The moment-to-moment processing of information by the nervous system involves the propagation and interaction of electrical and chemical signals that are distributed in space and time. Biologically realistic modeling is needed to test hypotheses about the mechanisms that govern these signals and how nervous system function emerges from the operation of these mechanisms. The NEURON simulation program provides a powerful and flexible environment for implementing such models of individual neurons and small networks of neurons. It is particularly useful when membrane potential is nonuniform and membrane currents are complex. We present the basic ideas that would help informed users make the most efficient use of NEURON.

1 Introduction

NEURON (Hines, 1984, 1989, 1993, 1994) provides a powerful and flexible environment for implementing biologically realistic models of electrical and chemical signaling in neurons and networks of neurons. This article describes the concepts and strategies that have guided the design and implementation of this simulator, with emphasis on those features that are particularly relevant to its most efficient use.

1.1 The Problem Domain. Information processing in the brain results from the spread and interaction of electrical and chemical signals within and among neurons. This involves nonlinear mechanisms that span a wide range of spatial and temporal scales (Carnevale & Rosenthal, 1992) and are constrained to operate within the intricate anatomy of neurons and their interconnections. Consequently the equations that describe brain mechanisms generally do not have analytical solutions, and intuition is not a reliable guide to understanding the working of the cells and circuits of the brain. Furthermore, these nonlinearities and spatiotemporal complexities are quite unlike those that are encountered in most nonbiological systems,
so the utility of many quantitative and qualitative modeling tools that were
developed without taking these features into consideration is severely lim-
ited.

NEURON is designed to address these problems by enabling both the
convenient creation of biologically realistic quantitative models of brain
mechanisms and the efficient simulation of the operation of these mecha-
nisms. In this context the term biological realism does not mean “infinitely
detailed.” Instead, it means that the choice of which details to include in
the model and which to omit are at the discretion of the investigator who
constructs the model, and not forced by the simulation program.

To the experimentalist, NEURON offers a tool for cross-validating data,
estimating experimentally inaccessible parameters, and deciding whether
known facts account for experimental observations. To the theoretician, it
is a means for testing hypotheses and determining the smallest subset of
anatomical and biophysical properties that is necessary and sufficient to
account for particular phenomena. To the student in a laboratory course,
it provides a vehicle for illustrating and exploring the operation of brain
mechanisms in a simplified form that is more robust than the typical “wet
lab” experiment. For the experimentalist, theoretician, and student alike, a
powerful simulation tool such as NEURON can be an indispensable aid to
developing the insight and intuition that is needed if one is to discover the
order hidden within the intricacy of biological phenomena, the order that
transcends the complexity of accident and evolution.

1.2 Experimental Advances and Quantitative Modeling. Experimental
advances drive and support quantitative modeling. Over the past two
decades, the field of neuroscience has seen striking developments in experi-
mental techniques, among them the following:

- High-quality electrical recording from neurons in vitro and in vivo
  using patch clamp.
- Multiple impalements of visually identified cells.
- Simultaneous intracellular recording from paired pre- and postsynap-
tic neurons.
- Simultaneous measurement of electrical and chemical signals.
- Multisite electrical and optical recording.
- Quantitative analysis of anatomical and biophysical properties from
  the same neuron.
- Photolesioning of cells.
- Photorelease of caged compounds for spatially precise chemical stim-
  ulation.
• New drugs such as channel blockers and receptor agonists and antagonists.
• Genetic engineering of ion channels and receptors.
• Analysis of messenger RNA and biophysical properties from the same neuron.
• “Knockout” mutations.

These and other advances are responsible for impressive progress in the definition of the molecular biology and biophysics of receptors and channels; the construction of libraries of identified neurons and neuronal classes that have been characterized anatomically, pharmacologically, and biophysically; and the analysis of neuronal circuits involved in perception, learning, and sensorimotor integration.

The result is a data avalanche that catalyzes the formulation of new hypotheses of brain function, while at the same time serving as the empirical basis for the biologically realistic quantitative models that must be used to test these hypotheses. Following are some examples from the large list of topics that have been investigated through the use of such models:

• The cellular mechanisms that generate and regulate chemical and electrical signals (Destexhe, Contreras, Steriade, Sejnowski, & Huguenard, 1996; Jaffe et al., 1994).
• Drug effects on neuronal function (Lytton & Sejnowski, 1992).
• Presynaptic (Lindgren & Moore, 1989) and postsynaptic (Destexhe & Sejnowski, 1995; Traynelis, Silver, & Cull-Candy, 1993) mechanisms underlying communication between neurons.
• Integration of synaptic inputs (Bernander, Douglas, Martin, & Koch, 1991; Cauller & Connors, 1992).
• Action potential initiation and conduction (Hausser, Stuart, Racca, & Sakmann, 1995; Hines & Shrager, 1991; Mainen, Joerges, Huguenard, & Sejnowski, 1995).
• Cellular mechanisms of learning (Brown, Zador, Mainen, & Claiborne, 1992; Tsai, Carnevale, & Brown, 1994a).
• Cellular oscillations (Destexhe, Babloyantz, & Sejnowski, 1993a; Lytton, Destexhe, & Sejnowski, 1996).
• Thalamic networks (Destexhe, McCormick, & Sejnowski, 1993b; Destexhe, Contreras, Sejnowski, & Steriade, 1994).
• Neural information encoding (Hsu et al., 1993; Mainen & Sejnowski, 1995; Softky, 1994).
2 Overview of NEURON

NEURON is intended to be a flexible framework for handling problems in which membrane properties are spatially inhomogeneous and where membrane currents are complex. Since it was designed specifically to simulate the equations that describe nerve cells, NEURON has three important advantages over general-purpose simulation programs. First, the user is not required to translate the problem into another domain, but instead is able to deal directly with concepts that are familiar at the neuroscience level. Second, NEURON contains functions that are tailored specifically for controlling the simulation and graphing the results of real neurophysiological problems. Third, its computational engine is particularly efficient because of the use of special methods and tricks that take advantage of the structure of nerve equations (Hines, 1984; Mascagni, 1989).

The general domain of nerve simulation, however, is still too large for any single program to deal optimally with every problem. In practice, each program has its origin in a focused attempt to solve a restricted class of problems. Both speed of simulation and the ability of the user to maintain conceptual control degrade when any program is applied to problems outside the class for which it is best suited.

NEURON is computationally most efficient for problems that range from parts of single cells to small numbers of cells in which cable properties play a crucial role. In terms of conceptual control, it is best suited to tree-shaped structures in which the membrane channel parameters are approximated by piecewise linear functions of position. Two classes of problems for which it is particularly useful are those in which it is important to calculate ionic concentrations and those where one needs to compute the extracellular potential just next to the nerve membrane. It is especially capable for investigating new kinds of membrane channels since they are described using a high-level neuron model description language (NMODL) (Moore & Hines, 1996), which allows the expression of models in terms of kinetic schemes or sets of simultaneous differential and algebraic equations. To maintain efficiency, user-defined mechanisms in NMODL are automatically translated into C, compiled, and linked into the rest of NEURON.

The flexibility of NEURON comes from a built-in object-oriented interpreter, which is used to define the morphology and membrane properties of neurons, control the simulation, and establish the appearance of a graphical interface. The default graphical interface is suitable for exploratory simulations involving the setting of parameters, control of voltage and current stimuli, and graphing variables as a function of time and position.

Simulation speed is excellent since membrane voltage is computed by an implicit integration method optimized for branched structures (Hines, 1984). The performance of NEURON degrades very slowly with increased complexity of morphology and membrane mechanisms, and it has been applied to very large network models: $10^4$ cells with six compartments each,
for a total of $10^6$ synapses in the net (T. Sejnowski, personal communication, March 29, 1996).

3 Mathematical Basis

Strategies for numerical solution of the equations that describe chemical and electrical signaling in neurons have been discussed in many places. Elsewhere we have briefly presented an intuitive rationale for the most commonly used methods (Hines & Carnevale, 1995). Here we start from this base and proceed to address those aspects that are most pertinent to the design and application of NEURON.

3.1 The Cable Equation. The application of cable theory to the study of electrical signaling in neurons has a long history, which is briefly summarized elsewhere (Rall, 1989). The basic computational task is to solve numerically the cable equation

$$\frac{\partial V}{\partial t} + I(V, t) = \frac{\partial^2 V}{\partial x^2},$$

(3.1)

which describes the relationship between current and voltage in a one-dimensional cable. The branched architecture typical of most neurons is incorporated by combining equations of this form with appropriate boundary conditions.

Spatial discretization of this partial differential equation is equivalent to reducing the spatially distributed neuron to a set of connected compartments. The earliest example of a multicompartmental approach to the analysis of dendritic electrotonus was provided by Rall (1964).

Spatial discretization produces a family of ordinary differential equations of the form

$$c_j \frac{dv_j}{dt} + i_{ion_j} = \sum_k \frac{v_k - v_j}{r_{jk}}.$$

(3.2)

Equation 3.2 is a statement of Kirchhoff’s current law, which asserts that net transmembrane current leaving the $j$th compartment must equal the sum of axial currents entering this compartment from all sources (see Figure 1). The left-hand side of equation 3.2 is the total membrane current, which is the sum of capacitive and ionic components. The capacitive component is $c_j \frac{dv_j}{dt}$, where $c_j$ is the membrane capacitance of the compartment. The ionic component $i_{ion}$ includes all currents through ionic channel conductances. The right-hand side of equation 3.2 is the sum of axial currents that enter this compartment from its adjacent neighbors. Currents injected through a microelectrode would be added to the right-hand side. The sign conventions for current are as follows: outward transmembrane current is positive; axial
current flow into a region is positive; positive injected current drives $v_j$ in a positive direction.

Equation 3.2 involves two approximations. First, axial current is specified in terms of the voltage drop between the centers of adjacent compartments. The second approximation is that spatially varying membrane current is represented by its value at the center of each compartment. This is much less drastic than the often-heard statement that a compartment is assumed to be “isopotential.” It is far better to picture the approximation in terms of voltage varying linearly between the centers of adjacent compartments. Indeed, the linear variation in voltage is implicit in the usual description of a cable in terms of discrete electrical equivalent circuits.

If the compartments are of equal size, it is easy to use Taylor’s series to show that both of these approximations have errors proportional to the square of compartment length. Thus, replacing the second partial derivative by its central difference approximation introduces errors proportional to $\Delta x^2$, and doubling the number of compartments reduces the error by a factor of four.

It is often not convenient for the size of all compartments to be equal. Unequal compartment size might be expected to yield simulations that are only first-order accurate. However, comparison of simulations in which unequal compartments are halved or quartered in size generally reveals a
second-order reduction of error. A rough rule of thumb is that simulation error is proportional to the square of the size of the largest compartment.

The first of two special cases of equation 3.2 that we wish to discuss allows us to recover the usual parabolic differential form of the cable equation. Consider the interior of an unbranched cable with constant diameter. The axial current consists of two terms involving compartments with the natural indices \( j-1 \) and \( j+1 \), that is,

\[
e_j \frac{dv_j}{dt} + i_{ion,j} = \frac{v_{j-1} - v_j}{r_{j-1,j}} + \frac{v_{j+1} - v_j}{r_{j+1,j}}.
\]

If the compartments have the same length \( \Delta x \) and diameter \( d \), then the capacitance of a compartment is \( C_m \pi d \Delta x \), and the axial resistance is \( R_a \Delta x / \pi (d/2)^2 \). \( C_m \) is called the specific capacitance of the membrane, which is generally taken to be 1 \( \mu F/cm^2 \). \( R_a \) is the axial resistivity, which has different reported values for different cell classes (e.g., 35.4 \( \Omega \) cm for squid axon). Equation 3.2 then becomes

\[
C_m \frac{dv_j}{dt} + i_j = \frac{d}{4R_a} \frac{v_{j+1} - 2v_j + v_{j-1}}{\Delta x^2},
\]

where we have replaced the total ionic current \( i_{ion} \) with the current density \( i_j \). The right-hand term, as \( \Delta x \to 0 \), is just \( \partial^2 v / \partial x^2 \) at the location of the now-infinitesimal compartment \( j \).

The second special case of equation 3.2 allows us to recover the boundary condition. This is an important exercise since naive discretizations at the ends of the cable have destroyed the second-order accuracy of many simulations. Nerve boundary conditions are that no axial current flows at the end of the cable; the end is sealed. This is implicit in equation 3.2, where the right-hand side consists of only the single term \( (v_{j-1} - v_j) / r_{j-1,j} \) when compartment \( j \) lies at the end of an unbranched cable.

### 3.2 Spatial Discretization in a Biological Context: Sections and Segments

Every nerve simulation program solves for the longitudinal spread of voltage and current by approximating the cable equation as a series of compartments connected by resistors (see Figure 4 and equation 3.2). The sum of all the compartment areas is the total membrane area of the whole nerve. Unfortunately, it is usually not clear at the outset how many compartments should be used. Both the accuracy of the approximation and the computation time increase as the number of compartments used to represent the cable increases. When the cable is “short,” a single compartment can be made to represent the entire cable adequately. For long cables or highly branched structures, it may be necessary to use a large number of compartments.

This raises the question of how best to manage all the parameters that exist within these compartments. Consider membrane capacitance, which
has a different value in each compartment. Rather than specify the capacitance of each compartment individually, it is better to deal in terms of a single specific membrane capacitance that is constant over the entire cell and have the program compute the values of the individual capacitances from the areas of the compartments. Other parameters such as diameter or channel density may vary widely over short distances, so the graininess of their representation may have little to do with numerically adequate compartmentalization.

Although NEURON is a compartmental modeling program, the specification of biological properties (neuron shape and physiology) has been separated from the numerical issue of compartment size. What makes this possible is the notion of a section, which is a continuous length of unbranched cable. Although each section is ultimately discretized into compartments, values that can vary with position along the length of a section are specified in terms of a continuous parameter that ranges from 0 to 1 (normalized distance). In this way, section properties are discussed without regard to the number of segments used to represent it. This makes it easy to trade off between accuracy and speed, and enables convenient verification of the numerical correctness of simulations.

Sections are connected together to form any kind of branched tree structure. Figure 2 illustrates how sections are used to represent biologically significant anatomical features. The top of this figure is a cartoon of a neuron with a soma that gives rise to a branched dendritic tree and an axon hillock connected to a myelinated axon. Each biologically significant component of this cell has its counterpart in one of the sections of the NEURON model, as shown in the bottom of Figure 2: the cell body (Soma), axon hillock (AH), myelinated internodes (Ii), nodes of Ranvier (Ni), and dendrites (Di).

Sections allow this kind of functional and anatomical parcellation of the cell to remain foremost in the mind of the person who constructs and uses a NEURON model.

To accommodate requirements for numerical accuracy, NEURON represents each section by one or more segments of equal length (see Figures 3 and 4). The number of segments is specified by the parameter nseg, which can have a different value for each section.

At the center of each segment is a node, the location where the internal voltage of the segment is defined. The transmembrane currents over the entire surface area of a segment are associated with its node. The nodes of adjacent segments are connected by resistors.

It is crucial to realize that the location of the second-order correct voltage is not at the edge of a segment but rather at its center, that is, at its node. This is the discretization method employed by NEURON. To allow branching and injection of current at the precise ends of a section while maintaining second-order correctness, extra voltage nodes that represent compartments with zero area are defined at the section ends. It is possible to achieve second-order accuracy with sections whose end nodes have
Figure 2: (Top) Cartoon of a neuron indicating the approximate boundaries between biologically significant structures. The left-hand side of the cell body (Soma) is attached to an axon hillock (AH) that drives a myelinated axon (myelinated internodes $I_i$ alternating with nodes of Ranvier $N_i$). From the right-hand side of the cell body originates a branched dendritic tree ($D_i$). (Bottom) How sections would be employed in a NEURON model to represent these structures.

nonzero area compartments. However, the areas of these terminal compartments would have to be exactly half that of the internal compartments, and extra complexity would be imposed on administration of channel density at branch points.

Based on the position of the nodes, NEURON calculates the values of internal model parameters such as the average diameter, axial resistance, and compartment area that are assigned to each segment. Figures 3 and 4 show how an unbranched portion of a neuron, called a neurite (see Figure 3A), is represented by a section with one or more segments. Morphometric analysis generates a series of diameter measurements whose centers lie on the midline of the neurite (the thin axial line in Figure 3B). These measurements and the path lengths between their centers are the dimensions of the section, which can be regarded as a chain of truncated cones or frusta (see Figure 3C).

Distance along the length of a section is discussed in terms of the normalized position parameter $x$. That is, one end of the section corresponds to
Figure 3: (A) Cartoon of an unbranched neurite (thick lines) that is to be represented by a section in a NEURON model. Computer-assisted morphometry generates a file that stores successive measurements of diameter (thin circles) centered at x, y, z coordinates (thin crosses). (B) Each adjacent pair of diameter measurements (thick circles) becomes the parallel faces of a truncated cone or frustum, the height of which is the distance between the measurement locations. The outline of each frustum is shown with thin lines, and a thin centerline passes through the central axis of the chain of solids. (C) The centerline has been straightened so the faces of adjacent frusta are flush with each other. The scale underneath the figure shows the distance along the midline of the section in terms of the normalized position parameter x. The vertical dashed line at x = 0.5 divides the section into two halves of equal length. (D) Electrical equivalent circuit of the section as represented by a single segment (nseg = 1). The open rectangle includes all mechanisms for ionic (noncapacitive) transmembrane currents.

x = 0 and the other end to x = 1. In Figure 3C these locations are depicted as being on the left- and right-hand ends of the section. The locations of the nodes and the boundaries between segments are conveniently specified
Figure 4: How the neurite of Figure 3 would be represented by a section with two segments ($n_{seg} = 2$). Now the electrical equivalent circuit (bottom) has two nodes. The membrane properties attached to the first and second nodes are based on neurite dimensions and biophysical parameters over the $x$ intervals $[0, 0.5]$ and $[0.5, 1]$, respectively. The three axial resistances are computed from the cytoplasmic resistivity and neurite dimensions over the $x$ intervals $[0, 0.25]$, $[0.25, 0.75]$, and $[0.75, 1]$. In terms of this normalized position parameter. In general, a section has $n_{seg}$ segments that are demarcated by evenly spaced boundaries at intervals of $1/n_{seg}$. The nodes at the centers of these segments are located at $x = (2i - 1)/2n_{seg}$ where $i$ is an integer in the range $[1, n_{seg}]$. As we shall see later, $x$ is also used in specifying model parameters or retrieving state variables that are a function of position along a section (see section 4.4).

The special importance of $x$ and $n_{seg}$ lies in the fact that they free the user from having to keep track of the correspondence between segment number and position on the nerve. In early versions of NEURON, all nerve properties were stored in vector variables where the vector index was the segment number. Changing the number of segments was an error-prone and laborious process that demanded a remapping of the relationship between the user’s mental image of the biologically important features of the model, on the one hand, and the implementation of this model in a digital computer, on the other. The use of $x$ and $n_{seg}$ insulates the user from the most inconvenient aspects of such low-level details.
When $n_{\text{seg}} = 1$ the entire section is lumped into a single compartment. This compartment has only one node, which is located midway along its length, at $x = 0.5$ (see Figures 3C and D). The integral of the surface area over the entire length of the section ($0 \leq x \leq 1$) is used to calculate the membrane properties associated with this node. The values of the axial resistors are determined by integrating the cytoplasmic resistivity along the paths from the ends of the section to its midpoint (dashed line in Figure 3C). The left- and right-hand axial resistances of Figure 3D are evaluated over the $x$ intervals $[0, 0.5]$ and $[0.5, 1]$, respectively.

Figure 4 shows what happens when $n_{\text{seg}} = 2$. Now NEURON breaks the section into two segments of equal length that correspond to $x$ intervals $[0, 0.5]$ and $[0.5, 1]$. The membrane properties over these intervals are attached to the nodes at 0.25 and 0.75, respectively. The three axial resistors $R_{i1}, R_{i2},$ and $R_{i3}$ are determined by integrating the path resistance over the $x$ intervals $[0, 0.25], [0.25, 0.75],$ and $[0.75, 1]$.

3.3 Integration Methods. Spatial discretization reduced the cable equation, a partial differential equation with derivatives in space and time, to a set of ordinary differential equations with first-order derivatives in time. Selection of an integration method to solve these equations is guided by concerns of stability, accuracy, and efficiency (Hines & Carnevale, 1995).

NEURON offers the user a choice of two stable implicit integration methods: backward Euler and a variant of Crank-Nicholson (C-N). Backward Euler is the default because of its robust numerical stability properties. Backward Euler can be used with extremely large time steps in order to find the steady-state solution for a linear (“passive”) system. It produces good qualitative results even with large time steps, and it works even if some or all of the equations are strictly algebraic relations among states.

A more accurate method for small time steps is available by setting the global parameter $\text{secondorder}$ to 2. NEURON then uses a variant of the C-N method, in which numerical error is proportional to $\Delta t^2$.

Both of these are implicit integration methods, in which all current balance equations must be solved simultaneously. The backward Euler algorithm does not resort to iteration to deal with nonlinearities, since its numerical error is proportional to $\Delta t$ anyway. The special feature of the C-N variant is its use of a staggered time step algorithm to avoid iteration of nonlinear equations (see section 3.3.1). This converts the current balance part of the problem to one that requires only the solution of simultaneous linear equations.

Although the C-N method is formally stable, it is sometimes plagued by spurious large-amplitude oscillations (see Hines and Carnevale, 1995, Figure 7). This occurs when $\Delta t$ is too large, as may occur in models that involve fast voltage clamps or have compartments coupled by very small resistances. However, C-N is safe in most situations, and it can be much more efficient than backward Euler for a given accuracy.
These two methods are almost identical in terms of computational cost per time step (see section 3.3.1). Since the current balance equations have the structure of a tree (there are no current loops), direct gaussian elimination is optimal for their solution (Hines, 1984). This takes exactly the same number of computer operations as would be required for an unbranched cable with the same number of compartments.

The best way to determine the method of choice for any particular problem is to compare both methods with several values of $\Delta t$ to see which allows the largest $\Delta t$ consistent with the desired accuracy. In performing such trials, one must remember that the stability properties of a simulation depend on the entire system that is being modeled. Because of interactions between the “biological” components and any “nonbiological” elements, such as stimulators or voltage clamps, the time constants of the entire system may be different from those of the biological components alone. A current source (perfect current clamp) does not affect stability because it does not change the time constants. Any other signal source imposes a load on the compartment to which it is attached, changing the time constants and potentially requiring use of a smaller time step to avoid numerical oscillations in the C-N method. The more closely a signal source approximates a voltage source (perfect voltage clamp), the greater this effect will be.

3.3.1 Efficiency. Nonlinear equations generally need to be solved iteratively to maintain second-order correctness. However, voltage-dependent membrane properties, which are typically formulated in analogy to Hodgkin-Huxley (HH) type channels, allow the cable equation to be cast in a linear form, still second order correct, that can be solved without iterations. A direct solution of the voltage equations at each time step $t \rightarrow t + \Delta t$ using the linearized membrane current $I(V, t) = G \cdot (V - E)$ is sufficient as long as the slope conductance $G$ and the effective reversal potential $E$ are known to second order at time $t + 0.5\Delta t$. HH type channels are easy to solve at $t + 0.5\Delta t$ since the conductance is a function of state variables, which can be computed using a separate time step that is offset by $0.5\Delta t$ with respect to the voltage equation time step. That is, to integrate a state from $t - 0.5\Delta t$ to $t + 0.5\Delta t$, we require only a second-order correct value for the voltage-dependent rates at the midpoint time $t$.

Figure 5 contrasts this approach with the common technique of replacing nonlinear coefficients by their values at the beginning of a time step. For HH equations in a single compartment, the staggered time grid approach converts four simultaneous nonlinear equations at each time step to four independent linear equations that have the same order of accuracy at each time step. Since the voltage-dependent rates use the voltage at the midpoint of the integration step, integration of channel states can be done analytically in just a single addition and multiplication operation and two table-lookup operations. Although this efficient scheme achieves second-order accuracy,
Figure 5: The equations shown at the top of the figure are computed using the Crank-Nicholson method. (Top) $x(t+\Delta t)$ and $y(t+\Delta t)$ are determined using their values at time $t$. (Bottom) Staggered time steps yield decoupled linear equations. $y(t + \Delta t/2)$ is determined using $x(t)$, after which $x(t + \Delta t)$ is determined using $y(t + \Delta t/2)$.

The trade-off is that the tables depend on the value of the time step and must be recomputed whenever the time step changes.

Neuronal architecture can also be exploited to increase computational efficiency. Since neurons generally have a branched tree structure with no loops, the number of arithmetic operations required to solve the cable equation by gaussian elimination is exactly the same as for an unbranched cable with the same number of compartments. That is, we need only $O(N)$ arith-
metic operations for the equations that describe \( N \) compartments connected in the form of a tree, even though standard gaussian elimination generally takes \( O(N^3) \) operations to solve \( N \) equations in \( N \) unknowns. The tremendous efficiency increase results from the fact that in a tree, one can always find a leaf compartment \( i \) that is connected to only one other compartment \( j \), so that the equation for compartment \( i \) (see equation 3.3a) involves only the voltages in compartments \( i \) and \( j \), and the voltage in leaf compartment \( i \) is involved only in the equations for compartments \( i \) and \( j \) (see equations 3.3a and 3.3b):

\[
a_{ii}V_i + a_{ij}V_j = b_i \tag{3.3a}
\]

\[
a_{ji}V_i + a_{jj}V_j + \text{[terms from other compartments]} = b_j. \tag{3.3b}
\]

Using equation 3.3a to eliminate the \( V_i \) term from equation 3.3b, which requires \( O(1) \) (instead of \( N \) ) operations, gives equation 3.4 and leaves \( N - 1 \) equations in \( N - 1 \) unknowns.

\[
a'_{jj}V_j + \text{[terms from other compartments]} = b'_j \tag{3.4}
\]

where

\[
a'_{jj} = a_{jj} - \frac{a_{ij}a_{ji}}{a_{ii}}
\]

and

\[
b'_j = b_j - \frac{b_i a_{ij}}{a_{ii}}.
\]

This strategy can be applied until there is only one equation in one unknown.

Assume that we know the solution to these \( N - 1 \) equations, and in particular that we know \( V_j \). Then we can find \( V_i \) from equation 3.3a with \( O(1) \) step. Therefore, the effort to solve these \( N \) equations is \( O(1) \) plus the effort needed to solve \( N - 1 \) equations. The number of operations required is independent of the branching structure, so a tree of \( N \) compartments uses exactly the same number of arithmetic operations as a one-dimensional cable of \( N \) compartments.

Efficient gaussian elimination requires an ordering that can be found by a simple algorithm: choose the equation with the current minimum number of terms as the equation to use in the elimination step. This minimum degree ordering algorithm is commonly employed in standard sparse matrix solver packages. For example, NEURON’s Matrix class uses the matrix library written by Stewart and Leyk (1994). This and many other sparse matrix packages are freely available on the Internet at http://www.netlib.org.
4 The Neuron Simulation Environment

No matter how powerful and robust its computational engine may be, the real utility of any software tool depends largely on its ease of use. Therefore a great deal of effort has been invested in the design of the simulation environment provided by NEURON. In this section we first briefly consider general aspects of the high-level language used for writing NEURON programs. Then we turn to an example of a model of a nerve cell to introduce specific aspects of the user environment, after which we cover these features more thoroughly.

4.1 The hoc Interpreter. NEURON incorporates a programming language based on hoc, a floating-point calculator with C-like syntax described by Kernighan and Pike (1984). This interpreter has been extended by the addition of object-oriented syntax (not including polymorphism or inheritance) that can be used to implement abstract data types and data encapsulation. Other extensions include functions that are specific to the domain of neural simulations and functions that implement a graphical user interface (see below). With hoc one can quickly write short programs that meet most problem-specific needs. The interpreter is used to execute simulations, customize the user interface, optimize parameters, analyze experimental data, calculate new variables such as impulse propagation velocity, and so forth.

NEURON simulations are not subject to the performance penalty often associated with interpreted (as opposed to compiled) languages because computationally intensive tasks are carried out by highly efficient precompiled code. Some of these tasks are related to integration of the cable equation, and others are involved in the emulation of biological mechanisms that generate and regulate chemical and electrical signals.

NEURON provides a built-in implementation of the microemacs text editor. Since the choice of a programming editor is highly personal, NEURON will also accept hoc code in the form of straight ASCII files created with any other editor.

4.2 A Specific Example. In the following example we show how NEURON might be used to model the cell in the top of Figure 6. Comments in the hoc code are preceded by double slashes (//), and code blocks are enclosed in curly brackets ({}).

4.2.1 First Step: Establish Model Topology. One very important feature of NEURON is that it allows the user to think about models in terms that are familiar to the neurophysiologist, keeping numerical issues (e.g., number of spatial segments) entirely separate from the specification of morphology and biophysical properties. This separation is achieved through the use of one-dimensional cable “sections” as the basic building block from which model cells are constructed. These sections can be connected together to
Figure 6: (Top) Cartoon of a neuron with a soma, three dendrites, and an unmyelinated axon (not to scale). The diameter of the spherical soma is 50 µm. Each dendrite is 200 µm long and tapers uniformly along its length from 10 µm diameter at its site of origin on the soma, to 3 µm at its distal end. The unmyelinated cylindrical axon is 1000 µm long and has a diameter of 1 µm. An electrode (not shown) is inserted into the soma for intracellular injection of a stimulating current. (Bottom) Topology of a NEURON model that represents this cell (see text for details).

form any kind of branched cable and endowed with properties that may vary with position along their length.

The idealized neuron in Figure 6 has several anatomical features whose existence and spatial relationships we want the model to include: a cell body (soma), three dendrites, and an unmyelinated axon. The following hoc code sets up the basic topology of the model:

```hoc
create soma, axon, dendrite[3]
connect axon(0), soma(0)
for i=0,2 { connect dendrite[i](0), soma[i] }
```

The program starts by creating named sections that correspond to the important anatomical features of the cell. These sections are attached to each other using connect statements. As noted previously, each section has a normalized position parameter \( x \), which ranges from 0 at one end to 1 at the other. Because the axon and dendrites arise from opposite sides of the cell body, they are connected to the 0 and 1 ends of the soma section (see the
bottom of Figure 6). A child section can be attached to any location on the parent, but attachment at locations other than 0 or 1 is generally employed only in special cases such as spines on dendrites.

4.2.2 Second Step: Assign Anatomical and Biophysical Properties. Next we set the anatomical and biophysical properties of each section. Each section has its own segmentation, length, and diameter parameters, so it is necessary to indicate which section is being referenced. There are several ways to declare which is the currently accessed section, but here the most convenient is to precede blocks of statements with the appropriate section name:

```plaintext
// specify anatomical and biophysical properties
soma {
    nseg = 1 // compartmentalization parameter
    L = 50 // [µm] length
    diam = 50 // [µm] diameter
    insert hh // standard Hodgkin-Huxley
        // currents
    gnabar_hh = 0.5*0.120 // max HH sodium conductance
}
axon {
    nseg = 20
    L = 1000
    diam = 1
    insert hh
}
for i=0,2 dendrite[i] {
    nseg = 5
    L = 200
    diam(0:1) = 10:3 // dendritic diameter tapers
        // along its length
    insert pas // standard passive current
    e_pas = -65 // [mV] equilibrium potential
        // for passive current
    g_pas = 0.001 // [siemens/cm²] conductance
        // for passive current
}
```

The fineness of the spatial grid is determined by the compartmentalization parameter nseg. Here the soma is lumped into a single compartment (nseg = 1), while the axon and each of the dendrites are broken into several subcompartments (nseg = 20 and 5, respectively).

In this example, we specify the geometry of each section by assigning values directly to section length and diameter. This creates a stylized model. Alternatively, one can use the 3D method, in which NEURON computes
section length and diameter from a list of $(x, y, z, \text{diam})$ measurements (see section 4.5).

Since the axon is a cylinder, the corresponding section has a fixed diameter along its entire length. The spherical soma is represented by a cylinder with the same surface area as the sphere. The dimensions and electrical properties of the soma are such that its membrane will be nearly isopotential, so the cylinder approximation is not a significant source of error. If chemical signals such as intracellular ion concentrations were important in this model, it would be necessary to approximate not only the surface area but also the volume of the soma.

Unlike the axon, the dendrites become progressively narrower with distance from the soma. Furthermore, unlike the soma, they are too long to be lumped into a single compartment with constant diameter. The taper of the dendrites is accommodated by assigning a sequence of decreasing diameters to their segments. This is done through the use of range variables, which are discussed in section 4.4.

In this model the soma and axon contain HH sodium, potassium, and leak channels (Hodgkin & Huxley, 1952), while the dendrites have constant, linear (“passive”) ionic conductances. The insert statement assigns the biophysical mechanisms that govern electrical signals in each section. Particular values are set for the density of sodium channels on the soma ($g_{nabar\_hh}$) and for the ionic conductance and equilibrium potential of the passive current in the dendrites ($g_{\text{pas}}$ and $e_{\text{pas}}$). More information about membrane mechanisms is presented in section 4.6.

4.2.3 Third Step: Attach Stimulating Electrodes. This code emulates the use of an electrode to inject a stimulating current into the soma by placing a current pulse stimulus in the middle of the soma section. The stimulus starts at $t = 1$ ms, lasts for 0.1 ms, and has an amplitude of 60 nA.

```plaintext
objref stim
soma stim = new IClamp(0.5) // put it in middle of soma
stim.del = 1           // [ms] delay
stim.dur = 0.1         // [ms] duration
stim.amp = 60          // [nA] amplitude
```

The stimulating electrode is an example of a point process. Point processes are discussed in more detail in section 4.6.

4.2.4 Fourth Step: Control Simulation Time Course. At this point all model parameters have been specified. All that remains is to define the simulation parameters, which govern the time course of the simulation, and write some code that executes the simulation.

This is generally done in two procedures. The first procedure initializes the membrane potential and the states of the inserted mechanisms (channel states, ionic concentrations, extracellular potential next to the membrane).
The second procedure repeatedly calls the built-in single-step integration function `fadvance()` and saves, plots, or computes functions of the desired output variables at each step. In this procedure it is possible to change the values of model parameters during a run.

```plaintext
dt = 0.05 // [ms] integration time step
tstop = 5 // [ms]
finitialize(-65) // initialize membrane potential, // state variables, and time
proc integrate() {
    print t, soma.v(0.5) // show starting time // and initial somatic // membrane potential
    while (t < tstop) {
        fadvance() // advance solution by dt // function calls to save or plot results // would go here
        print t, soma.v(0.5) // show present time // and somatic // membrane potential
        // statements that change model parameters // would go here
    }
}
```

The built-in function `finitialize()` initializes time \( t \) to 0, membrane potential \( v \) to −65 mV throughout the model, and the HH state variables \( m, n, \) and \( h \) to their steady-state values at \( v = -65 \text{ mV} \). Initialization can also be performed with a user-written routine if there are special requirements that `finitialize()` cannot accommodate, such as nonuniform membrane potential.

Both the integration time step \( dt \) and the solution time \( t \) are global variables. For this example \( dt = 50 \mu s \). The `while(){}` statement repeatedly calls `fadvance()`, which integrates the model equations over the interval \( dt \) and increments \( t \) by \( dt \) on each call. For this example, the time and somatic membrane potential are displayed at each step. This loop exits when \( t \geq tstop \).

When this program is first processed by the NEURON interpreter, the model is set up and initiated, but the `integrate()` procedure is not executed. When the user enters an `integrate()` statement in the NEURON interpreter window, the simulation advances for 5 ms using 50 \( \mu s \) time steps.

### 4.3 Section Variables.

Three parameters apply to the section as a whole: cytoplasmic resistivity \( R_a \) (\( \Omega \text{ cm} \)), the section length \( L \), and the compartmentalization parameter \( nseg \). The first two are ordinary in the sense that they
do not affect the structure of the equations that describe the model. Note that
the hoc code specifies values for \( L \) but not for \( Ra \). This is because each sec-
tion in a model is likely to have a different length, whereas the cytoplasm
(and therefore \( Ra \)) is usually assumed to be uniform throughout the cell.
The default value of \( Ra \) is 35.4 \( \Omega \) cm, which is appropriate for invertebrate
neurons. Like \( L \) it can be assigned a new value in any or all sections (e.g.,
\( \sim 200 \) \( \Omega \) cm for mammalian neurons).

The user can change the compartmentalization parameter \( nseg \) without
having to modify any of the statements that set anatomical or biophysical
properties. However, if parameters vary with position in a section, care must
be taken to ensure that the model incorporates the spatial detail inherent in
the parameter description.

4.4 Range Variables. Like dendritic diameter in our example, most cel-
lar properties are functions of the position parameter \( x \). NEURON has
special provisions for dealing with these properties, which are called range
variables. Other examples of range variables are the membrane potential \( v \) and ionic conductance parameters such as the maximum HH sodium
conductance \( g_{nabar, hh} \) (siemens/cm\(^2\)).

Range variables enable the user to separate property specification from
segment number. A range variable is assigned a value in one of two ways.
The simpler and more common is as a constant. For example, the statement
\( \text{axon.diam} = 10 \) asserts that the diameter of the axon is uniform over its
entire length.

The syntax for a property that changes along a length of a section is
\( \text{rangevar}(xmin: xmax) = e1 : e2 \). The four italicized symbols are expres-
sions, with \( e1 \) and \( e2 \) being the values of the property at \( xmin \) and \( xmax \),
respectively. The position expressions must meet the constraint \( 0 \leq xmin \leq
xmax \leq 1 \). Linear interpolation is used to assign the values of the property at
the segment centers that lie in the position range \([xmin, xmax] \). In this man-
ner a continuously varying property can be approximated by a piecewise
linear function. If the range variable is diameter, neither \( e1 \) nor \( e2 \) should be
0, or corresponding axial resistance will be infinite.

In our model neuron, the simple dendritic taper is specified by \( \text{diam}(0:1)
= 10:3 \) and \( nseg = 5 \). This results in five segments that have centers at
\( x = 0.1, 0.3, 0.5, 0.7, \) and 0.9 and diameters of 9.3, 7.9, 6.5, 5.1, and 3.7,
respectively.

The value of a range variable at the center of a segment can appear in any
expression using the syntax \( \text{rangevar}(x) \) in which \( 0 \leq x \leq 1 \). The value
returned is the value at the center of the segment containing \( x \), not the linear
interpolation of the values stored at the centers of adjacent segments. If the
parentheses are omitted, the position defaults to a value of 0.5 (middle of
the section).

A special form of the for statement is available: \( \text{for (var) stmt.} \) For
each value of the normalized position parameter \( x \) that defines the center
of each segment in the selected section (along with positions 0 and 1), this statement assigns \texttt{var} that value and executes the \texttt{stmt}. This hoc code would print the membrane potential as a function of physical position (in \( \mu \text{m} \)) along the axon:

\begin{verbatim}
axon for (x) print x*L, v(x)
\end{verbatim}

4.5 Specifying Geometry: Stylized versus 3D. There are two ways to specify section geometry. Our example uses the stylized method, which simply assigns values to section length and diameter. This is most appropriate when cable length and diameter are authoritative and 3D shape is irrelevant.

If the model is based on anatomical reconstruction data (quantitative morphometry) or if 3D visualization is paramount, it is better to use the 3D method. This approach keeps the anatomical data in a list of \((x, y, z, \text{diam})\) “points.” The first point is associated with the end of the section that is connected to the parent (this is not necessarily the zero end), and the last point is associated with the opposite end. There must be at least two points per section, and they should be ordered in terms of monotonically increasing arc length. This pt3d list, which is the authoritative definition of the shape of the section, automatically determines the length and diameter of the section.

When the pt3d list is nonempty, the shape model used for a section is a sequence of frusta. The pt3d points define the locations and diameters of the ends of these frusta. The effective area, diameter, and resistance of each segment are computed from this sequence of points by trapezoidal integration along the segment length. This takes into account the extra area introduced by diameter changes; even degenerate cones of 0 length can be specified (i.e., two points with same coordinates but different diameters), which add area but not length to the section. No attempt is made to deal with the effects of centroid curvature on surface area. The number of 3D points used to describe a shape has nothing to do with \texttt{nseg} and does not affect simulation speed.

4.6 Density Mechanisms and Point Processes. The \texttt{insert} statement assigns biophysical mechanisms, which govern electrical and (if present) chemical signals, to a section. Many sources of electrical and chemical signals are distributed over the membrane of the cell. These density mechanisms are described in terms of current per unit area and conductance per unit area; examples include voltage-gated ion channels such as the HH currents. However, density mechanisms are not the most appropriate representation of all signal sources. Synapses and electrodes are best described in terms of localized absolute current in nanoamperes and conductance in microsiemens. These are called point processes.
An object syntax is used to manage the creation, insertion, attributes, and destruction of point processes. For example, a current clamp (electrode for injecting a current) is created by declaring an object variable and assigning it a new instance of the IClamp object class. When a point process is no longer referenced by any object variable, the point process is removed from the section and destroyed. In our example, redeclaring stim with the statement objref stim would destroy the pulse stimulus, since no other object variable is referencing it.

The location of a point process can be changed with no effect on its other attributes. In our example, the statement dendrite[2] stim.loc(1) would move the current stimulus to the distal end of the third dendrite.

Many user-defined density mechanisms and point processes can be simultaneously present in each compartment of a neuron. One important difference between density mechanisms and point processes is that any number of the same kind of point process can exist at the same location.

User-defined density mechanisms and point processes can be linked into NEURON using the model description language NMODL. This lets the user focus on specifying the equations for a channel or ionic process without regard to its interactions with other mechanisms. The NMODL translator then constructs the appropriate C program, which is compiled and becomes available for use in NEURON. This program properly and efficiently computes the total current of each ionic species used, as well as the effect of that current on ionic concentration, reversal potential, and membrane potential.

An extensive discussion of NMODL is beyond the scope of this article, but its major advantages can be listed succinctly.

1. Interface details to NEURON are handled automatically, and there are a great many such details. NEURON needs to know that model states are range variables and which model parameters can be assigned values and evaluated from the interpreter. Point processes need to be accessible via the interpreter object syntax, and density mechanisms need to be added to a section when the “insert” statement is executed. If two or more channels use the same ion at the same place, the individual current contributions need to be added together to calculate a total ionic current.

2. Consistency of units is ensured.

3. Mechanisms described by kinetic schemes are written with a syntax in which the reactions are clearly apparent. The translator provides tremendous leverage by generating a large block of C code that calculates the analytic Jacobian and the state fluxes.

4. There is often a great increase in clarity since statements are at the model level instead of the C programming level and are independent of the numerical method. For example, sets of differential and non-
linear simultaneous equations are written using an expression syntax such as
\[ x' = f(x, y, t) \]
\[ g(x, y) = h(x, y) \]
where the prime refers to the derivative with respect to time (multiple primes such as \(x''\) refer to higher derivatives), and the tilde introduces an algebraic equation. The algebraic portion of such systems of equations is solved by Newton’s method, and a variety of methods are available for solving the differential equations, such as Runge-Kutta or backward Euler.

5. Function tables can be generated automatically for efficient computation of complicated expressions.

6. Default initialization behavior of a channel can be specified.

4.7 Graphical Interface. The user is not limited to operating within the traditional code-based command-mode environment. Among its many extensions to \texttt{hoc}, NEURON includes functions for implementing a fully graphical, windowed interface. Through this interface, and without having to write any code at all, the user can effortlessly create and arrange displays of menus, parameter value editors, graphs of parameters and state variables, and views of the model neuron. Anatomical views, called space plots, can be explored, revealing what mechanisms and point processes are present and where they are located.

The purpose of NEURON’s graphical interface is to promote a match between what the user thinks is inside the computer and what is actually there. These visualization enhancements are a major aid to maintaining conceptual control over the simulation because they provide immediate answers to questions about what is being represented in the computer.

The interface has no provision for constructing neuronal topology, a conscious design choice based on the strong likelihood that a graphical toolbox for building neuronal topologies would find little use. Small models with simple topology are so easily created in \texttt{hoc} that a graphical topology editor is unnecessary. More complex models are too cumbersome to deal with using a graphical editor. It is best to express the topological specifications of complex stereotyped models through algorithms, written in \texttt{hoc}, that generate the topology automatically. Biologically realistic models often involve hundreds or thousands of sections, whose dimensions and interconnections are contained in large data tables generated by hours of painstaking quantitative morphometry. These tables are commonly read by \texttt{hoc} procedures that in turn create and connect the required sections without operator intervention.

The basic features of the graphical interface and how to use it to monitor and control simulations are discussed elsewhere (Moore & Hines, 1996).
However, several sophisticated analysis and simulation tools that have special utility for nerve simulation are worthy of mention:

- The Function Fitter optimizes a parameterized mathematical expression to minimize the least squared difference between the expression and data.

- The Run Fitter allows one to optimize several parameters of a complete neuron model to experimental data. This is most useful in the context of voltage clamp data that are contaminated by incomplete space clamp or models that cannot be expressed in closed form, such as kinetic schemes for channel conductance.

- The Electrotonic Workbench plots small signal input and transfer impedance and voltage attenuation as functions of space and frequency (Carnevale, Tsai, & Hines, 1996). These plots include the neuromorphic (Carnevale, Tsai, Claiborne, & Brown, 1995) and L versus x (O’Boyle, Barrevalle, Carnevale, Claiborne, & Brown, 1996) renderings of the electrotonic transformation (Brown et al., 1992; Tsai, Carnevale, Claiborne, & Brown, 1994; Zador, Agmon-Šnir, & Segev, 1995). By revealing the effectiveness of signal transfer, the Workbench quickly provides insight into the “functional shape” of a neuron.

All interactions with these and other tools take place in the graphical interface, and no interpreter programming is needed to use them. However, they are constructed entirely within the interpreter and can be modified when special needs require.

4.8 Object-Oriented Syntax.

4.8.1 Neurons. It is often convenient to deal with groups of sections that are related. Therefore NEURON provides a data class called a SectionList that can be used to identify subsets of sections. Section lists fit nicely with the “regular expression” method of selecting sections, used in earlier implementations of NEURON, in that the section list is easily constructed by using regular expressions to add and delete sections. After the list is constructed it is available for reuse, and it is much more efficient to loop over the sections in a section list than to pick out the sections accepted by a combination of regular expressions. This code

```object
objref alldend
alldend = new SectionList()
forsec "dend" alldend.append()
forsec alldend print secname()
```

forms a list of all the sections whose names contain the string “dend” and then iterates over the list, printing the name of each section in it. For the example program presented here, this would generate the following output
in the NEURON interpreter window,

```plaintext
dendrite[0]
dendrite[1]
dendrite[2]
```

although in this very simple example it would clearly have been easy enough to loop over the array of dendrites directly, for example:

```plaintext
for i = 0,2 {
    dendrite[i] print secname()
}
```

4.8.2 Networks. To help the user manage very large simulations, the interpreter syntax has been extended to facilitate the construction of hierarchical objects. This is illustrated by the following code fragment, which specifies a pattern for a simple stylized neuron consisting of three dendrites connected to one end of a soma and an axon connected to the other end:

```plaintext
begintemplate Cell1
  public soma, dendrite, axon
  create soma, dendrite[3], axon
  proc init() {
      for i=0,2 connect dendrite[i](0), soma(0)
      connect axon(0), soma(1)
      axon insert hh
  }
endtemplate Cell1
```

Whenever a new instance of this pattern is created, the `init()` procedure automatically connects the `soma`, `dendrite`, and `axon` sections together. A complete pattern would also specify default membrane properties, as well as the number of segments for each section.

Names that can be referenced outside the pattern are listed in the `public` statement. In this case, since `init` is not in the list, the user could not reinitialize by calling the `init()` procedure. Public names are referenced through a dot notation.

The particular benefit of using templates (“classes” in standard object-oriented terminology) is the fact that they can be employed to create any number of instances of a pattern. For example,

```plaintext
objref cell[10][10]
for i=0,9 for j=0,9 cell[i][j]=new Cell1()
```

creates an array of one hundred objects of type `Cell1` that can be referenced individually by the object variable `cell`. In this example, `cell[4][5].axon.gnabar.hh(0.5)` is the value of the maximum HH sodium conductance in the middle of the axon of `cell[4][5]`.
As this example implies, templates offer a natural syntax for the creation of networks. However, it is entirely up to the user to organize the templates logically so that they appropriately reflect the structure of the problem. Generally any given structural organization can be viewed as a hierarchy of container classes, such as cells, microcircuits, layers, or networks. The important issue is how much effort is required for the concrete network representation to support a range of logical views of the same abstract network. A logical view that organizes the cells differently may not be easy to compute if the network is built as an elaborate hierarchy. This kind of pressure tends to encourage relatively flat organizations that make it easier to implement functions that search for specific information. The bottom line is that network simulation design remains an ad hoc process that requires careful programming judgment.

One very important class of logical views that are not generally organiz-able as a hierarchy are those of synaptic organization. In connecting cells with synapses, one is often driven to deal with general graphs, which is to say no structure at all.

In addition to the notions of classes and objects (a synapse is an object with a pre- and a postsynaptic logical connection) the interpreter offers one other fundamental language feature that can be useful in dealing with objects that are collections of other objects: the notion of iterators, taken from the Sather programming language (Murer, Omohundro, Stoutamire, & Szwerski, 1996). This is a separation of the process of iteration from that of “what is to be done for each item.” If a programmer implements one or more iterators in a collection class, the user of the class does not need to know how the class indexes its items. Instead the class will return each item in turn for execution in the context of the loop body. This allows the user to write

```java
for layer1.synapses(syn, type) {
    // statements that manipulate the object
    // reference named "syn" (The iterator
    // causes "syn" to refer, in turn, to
    // each synapse of a certain type in
    // the layer1 object)
}
```

without being aware of the possibly complicated process of picking out these synapses from the layer (that is the responsibility of the author of the class of which layer1 is an instance).

It is to be sadly emphasized that these kinds of language features, though very useful, do not impose any policy with regard to the design decisions that users must make in building their networks. Different programmers express very different designs on the same language base, with the consequence that it is more often than not infeasible to reconcile slightly different representations of even very similar concepts.
An example of a useful way to deal uniformly with the issue of synaptic connectivity is the policy implemented in NEURON by Lytton (1996). This implementation uses the normal NMODL methodology to define a synaptic conductance model and enclose it within a framework that manages network connectivity.

5 Summary

The recent striking expansion in the use of simulation tools in the field of neuroscience has been encouraged by the rapid growth of quantitative observations that both stimulate and constrain the formulation of new hypotheses of neuronal function, enabled by the availability of ever-increasing computational power at low cost. These factors have motivated the design and implementation of NEURON, the goal of which is to provide a powerful and flexible environment for simulations of individual neurons and networks of neurons. NEURON has special features that accommodate the complex geometry and nonlinearities of biologically realistic models, without interfering with its ability to handle more speculative models that involve a high degree of abstraction.

As we note in this article, one particularly advantageous feature is that the user can specify the physical properties of a cell without regard for the strictly computational concern of how many compartments are employed to represent each of the cable sections. In a future publication, we will examine how the NMODL translator is used to define new membrane channels and calculate ionic concentration changes. Another will describe the Vector class. In addition to providing very efficient implementations of frequently needed operations on lists of numbers, the vector class offers a great deal of programming leverage, especially in the management of network models.


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