Python for Education: Computational Methods for Nonlinear Systems

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(Dated: October 26, 2018)

We describe a novel, interdisciplinary, computational methods course that uses Python and associated numerical and visualization libraries to enable students to implement simulations for a number of different course modules. Problems in complex networks, biomechanics, pattern formation, and gene regulation are highlighted to illustrate the breadth and flexibility of Python-powered computational environments.

Introduction

Computational science and engineering (CSE) involves the integration of a number of different techniques, requiring expertise in data structures, algorithms, numerical analysis, programming methodologies, simulation, visualization, data analysis, and performance optimization. The CSE community has embraced Python as a platform for attacking a wide variety of research problems, in part because of Python’s support for easily gluing together tools from different domains to solve complex problems. Teaching the theory and practice of CSE requires touching on all the subjects mentioned above, in the context of important and interesting scientific problems. Many of the same advantages that Python brings to CSE research make it useful for teaching too. Traditionally, courses have tended to focus more narrowly on particular aspects of CSE, such as numerical analysis, algorithms, or high-performance computing. In developing a new, broadly focused laboratory course in computational science, engineering and biology, we have sought to introduce students to the wide swath of techniques necessary to do effective research in CSE. Python and its many batteries serve remarkably well in this endeavor.

Computational methods for nonlinear systems is a graduate computational science laboratory course jointly developed and taught by us. We initiated course development in the summer of 2004 to support the curricular needs of the Cornell IGERT program in nonlinear systems, a broad and interdisciplinary graduate fellowship program aimed at introducing theoretical and computational techniques developed in the study of nonlinear and complex systems to a range of fields. The focal themes of the IGERT program span a number of areas - including complex networks, biological locomotion and manipulation, pattern formation, and gene regulation - broadly interpreted in the context of complex systems and nonlinear dynamics. These themes form the core of our course curriculum, augmented with other problems of interest arising in the fields of statistical mechanics, applied mathematics, and computer science.

The format of the course is somewhat unusual. As a computational laboratory course, it provides relatively little in the way of lectures: we prefer to have students learn by doing, rather than having us tell them how to do things. The course is autonomous, modular, and self-paced: students choose computational modules to work on from a large (and hopefully growing) suite of those available, and then proceed to implement relevant simulations and analyses as laid out in the exercise. We provide hints files to help the students along: these consist of documented skeletal code that the students are meant to flesh out. (In practice, we develop a module ourselves, document each of the relevant pieces using Python’s doctstrings, and then replace all the code bodies with the Python keyword pass so that the students can repopulate those code bodies themselves.) We have written several different visualization tools to provide visual feedback. We find these help to engage the students in new problems and are useful in code debugging.

Python is a useful language for teaching for several reasons (even though most of our incoming students have had no previous experience with Python). Its clean syntax enables students to learn the language quickly, and allows us to provide concise programming hints in our documented code fragments. Python’s dynamic typing and high-level, built-in datatypes enables students to get programs working quickly, without having to struggle with type declarations and compile-link-run loops. Since Python is interpreted, students can learn the language by executing and analyzing individual commands, and we can help them debug their programs by working with them in the interpreter.

One of the other key advantages that Python brings to scientific computing is the availability of many packages supporting numerical algorithms and visualization. While some of our exercises require development of algorithms from scratch, others rely on established numerical routines implemented in third-party libraries. It is of course important to understand the fundamentals of algorithms, error analysis, and algorithmic complexity, but it is also useful to know when and how to use existing solutions that have been developed by others. We make heavy use of the numpy$^1$ and scipy$^2$ packages, for construction of efficient arrays and for access to routines for generation of random numbers, integration of ordinary differential equations, root-finding, computation of eigenvalues, etc. We use matplotlib$^3$ for x-y plotting and histograms. We have written several visualization modules that we provide to students, based on the Python Imaging Library (PIL)$^4$, using PIL’s ImageDraw module to place graphics primitives within an image,
FIG. 1: Node and edge betweenness in a model of small-world networks. Nodes (red dots) are connected by undirected edges (black lines). Betweenness measures how central each node and edge is to the shortest network paths connecting any two nodes. In this plot, node diameter and edge thickness are proportional to node and edge betweenness, respectively. (Our simple graph visualization tool uses the Python Imaging Library.)

and the ImageTk module to paste an image into a Tk window for real-time animation. We recommend the use of the ipython interpreter, which facilitates exploration by students. And we have used VPython to generate three-dimensional animations to accompany some of our modules.

Course modules

There are too many course modules to describe in detail here, and we refer interested readers to our course website for information on all the modules, as well as access to problems, hints, and answers. (Many of the exercises have also been incorporated into a new textbook written by one of us.) Here, we highlight a few of the modules, in order to illustrate both the breadth of science that can be usefully taught with Python and variety of tools and techniques that Python can bring to bear on such problems.

Small world networks

The study of complex networks has flourished over the last several years as researchers have discovered commonalities among networked structures that arise in diverse fields such as biology, ecology, sociology, and computer science. An interesting property found in many complex networks is exemplified in the popular notion of “six degrees of separation”, which suggests that any two people on earth are connected through at most roughly five intermediate acquaintances. Duncan Watts and Steve Strogatz developed a simple model of random networks that demonstrate this “small-world” property. Our course module enables students to construct small-world networks and to examine how the average path length connecting two nodes decreases rapidly as random, long-range bonds are introduced into a network consisting initially of only short-ranged bonds (Figure 1).

Computationally, this module introduces students to data structures for the representation of undirected graphs, object-oriented encapsulation of those data structures, and graph traversal algorithms. Python makes the development of an undirected graph data structure exceedingly simple, a point made long ago by Python creator Guido van Rossum in one of his early essays on Python. In an undirected graph, nodes are connected to other nodes by edges. A simple way to implement this is to combine the two cornerstones of container-based programming in Python: lists and dictionaries. In our UndirectedGraph class, a dictionary of network neighbor connections (a neighbor dictionary) maps a node identifier to a list of other nodes to which the reference node is connected. Because the graph edges are undirected, we duplicate the connection information for each node: if an edge is added connecting node 1 and node 2, the neighbor dictionary must be updated so that node 2 is added to node 1’s list of neighbors, and vice versa.

We can of course hide the details of adding edges inside an AddEdge method defined on an UndirectedGraph class:

class UndirectedGraph:
    # ...
    def AddEdge(self, node1, node2):
        self.AddNode(node1)
In the small-world networks exercise, we choose to label nodes simply by integers, but Python’s dynamic typing does not require this. If we were playing the “Kevin Bacon game” of searching for shortest paths in actor collaboration networks, we could use our code above to build a graph connecting names of actors (encoded as strings). This dynamic typing allows for significant code reuse (as described below in the section on Percolation). And it is worth mentioning that, while our `UndirectedGraph` class is exceedingly simple and built to support only the analyses relevant to our course module, the same basic principles are at work in a much more comprehensive, Python-based, graph construction and analysis package - named NetworkX - that has been developed at Los Alamos National Labs. [12]

Percolation

Percolation is the study of how objects become connected (or disconnected) as they are randomly wired together (or cut apart). Percolation is an important and classic problem in the study of phase transitions, and has practical relevance as well: considerable interest over the years in percolation phenomena has come from the oil and gas industry, for example, where one is interested in extracting a fluid through a network of pores in rock.

Although percolation is traditionally studied on regular lattices, it is a problem more generally applicable to arbitrary networks, and in fact, we are able to reuse some of the code developed in the small-world networks module to support the study of percolation. As noted above, Python’s dynamic typing makes our definition of a node in a graph very flexible; in a percolation problem on a lattice, we can reuse our `UndirectedGraph` class described previously by making node identifiers be lattice index tuples \((i, j)\). We can thus easily make an instance of bond percolation on a 2D square lattice of size \(L\) (with periodic boundary conditions) and bond fraction \(p\):

```python
def MakeSquareBondPercolation(L, p):
    g = UndirectedGraph()
    for i in range(L):
        for j in range(L):
            g.AddNode((i, j))
            if random.random() < p:
                g.AddEdge((i, j), ((i+1)%L, j))
            if random.random() < p:
                g.AddEdge((i, j), (i, (j+1)%L))
    return g
```

Instances of percolation networks generated by this procedure are illustrated in Figure 2. Students use breadth-first search to identify all connected clusters in such a network, and our PIL-based visualization tool colors each separate cluster distinctly, taking as input all nodes of each cluster.

The concept of universality of phase transitions is also introduced: despite their microscopic differences, site-percolation on a 2D hexagonal lattice and bond-percolation on a 2D square lattice are indistinguishable from each other on long length scales, exhibiting the same critical behavior (e.g., scaling exponents). Scaling collapses are a useful construct for revealing the universality of phase transitions, and typically involve transforming the \(x\) and \(y\) axes in specified ways to get disparate data sets to “collapse” onto one universal scaling form. With Python, we can support such scaling collapses very flexibly by using the built-in `eval()` function that evaluates expressions encoded as strings. Rather than hard-coding particular functional forms for scaling collapses, arbitrary mathematical expressions can be simply encoded and evaluated.

Biomechanics: The Walker

Research in biomechanics aims to understand how living beings move, and robotics and prosthetics are two important technological areas that can benefit from advances in the field. While much research in robotics is focused on active sensing and control, Andy Ruina and collaborators have been interested in passive biolocomotive systems, which are more properly understood as dynamical systems than as control systems. The “simplest walking model” of Garcia et al. [13] provides the basis of our Walker module. This model consists of a pair of legs connected at the hip.
(a double pendulum), walking down an inclined ramp under the influence of gravity, with a heelstrike that imparts angular momentum to the Walker as the swing leg strikes the floor and becomes the stance leg. As a warmup, students integrate the equation of motion for a single pendulum under gravity, and compute the period of the motion as a function of the initial pendulum angle.

The Pendulum and Walker modules introduce several important scientific and computational aspects. Ordinary differential equations (ODEs) describing the time evolution of the Pendulum and Walker need to be integrated forward in time. In the context of the simpler Pendulum, we highlight the properties of accuracy, fidelity, and stability in numerical integration, having students explore errors introduced by a finite time step $\Delta t$. We also highlight the need for event detection in many numerical integration problems. In the Walker, for example, a heelstrike occurs when the swing leg hits the floor. Accurately solving for the heelstrike collision involves transforming to a new set of integration variables, where an appropriate combination of the pendulum angles becomes the independent variable, and time a dependent variable. We then integrate backwards in angle to find the time at which the heelstrike occurred. We use the `scipy.integrate.odeint` function to execute these integrations, providing a function $dydt$ that evaluates the instantaneous time derivative of the Walker state vector field $\dot{\mathbf{y}}$ and a function $dzdc$ that evaluates the instantaneous time derivative of the transformed system for heelstrike detection (where the independent variable is the “collision variable” $c = \phi - 2\theta$).

```python
def dydt(self, y,t):
    theta, thetaDot, phi, phiDot = y
    thetaDotdot = scipy.sin(theta-self.gamma)
    phiDotdot = thetaDotdot + \
        (thetaDot**2)*sin(phi)-cos(theta-self.gamma)*sin(phi)
    return [thetaDot,thetaDotdot,phiDot,phiDotdot]

self.trajectory = scipy.integrate.odeint(self.dydt, self.GetStateVector(),
    timepoints)

def dzdc(self, z, c):
    theta,thetaDot,phi,phiDot,t = z
    y = array([theta, thetaDot, phi, phiDot])
    thetaDot, thetaDotdot, phiDot, phiDotdot = self.dydt(y, t)
    cDot = phiDot - 2.*thetaDot
    return [thetaDot/cDot,thetaDotdot/cDot,phiDot/cDot,phiDotdot/cDot,cDot, 1./cDot]

z = scipy.integrate.odeint(self.dzdc, [y[0],y[1],y[2],y[3],t],
    scipy.array([self.CollisionCondition(), 0.1]))
```
FIG. 3: Snapshot in the perambulation of the Walker. The model consists of a pair of coupled pendula (legs) walking down a ramp. The stance leg (red) remains fixed with respect to the floor, while the swing leg (orange) swings forward. Once the swing leg hits the floor ahead of the stance leg (heelstrike), the two legs switch roles. Real-time animation of the Walker is accomplished using VPython.

The Walker exhibits an interesting period-doubling route to chaos as the slope of the inclined ramp is increased. Simple periodic walking is stable for small ramp angles, but becomes unstable to a period-two gait at a critical angle. The period-two orbit bifurcates to a period-four gait, etc., with increasing angle, culminating in a chaotic walk. (The chaos is, however, remarkably subtle, as is also true in systems like dripping faucets.) A snapshot of the Walker is shown in Figure 3. Other modules in our course enable students to study in considerable more detail these sorts of period-doubling bifurcations and chaotic dynamics in iterated one-dimensional maps.

Pattern formation in cardiac dynamics

Pattern formation is ubiquitous in spatially-extended nonequilibrium systems. Many patterns involve regular, periodic phenomena in space and time, but equally important are localized coherent structures that break or otherwise interrupt these periodic structures. Patterns lie at the root of much activity in living tissues: the regular beating of the human heart is perhaps our most familiar reminder of the spatiotemporal rhythmicity of biological patterns. Cardiac tissue is an excitable medium: rhythmic voltage pulses, initiated by the heart’s pacemaker cells (in the sinoatrial node), spread as a wave through the rest of the heart inducing the heart muscle to contract, thereby pumping blood in a coherent fashion. In some situations, however, this regular beating can become interrupted by the presence of spiral waves in the heart’s electrical activity (see Figure 4). These spiral waves generate voltage pulses on their own, disrupting the coordinated rhythm of the normal heart, leading to cardiac arrhythmia. A simple model of cardiac dynamics - the two-dimensional FitzHugh-Nagumo equations[14, 15] - is introduced in this course module, which we developed in conjunction with Niels Otani. The FitzHugh-Nagumo model describes the coupled time evolution of two fields, the transmembrane potential $V$ and the recovery variable $W$ (given parameters $\epsilon$, $\gamma$ and $\beta$):

\[
\frac{\partial V}{\partial t} = \nabla^2 V + \frac{1}{\epsilon} (V - V^3/3 - W) \quad \frac{\partial W}{\partial t} = \epsilon (V - \gamma W + \beta)
\]

Fixed point solutions to the FitzHugh-Nagumo equations are found by root-finding, which we accomplish using the `brentq` function in scipy:

```python
def FindFixedPoint(gamma, beta):
    f = lambda v, gamma, beta: (v-(v**3)/3.)-(1./gamma)*(v+beta)
    vstar = scipy.optimize.brentq(f, -2., 2., args=(gamma, beta))
    wstar = (1./gamma)*(vstar+beta)
    return vstar, wstar
```

We also introduce students to finite-difference techniques for computing spatial derivatives in the solution of partial differential equations (PDEs). Numpy arrays are used to represent the $V$ and $W$ fields of the FitzHugh-Nagumo model, and an important operation is the computation of the laplacian of the voltage field, $\nabla^2 V(x, y)$. We introduce the stencil notation for characterizing finite-difference approximations to $\nabla^2 V$, and use a combination of array arithmetic and array slicing to compactly and efficiently compute the derivative on the interior (non-boundary) cells of the
FIG. 4: Snapshots in the time evolution of the FitzHugh-Nagumo model of cardiac dynamics. The transmembrane voltage $V$ is depicted via a grayscale map (higher voltages in lighter grays). Spiral waves in the voltage field can lead to cardiac arrhythmias by disrupting the normal periodic rhythm generated by the sinoatrial node. (Right) Users can administer local voltage pulses (white rectangle) to trigger spiral wave formation or to shock the arrhythmic heart back to a normal beating state.

simulation domain. Students are asked to implement two different approximations to the laplacian operator (a five-point and nine-point stencil), and compare their effects on the detailed form of propagating electrical waves. The computation of the five-point stencil is shown here:

```python
def del2_5(a, dx):
    del2 = scipy.zeros(a.shape, float)
    del2[1:-1, 1:-1] = (a[1:-1,2:] + a[1:-1,:-2] + a[2:,1:-1] + a[:-2,1:-1] - 4.*a[1:-1,1:-1])/(dx*dx)
    return del2
```

We provide an animation tool that we have written, based on PIL and Tkinter, that enables students to update the display of the voltage field $V$ at every time step, and to use the mouse to introduce local “shocks” to the system. (See Figure 4.) These shocks are both useful in initiating spiral waves and in resetting the global electrical state of the system as a defibrillator might do. Optional extensions to the module, developed by our collaborator Otani, allow for simulations of spontaneous pacemakers, dead regions of tissue, and more complex heart-chamber geometries, by letting the various parameters of the model become spatially-varying fields themselves (again implemented via numpy arrays).

**Gene regulation and the Repressilator**

Gene regulation describes a set of processes by which the expression of genes within a living cell - i.e., their transcription to messenger RNA and ultimately their translation to protein - is controlled. While modern genome sequencing has provided great insights into the constituent parts (genes and proteins) of many organisms, much less is known about how those parts of turned on and off and mixed and matched in different contexts: how is that a brain cell and a hair cell, for example, can derive from the same genomic blueprint but have such different properties?

The Repressilator is a relatively simple synthetic gene regulatory network developed by Michael Elowitz and Stan Leibler\[16\]. Its name derives from its use of three repressor proteins arranged to form a biological oscillator: these three repressors act in a manner akin to the “rock-paper-scissors” game where TetR inhibits λ CI, which in turn inhibits LacI, which in turn inhibits TetR. A snapshot in the time evolution of the Repressilator is shown in Figure 5.

One of the important scientific and computational features that we emphasize in this module are the differences between stochastic and deterministic representations of chemical reaction networks. (We first introduce these concepts in a warmup exercise, Stochastic Cells, in which students simulate a much simpler biochemical network: one
FIG. 5: Snapshot in the stochastic time evolution of the Repressilator. The state for this model consists of 15 components: 3 protein concentrations (back row), 3 mRNA concentrations (middle row), and 3 sets of promoter-binding states (front row): each promoter can be either unbound, singly-bound, or doubly-bound. At this instant, TetR (red) concentrations are high, leading to suppression of λ cI (yellow). Since λ cI is low, however, LacI (green) concentrations are allowed to grow. This will lead to the eventual suppression of TetR.

representation the binding and unbinding of two monomer molecules $M$ to form a single dimer $D$: $M + M \leftrightarrow D$.) We introduce students to Petri nets as a graphical notation for encoding such networks, and then have them, from the underlying Petri net representation, (a) synthesize differential equations describing the deterministic time evolution of the system, and (b) implement the Gillespie algorithm (a form of continuous time Monte Carlo) for stochastic simulation.\[17\] Gillespie’s “direct method” involves choosing a particular reaction and reaction time based on the instantaneous reaction rates. For the Repressilator, this can be done quite compactly using array operations within numpy/scipy:

class StochasticRepressilator (Repressilator):
    # ...
    def Step(self, dtmax):
        self.ComputeReactionRates()
        total_rate = sum(self.rates)
        # get exponentially distributed time
        ran_time = -scipy.log(1.-random.random())/total_rate
        if ran_time > dtmax:
            return dtmax
        # get uniformly drawn rate in interval defined by total_rate
        ran_rate = total_rate*random.random()
        # find interval corresponding to random rate
        reac_index = len(self.rates) - sum(scipy.cumsum(self.rates) > ran_rate)
        reaction = self.reactions[reac_index]
        # execute specified reaction
        for chem, dchem in reaction.stoichiometry.items():
            chem.amount += dchem
        # return time at which reaction takes place
        return ran_time

Other modules

Our course consists of a number of other modules which we can only mention in passing here. As noted, there is a suite of problems introducing various aspects of chaos and bifurcations in iterated maps. There is also a suite of small modules exploring properties of random walks and extremal statistics. We have two exercises examining connections between statistical mechanics and computational complexity, by probing the nature of phase transitions in NP-complete problems such as 3SAT. A random matrix theory module examines the nature of universality of eigenvalue distributions, and two other modules explore the thermodynamics of large collective systems (the Ising model of simple magnets, and the molecular dynamics of large numbers of atoms).
We continue to look for new problems to add to this collection, and for collaborators interested in contributing their scientific and computational expertise to this endeavor. (Please contact us if you have ideas for interesting modules.) Our goal is to provide a hands-on introduction in scientific computing, and it is our hope that this course can help serve a number of educational objectives in the part of a larger curriculum in computational science and engineering.

Acknowledgments

We thank our colleagues who have helped us develop computational modules and have given us useful feedback: Steve Strogatz, Andy Ruina, Niels Otani, Bart Selman, Carla Gomes, and John Guckenheimer. We also thank all the students who have taken our course and have helped us work the bugs out of exercises and solutions. Funding from the NSF IGERT program (award NSF DGE-0333366) and NSF DMR-0218475 helped support some initial development of course modules.

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