cos(x**2 + y**2) / (1 + x**2 + y**2)

grid = np.linspace(-3, 3, 5000)
x, y = np.meshgrid(grid, grid)

%timeit np.max(f(x, y))
```

```
842 ms \pm 8.59 ms per loop (mean \pm std. dev. of 7 runs, 1 loop each)
```

If you have a system monitor such as http (Linux/Mac) or perfmon (Windows), then try running this and then observing the load on your CPUs.

(You will probably need to bump up the grid size to see large effects.)

At least on our machine, the output shows that the operation is successfully distributed across multiple threads.

This is one of the reasons why the vectorized code above is fast.

A Comparison with Numba

To get some basis for comparison for the last example, let's try the same thing with Numba.

In fact there is an easy way to do this, since Numba can also be used to create custom $\underline{\text{ufuncs}}$ with the $\underline{\text{@vectorize}}$ decorator.

```
from numba import vectorize
@vectorize
def f_vec(x, y):
    return np.cos(x**2 + y**2) / (1 + x**2 + y**2)
np.max(f_vec(x, y)) # Run once to compile
```

0.9999992797121728

```
%timeit np.max(f_vec(x, y))
```

```
327 ms \pm 5.66 ms per loop (mean \pm std. dev. of 7 runs, 1 loop each)
```

At least on our machine, the difference in the speed between the Numba version and the vectorized NumPy version shown above is not large.

But there's quite a bit going on here so let's try to break down what is happening.

Both Numba and NumPy use efficient machine code that's specialized to these floating point operations.

However, the code NumPy uses is, in some ways, less efficient.

The reason is that, in NumPy, the operation np.cos(x**2 + y**2) / (1 + x**2 + y**2) generates several intermediate arrays.

For example, a new array is created when x**2 is calculated.

The same is true when y**2 is calculated, and then x**2 + y**2 and so on.

Numba avoids creating all these intermediate arrays by compiling one function that is specialized to the entire operation.

But if this is true, then why isn't the Numba code faster?

The reason is that NumPy makes up for its disadvantages with implicit multithreading, as we've just discussed.

Multithreading a Numba Ufunc

Can we get both of these advantages at once?

In other words, can we pair

- the efficiency of Numba's highly specialized JIT compiled function and
- the speed gains from parallelization obtained by NumPy's implicit multithreading?

It turns out that we can, by adding some type information plus target='parallel'.

```
@vectorize('float64(float64, float64)', target='parallel')
def f_vec(x, y):
    return np.cos(x**2 + y**2) / (1 + x**2 + y**2)
np.max(f_vec(x, y)) # Run once to compile
```

0.9999992797121728

```
%timeit np.max(f_vec(x, y))
```

```
114 ms \pm 2.25 ms per loop (mean \pm std. dev. of 7 runs, 10 loops each)
```

Now our code runs significantly faster than the NumPy version.

Multithreaded Loops in Numba

We just saw one approach to parallelization in Numba, using the parallel flag in @vectorize.

This is neat but, it turns out, not well suited to many problems we consider.

Fortunately, Numba provides another approach to multithreading that will work for us almost everywhere parallelization is possible.

To illustrate, let's look first at a simple, single-threaded (i.e., non-parallelized) piece of code.

The code simulates updating the wealth $\boldsymbol{w_t}$ of a household via the rule

$$w_{t+1} = R_{t+1} s w_t + y_{t+1}$$

- R is the gross rate of return on assets
- s is the savings rate of the household and
- y is labor income.

We model both R and y as independent draws from a lognormal distribution.

Here's the code:

Let's have a look at how wealth evolves under this rule.

```
fig, ax = plt.subplots()
T = 100
w = np.empty(T)
w[0] = 5
for t in range(T-1):
    w[t+1] = h(w[t])
ax.plot(w)
ax.set_xlabel('$t$', fontsize=12)
ax.set_ylabel('$w_{t}$', fontsize=12)
plt.show()

.../_images/parallelization_10_0.png
```

Now let's suppose that we have a large population of households and we want to know what median wealth will be.

This is not easy to solve with pencil and paper, so we will use simulation instead.

In particular, we will simulate a large number of households and then calculate median wealth for this group.

Suppose we are interested in the long-run average of this median over time.

It turns out that, for the specification that we've chosen above, we can calculate this by taking a one-period snapshot of what has happened to median wealth of the group at the end of a long simulation.

 $Moreover, provided \ the \ simulation \ period \ is \ long \ enough, initial \ conditions \ don't \ matter.$

 $\bullet~$ This is due to something called ergodicity, which we will discuss $\underline{\mathsf{later}\ \mathsf{on}}.$

So, in summary, we are going to simulate 50,000 households by

- 1. arbitrarily setting initial wealth to 1 and
- 2. simulating forward in time for 1,000 periods.

Then we'll calculate median wealth at the end period.

Here's the code:

```
@njit
def compute_long_run_median(w0=1, T=1000, num_reps=50_000):
    obs = np.empty(num_reps)
    for i in range(num_reps):
        w = w0
        for t in range(T):
              w = h(w)
        obs[i] = w

return np.median(obs)
```

Let's see how fast this runs:

```
%%time compute_long_run_median()

CPU times: user 4.85 s, sys: 10.7 ms, total: 4.86 s
Wall time: 4.86 s
```

To speed this up, we're going to parallelize it via multithreading.

To do so, we add the parallel=True flag and change range to prange:

```
from numba import prange
@njit(parallel=True)
def compute_long_run_median_parallel(w0=1, T=1000, num_reps=50_000):

   obs = np.empty(num_reps)
   for i in prange(num_reps):
        w = w0
        for t in range(T):
            w = h(w)
        obs[i] = w

return np.median(obs)
```

Let's look at the timing:

1.8380439509863367

```
%%time compute_long_run_median_parallel()

CPU times: user 6.9 s, sys: 9.74 ms, total: 6.91 s Wall time: 1.27 s

1.842278795929991
```

The speed-up is significant.

A Warning

Parallelization works well in the outer loop of the last example because the individual tasks inside the loop are independent of each other.

If this independence fails then parallelization is often problematic.

For example, each step inside the inner loop depends on the last step, so independence fails, and this is why we use ordinary range instead of prange.

When you see us using prange in later lectures, it is because the independence of tasks holds true.

When you see us using ordinary range in a jitted function, it is either because the speed gain from parallelization is small or because independence fails.

Exercises

Exercise 1

In <u>an earlier exercise</u>, we used Numba to accelerate an effort to compute the constant π by Monte Carlo.

Now try adding parallelization and see if you get further speed gains. \\

You should not expect huge gains here because, while there are many independent tasks (draw point and test if in circle), each one has low execution time.

Generally speaking, parallelization is less effective when the individual tasks to be parallelized are very small relative to total execution time.

This is due to overheads associated with spreading all of these small tasks across multiple CPUs.

Nevertheless, with suitable hardware, it is possible to get nontrivial speed gains in this evercise

For the size of the Monte Carlo simulation, use something substantial, such as $n=100_000_000$.

Solutions

Exercise 1

Here is one solution:

```
from random import uniform
@njit(parallel=True)
def calculate_pi(n=1_000_000):
    count = 0
    for i in prange(n):
        u, v = uniform(0, 1), uniform(0, 1)
        d = np.sqrt((u - 0.5)**2 + (v - 0.5)**2)
        if d < 0.5:
            count += 1

    area_estimate = count / n
    return area_estimate * 4 # dividing by radius**2</pre>
```

Now let's see how fast it runs:

```
%time calculate_pi()

CPU times: user 367 ms, sys: 1.71 ms, total: 369 ms
Wall time: 344 ms

3.139692

%time calculate_pi()

CPU times: user 26 ms, sys: 152 µs, total: 26.2 ms
Wall time: 3.39 ms

3.143128
```

By switching parallelization on and off (selecting True or False in the @njit annotation), we can test the speed gain that multithreading provides on top of JIT compilation.

On our workstation, we find that parallelization increases execution speed by a factor of 2 or 3.

(If you are executing locally, you will get different numbers, depending mainly on the number of CPUs on your machine.)

Pandas

In addition to what's in Anaconda, this lecture will need the following libraries:

```
!pip install ——upgrade pandas—datareader
```

Overview

Pandas is a package of fast, efficient data analysis tools for Python.

Its popularity has surged in recent years, coincident with the rise of fields such as data science and machine learning.

Here's a popularity comparison over time against STATA, SAS, and \underline{dplyr} courtesy of Stack Overflow Trends

Just as NumPy provides the basic array data type plus core array operations, pandas

- 1. defines fundamental structures for working with data and
- 2. endows them with methods that facilitate operations such as
- reading in data
- · adjusting indices
- · working with dates and time series
- sorting, grouping, re-ordering and general data munging2
- dealing with missing values, etc., etc.

More sophisticated statistical functionality is left to other packages, such as <u>statsmodels</u> and <u>scikit-learn</u>, which are built on top of pandas.

This lecture will provide a basic introduction to pandas.

Throughout the lecture, we will assume that the following imports have taken place

```
import pandas as pd
import numpy as np
import matplotlib.pyplot as plt
%matplotlib inline
import requests
```

Series

Two important data types defined by pandas are Series and DataFrame.

You can think of a Series as a "column" of data, such as a collection of observations on a single variable.

A DataFrame is an object for storing related columns of data.

Let's start with Series

```
s = pd.Series(np.random.randn(4), name='daily returns')

0 -1.321149
1 -0.547052
2 2.247000
3 -0.244340
Name: daily returns, dtype: float64
```

Here you can imagine the indices 0, 1, 2, 3 as indexing four listed companies, and the values being daily returns on their shares.

Pandas Series are built on top of NumPy arrays and support many similar operations

```
s * 100

0 -132.114919

1 -54.705207

2 224.699963

3 -24.433987

Name: daily returns, dtype: float64

np.abs(s)

0 1.321149

1 0.547052

2 2.247000

3 0.244340

Name: daily returns, dtype: float64
```

Not only do they have some additional (statistically oriented) methods

```
s.describe()
          4.000000
count
mean
          0.033615
std
          1.543685
min
         -1.321149
        -0.740576
-0.395696
25%
50%
          0.378495
75%
          2.247000
max
Name: daily returns, dtype: float64
```

But their indices are more flexible

```
s.index = ['AMZN', 'AAPL', 'MSFT', 'G00G']
s

AMZN    -1.321149
AAPL    -0.547052
MSFT     2.247000
G00G     -0.244340
Name: daily returns, dtype: float64
```

Viewed in this way, Series are like fast, efficient Python dictionaries (with the restriction that the items in the dictionary all have the same type—in this case, floats).

In fact, you can use much of the same syntax as Python dictionaries

```
s['AMZN']

-1.3211491865776104

s['AMZN'] = 0
s

AMZN      0.000000
AAPL      -0.547052
MSFT      2.247000
GOOG      -0.244340
Name: daily returns, dtype: float64

'AAPL' in s
```

DataFrames

True

While a Series is a single column of data, a DataFrame is several columns, one for each variable.

In essence, a DataFrame in pandas is analogous to a (highly optimized) Excel spreadsheet.

Thus, it is a powerful tool for representing and analyzing data that are naturally organized into rows and columns, often with descriptive indexes for individual rows and individual columns.

Let's look at an example that reads data from the CSV file pandas/data/test_pwt.csv that can be downloaded $\underline{\text{here}}$.

Here's the content of test_pwt.csv

```
"country", "country isocode", "year", "POP", "XRAT", "tcgdp", "cc", "cg"
"Argentina", "ARG", "2000", "37335.653", "0.9995", "295072.21869", "75.716805379"
,"5.5788042896"
"Australia", "AUS", "2000", "19053.186", "1.72483", "541804.6521", "67.759025993"
,"6.720975332"
"India", "IND", "2000", "1006300.297", "44.9416", "1728144.3748", "64.575551328", "14.072205773"
"Israel", "ISR", "2000", "6114.57", "4.07733", "129253.89423", "64.436450847", "10.266688415"
"Malawi", "MWI", "2000", "11801.505", "59.543808333", "5026.2217836", "74.707624181", "11.658954494"
"South Africa", "ZAF", "2000", "45064.098", "6.93983", "227242.36949", "72.718710427", "5.7265463933"
"United States", "USA", "2000", "282171.957", "1", "9898700", "72.347054303", "6.0324539789"
"Uruguay", "URY", "2000", "3219.793", "12.099591667", "25255.961693", "78.978740282", "5.108067988"
```

Supposing you have this data saved as test_pwt.csv in the present working directory (type %pwd in Jupyter to see what this is), it can be read in as follows:

```
df = pd.read_csv('https://raw.githubusercontent.com/QuantEcon/lecture-
source-py/master/source/_static/lecture_specific/pandas/data/test_pwt.csv')
type(df)
```

pandas.core.frame.DataFrame

C	f							
	country	country isocode	year	POP	XRAT	tcgdp	сс	cg
0	Argentina	ARG	2000	37335.653	0.999500	2.950722e+05	75.716805	5.578804
1	Australia	AUS	2000	19053.186	1.724830	5.418047e+05	67.759026	6.720098
2	India	IND	2000	1006300.297	44.941600	1.728144e+06	64.575551	14.072206
3	Israel	ISR	2000	6114.570	4.077330	1.292539e+05	64.436451	10.266688
4	Malawi	MWI	2000	11801.505	59.543808	5.026222e+03	74.707624	11.658954
5	South Africa	ZAF	2000	45064.098	6.939830	2.272424e+05	72.718710	5.726546
6	United States	USA	2000	282171.957	1.000000	9.898700e+06	72.347054	6.032454
7	Uruguay	URY	2000	3219.793	12.099592	2.525596e+04	78.978740	5.108068

We can select particular rows using standard Python array slicing notation

C	df[2:5]							
	country	country isocode	year	POP	XRAT	tcgdp	сс	cg
2	India	IND	2000	1006300.297	44.941600	1.728144e+06	64.575551	14.072206
3	Israel	ISR	2000	6114.570	4.077330	1.292539e+05	64.436451	10.266688
4	Malawi	MWI	2000	11801.505	59.543808	5.026222e+03	74.707624	11.658954

To select columns, we can pass a list containing the names of the desired columns represented as strings

df[['country', 'tcgdp']]

	country	tcgdp
0	Argentina	2.950722e+05
1	Australia	5.418047e+05
2	India	1.728144e+06
3	Israel	1.292539e+05
4	Malawi	5.026222e+03
5	South Africa	2.272424e+05
6	United States	9.898700e+06
7	Uruguay	2.525596e+04

To select both rows and columns using integers, the iloc attribute should be used with the format.iloc[rows, columns]

df.iloc[2:5, 0:4]

	country	country isocode	year	POP
2	India	IND	2000	1006300.297
3	Israel	ISR	2000	6114.570
4	Malawi	MWI	2000	11801.505

To select rows and columns using a mixture of integers and labels, the loc attribute can be used in a similar way

df.loc[df.index[2:5], ['country', 'tcgdp']]

	country	tcgdp
2	India	1.728144e+06
3	Israel	1.292539e+05
4	Malawi	5.026222e+03

Let's imagine that we're only interested in population (POP) and total GDP (tcgdp).

One way to strip the data frame df down to only these variables is to overwrite the dataframe using the selection method described above

u	•		
	country	POP	tcgdp
0	Argentina	37335.653	2.950722e+05
1	Australia	19053.186	5.418047e+05
2	India	1006300.297	1.728144e+06
3	Israel	6114.570	1.292539e+05
4	Malawi	11801.505	5.026222e+03
5	South Africa	45064.098	2.272424e+05
6	United States	282171.957	9.898700e+06
7	Uruguay	3219.793	2.525596e+04

Here the index 0, 1, \dots , 7 is redundant because we can use the country names as an index.

To do this, we set the index to be the country variable in the dataframe

df = df.set_index('country')

a i		
	POP	tcgdp
country		
Argentina	37335.653	2.950722e+05
Australia	19053.186	5.418047e+05
India	1006300.297	1.728144e+06
Israel	6114.570	1.292539e+05
Malawi	11801.505	5.026222e+03
South Africa	45064.098	2.272424e+05
United States	282171.957	9.898700e+06
Uruguay	3219.793	2.525596e+04

Let's give the columns slightly better names

```
df.columns = 'population', 'total GDP'
```

	population	total GDP
country		
Argentina	37335.653	2.950722e+05
Australia	19053.186	5.418047e+05
India	1006300.297	1.728144e+06
Israel	6114.570	1.292539e+05
Malawi	11801.505	5.026222e+03
South Africa	45064.098	2.272424e+05
United States	282171.957	9.898700e+06
Uruguay	3219.793	2.525596e+04

Population is in thousands, let's revert to single units

```
df['population'] = df['population'] * 1e3
df
```

	population	total GDP
country		
Argentina	3.733565e+07	2.950722e+05
Australia	1.905319e+07	5.418047e+05
India	1.006300e+09	1.728144e+06
Israel	6.114570e+06	1.292539e+05
Malawi	1.180150e+07	5.026222e+03
South Africa	4.506410e+07	2.272424e+05
United States	2.821720e+08	9.898700e+06
Uruguay	3.219793e+06	2.525596e+04

Next, we're going to add a column showing real GDP per capita, multiplying by 1,000,000 as we go because total GDP is in millions

```
df['GDP percap'] = df['total GDP'] * 1e6 / df['population']
df
```

	population	total GDP	GDP percap
country			
Argentina	3.733565e+07	2.950722e+05	7903.229085
Australia	1.905319e+07	5.418047e+05	28436.433261
India	1.006300e+09	1.728144e+06	1717.324719
Israel	6.114570e+06	1.292539e+05	21138.672749
Malawi	1.180150e+07	5.026222e+03	425.896679
South Africa	4.506410e+07	2.272424e+05	5042.647686
United States	2.821720e+08	9.898700e+06	35080.381854
Uruguay	3.219793e+06	2.525596e+04	7843.970620

One of the nice things about pandas DataFrame and Series objects is that they have methods for plotting and visualization that work through Matplotlib.

For example, we can easily generate a bar plot of GDP per capita $\,$

```
ax = df['GDP percap'].plot(kind='bar')
ax.set_xlabel('country', fontsize=12)
ax.set_ylabel('GDP per capita', fontsize=12)
plt.show()

..../_images/pandas_21_0.png
```

At the moment the data frame is ordered alphabetically on the countries—let's change it to GDP per capita

```
df = df.sort_values(by='GDP percap', ascending=False)
df
```

	population	total GDP	GDP percap
country			
United States	2.821720e+08	9.898700e+06	35080.381854
Australia	1.905319e+07	5.418047e+05	28436.433261
Israel	6.114570e+06	1.292539e+05	21138.672749
Argentina	3.733565e+07	2.950722e+05	7903.229085
Uruguay	3.219793e+06	2.525596e+04	7843.970620
South Africa	4.506410e+07	2.272424e+05	5042.647686
India	1.006300e+09	1.728144e+06	1717.324719
Malawi	1.180150e+07	5.026222e+03	425.896679

Plotting as before now yields

```
ax = df['GDP percap'].plot(kind='bar')
ax.set_xlabel('country', fontsize=12)
ax.set_ylabel('GDP per capita', fontsize=12)
plt.show()

././_images/pandas_23_0.png
```

On-Line Data Sources

Python makes it straightforward to query online databases programmatically.

An important database for economists is $\underline{\mathsf{FRED}}-\mathsf{a}$ vast collection of time series data maintained by the St. Louis Fed.

For example, suppose that we are interested in the <u>unemployment rate</u>.

Via FRED, the entire series for the US civilian unemployment rate can be downloaded directly by entering this URL into your browser (note that this requires an internet connection)

```
https://research.stlouisfed.org/fred2/series/UNRATE/downloaddata/UNRATE.csv
```

(Equivalently, click here:

https://research.stlouisfed.org/fred2/series/UNRATE/downloaddata/UNRATE.csv)

This request returns a CSV file, which will be handled by your default application for this class of files.

Alternatively, we can access the CSV file from within a Python program.

This can be done with a variety of methods.

We start with a relatively low-level method and then return to pandas.

Accessing Data with requests

One option is to use $\underline{requests}$, a standard Python library for requesting data over the Internet.

To begin, try the following code on your computer

```
r =
requests.get('http://research.stlouisfed.org/fred2/series/UNRATE/download
data/UNRATE.csv')
```

If there's no error message, then the call has succeeded.

If you do get an error, then there are two likely causes

- 1. You are not connected to the Internet hopefully, this isn't the case.
- Your machine is accessing the Internet through a proxy server, and Python isn't aware of this.

In the second case, you can either

- switch to another machine
- solve your proxy problem by reading the documentation

Assuming that all is working, you can now proceed to use the source object returned by the

requests.get('http://research.stlouisfed.org/fred2/series/UNRATE/downloaddat a/UNRATE.csv')

```
url =
'http://research.stlouisfed.org/fred2/series/UNRATE/downloaddata/UNRATE.c
sv'
source = requests.get(url).content.decode().split("\n")
source[0]
```

'DATE, VALUE\r'

```
source[1]
```

'1948-01-01,3.4\r'

```
source[2]
```

'1948-02-01,3.8\r'

We could now write some additional code to parse this text and store it as an array.

But this is unnecessary — pandas' read_csv function can handle the task for us.

We use parse_dates=True so that pandas recognizes our dates column, allowing for simple date filtering

```
data = pd.read_csv(url, index_col=0, parse_dates=True)
```

The data has been read into a pandas DataFrame called data that we can now manipulate in the usual way

```
type(data)
```

pandas.core.frame.DataFrame

data.head() # A useful method to get a quick look at a data frame

	VALUE
DATE	
1948-01-01	3.4
1948-02-01	3.8
1948-03-01	4.0
1948-04-01	3.9
1948-05-01	3.5

```
pd.set_option('precision', 1)
data.describe() # Your output might differ slightly
```

	VALUE
count	866.0
mean	5.7
std	1.6
min	2.5
25%	4.5
50%	5.5
75%	6.8
max	10.8

We can also plot the unemployment rate from 2006 to 2012 as follows

```
ax = data['2006':'2012'].plot(title='US Unemployment Rate', legend=False)
ax.set_xlabel('year', fontsize=12)
ax.set_ylabel('%', fontsize=12)
plt.show()
```

Note that pandas offers many other file type alternatives.

Pandas has <u>a wide variety</u> of top-level methods that we can use to read, excel, json, parquet or plug straight into a database server.

Using pandas_datareader to Access Data

The maker of pandas has also authored a library called pandas_datareader that gives programmatic access to many data sources straight from the Jupyter notebook.

While some sources require an access key, many of the most important (e.g., FRED, <u>OECD</u>, <u>EUROSTAT</u> and the World Bank) are free to use.

For now let's work through one example of downloading and plotting data — this time from the World Bank.

The World Bank collects and organizes data on a huge range of indicators.

For example, here's some data on government debt as a ratio to GDP.

The next code example fetches the data for you and plots time series for the US and Australia

```
from pandas_datareader import wb

govt_debt = wb.download(indicator='GC.DDD.TOTL.GD.ZS', country=['US',
'AU'], start=2005, end=2016).stack().unstack(0)
ind = govt_debt.index.droplevel(-1)
govt_debt.index = ind
ax = govt_debt.plot(lw=2)
ax.set_xlabel('year', fontsize=12)
plt.title("Government Debt to GDP (%)")
plt.show()
```

==../../_images/pandas_55_1.pmg

The <u>documentation</u> provides more details on how to access various data sources.

Exercises

Exercise 1

With these imports:

```
import datetime as dt
from pandas_datareader import data
```

Write a program to calculate the percentage price change over 2019 for the following shares:

Here's the first part of the program

Complete the program to plot the result as a bar graph like this one:

Exercise 2

Using the method read_data introduced in <u>Exercise 1</u>, write a program to obtain year-on-year percentage change for the following indices:

Complete the program to show summary statistics and plot the result as a time series graph like this one:

Solutions

Exercise 1

There are a few ways to approach this problem using Pandas to calculate the percentage change.

First, you can extract the data and perform the calculation such as:

```
p1 = ticker.iloc[0]  #Get the firs
p2 = ticker.iloc[-1]  #Get the last
price_change = (p2 - p1) / p1 * 100
price_change
                                      #Get the first set of prices as a Series
#Get the last set of prices as a Series
INTC
              27.1
MSFT
              56.0
IBM
              16.3
BHP
             14.3
TM
              20.9
AAPL
              85.9
AMZN
             20.1
               0.6
BA
QCOM
K0
              17.9
G00G
             27.8
39.6
SNE
PTR
             -17.4
dtype: float64
```

Alternatively you can use an inbuilt method pct_change and configure it to perform the correct calculation using periods argument.

```
change = ticker.pct_change(periods=len(ticker)-1, axis='rows')*100
price_change = change.iloc[-1]
price_change
```

```
INTC
MSFT
         56.0
IBM
        16.3
BHP
         20.9
TM
AAPL
AMZN
BA
         0.6
QCOM
        53.7
17.9
K0
GOOG
        27.8
SNE
        39.6
PTR
        -17.4
Name: 2019-12-31 00:00:00, dtype: float64
```

Then to plot the chart

```
price_change.sort_values(inplace=True)
price_change = price_change.rename(index=ticker_list)
fig, ax = plt.subplots(figsize=(10,8))
ax.set_xlabel('stock', fontsize=12)
ax.set_ylabel('percentage change in price', fontsize=12)
price_change.plot(kind='bar', ax=ax)
plt.show()

.../_images/pandas_40_0.png
```

Exercise 2

Following the work you did in <u>Exercise 1</u>, you can query the data using read_data by updating the start and end dates accordingly.

```
indices_data = read_data(
    indices_list,
        start=dt.datetime(1928, 1, 2),
        end=dt.datetime(2020, 12, 31)
)
```

Then, extract the first and last set of prices per year as DataFrames and calculate the yearly returns such as:

```
yearly_returns = pd.DataFrame()
for index, name in indices_list.items():
    p1 = indices_data.groupby(indices_data.index.year)[index].first() #
Get the first set of returns as a DataFrame
    p2 = indices_data.groupby(indices_data.index.year)[index].last() #
Get the last set of returns as a DataFrame
    returns = (p2 - p1) / p1
    yearly_returns[name] = returns

yearly_returns
```

,					
	S&P 500	NASDAQ	Dow Jones	Nikkei	
Date					
1928	3.7e-01	NaN	NaN	NaN	
1929	-1.4e-01	NaN	NaN	NaN	
1930	-2.8e-01	NaN	NaN	NaN	
1931	-4.9e-01	NaN	NaN	NaN	
1932	-8.5e-02	NaN	NaN	NaN	
•••					
2016	1.1e-01	9.8e-02	1.5e-01	3.6e-02	
2017	1.8e-01	2.7e-01	2.4e-01	1.6e-01	
2018	-7.0e-02	-5.3e-02	-6.0e-02	-1.5e-01	
2019	2.9e-01	3.5e-01	2.2e-01	2.1e-01	
2020	-1.9e-01	-1.4e-01	-2.3e-01	-1.8e-01	

93 rows × 4 columns

Next, you can obtain summary statistics by using the method ${\tt describe}.$

```
yearly_returns.describe()
```

	S&P 500	NASDAQ	Dow Jones	Nikkei
count	9.3e+01	5.0e+01	3.6e+01	5.6e+01
mean	7.3e-02	1.2e-01	8.9e-02	7.4e-02
std	1.9e-01	2.5e-01	1.5e-01	2.4e-01
min	-4.9e-01	-4.0e-01	-3.3e-01	-4.0e-01
25%	-6.5e-02	-3.0e-02	4.2e-03	-1.0e-01
50%	9.9e-02	1.4e-01	8.9e-02	7.5e-02
75%	2.0e-01	2.7e-01	2.2e-01	2.1e-01
max	4.6e-01	8.4e-01	3.3e-01	9.2e-01

Then, to plot the chart

Footnotes

2 Wikipedia defines munging as cleaning data from one raw form into a structured, purged one.

Advanced Python Programming

This part provides a look at more advanced concepts in Python programming.

Writing Good Code

Overview

When computer programs are small, poorly written code is not overly costly.

But more data, more sophisticated models, and more computer power are enabling us to take on more challenging problems that involve writing longer programs.

For such programs, investment in good coding practices will pay high returns.

The main payoffs are higher productivity and faster code.

In this lecture, we review some elements of good coding practice.

We also touch on modern developments in scientific computing — such as just in time compilation — and how they affect good program design.

An Example of Poor Code

Let's have a look at some poorly written code.

The job of the code is to generate and plot time series of the simplified Solow model

$$k_{t+1} = sk_t^{\alpha} + (1 - \delta)k_t, \quad t = 0, 1, 2, \dots$$
 (8)

Here

- k_t is capital at time t and
- s, α, δ are parameters (savings, a productivity parameter and depreciation)

For each parameterization, the code

```
1. sets k_0=1
2. iterates using (8) to produce a sequence k_0,k_1,k_2\ldots,k_T
3. plots the sequence
```

The plots will be grouped into three subfigures.

In each subfigure, two parameters are held fixed while another varies

```
import numpy as np
import matplotlib.pyplot as plt
%matplotlib inline
# Allocate memory for time series
k = np.empty(50)
fig, axes = plt.subplots(3, 1, figsize=(6, 14))
# Trajectories with different \alpha
\delta = 0.1
s = 0.4
\alpha = (0.25, 0.33, 0.45)
for j in range(3):
      k[0] = 1
      for t in range(49):
      axes[0].grid(lw=0.2)
axes[0].set_ylim(0, 18)
axes[0].set_xlabel('time')
axes[0].set_ylabel('capital')
axes[0] legend(loc='upper left', frameon=True)
# Trajectories with different s
\delta = 0.1
\alpha = 0.33
s = (0.3, 0.4, 0.5)
for j in range(3):
      k[0] = 1
      for t in range(49):
      k[t+1] = s[j] * k[t] **\alpha + (1 - \delta) * k[t]

axes[1].plot(k, 'o-', label=rf"$\alpha = {\alpha},\; s = {s[j]},\; \delta=
axes[1].grid(lw=0.2)
axes[1].set_xlabel('time')
axes[1].set_ylabel('capital')
axes[1].set_ylim(0, 18)
axes[1].legend(loc='upper left', frameon=True)
# Trajectories with different \delta
\delta = (0.05, 0.1, 0.15)
\alpha = 0.33
s = 0.4
for j in range(3):
      k[0] = 1
      for t in range(49): k[t+1] = s * k[t] **\alpha + (1 - \delta[j]) * k[t] axes[2].plot(k, 'o-', label=rf"$\alpha = {\alpha},\; s = {s},\; \delta=
{δ[j]}$")
axes[2].set_ylim(0, 18)
axes[2].set_xlabel('time')
axes[2].set_ylabel('capital')
axes[2].grid(lw=0.2)
axes[2].legend(loc='upper left', frameon=True)
plt.show()
......_images/writing_good_code_0_0.png
```

True, the code more or less follows <u>PEP8</u>.

At the same time, it's very poorly structured.

Let's talk about why that's the case, and what we can do about it.

Good Coding Practice

There are usually many different ways to write a program that accomplishes a given task.

For small programs, like the one above, the way you write code doesn't matter too much.

But if you are ambitious and want to produce useful things, you'll write medium to large programs too.

In those settings, coding style matters a great deal.

Fortunately, lots of smart people have thought about the best way to write code.

Here are some basic precepts.

Don't Use Magic Numbers

If you look at the code above, you'll see numbers like 50 and 49 and 3 scattered through the code.

These kinds of numeric literals in the body of your code are sometimes called "magic numbers".

This is not a compliment.

While numeric literals are not all evil, the numbers shown in the program above should certainly be replaced by named constants.

For example, the code above could declare the variable time_series_length = 50.

Then in the loops, 49 should be replaced by time_series_length - 1.

The advantages are:

- the meaning is much clearer throughout
- to alter the time series length, you only need to change one value

Don't Repeat Yourself

The other mortal sin in the code snippet above is repetition.

Blocks of logic (such as the loop to generate time series) are repeated with only minor changes.

This violates a fundamental tenet of programming: Don't repeat yourself (DRY).

• Also called DIE (duplication is evil).

Yes, we realize that you can just cut and paste and change a few symbols.

But as a programmer, your aim should be to ${\bf automate}$ repetition, ${\bf not}$ do it yourself.

More importantly, repeating the same logic in different places means that eventually one of them will likely be wrong.

If you want to know more, read the excellent summary found on $\underline{\text{this page}}.$

We'll talk about how to avoid repetition below.

Minimize Global Variables

Sure, global variables (i.e., names assigned to values outside of any function or class) are convenient.

Rookie programmers typically use global variables with abandon — as we once did ourselves.

But global variables are dangerous, especially in medium to large size programs, since

- they can affect what happens in any part of your program
- they can be changed by any function

This makes it much harder to be certain about what some small part of a given piece of code actually commands.

Here's a useful discussion on the topic.

While the odd global in small scripts is no big deal, we recommend that you teach yourself to avoid them.

(We'll discuss how just below).

JIT Compilation

For scientific computing, there is another good reason to avoid global variables.

As <u>we've seen in previous lectures</u>, JIT compilation can generate excellent performance for scripting languages like Python.

But the task of the compiler used for JIT compilation becomes harder when global variables are present.

Put differently, the type inference required for JIT compilation is safer and more effective when variables are sandboxed inside a function.

Use Functions or Classes

Fortunately, we can easily avoid the evils of global variables and WET code.

• WET stands for "we enjoy typing" and is the opposite of DRY.

We can do this by making frequent use of functions or classes.

In fact, functions and classes are designed specifically to help us avoid shaming ourselves by repeating code or excessive use of global variables.

Which One, Functions or Classes?

Both can be useful, and in fact they work well with each other.

We'll learn more about these topics over time.

(Personal preference is part of the story too)

What's really important is that you use one or the other or both.

Revisiting the Example

Here's some code that reproduces the plot above with better coding style.

```
from itertools import product

def plot_path(ax, αs, s_vals, δs, time_series_length=50):
    """

    Add a time series plot to the axes ax for all given parameters.
    """

    k = np.empty(time_series_length)

    for (α, s, δ) in product(αs, s_vals, δs):
        k[0] = 1
        for t in range(time_series_length=1):
              k[t+1] = s * k[t]**α + (1 - δ) * k[t]
        ax.plot(k, 'o-', label=rf"$\alpha = {α},\; s = {s},\; \delta = {δ}$")

    ax.set_xlabel('time')
    ax.set_ylabel('capital')
    ax.set_ylam(0, 18)
    ax.legend(loc='upper left', frameon=True)

fig, axes = plt.subplots(3, 1, figsize=(6, 14))

# Parameters (αs, s_vals, δs)
set_one = ([0.25, 0.33, 0.45], [0.4], [0.1])
set_two = ([0.33], [0.3, 0.4, 0.5], [0.1])
set_three = ([0.33], [0.4], [0.05, 0.1, 0.15])

for (ax, params) in zip(axes, (set_one, set_two, set_three)):
    αs, s_vals, δs = params
    plot_path(ax, αs, s_vals, δs)

plt.show()
```

../../_images/writing_good_code_1_0.png

If you inspect this code, you will see that

- it uses a function to avoid repetition.
- Global variables are quarantined by collecting them together at the end, not the start of the program.
- Magic numbers are avoided.
- The loop at the end where the actual work is done is short and relatively simple.

Exercises

Exercise 1

Here is some code that needs improving.

It involves a basic supply and demand problem.

Supply is given by

$$q_s(p) = \exp(\alpha p) - \beta.$$

The demand curve is

$$q_d(p) = \gamma p^{-\delta}.$$

The values α , β , γ and δ are **parameters**

The equilibrium p^* is the price such that $q_d(p) = q_s(p)$.

We can solve for this equilibrium using a root finding algorithm. Specifically, we will find the p such that h(p)=0, where

$$h(p) := q_d(p) - q_s(p)$$

This yields the equilibrium price p^* . From this we get the equilibrium price by $q^* = q_s(p^*)$

The parameter values will be

- $\alpha = 0.1$
- $\beta = 1$
- $\gamma = 1$
- $\delta = 1$

```
from scipy.optimize import brentq

# Compute equilibrium
def h(p):
    return p**(-1) - (np.exp(0.1 * p) - 1) # demand - supply

p_star = brentq(h, 2, 4)
q_star = np.exp(0.1 * p_star) - 1

print(f'Equilibrium price is {p_star: .2f}')
print(f'Equilibrium quantity is {q_star: .2f}')
```

Equilibrium price is 2.93 Equilibrium quantity is 0.34

Let's also plot our results.

```
# Now plot
grid = np.linspace(2, 4, 100)
fig, ax = plt.subplots()

qs = np.exp(0.1 * grid) - 1
qd = grid**(-1)

ax.plot(grid, qd, 'b-', lw=2, label='demand')
ax.plot(grid, qs, 'g-', lw=2, label='supply')
ax.set_xlabel('price')
ax.set_ylabel('price')
ax.set_ylabel('quantity')
ax.legend(loc='upper center')
plt.show()
```

We also want to consider supply and demand shifts.

For example, let's see what happens when demand shifts up, with γ increasing to 1.25:

```
# Compute equilibrium
def h(p):
    return 1.25 * p**(-1) - (np.exp(0.1 * p) - 1)

p_star = brentq(h, 2, 4)
q_star = np.exp(0.1 * p_star) - 1

print(f'Equilibrium price is {p_star: .2f}')
print(f'Equilibrium quantity is {q_star: .2f}')
```

Equilibrium price is 3.25 Equilibrium quantity is 0.38

```
# Now plot
p_grid = np.linspace(2, 4, 100)
fig, ax = plt.subplots()

qs = np.exp(0.1 * p_grid) - 1
qd = 1.25 * p_grid**(-1)

ax.plot(grid, qd, 'b-', lw=2, label='demand')
ax.plot(grid, qs, 'g-', lw=2, label='supply')

ax.set_xlabel('price')
ax.set_ylabel('price')
ax.set_ylabel('quantity')
ax.legend(loc='upper center')
plt.show()
```

.../../_images/writing_good_code_5_0.png

Now we might consider supply shifts, but you already get the idea that there's a lot of repeated code here.

Refactor and improve clarity in the code above using the principles discussed in this lecture.

Solutions

Exercise 1

Here's one solution, that uses a class:

```
class Equilibrium:
          def qs(self, p):
          return np.exp(self.\alpha * p) - self.\beta
     def qd(self, p):
    return self.γ * p**(-self.δ)
     def compute_equilibrium(self):
          def h(p):
               return self.qd(p) - self.qs(p)
          p_star = brentq(h, 2, 4)
q_star = np.exp(self.α * p_star) - self.β
          print(f'Equilibrium price is {p_star: .2f}')
print(f'Equilibrium quantity is {q_star: .2f}')
     def plot_equilibrium(self):
          # Now plot
grid = np.linspace(2, 4, 100)
fig, ax = plt.subplots()
          ax.plot(grid, self.qd(grid), 'b-', lw=2, label='demand') ax.plot(grid, self.qs(grid), 'g-', lw=2, label='supply')  
          ax.set_xlabel('price')
ax.set_ylabel('quantity')
          ax.legend(loc='upper center')
          plt.show()
```

Let's create an instance at the default parameter values.

```
eq = Equilibrium()
```

Now we'll compute the equilibrium and plot it.

```
eq.compute_equilibrium()

Equilibrium price is 2.93
Equilibrium quantity is 0.34
```

eq.plot_equilibrium()

.../_images/writing_good_code_9_0.png

One of the nice things about our refactored code is that, when we change parameters, we don't need to repeat ourselves:

```
eq.γ = 1.25

eq.compute_equilibrium()

Equilibrium price is 3.25
Equilibrium quantity is 0.38

eq.plot_equilibrium()

././_images/writing_good_code_12_0.png
```

More Language Features

Overview

Tip

With this last lecture, our advice is to **skip it on first pass**, unless you have a burning desire to read it.

It's here

- 1. as a reference, so we can link back to it when required, and
- 2. for those who have worked through a number of applications, and now want to learn more about the Python language

A variety of topics are treated in the lecture, including generators, exceptions and descriptors.

Iterables and Iterators

We've <u>already said something</u> about iterating in Python.

Now let's look more closely at how it all works, focusing in Python's implementation of the for loop.

Iterators

Iterators are a uniform interface to stepping through elements in a collection.

Here we'll talk about using iterators—later we'll learn how to build our own.

Formally, an *iterator* is an object with a __next__ method.

For example, file objects are iterators .

To see this, let's have another look at the <u>US cities data</u>, which is written to the present working directory in the following cell

```
%%file us_cities.txt
new york: 8244910
los angeles: 3819702
chicago: 2707120
houston: 2145146
philadelphia: 1536471
phoenix: 1469471
san antonio: 1359758
san diego: 1326179
dallas: 1223229
```

Overwriting us_cities.txt

```
f = open('us_cities.txt')
f.__next__()
```

'new york: 8244910\n'

```
f.__next__()
```

'los angeles: 3819702\n'

We see that file objects do indeed have a __next__ method, and that calling this method returns the next line in the file.

The next method can also be accessed via the builtin function $\mathsf{next}()$, which directly calls this method

```
next(f)
'chicago: 2707120\n'
```

The objects returned by enumerate() are also iterators

```
e = enumerate(['foo', 'bar'])
next(e)
```

(0, 'foo')

```
next(e)
(1, 'bar')
```

as are the reader objects from the csv module.

Let's create a small csv file that contains data from the NIKKEI index

```
%%file test_table.csv
Date,Open,High,Low,Close,Volume,Adj Close
2009-05-21,9280.35,9286.35,9189.92,9264.15,133200,9264.15
2009-05-20,9372.72,9399.40,9311.61,9344.64,143200,9344.64
2009-05-19,9172.56,9326.75,9166.97,9290.29,167000,9290.29
2009-05-18,9167.05,9167.82,8997.74,9038.69,147800,9038.69
2009-05-15,9150.21,9272.08,9140.90,9265.02,172000,9265.02
2009-05-14,9212.30,9223.77,9052.41,9093.73,169400,9093.73
2009-05-13,9305.79,9379.47,9278.89,9340.49,176000,9340.49
2009-05-12,9358.25,9389.61,9298.61,9298.61,188400,9298.61
2009-05-11,9460.72,9503.91,9342.75,9451.98,230800,9451.98
2009-05-00,9351.40,9464.43,9349.57,9432.83,220200,9432.83
```

Overwriting test_table.csv

```
from csv import reader

f = open('test_table.csv', 'r')
nikkei_data = reader(f)
next(nikkei_data)
```

```
['Date', 'Open', 'High', 'Low', 'Close', 'Volume', 'Adj Close']
```

```
next(nikkei_data)
```

```
['2009-05-21', '9280.35', '9286.35', '9189.92', '9264.15', '133200', '9264.15']
```

Iterators in For Loops

All iterators can be placed to the right of the in keyword in for loop statements.

In fact this is how the for loop works: If we write

```
for x in iterator:
     <code block>
```

then the interpreter

- calls iterator.___next___() and binds x to the result
- executes the code block
- repeats until a StopIteration error occurs

So now you know how this magical looking syntax works

```
f = open('somefile.txt', 'r')
for line in f:
    # do something
```

The interpreter just keeps

- 1. calling f.__next__() and binding line to the result
- 2. executing the body of the loop

This continues until a StopIteration error occurs.

Iterables

You already know that we can put a Python list to the right of in in a for loop

```
for i in ['spam', 'eggs']:
    print(i)

spam
eggs
```

So does that mean that a list is an iterator?

The answer is no

```
x = ['foo', 'bar']
type(x)
```

list

```
next(x)
```

TypeError Traceback (most recent call last)
<ipython-input-12-92de4e9f6b1e> in <module>
----> 1 next(x)

TypeError: 'list' object is not an iterator

So why can we iterate over a list in a for loop?

The reason is that a list is iterable (as opposed to an iterator).

Formally, an object is iterable if it can be converted to an iterator using the built-in function iter().

Lists are one such object

```
x = ['foo', 'bar']
 type(x)
 list
 y = iter(x)
 type(y)
 list_iterator
 next(y)
  'foo'
 next(y)
  'bar'
 next(y)
 StopIteration
                                                  Traceback (most recent call
 <ipython-input-17-81e57198612b> in <module>
----> 1 next(y)
 StopIteration:
Many other objects are iterable, such as dictionaries and tuples.
Of course, not all objects are iterable
 iter(42)
 TypeError last)
                                                  Traceback (most recent call
 <ipython-input-18-ef50b48e4398> in <module>
---> 1 iter(42)
 TypeError: 'int' object is not iterable
To conclude our discussion of for loops
   • for loops work on either iterators or iterables.
   • In the second case, the iterable is converted into an iterator before the loop starts.
Iterators and built-ins
Some built-in functions that act on sequences also work with iterables
   • max(),min(),sum(),all(),any()
For example
```

```
x = [10, -10]
max(x)
```

10

```
y = iter(x)
type(y)
```

list_iterator

```
max(y)
```

One thing to remember about iterators is that they are depleted by use

```
x = [10, -10]
y = iter(x)
max(y)
```

10

Names and Name Resolution

Variable Names in Python

Consider the Python statement

```
x = 42
```

We now know that when this statement is executed, Python creates an object of type int in your computer's memory, containing

- the value 42
- some associated attributes

But what is x itself?

In Python, x is called a *name*, and the statement x = 42 *binds* the name x to the integer object we have just discussed.

Under the hood, this process of binding names to objects is implemented as a dictionary—more about this in a moment.

There is no problem binding two or more names to the one object, regardless of what that object is

```
def f(string):  # Create a function called f
  print(string)  # that prints any string it's passed

g = f
id(g) == id(f)
```

True

```
g('test')
```

test

In the first step, a function object is created, and the name f is bound to it.

After binding the name ${\tt g}$ to the same object, we can use it anywhere we would use {\tt f}.

What happens when the number of names bound to an object goes to zero?

Here's an example of this situation, where the name \times is first bound to one object and then rebound to another

```
x = 'foo'
id(x)
```

4518185008

```
x = 'bar' # No names bound to the first object
```

What happens here is that the first object is garbage collected.

In other words, the memory slot that stores that object is deallocated, and returned to the operating system.

Namespaces

Recall from the preceding discussion that the statement

```
x = 42
```

binds the name x to the integer object on the right-hand side.

We also mentioned that this process of binding \mathbf{x} to the correct object is implemented as a dictionary.

This dictionary is called a *namespace*.

Definition: A namespace is a symbol table that maps names to objects in memory.

 $Python \ uses \ multiple \ names paces, creating \ them \ on \ the \ fly \ as \ necessary \ .$

For example, every time we import a module, Python creates a namespace for that module.

To see this in action, suppose we write a script math2.py with a single line

```
%%file math2.py
pi = 'foobar'
```

Overwriting math2.py

Now we start the Python interpreter and import it

```
import math2
```

Next let's import the $\operatorname{{\it math}}$ module from the standard library

```
import math
```

Both of these modules have an attribute called pi

```
math.pi
```

3.141592653589793

```
math2.pi
```

'foobar'

These two different bindings of pi exist in different namespaces, each one implemented as a dictionary.

We can look at the dictionary directly, using module_name.__dict__

```
import math
math.__dict__.items()
```

import math2

math2.__dict__.items()

dict_items([('__name__', 'math2'), ('__doc__', None), ('__package__', ''),
 ('__loader__', <_frozen_importlib_external.SourceFileLoader object at
 0x10f389dc0>), ('__spec__', ModuleSpec(name='math2', loader=
 <_frozen_importlib_external.SourceFileLoader_object at 0x10f389dc0>, origin='/Users/junaluzi/Documents/ExecutableBookProject/quantecon-example/math2.py')), ('__file__', '/Users/junaluzi/Documents/ExecutableBookProject/quanteconr/ose1s/junatu21/bocuments/ExecutableBookProject/quantecon-example/math2.py'), ('_cached_', '/Users/junatu21/Documents/ExecutableBookProject/quantecon-example/_pycache__/math2.cpython-38.pyc'), ('__builtins_', {'__ibuiltins', '__doc__': "Built-in functions, exceptions, and other 'builtins', '_doc_': 'Built-in functions, exceptions, and other objects.\nNnOteworthy: None is the `nil' object; Ellipsis represents ...' in slices.", '_package__': '', '_loader__': <class '_frozen_importlib.BuiltinImporter'>, '_spec__': ModuleSpec(name='builtins', loader=<class '_frozen_importlib.BuiltinImporter'>), '_build_class__': <built-in function _build_class__, '_import__': <built-in function _import_>, 'abs': <built-in function abs>, 'all': <built-in function all>, 'any': <built-in function any>, 'ascii': <built-in function ascii>, 'bin': <built-in function bin>, 'breakpoint': <built-in function bin>, 'breakpoint': <built-in function corp>, 'callable': <built-in function callable>, 'chr': <built-in function delattr>, 'dir': <built-in function dir>, 'divmod': <built-in function divmod>, 'eval': <built-in function eval>, 'exec': <built-in function exec>, 'format': <built-in function globals>, 'hasattr': <built-in function getattr>, 'globals': <built-in function globals>, 'hasattr': <built-in function getattr>, 'globals': <built-in function globals>, 'hasattr': <built-in function hasattr; duilt-in function hasattr; <built-in function hasattr>, 'hash': <built-in function hasa>, 'hex': <built-in function hex>, 'id': <built-in function id>, 'input': <built-in func function hex>, 'id': <built-in function id>, 'input': <bundant with the control of the control o 'NotImplementedError': <class 'NotImplementedError'>, 'NameError': <class 'NameError'>, 'UnboundLocalError': <class 'UnboundLocalError'>, 'AttributeError': <class 'AttributeError'>, 'SyntaxError': <class 'SyntaxError'>, 'IndentationError': <class 'IndentationError'>, 'TabError': <class 'IndentationError'>, 'IndexError'>, 'LookupError': <class 'LookupError'>, 'IndexError': <class 'IndexError'>, 'KeyError': <class 'KeyError'>, 'ValueError': <class 'ValueError'>, 'UnicodeError': <class ValueError': <ctass ValueError': <ctass 'UnicodeError's, UnicodeError's, 'UnicodeError's, 'UnicodeError's, 'UnicodeError': <ctass 'UnicodeError's, 'UnicodeTranslateError'; <ctass 'UnicodeTranslateError's, 'ArithmeticError': <ctass 'ArithmeticError': <ctass 'ArithmeticError': <ctass 'ArithmeticError': <ctass 'ArithmeticError'; 'FloatingPointError': <ctass 'FloatingPointError's, 'OverflowError': <ctass 'OverflowError': <ctass 'Initial 'Initia 'UnicodeError'>, 'UnicodeEncodeError': <class 'UnicodeEncodeError'>, 'ChildProcessError': <class 'ChildProcessError'>,
'ConnectionAbortedError': <class 'ConnectionAbortedError'>,
'ConnectionRefusedError': <class 'ConnectionRefusedError'>,
'ConnectionResetError': <class 'ConnectionResetError'>, 'FileExistsError':
<class 'FileExistsError'>, 'FileNotFoundError': <class 'IsaDirectoryError': <class 'IsaDirectoryError'>,
'NotADirectoryError': <class 'NotADirectoryError'>, 'InterruptedError':
<class 'InterruptedError'>, 'PermissionError': <class 'PermissionError'>,
'ProcessLookupError': <class 'ProcessLookupError'>, 'TimeoutError': <class 'TimeoutError'>, 'open':

'TimeoutError'>, 'open':

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*bound method InteractiveShell.get_ipython of <ipykernel.zmqshell.ZMQInteractiveShell object at 0x10f212e50>>}), ('pi', 'foobar')])

As you know, we access elements of the namespace using the dotted attribute notation

```
math.pi

3.141592653589793

In fact this is entirely equivalent to math.__dict__['pi']
```

math.__dict__['pi'] == math.pi

True

Viewing Namespaces

As we saw above, the math namespace can be printed by typing math. __dict__.

Another way to see its contents is to type vars (math)

If you just want to see the names, you can type

dir(math)[0:10]

Notice the special names <u>__doc__</u> and <u>__name__</u>.

These are initialized in the namespace when any module is imported

- __doc__ is the doc string of the module
- __name__ is the name of the module

```
print(math.__doc__)
```

This module provides access to the mathematical functions defined by the $\ensuremath{\mathsf{C}}$ standard.

```
math.__name__
'math'
```

Interactive Sessions

In Python, all code executed by the interpreter runs in some module.

What about commands typed at the prompt?

These are also regarded as being executed within a module - in this case, a module called $_$ main $_$.

To check this, we can look at the current module name via the value of $_$ name $_$ given at the prompt

```
print(__name__)
__main__
```

When we run a script using IPython's run command, the contents of the file are executed as part of __main__ too.

To see this, let's create a file mod.py that prints its own __name__ attribute

```
%%file mod.py
print(__name__)
```

Overwriting mod.py

Now let's look at two different ways of running it in IPython

```
import mod # Standard import

mod

%run mod.py # Run interactively
__main__
```

In the second case, the code is executed as part of __main__, so __name__ is equal to __main__.

```
To see the contents of the namespace of \_main\_ we use vars() rather than vars(\_main\_).
```

If you do this in IPython, you will see a whole lot of variables that IPython needs, and has initialized when you started up your session.

If you prefer to see only the variables you have initialized, use whos

```
x = 2
y = 3
import numpy as np
%whos
```

Variable	Туре	Data/Info			
e	enumerate	<pre><enumerate at<="" object="" pre=""></enumerate></pre>			
0x10f383440>					
f	function	<function 0x10f34bca0="" at="" f=""></function>			
g i	function	<function 0x10f34bca0="" at="" f=""></function>			
i	str	eggs			
math	module	<pre><module 'math'="" from<="" pre=""></module></pre>			
'/Use<>th.cpython-38-darwin.so'>					
math2	module	<module 'math2'="" from<="" td=""></module>			
'/Us<>ntecon-example/math2.py'>					
mod	module	<module 'mod'="" from<="" td=""></module>			
'/User<>uantecon-example/mod.py'>					
nikkei data		< csv.reader object at			
0x10f364ba0>					
np	module	<module 'numpy'="" from<="" td=""></module>			
'/Us<>kages/numpy/initpy'>					
reader	builtin function or method	<pre><built-in function="" reader=""></built-in></pre>			
X	int	2			
V	int	3			
У	THE	J			

The Global Namespace

Python documentation often makes reference to the "global namespace".

The global namespace is the namespace of the module currently being executed.

For example, suppose that we start the interpreter and begin making assignments .

We are now working in the module $_$ main $_$, and hence the namespace for $_$ main $_$ is the global namespace.

Next, we import a module called amodule

```
import amodule
```

At this point, the interpreter creates a namespace for the module amodule and starts executing commands in the module.

While this occurs, the namespace amodule. __dict__ is the global namespace.

Once execution of the module finishes, the interpreter returns to the module from where the import statement was made.

In this case it's __main__, so the namespace of __main__ again becomes the global namespace.

Local Namespaces

Important fact: When we call a function, the interpreter creates a *local namespace* for that function, and registers the variables in that namespace.

The reason for this will be explained in just a moment.

Variables in the local namespace are called *local variables*.

After the function returns, the namespace is deallocated and lost.

While the function is executing, we can view the contents of the local name space with locals ().

For example, consider

```
def f(x):
    a = 2
    print(locals())
    return a * x
```

Now let's call the function

```
f(1)
{'x': 1, 'a': 2}
2
```

You can see the local namespace of f before it is destroyed.

```
The __builtins__ Namespace
```

We have been using various built-in functions, such as $\max()$, dir(), str(), list(), len(), range(), type(), etc.

How does access to these names work?

- These definitions are stored in a module called __builtin__.
- They have there own namespace called __builtins__.

```
dir()[0:10]

['In', 'Out', '_', '_11', '_13', '_14', '_15', '_16', '_19', '_2']

dir(__builtins__)[0:10]

['ArithmeticError',
    'AssertionError',
    'BaseException',
    'Blocking10Error',
    'BrokenPipeError',
    'BufferError',
    'BytesWarning',
    'ConnectionAbortedError']
```

We can access elements of the namespace as follows

```
__builtins__.max
<function max>
```

Tanceron max

But __builtins__ is special, because we can always access them directly as well

```
max
<function max>

_builtins_.max == max
```

True

The next section explains how this works \dots

Name Resolution

Namespaces are great because they help us organize variable names.

(Type import this at the prompt and look at the last item that's printed)

However, we do need to understand how the Python interpreter works with multiple name spaces .

At any point of execution, there are in fact at least two namespaces that can be accessed directly.

("Accessed directly" means without using a dot, as in pi rather than math.pi)

These namespaces are

- The global namespace (of the module being executed)
- The builtin namespace

If the interpreter is executing a function, then the directly accessible namespaces are

- The local namespace of the function
- The global namespace (of the module being executed)
- The builtin namespace

Sometimes functions are defined within other functions, like so

```
def f():
    a = 2
    def g():
        b = 4
        print(a * b)
    g()
```

Here ${\bf f}$ is the ${\it enclosing function}$ for ${\bf g},$ and each function gets its own namespaces.

Now we can give the rule for how namespace resolution works:

The order in which the interpreter searches for names is

- 1. the local namespace (if it exists)
- 2. the hierarchy of enclosing namespaces (if they exist)
- 3. the global namespace
- 4. the builtin namespace

If the name is not in any of these namespaces, the interpreter raises a NameError.

This is called the **LEGB rule** (local, enclosing, global, builtin).

Here's an example that helps to illustrate.

Consider a script test.py that looks as follows

```
%*file test.py
def g(x):
    a = 1
    x = x + a
    return x

a = 0
y = g(10)
print("a = ", a, "y = ", y)
```

Overwriting test.py

What happens when we run this script?

```
%run test.py

a = 0 y = 11
```

```
x
```

First,

- The global namespace {} is created.
- The function object is created, and g is bound to it within the global namespace.
- The name a is bound to 0, again in the global namespace.

Next g is called via y = g(10), leading to the following sequence of actions

- The local namespace for the function is created.
- Local names x and a are bound, so that the local namespace becomes {'x': 10, 'a': 1}.
- Statement x = x + a uses the local a and local x to compute x + a, and binds local name x to the result.
- This value is returned, and y is bound to it in the global namespace.
- Local x and a are discarded (and the local namespace is deallocated).

Note that the global a was not affected by the local a.

Mutable Versus Immutable Parameters

This is a good time to say a little more about mutable vs immutable objects.

Consider the code segment

```
def f(x):
    x = x + 1
    return x

x = 1
print(f(x), x)
```

2 1

We now understand what will happen here: The code prints 2 as the value of f(x) and 1 as the value of x.

First f and x are registered in the global namespace.

The call f(x) creates a local namespace and adds x to it, bound to 1.

Next, this local x is rebound to the new integer object 2, and this value is returned.

None of this affects the global x.

However, it's a different story when we use a **mutable** data type such as a list

```
def f(x):
    x[0] = x[0] + 1
    return x

x = [1]
print(f(x), x)
```

[2]

[2]

This prints [2] as the value of f(x) and same for x.

Here's what happens

- f is registered as a function in the global namespace
- x bound to [1] in the global namespace
- The call f(x)
 - o Creates a local namespace
 - Adds x to local namespace, bound to [1]
 - The list [1] is modified to [2]
 - o Returns the list [2]
 - o The local namespace is deallocated, and local x is lost
- Global x has been modified

Handling Errors

Sometimes it's possible to anticipate errors as we're writing code.

For example, the unbiased sample variance of sample y_1, \ldots, y_n is defined as

$$s^2 := \frac{1}{n-1} \sum_{i=1}^{n} (y_i - \bar{y})^2$$
 $\bar{y} = \text{sample mean}$

This can be calculated in NumPy using np.var.

But if you were writing a function to handle such a calculation, you might anticipate a divide-by-zero error when the sample size is one.

One possible action is to do nothing — the program will just crash, and spit out an error message.

But sometimes it's worth writing your code in a way that anticipates and deals with runtime errors that you think might arise.

Why?

- Because the debugging information provided by the interpreter is often less useful than the information on possible errors you have in your head when writing code.
- Because errors causing execution to stop are frustrating if you're in the middle of a large computation.
- Because it's reduces confidence in your code on the part of your users (if you are writing for others).

Assertions

A relatively easy way to handle checks is with the assert keyword.

For example, pretend for a moment that the ${\sf np.var}$ function doesn't exist and we need to write our own

```
def var(y):
    n = len(y)
    assert n > 1, 'Sample size must be greater than one.'
    return np.sum((y - y.mean())**2) / float(n-1)
```

If we run this with an array of length one, the program will terminate and print our error message

```
var([1])
```

The advantage is that we can

- fail early, as soon as we know there will be a problem
- supply specific information on why a program is failing

Handling Errors During Runtime

The approach used above is a bit limited, because it always leads to termination.

Sometimes we can handle errors more gracefully, by treating special cases.

Let's look at how this is done.

Exceptions

Here's an example of a common error type

```
def f:
```

Since illegal syntax cannot be executed, a syntax error terminates execution of the program.

1 / 0

ZeroDivisionError Traceback (most recent call

last)

Traceback (most recent call

<ipython-input-61-bc757c3fda29> in <module> --> 1 1 / 0

ZeroDivisionError: division by zero

Here's another

```
x1 = y1
```

NameError last) <ipython-input-62-a7b8d65e9e45> in <module> -> 1 x1 = y1

NameError: name 'y1' is not defined

And another

```
'foo' + 6
```

TypeError Traceback (most recent call <ipython-input-63-216809d6e6fe> in <module>
----> 1 'foo' + 6

TypeError: can only concatenate str (not "int") to str

And another

```
X = []
X = X[0]
```

 ${\tt IndexError}$ Traceback (most recent call last) <ipython-input-64-082a18d7a0aa> in <module> 1 X = []---> 2 x = X[0]

On each occasion, the interpreter informs us of the error type

• NameError, TypeError, IndexError, ZeroDivisionError, etc.

In Python, these errors are called exceptions.

IndexError: list index out of range

Catching Exceptions

We can catch and deal with exceptions using try - except blocks.

Here's a simple example

```
def f(x):
    try:
       return 1.0 / x
    except ZeroDivisionError:
       print('Error: division by zero. Returned None')
    return None
```

When we call f we get the following output

f(2)

```
f(0)
```

Error: division by zero. Returned None

```
f(0.0)
```

Error: division by zero. Returned None

The error is caught and execution of the program is not terminated.

Note that other error types are not caught.

If we are worried the user might pass in a string, we can catch that error too

```
def f(x):
    try:
        return 1.0 / x
    except ZeroDivisionError:
        print('Error: Division by zero. Returned None')
    except TypeError:
        print('Error: Unsupported operation. Returned None')
    return None
```

Here's what happens

```
f(2)
```

0.5

```
f(0)
```

Error: Division by zero. Returned None

```
f('foo')
```

Error: Unsupported operation. Returned None

If we feel lazy we can catch these errors together

```
def f(x):
    try:
        return 1.0 / x
    except (TypeError, ZeroDivisionError):
        print('Error: Unsupported operation. Returned None')
    return None
```

Here's what happens

```
f(2)
```

0.5

```
f(0)
```

Error: Unsupported operation. Returned None

```
f('foo')
```

Error: Unsupported operation. Returned None

If we feel extra lazy we can catch all error types as follows

```
def f(x):
    try:
        return 1.0 / x
    except:
        print('Error. Returned None')
    return None
```

Decorators and Descriptors

Let's look at some special syntax elements that are routinely used by Python developers.

You might not need the following concepts immediately, but you will see them in other people's code.

Hence you need to understand them at some stage of your Python education.

Decorators

Decorators are a bit of syntactic sugar that, while easily avoided, have turned out to be popular.

It's very easy to say what decorators do.

On the other hand it takes a bit of effort to explain why you might use them.

An Example

Suppose we are working on a program that looks something like this

```
import numpy as np
def f(x):
    return np.log(np.log(x))

def g(x):
    return np.sqrt(42 * x)

# Program continues with various calculations using f and g
```

Now suppose there's a problem: occasionally negative numbers get fed to ${\tt f}$ and ${\tt g}$ in the calculations that follow.

If you try it, you'll see that when these functions are called with negative numbers they return a NumPy object called nan .

This stands for "not a number" (and indicates that you are trying to evaluate a mathematical function at a point where it is not defined).

Perhaps this isn't what we want, because it causes other problems that are hard to pick up later on.

Suppose that instead we want the program to terminate whenever this happens, with a sensible error message.

This change is easy enough to implement

```
import numpy as np

def f(x):
    assert x >= 0, "Argument must be nonnegative"
    return np.log(np.log(x))

def g(x):
    assert x >= 0, "Argument must be nonnegative"
    return np.sqrt(42 * x)

# Program continues with various calculations using f and g
```

Notice however that there is some repetition here, in the form of two identical lines of code.

Repetition makes our code longer and harder to maintain, and hence is something we try hard to avoid.

Here it's not a big deal, but imagine now that instead of just f and g, we have 20 such functions that we need to modify in exactly the same way.

This means we need to repeat the test logic (i.e., the assert line testing nonnegativity) 20 times.

The situation is still worse if the test logic is longer and more complicated.

In this kind of scenario the following approach would be neater

```
import numpy as np

def check_nonneg(func):
    def safe_function(x):
        assert x >= 0, "Argument must be nonnegative"
        return func(x)
    return safe_function

def f(x):
    return np.log(np.log(x))

def g(x):
    return np.sqrt(42 * x)

f = check_nonneg(f)
    g = check_nonneg(g)
# Program continues with various calculations using f and g
```

This looks complicated so let's work through it slowly.

To unravel the logic, consider what happens when we say $f = \text{check_nonneg}(f)$.

This calls the function check_nonneg with parameter func set equal to f.

Now check_nonneg creates a new function called safe_function that verifies x as nonnegative and then calls func on it (which is the same as f).

Finally, the global name f is then set equal to safe_function.

Now the behavior of f is as we desire, and the same is true of g.

At the same time, the test logic is written only once.

Enter Decorators

The last version of our code is still not ideal.

For example, if someone is reading our code and wants to know how f works, they will be looking for the function definition, which is

```
def f(x):
    return np.log(np.log(x))
```

They may well miss the line $f = check_nonneg(f)$.

For this and other reasons, decorators were introduced to Python.

With decorators, we can replace the lines

```
def f(x):
    return np.log(np.log(x))

def g(x):
    return np.sqrt(42 * x)

f = check_nonneg(f)
g = check_nonneg(g)
```

with

```
@check_nonneg
def f(x):
    return np.log(np.log(x))
@check_nonneg
def g(x):
    return np.sqrt(42 * x)
```

These two pieces of code do exactly the same thing.

If they do the same thing, do we really need decorator syntax?

Well, notice that the decorators sit right on top of the function definitions.

Hence anyone looking at the definition of the function will see them and be aware that the function is modified.

In the opinion of many people, this makes the decorator syntax a significant improvement to the language.

Descriptors

Descriptors solve a common problem regarding management of variables.

To understand the issue, consider a Car class, that simulates a car.

Suppose that this class defines the variables miles and kms, which give the distance traveled in miles and kilometers respectively.

A highly simplified version of the class might look as follows

```
class Car:
    def __init__(self, miles=1000):
        self.miles = miles
        self.kms = miles * 1.61
# Some other functionality, details omitted
```

One potential problem we might have here is that a user alters one of these variables but not the other

```
car = Car()
car.miles

1000

car.kms

1610.0

car.miles = 6000
car.kms
```

1610.0

In the last two lines we see that miles and kms are out of sync.

What we really want is some mechanism whereby each time a user sets one of these variables, *the other is automatically updated*.

A Solution

In Python, this issue is solved using descriptors.

A descriptor is just a Python object that implements certain methods.

These methods are triggered when the object is accessed through dotted attribute notation.

The best way to understand this is to see it in action.

Consider this alternative version of the Car class

```
class Car:

def __init__(self, miles=1000):
    self._miles = miles
    self._kms = miles * 1.61

def set_miles(self, value):
    self._miles = value
    self._kms = value * 1.61

def set_kms(self, value):
    self._kms = value
    self._kms = value
    self._kms = value / 1.61

def get_miles(self):
    return self._miles

def get_kms(self):
    return self._kms

miles = property(get_miles, set_miles)
    kms = property(get_kms, set_kms)
```

First let's check that we get the desired behavior

```
car = Car()
car.miles
```

1000

```
car.miles = 6000 car.kms
```

9660.0

Yep, that's what we want - car.kms is automatically updated.

How it Works

The names _miles and _kms are arbitrary names we are using to store the values of the variables.

The objects miles and kms are *properties*, a common kind of descriptor.

The methods get_miles, set_miles, get_kms and set_kms define what happens when you get (i.e. access) or set (bind) these variables

• So-called "getter" and "setter" methods.

The builtin Python function property takes getter and setter methods and creates a property.

For example, after car is created as an instance of Car, the object car.miles is a property.

Being a property, when we set its value via car.miles = 6000 its setter method is triggered — in this case set_miles.

Decorators and Properties

These days its very common to see the $\ensuremath{\mathsf{property}}$ function used via a decorator.

Here's another version of our Car class that works as before but now uses decorators to set up the properties

```
class Car:

def __init__(self, miles=1000):
    self._miles = miles
    self._kms = miles * 1.61

@property
def miles(self):
    return self._miles

@property
def kms(self):
    return self._kms

@miles.setter
def miles(self, value):
    self._miles = value
    self._kms = value * 1.61

@kms.setter
def kms(self, value):
    self._kms = value * 1.61
```

We won't go through all the details here.

For further information you can refer to the $\underline{\text{descriptor documentation}}$.

Generators

A generator is a kind of iterator (i.e., it works with a next function).

We will study two ways to build generators: generator expressions and generator functions.

Generator Expressions

The easiest way to build generators is using *generator expressions*.

Just like a list comprehension, but with round brackets.

Here is the list comprehension:

```
singular = ('dog', 'cat', 'bird')
type(singular)

tuple

plural = [string + 's' for string in singular]
plural

['dogs', 'cats', 'birds']

type(plural)

list
```

And here is the generator expression

'cats'

```
singular = ('dog', 'cat', 'bird')
plural = (string + 's' for string in singular)
type(plural)

generator

next(plural)

'dogs'
```

```
next(plural)
```

'birds'

285

Since sum() can be called on iterators, we can do this

```
sum((x * x for x in range(10)))
```

The function sum() calls next() to get the items, adds successive terms.

In fact, we can omit the outer brackets in this case

```
sum(x * x for x in range(10))
285
```

Generator Functions

The most flexible way to create generator objects is to use generator functions.

Let's look at some examples.

Example 1

Here's a very simple example of a generator function

```
def f():
    yield 'start'
    yield 'middle'
    yield 'end'
```

It looks like a function, but uses a keyword yield that we haven't met before.

Let's see how it works after running this code

```
type(f)

function

gen = f()
gen

<generator object f at 0x117ce5350>

next(gen)

'start'

next(gen)

'middle'

next(gen)

'end'

next(gen)
```

 ${\tt StopIteration}$ Traceback (most recent call <ipython-input-107-6e72e47198db> in <module>
----> 1 next(gen) StopIteration:

The generator function f() is used to create generator objects (in this case gen).

Generators are iterators, because they support a next method.

The first call to next (gen)

- Executes code in the body of f() until it meets a yield statement.
- Returns that value to the caller of next (gen).

The second call to next (gen) starts executing from the next line

```
def f():
                 'start'
'middle'
'end'
      yield
yield
yield
                                # This line!
```

and continues until the next yield statement.

At that point it returns the value following yield to the caller of next(gen), and so on.

When the code block ends, the generator throws a StopIteration error.

Example 2

Our next example receives an argument x from the caller

```
def g(x):
    while x < 100:
       yield x
        x = x * x
```

```
Let's see how it works
 g
 <function __main__.g(x)>
 gen = g(2)
type(gen)
 generator
 next(gen)
 next(gen)
 4
```

```
next(gen)
```

16

```
next(gen)
```

```
StopIteration
                                          Traceback (most recent call
<ipython-input-115-6e72e47198db> in <module>
   -> 1 next(gen)
```

The call gen = g(2) binds gen to a generator.

Inside the generator, the name x is bound to 2.

When we call next(gen)

• The body of g() executes until the line yield x, and the value of x is returned.

Note that value of x is retained inside the generator.

When we call next (gen) again, execution continues from where it left off

```
def g(x):
    while x < 100:
        yield x
        x = x * x # execution continues from here</pre>
```

When x < 100 fails, the generator throws a StopIteration error.

Incidentally, the loop inside the generator can be infinite

```
def g(x):
    while 1:
        yield x
        x = x * x
```

Advantages of Iterators

What's the advantage of using an iterator here?

Suppose we want to sample a binomial(n,0.5).

One way to do it is as follows

```
import random
n = 10000000
draws = [random.uniform(0, 1) < 0.5 for i in range(n)]
sum(draws)</pre>
```

5000587

But we are creating two huge lists here, range(n) and draws.

This uses lots of memory and is very slow.

If we make n even bigger then this happens

```
n = 100000000
draws = [random.uniform(0, 1) < 0.5 for i in range(n)]</pre>
```

We can avoid these problems using iterators.

Here is the generator function

```
def f(n):
    i = 1
    while i <= n:
        yield random.uniform(0, 1) < 0.5
        i += 1</pre>
```

Now let's do the sum

```
n = 10000000
draws = f(n)
draws
```

<generator object f at 0x117d05900>

```
sum(draws)
```

In summary, iterables

- avoid the need to create big lists/tuples, and
- provide a uniform interface to iteration that can be used transparently in for loops

Recursive Function Calls

This is not something that you will use every day, but it is still useful — you should learn it at some stage.

Basically, a recursive function is a function that calls itself.

For example, consider the problem of computing x_t for some t when

$$x_{t+1} = 2x_t, \quad x_0 = 1$$

Obviously the answer is 2^t .

We can compute this easily enough with a loop

```
def x_loop(t):
    x = 1
    for i in range(t):
        x = 2 * x
    return x
```

We can also use a recursive solution, as follows

```
def x(t):
    if t == 0:
        return 1
    else:
        return 2 * x(t-1)
```

What happens here is that each successive call uses it's own frame in the stack

- a frame is where the local variables of a given function call are held
- stack is memory used to process function calls
 - o a First In Last Out (FILO) queue

This example is somewhat contrived, since the first (iterative) solution would usually be preferred to the recursive solution.

We'll meet less contrived applications of recursion later on.

Exercises

Exercise 1

The Fibonacci numbers are defined by

$$x_{t+1} = x_t + x_{t-1}, \quad x_0 = 0, \ x_1 = 1$$

The first few numbers in the sequence are 0, 1, 1, 2, 3, 5, 8, 13, 21, 34, 55.

Write a function to recursively compute the *t*-th Fibonacci number for any *t*.

Exercise 2

Complete the following code, and test it using $\underline{\text{this csv file}}$, which we assume that you've put in your current working directory

```
def column_iterator(target_file, column_number):
    """A generator function for CSV files.
    When called with a file name target_file (string) and column number
    column_number (integer), the generator function returns a generator
    that steps through the elements of column column_number in file
    target_file.
    # put your code here

dates = column_iterator('test_table.csv', 1)

for date in dates:
    print(date)
```

Exercise 3

Suppose we have a text file numbers.txt containing the following lines

```
prices
3
8
7
21
```

Using try – except, write a program to read in the contents of the file and sum the numbers, ignoring lines without numbers.

Solutions

Exercise 1

Here's the standard solution

```
def x(t):
    if t == 0:
        return 0
    if t == 1:
        return 1
    else:
        return x(t-1) + x(t-2)
```

Let's test it

```
print([x(i) for i in range(10)])
[0, 1, 1, 2, 3, 5, 8, 13, 21, 34]
```

Exercise 2

One solution is as follows

```
Date 2009-05-21 2009-05-19 2009-05-18 2009-05-14 2009-05-12 2009-05-12 2009-05-11
```

Exercise 3

Let's save the data first

```
%file numbers.txt
prices
3
8
```

Overwriting numbers.txt

```
f = open('numbers.txt')

total = 0.0
for line in f:
    try:
        total += float(line)
    except ValueError:
        pass

f.close()
print(total)
```

39.0

Debugging

"Debugging is twice as hard as writing the code in the first place. Therefore, if you write the code as cleverly as possible, you are, by definition, not smart enough to debug it." – Brian Kernighan

Overview

Are you one of those programmers who fills their code with print statements when trying to debug their programs?

Hey, we all used to do that.

(OK, sometimes we still do that...)

But once you start writing larger programs you'll need a better system.

Debugging tools for Python vary across platforms, IDEs and editors.

Here we'll focus on Jupyter and leave you to explore other settings.

We'll need the following imports

```
import numpy as np
import matplotLib.pyplot as plt
%matplotLib inline
```

Let's consider a simple (and rather contrived) example

```
def plot_log():
    fig, ax = plt.subplots(2, 1)
    x = np.linspace(1, 2, 10)
    ax.plot(x, np.log(x))
    plt.show()
plot_log() # Call the function, generate plot
```

AttributeError Traceback (most recent call last)
<ipython-input-2-c32a2280f47b> in <module>
5 plt.show()
6
----> 7 plot_log() # Call the function, generate plot
<ipython-input-2-c32a2280f47b> in plot_log()
2 fig, ax = plt.subplots(2, 1)
3 x = np.linspace(1, 2, 10)
----> 4 ax.plot(x, np.log(x))
5 plt.show()

.../../_images/debugging_1_1.png

This code is intended to plot the \log function over the interval [1, 2].

But there's an error here: plt.subplots(2, 1) should be just plt.subplots().

AttributeError: 'numpy.ndarray' object has no attribute 'plot'

(The call plt.subplots (2, 1) returns a NumPy array containing two axes objects, suitable for having two subplots on the same figure)

The traceback shows that the error occurs at the method call ax.plot(x, np.log(x)).

The error occurs because we have mistakenly made ax a NumPy array, and a NumPy array has no plot method.

But let's pretend that we don't understand this for the moment.

We might suspect there's something wrong with ax but when we try to investigate this object, we get the following exception:

```
ax

NameError Traceback (most recent call last)
<ipython-input-3-b00e77935981> in <module>
----> 1 ax

NameError: name 'ax' is not defined
```

The problem is that ax was defined inside $plot_log()$, and the name is lost once that function terminates.

Let's try doing it a different way.

We run the first cell block again, generating the same error $% \left(1\right) =\left(1\right) \left(1\right)$

```
def plot_log():
    fig, ax = plt.subplots(2, 1)
    x = np.linspace(1, 2, 10)
    ax.plot(x, np.log(x))
    plt.show()
plot_log() # Call the function, generate plot
```

AttributeError: 'numpy.ndarray' object has no attribute 'plot'

```
.../../_images/debugging_3_1.png
```

But this time we type in the following cell block

```
%debug
```

You should be dropped into a new prompt that looks something like this

```
ipdb>
```

(You might see pdb\> instead)

Now we can investigate the value of our variables at this point in the program, step forward through the code, etc.

For example, here we simply type the name ax to see what's happening with this object:

It's now very clear that ax is an array, which clarifies the source of the problem.

To find out what else you can do from inside ipdb (or pdb), use the online help

```
ipdb> h
Documented commands (type help <topic>):
E0F
                                                         tbreak
                  continue
                           exit
                                         pdoc
                                                restart
                                                                  whatis
alias cl
                                   list
                                                         unalias
                 d
                           h
                                         pinfo
                                                return
                                                                  where
args
b
                 debua
                           heln
      clear
                                   n
                                          pp
                                                 run
                                                         unt
                           ignore next
       commands
                 disable
                                                         until
                                         q
break condition down
                                         quit
                                                step
                                                         up
                                   р
Miscellaneous help topics:
exec pdb
Undocumented commands:
retval rv
ipdb> h c
c(ont(inue))
Continue execution, only stop when a breakpoint is encountered.
```

Setting a Break Point

The preceding approach is handy but sometimes insufficient.

Consider the following modified version of our function above

```
def plot_log():
    fig, ax = plt.subplots()
    x = np.logspace(1, 2, 10)
    ax.plot(x, np.log(x))
    plt.show()

plot_log()
```

```
.../../_images/debugging_4_0.png
```

Here the original problem is fixed, but we've accidentally written np.logspace(1, 2, 10) instead of np.linspace(1, 2, 10).

Now there won't be any exception, but the plot won't look right.

To investigate, it would be helpful if we could inspect variables like \times during execution of the function.

To this end, we add a "break point" by inserting breakpoint() inside the function code block

```
def plot_log():
    breakpoint()
    fig, ax = plt.subplots()
    x = np.logspace(1, 2, 10)
    ax.plot(x, np.log(x))
    plt.show()

plot_log()
```

Now let's run the script, and investigate via the debugger

We used n twice to step forward through the code (one line at a time).

Then we printed the value of x to see what was happening with that variable.

To exit from the debugger, use ${\bf q}$.

Other Useful Magics

In this lecture, we used the %debug IPython magic.

There are many other useful magics:

- %precision 4 sets printed precision for floats to 4 decimal places
- %whos gives a list of variables and their values
- %quickref gives a list of magics

The full list of magics is <u>here</u>.

Troubleshooting

Note

This page is for readers experiencing errors when running the code from the lectures.

Fixing Your Local Environment

The basic assumption of the lectures is that code in a lecture should execute whenever

- 1. it is executed in a Jupyter notebook and
- 2. the notebook is running on a machine with the latest version of Anaconda Python.

You have installed Anaconda, haven't you, following the instructions in <u>this lecture</u>?

Assuming that you have, the most common source of problems for our readers is that their Anaconda distribution is not up to date.

Here's a useful article on how to update Anaconda.

Another option is to simply remove Anaconda and reinstall.

You also need to keep the external code libraries, such as **QuantEcon.py** up to date.

For this task you can either

- use pip install --upgrade quantecon on the command line, or
- execute !pip install \--upgrade quantecon within a Jupyter notebook.

If your local environment is still not working you can do two things.

First, you can use a remote machine instead, by clicking on the Launch Notebook icon available for each lecture



Second, you can report an issue, so we can try to fix your local set up.

We like getting feedback on the lectures so please don't hesitate to get in touch.

Reporting an Issue

One way to give feedback is to raise an issue through our <u>issue tracker</u>.

Please be as specific as possible. Tell us where the problem is and as much detail about your local set up as you can provide.

Another feedback option is to use our <u>discourse forum</u>.

Finally, you can provide direct feedback to contact@quantecon.org

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