Cellular/Molecular

How Multiple Conductances Determine Electrophysiological Properties in a Multicompartment Model

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Most neurons have large numbers of voltage- and time-dependent currents that contribute to their electrical firing patterns. Because these currents are nonlinear, it can be difficult to determine the role each current plays in determining how a neuron fires. The lateral pyloric (LP) neuron of the stomatogastric ganglion of decapod crustaceans has been studied extensively biophysically. We constructed ~600,000 versions of a four-compartment model of the LP neuron and distributed 11 different currents into the compartments. From these, we selected ~1300 models that match well the electrophysiological properties of the biological neuron. Interestingly, correlations that were seen in the expression of channel mRNA in biological studies were not found across the ~1300 admissible LP neuron models, suggesting that the electrical phenotype does not require these correlations. We used cubic fits of the function from maximal conductances to a series of electrophysiological properties to ask which conductances predominantly influence input conductance, resting membrane potential, resting spike rate, phasing of activity in response to rhythmic inhibition, and several other properties. In all cases, multiple conductances contribute to the measured property, and the combinations of currents that strongly influence each property differ. These methods can be used to understand how multiple currents in any candidate neuron interact to determine the cell’s electrophysiological behavior.

Introduction

The mixture of voltage- and calcium-dependent channels and their distribution across the surface of a neuron collectively determine the electrophysiological properties of the neuron, including its resting membrane potential, input impedance, firing rate in response to current, and the extent to which it exhibits neuronal behaviors such as bursting, plateauing, and postinhibitory rebound (Marder, 1998). For a neuronal network to function properly, the neurons in that network must exhibit network-appropriate behaviors.

Recent work has shown that although a neuron may behave similarly from animal to animal, the densities of voltage-gated channels can vary severalfold from animal to animal (Golowasch and Marder, 1992a; Golowasch et al., 2002; MacLean et al., 2005; Schulz et al., 2006, 2007). How, then, is reliable neuronal behavior maintained in the face of this variability? One possible answer is that although conductances are variable, they are variable in a correlated manner, and these correlations preserve the function of the neuron. This idea was given additional plausibility by the discovery that many channels are expressed in a correlated manner, and these correlations preserve the function of the neuron (Schulz et al., 2006, 2007).

We sought to examine the hypothesis that correlations between conductances are required to conserve the electrophysiological function of neurons. We generated a large population of models of a single identified neuron, the lateral pyloric (LP) neuron, found in the stomatogastric ganglion (STG) of the crab Cancer borealis. We generated this population by choosing model parameters at random and filtering out models that did not behave appropriately. We then measured a number of electrophysiological properties in the resulting models and looked for correlations between model parameters that were induced by our constraints on electrophysiological properties.

Historically, a number of different approaches have been used to assess the role of a given channel in a given electrophysiological property of a neuron. One such approach is pharmacological blockade (Hille, 2001). Although this can be done successfully, it presupposes both that a given drug’s cellular targets are known and that it is selective. Often, neither of these conditions is met. More recently, many have studied the effects of genetically altering or deleting currents on neuronal dynamics (Raman et al., 1997; Khaliq et al., 2003; MacLean et al., 2003; Swensen and Bean, 2005). Again, although these manipulations are often informative, their interpretation is complicated by compensatory mechanisms that occur during development. A third method is to use the dynamic clamp to add or subtract a current from a neuron (Prinz et al., 2004a).

The most common computational technique used to assess the role of a given parameter on the behavior of a neuron is sensitivity analysis (Butera et al., 1999; Hill et al., 2001, 2002; Jezzini et al., 2004). Here, we used a different approach to understand the extent to which combinations of neuronal conductances influence a number of different neuronal behaviors and compare this approach to sensitivity analysis.
Materials and Methods
Electrophysiology
Experiments were performed essentially as described previously (Gosso-
lard et al., 2004). All preparations had two intact superior esophageal
erves and included the esophageal ganglion, but they had from zero to
two intact inferior esophageal nerves.

Modeling voltage-gated currents
The LP neuron models contained 10 voltage- and calcium-gated cur-
rents. These were the delayed-rectifier potassium current (I_{Kd}); the tran-
sient potassium current (I_{Kt}); the calcium current (I_{Ca}); the calcium-
activated potassium current (I_{CaK}); the hyperpolarization-activated
inward current (I_{P}); the proctolin-activated current (I_{P}), a mixed inward
current that is activated by several neuromodulators, including proctolin
(Swensen and Marder, 2000); the fast sodium current (I_{Na}); the axonal
delayed-rectifier potassium current (I_{Kst}); and the axonal transient potas-
sium current (I_{Kst}). The I_{Kst} current was modeled as the sum of a fast and
a slow component (I_{Ks} and I_{Kst}), but the maximal conductances of these
currents were held in a fixed ratio (0.885:1), because they model a single
voltage-gated current as characterized physiologically. The model con-
tained two leak currents, I_{leak,ax} (the somatonic leak) and I_{leak,sk} (the
axonal leak conductance), both described by the usual formalism but
with different maximal conductances.

The mathematical description of most of these currents (I_{Kd}, I_{Kt}, I_{Ca},
I_{CaK}, I_{Na}, and I_{P}) was based on descriptions in previous work (Buchholtz
et al., 1992; Golowasch and Marder, 1992b), but with several revisions. In
cases, we tuned the half-activation voltages and kinetics of these cur-
rents to provide good fits to voltage-clamp data (Golowasch and Marder,
1992a; Turrigiano et al., 1995; Schulz et al., 2006). The axonal currents
(I_{Na}, I_{CaK}, and I_{Kst}) were based on those in the Connor-Stevens model of
crab walking leg axon (Connor et al., 1977), but the half-activation and
kinetic parameters of these currents were tuned to have the desired
current-clamp behavior in the baseline model (see below, Construction
of baseline model). Finally, there were three synaptic currents in the final
LP neuron models, representing synaptic inputs from other members of
the pyloric network: I_{synAB}, I_{synPD}, and I_{synPY}. They are described below.

All model currents except I_{Kst} were modeled as ohmic currents, de-
scribed by the following equation:

\[ I = g \cdot w \cdot (v - E), \]

where \( g \) is the maximal conductance, \( w \) is the product of the gating
variables raised to the appropriate powers, \( v \) is the membrane potential,
and \( E \) is the reversal potential. The reversal potential of each current in
the model depended on the permeant ions. The reversal potential was
+55 mV for sodium currents, -80 mV for potassium currents (including
I_{synPD}), -25 mV for I_{K}, -10 mV for I_{Ca}, and -70 mV for I_{synAB} and
I_{synSYN}.

The calcium current was modeled using the Goldman-Hodgkin-Katz
equation, and was given by the following:

\[ I_{Ca} = \frac{P_{Ca} d_{Ca}}{1 + \frac{[Ca^{2+}]}{K}}, \]  

where

\[ d_{Ca} = m_{Ca} h_{Ca} N_{A} q_{Ca} \frac{[Ca^{2+}]}{[Ca^{2+}]_{o}} \]

(3)

\[ \xi - \frac{z_{Ca} d_{Ca}}{kT} v, \]

(4)

\( P_{Ca} \) is the maximal calcium permeability, \( m_{Ca} \) is the calcium channel
activation, \( h_{Ca} \) is the calcium channel inactivation, \( N_{A} \) is Avogadro’s
number (6.022 \times 10^{23} \text{ mol}^{-1}), \( q_{Ca} \) is the elementary charge (1.602 \times
10^{-19} \text{ C}), \( z_{Ca} \) is the valence of a calcium ion (+2), \([Ca^{2+}]_{o}\) is the internal
 calcium concentration, \([Ca^{2+}]\) is the external calcium concentration
(13 mM), \( k \) is Boltzmann’s constant (1.381 \times 10^{-23} \text{ J/K}), and \( T \) is the
temperature in Kelvin (283.15K = 10°C).

In the model, we assumed that \( I_{Ca} \) and \( I_{CaK} \) channels were clustered
together, and each cluster was associated with a calcium microdomain
(Marrion and Tavalin, 1998; Fakler and Adelman, 2008). We know of no
evidence either for or against this hypothesis in the STG, but it seems to
be common in other systems (Fakler and Adelman, 2008). The maximal
permeability of \( I_{CaK} \) was related to the per-cluster maximum permeability
by the following:

\[ P_{Ca} = \rho_{max} \cdot \frac{P_{Ca}}{I_{CaK}}, \]  

and the maximal conductance of \( I_{CaK} \) was related to the per-cluster max-
imal conductance by the following:

\[ \frac{g_{max}}{I_{CaK}} = \frac{g_{Ca}}{P_{Ca}}, \]  

where \( \rho_{max} \) is the density of clusters (in clusters/nP), \( P_{Ca} \) is the per-cluster
maximal permeability of \( I_{Ca} \), and \( g_{max} \) is the per-cluster maximal con-
ductance of \( I_{CaK} \). We simulated the accumulation of Ca^{2+} in (an aver-
age) small volume associated with each microdomain. The equation gov-
erning intracellular calcium was as follows:

\[ \frac{d}{dt} \left( \frac{[Ca^{2+}]}{Ca} \right) = \frac{1}{z_{Ca} N_{A} V_{Ca} \rho_{max}} \left[ \frac{[Ca^{2+}]}{Ca} \right]_{i} - \frac{1}{z_{Ca} N_{A} V_{Ca} \rho_{max}} \left[ \frac{[Ca^{2+}]}{Ca} \right]_{o}, \]  

(7)

where \( [Ca^{2+}]_{i} \) is the steady-state calcium concentration in the absence
of calcium current (20 \muM), \( \tau_{Ca} \) is the time constant of calcium buffering
(70.4 ms), and \( V_{Ca} \) is the volume associated with each microdomain (6.49
\muM^3).

In generating our population of models, we picked values of \( P_{Ca} \) and
\( g_{CaK} \) at random and solved Equations 5 and 6 for \( \rho_{max} \) and \( g_{max} \), assum-
ing a fixed \( P_{Ca} \) of 1.167 \times 10^{-3} \muM/s. One consequence of this
scheme is that an increase in \( P_{Ca} \) alone will not change the dynamics
of calcium accumulation (because the calcium accumulation is done on a
per-microdomain basis, and increasing \( P_{Ca} \) increases the density of mi-
crodomains but not the dynamics within a microdomain) and therefore
will not change the dynamics of \( I_{CaK} \). Another approach would be to make
\( \rho_{max} \) and \( g_{max} \) all be independently tunable parameters, but this
seemed overly complex given the experimental uncertainties about the
mechanics of calcium accumulation in this system.

Equations for the steady-state (in)activation and time constants of
each voltage- and calcium-gated current are given in Table 1.

Modeling synaptic currents
During the pyloric rhythm, the LP neuron receives inhibition from three
main sources: the anterior burster (AB) neuron, the pyloric dilator (PD)
nurons, and the pyloric (PY) neurons. In the model, we approximated
these inputs by fixed rhythmic patterns of synaptic conductance.

The synapses from AB and PY to LP are glutamatergic, predominately
cholinergic, potassium-mediated, and slow (Eisen and Marder, 1982;
Marder and Eisen, 1984). The synapse from PD to LP, in contrast, is

\[ v_{APD}(t) = -62.6 + 11.039 \cos(2 \pi \cdot f_{0} t) + \]

\[ 5.199 \cos(2 \pi \cdot f_{0} t + 5.184) + 1.437 \cos(2 \pi \cdot 3 f_{0} t + 10.824) + \]

\[ 0.323 \cos(2 \pi \cdot 4 f_{0} t + 14.913) + 0.256 \cos(2 \pi \cdot 5 f_{0} t + 19.364) + \]

\[ 0.325 \cos(6 \pi \cdot 6 f_{0} t + 18.787) + 0.110 \cos(2 \pi \cdot 7 f_{0} t + 23.993), \]  

where \( v_{APD} \) is in millivolts, \( t \) is in seconds, and \( f_{0} \) is 1 Hz, the funda-
mental frequency of the simulated rhythm. Similarly, the presynaptic
voltage used for the PY synapse was as follows:

\[ v_{APY}(t) = -62.6 + 11.039 \cos(2 \pi \cdot f_{0} t) + \]

\[ 5.199 \cos(2 \pi \cdot f_{0} t + 5.184) + 1.437 \cos(2 \pi \cdot 3 f_{0} t + 10.824) + \]

\[ 0.323 \cos(2 \pi \cdot 4 f_{0} t + 14.913) + 0.256 \cos(2 \pi \cdot 5 f_{0} t + 19.364) + \]

\[ 0.325 \cos(6 \pi \cdot 6 f_{0} t + 18.787) + 0.110 \cos(2 \pi \cdot 7 f_{0} t + 23.993), \]  

where \( v_{APY} \) is in millivolts, \( t \) is in seconds, and \( f_{0} \) is 1 Hz, the funda-
mental frequency of the simulated rhythm.
Table 1. Equations governing the voltage dependence and kinetics of currents in the LP model

<table>
<thead>
<tr>
<th>Current</th>
<th>Form</th>
<th>Steady-State Activation</th>
<th>Time Constant of Activation</th>
<th>Steady-State Inactivation</th>
<th>Time Constant of Inactivation</th>
</tr>
</thead>
<tbody>
<tr>
<td>$I_{Na}$</td>
<td>$m^h$</td>
<td>$l_{gs}(v + 25) / 17$</td>
<td>$120 - 133.8 \cdot l_{gs}(v + 46.1) / (18.1)$</td>
<td>$114 - 100.1 \cdot l_{gs}(v + 1.8) / 4$</td>
<td>$1$</td>
</tr>
<tr>
<td>$I_{Na}$</td>
<td>$m^h$</td>
<td>$l_{gs}(v + 14.5) / (18.1)$</td>
<td>$3$</td>
<td>$l_{gs}(v + 68.1) / (4.5) / 4.8$</td>
<td>$1$</td>
</tr>
<tr>
<td>$I_{Na}$</td>
<td>$m^h$</td>
<td>$l_{gs}(v + 21.0) / 22.8$</td>
<td>$10.3 - 5.3 \cdot l_{gs}(v - 5.6) / 4$</td>
<td>$l_{gs}(v + 55.0) / (4.8) / 253.4 \cdot l_{gs}(v + 9.1) / 11.1$</td>
<td>$250.5 \cdot l_{gs}(v + 92) / 16$</td>
</tr>
<tr>
<td>$I_{Ca}$</td>
<td>$m^h$</td>
<td>$l_{gs}(v + 15.2) / 15.6$</td>
<td>$1.8 + 2.3 \cdot l_{gs}(v + 40.2) / 20.7$</td>
<td>$1 / (1 + [Ca] / 12.57)$</td>
<td>$0$</td>
</tr>
<tr>
<td>$I_{Na}$</td>
<td>$m^h$</td>
<td>$l_{gs}(v + 5.5) / 8$</td>
<td>$l_{gs}(v + 206.2) / 19.4397$</td>
<td>$l_{gs}(v + 206.2) / 19.4397$</td>
<td>$1 / (1 + [Ca] / 7.2)$</td>
</tr>
<tr>
<td>$I_{K_a}$</td>
<td>$m$</td>
<td>$l_{gs}(v + 84.3) / 6.4$</td>
<td>$46.9$</td>
<td>$l_{gs}(v + 29.7) / 19.4397$</td>
<td>$0$</td>
</tr>
<tr>
<td>$I_{K}$</td>
<td>$m$</td>
<td>$l_{gs}(v - v_{sat}) / 5$</td>
<td>$6$</td>
<td>$1 / (1 + [Ca] / 12.57)$</td>
<td>$0$</td>
</tr>
<tr>
<td>$I_{Na}$</td>
<td>$m^h$</td>
<td>$0.083^{+1} \cdot \exp \left( \frac{v + 48.73}{16.480} \right) + 0.333^{+1} \cdot \text{linoid} \left( \frac{v + 25.84}{9.155} \right)$</td>
<td>$0.083^{+1} \cdot \exp \left( \frac{v + 48.73}{16.480} \right)$</td>
<td>$0.333^{+1} \cdot \text{linoid} \left( \frac{v + 25.84}{9.155} \right)$</td>
<td>$1 / (1 + [Ca] / 10)$</td>
</tr>
<tr>
<td>$I_{K_a}$</td>
<td>$m^h$</td>
<td>$21.31^{+1} \cdot \exp \left( \frac{v + 47.3}{68.28} \right) + 0.639^{+1} \cdot \text{linoid} \left( \frac{v + 38.77}{8.535} \right)$</td>
<td>$21.31^{+1} \cdot \exp \left( \frac{v + 47.3}{68.28} \right)$</td>
<td>$0.639^{+1} \cdot \text{linoid} \left( \frac{v + 38.77}{8.535} \right)$</td>
<td>$1 / (1 + [Ca] / 10)$</td>
</tr>
<tr>
<td>$I_{Na}$</td>
<td>$m^h$</td>
<td>$0.0761^{+1} \cdot \exp \left( \frac{v + 102.41}{32.996} \right) / 29.98$</td>
<td>$0.710 + 5.453 \cdot \log \left( \frac{v - 5.98}{29.98} \right)$</td>
<td>$6.276 + 13.555 \cdot \log \left( \frac{v + 52.5}{16.550} \right)$</td>
<td>$1 / (1 + [Ca] / 12.57)$</td>
</tr>
</tbody>
</table>

The first column gives the name of the current, the second column gives the form of the gating factor (see Materials and Methods, Modeling voltage-gated currents), the third column gives the steady-state activation, the fourth column gives the time constant of activation, the fifth column gives the steady-state inactivation, and the sixth column gives the time constant of inactivation. Membrane potentials ($v$) are in millivolts, internal calcium concentration ([Ca$^2+$]) is in micromolars, and time constants ($\tau_a, \tau_h$) are in milliseconds. The function $\log(x)$ is equal to $\log(1 + \exp(-x))$, and $\text{linoid}(x)$ is equal to $x / (\exp(x) - 1)$. The parameter $v_{sat}$ was chosen at random when sampling the parameter space. In the baseline model, it had a value of $-45$ mV.

$$v_{asy}(t) = -57.6 + 11.425 \cos(2\pi \cdot f_0 \cdot t + 1.293) + 4.407 \cos(2\pi \cdot f_0 \cdot t + 0.503) + 1.581 \cos(2\pi \cdot f_0 \cdot t + 0.019) + 1.177 \cos(2\pi \cdot f_0 \cdot t + 0.095) + 0.786 \cos(2\pi \cdot f_0 \cdot t - 0.460) + 0.598 \cos(2\pi \cdot f_0 \cdot t - 7.240) + 0.305 \cos(2\pi \cdot f_0 \cdot t - 7.474). \quad (9)$$

For all synapses, the synaptic conductance was then given by the following equations:

$$g_{syn} = g_{syn} \cdot s \quad (10)$$

$$\tau_{syn} = \tau_{syn} \cdot s$$

where $g_{syn}$ is the synaptic conductance, $g_{syn}$ is the maximal synaptic conductance, $s$ is the fractional activation of the synapse, $\tau_{syn}$ is the synaptic time constant, $v_{pre}$ is the presynaptic voltage, $v_{thresh}$ is the synaptic threshold, and $v_{scale}$ sets the voltage sensitivity of the synapse. $s(x)$ is the Naka-Rushton function, given by $1 / (1 + x^{-2})$ for $x > 0$, and 0 otherwise.

For the AB and PD synapses, $v_{thresh} = -53$ mV and $v_{scale} = 3$ mV. For the AB and PY synapses, $v_{thresh} = 0$ ms, and for the PD synapse, $\tau_{syn} = 50$ ms.

**Construction of baseline model**

Once we had established reasonable descriptions of the intrinsic and synaptic currents in the LP neuron, we next wanted to establish a morphology for the model that mimicked the morphology of the biological LP and captured some rudimentary aspects of its electrophysiology. Additionally, we feared that a “brute force” search of possible morphologies and possible conductance distributions would be prohibitively time consuming. We therefore built a “baseline” model that was similar to LP in a number of ways: it had a morphology grossly consistent with the biological LP, it had spikes that were generated distally and were greatly attenuated in the soma, its input impedance was similar to that of LP (data not shown), its spike-frequency versus voltage curve was similar to that of LP, and it had voltage-gated currents similar to those of LP.

In developing the baseline model, we started with the simplest possible model and added complexity as needed to agree with the data. We eventually settled on a four-compartment model (Fig. 1) that combined a
good fit to the impedance over a range of frequencies (0.1–200 Hz), a reasonable spike height as recorded in the soma (−9 mV), and a reasonable resting membrane potential (approximately −40 mV).

We then added the reformulated voltage-gated conductances to the soma and neurite compartments of the four-compartment model (Fig. 1). Somatoneuritic currents (\(I_{\text{Na}}, I_{\text{Ca}}, I_{\text{K}}, I_{\text{K,Ca}}, I_{\text{leak}}\)) were present at uniform density in the soma and neurite compartments and were absent elsewhere. The synaptic currents were present at uniform density only in the neurite compartments, and the axonal currents (\(I_{\text{leak}}, I_{\text{Na}}, I_{\text{K}}, I_{\text{K,Ca}}\)) were present only in the axon compartment. All of these distributions are simplified versions of the true channel distributions in biological LP neurons (Hartline and Graubard, 1992; Baro and Harris-Warrick, 1998; Baro et al., 2000; French et al., 2004), but they are likely to be reasonable approximations at the level of detail we are interested in at present. The maximal conductances of most voltage-gated conductances were set to values close to those seen in the voltage-clamp recordings. The single exception to this was \(g_{\text{K,Ca}}\), which was set to a value −40% of the value seen in voltage-clamp recordings. We found that increasing \(g_{\text{K,Ca}}\) beyond this value led to endogenous bursting, which is not usually observed in the isolated LP in C. borealis. (It is not entirely clear why this is the case, but we suspect that the published voltage-clamp recordings were traces chosen for their large currents and that single LP neurons may not have all currents simultaneously large.)

Because the resulting model had physiological behavior that approximately matched that of the biological LP, with grossly reasonable maximal conductances, we declared this model an acceptable baseline model for later sampling of the parameter space. We note that the baseline model is not itself an admissible LP model. In particular, it bursts too late for later sampling of the parameter space. We note that the baseline conductances were zeroed, but the proctolin-activated conductance and the sodium conductance were not. We simulated this condition until the model came to apparent steady state and computed a number of features based on its steady-state waveform. These include the qualitative type of activity (silent, periodic spiker, aperiodic spiker, periodic nonspiker, or aperiodic nonspiker); the minimum, maximum, mean, and SD of the soma potential, axon potential, and somatic Ca\(^{2+}\) concentration; the number of spikes; the spike frequency; the mean spike duration; the mean and SD of the interspike intervals; the mean burst duration; the minimum, maximum, mean, and SD of the slow-wave somatic potential (the somatic potential if spikes are deleted, interpolated over, and the resulting waveform is low-pass filtered); the maximum spike height; and the cycle period. Some of these are only applicable for some qualitative activity types.

Third condition: periodic synaptic inhibition. Each model was subjected to a pattern of rhythmic synaptic inhibition that approximates what the LP neuron receives during the ongoing pyloric rhythm (see above, Modeling synaptic currents). In this condition, no conductances were zeroed. The synaptic currents were injected into the model until it reached apparent steady state, and the same scalar properties described above were computed based on the steady-state waveform in this condition. In more detail, the synaptic inputs were provided to each model for several cycles, using the same initial conditions as used for the spontaneous activity. After each cycle, the last cycle was compared with the penultimate cycle to determine whether the model had come to steady state. To be considered at steady state, four quantities had to be approximately equal on the last and the penultimate cycle: mean somatic voltage, SD of somatic voltage, mean somatic [Ca\(^{2+}\)], and SD of somatic [Ca\(^{2+}\)]. Voltages (both mean and SD) were considered “approximately equal” if they differed by <0.1 mV, and Ca\(^{2+}\) concentrations were considered approximately equal if they differed by <0.01 m-equiv.

Statistics
To test whether two parameters were significantly correlated in the population of 1304 LP models, we calculated the correlation coefficient (\(r\)) between the two parameters and used a permutation test to determine whether the calculated \(r\) differed significantly from zero. A significance level of \(\alpha = 0.001\) was used for each pair of parameters to facilitate comparison with Schulz et al. (2007).

Cubic fits
To develop a simple description of how a particular electrophysiological property depends on the model parameters, we fit cubic functions to a set of “data” consisting of a population of LP models. Each model had a set of parameters (maximal conductances, etc.) and a number of electrophysiological properties (resting membrane potential, slow-wave amplitude in response to rhythmic inhibition, etc.). One property at a time was fit, and each was treated as a scalar function of the parameters. To put all parameters on a common scale, each parameter was “z-scored” (the mean value of the parameter in the LP model population was subtracted off, and the SD was then divided out) before fits were performed.

A quadratic polynomial in 17 variables has 17 choose 2 coefficients (17 = 171 terms, each with an associated coefficient, and a full cubic model has 17 choose 3 coefficients).
17 + 3 choose 17 = 1140 terms/coefficients. A polynomial with a small number of coefficients will generally have good generalization performance but may not be able to fit the target function well. A polynomial with a large number of coefficients will be better able to fit the target but may suffer from overfitting and therefore not generalize well (Hastie et al., 2001). To have finer-grained control over the number of coefficients (and thus the fit-quality/generalization tradeoff), we used a subset of the 1140 possible terms, selected using a cross-validation procedure.

The LP models were first divided into a training set (90%) and a test set (10%). We used fivefold cross-validation on the training set to determine a set of terms that were likely to have an optimal fit on novel models. We varied the number of terms, starting with a single constant term, and added terms until we found a set of terms that gave the lowest cross-validation error. The next term to add was determined by temporarily adding all terms that could be obtained by increasing the degree of an existing term by 1 (not going beyond cubic terms), fitting the resulting polynomial to all of the training data, and retaining the most significant novel term (as determined by a t test). For this type of procedure, the cross-validation error at first decreases as the number of terms increases, but after a critical number of terms are added, the cross-validation error starts to increase. We then selected the simplest polynomial that had error not significantly worse than the minimum error, as assessed by a t test. For this type of procedure, the cross-validation error at first decreases as the number of terms increases, but after a critical number of terms are added, the cross-validation error starts to increase. We then selected the simplest polynomial that had error not significantly worse than the minimum error, as assessed by a t test. Each model was assessed in three different conditions. First, the LP neuron fires tonically from pyloric inputs, LP fires tonically in C. borealis (Fig. 2C) (Golowasch and Marder, 1992a). Previous work has shown that mRNA levels for a number of voltage-gated channels in LP are correlated (Schulz et al., 2007). Furthermore, for two of three channel types examined, these subunit mRNA levels correlate with the conductance of the channel (Schulz et al., 2006). To gain insight into whether these sorts of correlations might arise from electrophysiological constraints on LP, we constructed a population of multicompartment models of LP, in which maximal conductances were varied and their effects on electrophysiological properties were examined.

**Production of LP model population**

To generate the LP model population, we first developed a baseline model (see Materials and Methods) and then generated $\sim 6 \times 10^5$ models with randomized parameters in a large neighborhood around this baseline model (Fig. 3). For maximal conductances and maximal permeabilities, the range of possible values for each was from zero to approximately twice the value in the baseline model, so that each would span a realistic range. Maximal conductances are found to vary threefold to fourfold in this system (Golowasch and Marder, 1992a; Goldman et al., 2001; Golowasch et al., 2002; Schulz et al., 2006, 2007). Leak reversal potentials were varied over a 10 mV range centered at their baseline values, somewhat arbitrarily. The half-activation potential of the proctolin-activated conductance ($V_{1/2, pr}$) was varied over a 20 mV range centered at $-45$ mV, approximately consistent with the range observed experimentally (Golowasch and Marder, 1992b; Swensen and Marder, 2000) (Gouaillard, Taylor, Schulz, and Marder, unpublished observation). The parameters and their ranges are shown in Table 2. Each parameter was chosen from a uniform distribution within the given range, and each parameter was chosen independently of all other parameters. The capacitances and axial resistances of the model were fixed. This approach is similar to that taken in previous work (Foster et al., 1993; Goldman et al., 2001; Prinz et al., 2003, 2004b; Achard and De Schutter, 2006; Tobin et al., 2006; Hobbs and Hooper, 2008; Weaver and Wearne, 2008).

We chose 594,910 parameter sets in this way and performed simulations on each of the resulting models to measure their electrophysiological properties (see Materials and Methods). Each model was assessed in three different conditions. First, the

Figure 2. LP neuron morphology and activity. A, Two-photon laser-scanning confocal microscopy image of the LP neuron. B, Intracellular recording of the LP neuron during the ongoing pyloric rhythm. PD nerve (pdn) and pyloric nerve (pyn) recordings are included to show relative timing of the LP burst. Spikes on the pdn are from the two PD neurons, and the larger spikes on the pyn are from the PY neurons. C, Intracellular recording of the LP neuron when pyloric inputs have been blocked by applying $10^{-5}$M picrotoxin. The larger spikes on the A neuron play a critical role in the pyloric rhythm of decapod crustaceans. The LP neuron has a complex morphology (Fig. 2A, left gray bar), and it plays a critical role in the pyloric rhythm: during the first phase, it fires a burst of spikes (Fig. 2B). The phase at which the LP neuron fires is remarkably consistent from animal to animal, despite substantial variation in the pyloric frequency across animals (Bucher et al., 2005) (J. M. Gailllard, A. L. Taylor, D. J. Schulz, and E. Marder, unpublished observation). In the crab *C. borealis*, the waveform of the LP neuron reflects the three phases of the pyloric rhythm: during the first phase, when the pacemaker kernel is active (Fig. 2B, left gray bar), it is deeply hyperpolarized; in the second phase, it fires a burst of spikes; and in the third phase, it is inhibited by the PY neurons, but not as much as during the pacemaker activity (Fig. 2B, right gray bar). The bursting of LP is primarily network driven: when it is synaptically isolated from pyloric inputs, LP fires tonically in *C. borealis* (Fig. 2C) (Golowasch and Marder, 1992a).
model was subjected to a small voltage-clamp step. Second, the model was given no input, and its spontaneous behavior was observed. Third, a periodic synaptic input was delivered to mimic the pyloric inputs that LP receives during the ongoing pyloric rhythm. In the voltage-clamp condition, the model’s input conductance was measured. In the no-input condition, several properties were measured, including the model’s spontaneous activity pattern, its resting membrane potential, and its resting spike rate.

In the pyloric-like-input condition, measurements included the phase of burst onset, the phase of burst offset, the slow-wave amplitude, the peak slow-wave membrane potential, and the coefficient of variation of the interspike interval in the burst. All properties were automatically measured by the simulation code (see Materials and Methods).

We then examined this population of \( \sim 6 \times 10^5 \) models and selected only those with properties that were in a range that agreed with those of the biological LP neuron, using the bounds shown in Table 3.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Minimum</th>
<th>Maximum</th>
<th>Units</th>
</tr>
</thead>
<tbody>
<tr>
<td>Resting membrane potential (mV)</td>
<td>(-47.5)</td>
<td>(-32.5)</td>
<td></td>
</tr>
<tr>
<td>Resting spike rate (Hz)</td>
<td>13.1</td>
<td>30.6</td>
<td></td>
</tr>
<tr>
<td>Phase of burst onset (%)</td>
<td>32.0</td>
<td>44.0</td>
<td></td>
</tr>
<tr>
<td>Phase of burst offset (%)</td>
<td>61.7</td>
<td>74.9</td>
<td></td>
</tr>
<tr>
<td>Spike rate in burst (spikes/cycle)</td>
<td>16.3</td>
<td>30.2</td>
<td></td>
</tr>
<tr>
<td>Slow-wave amplitude (mV)</td>
<td>12.7</td>
<td>27.5</td>
<td></td>
</tr>
<tr>
<td>Peak slow-wave potential (mV)</td>
<td>(-47.5)</td>
<td>(-32.5)</td>
<td></td>
</tr>
<tr>
<td>ISI coefficient of variation in burst</td>
<td>0.2</td>
<td>0.25</td>
<td></td>
</tr>
</tbody>
</table>

In most cases, the bounds were chosen to contain the central \( \sim 85\% \) of the experimental data points (see Fig. 4, dashed lines). Resting spike rate was higher than those seen in experiments in which the LP neuron is isolated from pyloric network inputs, but similar to the spike rates within an LP burst. Input conductance was restricted to a range generally lower than previously reported experimentally (because the impedance measurements (data not shown) indicated a DC impedance generally larger than those reported previously), but the ratio between highest and lowest allowed values was similar to the ratio between the upper and lower bound of the central 85% of the experimentally measured values (Golowasch and Marder, 1992a).

Filtering the population of \( \sim 6 \times 10^5 \) models using these criteria yielded 1304 acceptable models of LP. Traces of model LP activity look reasonable, as shown in Figure 3. Both the spontaneous activity of these models (compare Figs. 2C, 3A) and their activity in response to pyloric-like input (compare Figs. 2B, 3B) look similar to recordings of biological LP neurons. In response to pyloric-like inputs, the model LPs generally displayed a burst phase, a phase in which they were mildly hyperpolarized (during which LP receives PY inhibition), and a phase of deep hyperpolarization (during which LP receives pacemaker inhibition). These three phases are present in recordings from biological LP neurons (Fig. 2B). Although strongly constrained electrophysiologically, the parameters of these models displayed wide variability (Fig. 3C) (also see below).

Histograms of the constrained properties are generally similar between model and experiment (Fig. 4). Consistent with this, the resting membrane potential (data not shown) was similar to that of the biological LPs (experiment: \(-46.8 \pm 6.5\) mV, \( n = 5 \); model: \(-44.1 \pm 2.1\) mV, \( n = 1304 \); mean \( \pm SD \)). Two additional properties that we did not explicitly constrain at this stage were also consistent with the biology. The height of spikes as measured in the soma was similar between model and experiment (experiment: \(7.7 \pm 1.8\) mV, \( n = 5 \); model: \(8.0 \pm 0.6\) mV, \( n = 1304 \); mean \( \pm SD \)). Also, the model impedances all matched the qualitative shape of the experimental impedance from 0.1 to 200 Hz (data not shown).

Weak correlations between pairs of parameters in the population of LP models

We examined whether the constraints imposed on the electrophysiological behavior of model LP neurons was reflected in the distributions of model parameters. Histograms of the model parameters for the 1304 models showed wide variability in most parameters (Fig. 5A). For most parameters, models could be found that spanned the sampled range. That is, when the range was binned into 10 bins, at least one model was found in all bins. Other parameters had restricted ranges in the LP population. The most strongly restricted parameter was \( g_{\text{synPY}} \) (which was restricted to about half the sampled range). \( g_{\text{KdA}} \) and \( g_{\text{synPY}} \) also had restricted ranges.

Does the population of 1304 model LP neurons exhibit any correlations between pairs of model parameters? This was of particular interest because previous experimental results demonstrated correlations between different voltage-gated conductances, some biophysically and some at the level of channel mRNA values (MacLean et al., 2005; Schulz et al., 2006, 2007). Therefore, we calculated correlation coefficients for each pair of parameters and tested these for significance (Fig. 5B) (see Materials and Methods). We found that correlations between different parameters in the LP population were generally weak (Fig. 5B) (red and blue backgrounds show significant correlations at \( \alpha = \)...
Although a rather stringent significance level of $\alpha = 0.001$ was used for individual comparisons (and to facilitate comparison to Schulz et al. [2007]), this corresponds to a family-wise significance level of $\alpha = 0.13$ ($n = 136$ comparisons, assuming comparisons are independent), which is rather generous. Of the 136 pairs, 126 (93%) had estimated correlation coefficients between $-0.2$ and $+0.2$. The strongest correlation was between $g_{Na}$ and $g_{Kda}$, which had a correlation coefficient ($r$) of $0.49 \pm 0.04$ [95% confidence interval (CI)].

To determine whether the observed correlations would likely be detected using typical experimental techniques, we picked a random subset of 20 models and tested for significant correlations (this time using a $t$ test, rather than the more computationally expensive permutation test). $n = 20$ was chosen because it is a reasonable value of $n$ for these types of experiments. This was repeated 2000 times, each time using a different random subset. A significant correlation (at $\alpha = 0.001$) between $g_{Na}$ and $g_{Kda}$ was found in 12.5 ± 1.5% (95% CI) of trials, and all other pairs were lower than this. Of the 136 pairs examined, 131 (96%) had a chance of discovery of <1%. Together, these results imply that the constraints imposed on the electrophysiological properties of the model LP neurons were not sufficient to induce the strong pairwise correlations observed experimentally. This implies that, at least in the model, strong correlations like those measured experimentally are not required for a population to have the electrophysiological properties of an LP neuron.

The LP models are part of a single “island” in parameter space

We were curious to know whether all of the LP models were part of a single connected island of LP cells in parameter space. That is, is it possible to continuously vary the parameters of one LP model and thereby transform it into any other LP model, and to do so in such a way that all intermediate models are also LP models? We cannot prove mathematically that this is the case, but we have strong circumstantial evidence that all of the LP models are part of a single connected volume in parameter space.

To establish this, we drew a line segment between pairs of models in parameter space and tested four equally spaced points on this line to see whether they also satisfied the criteria for an LP model. If they did, we considered the two models to be connected. In addition, if two models are each connected to a third model, then we considered the original two models to be connected. By performing this test on many pairs of models, we were able to establish that all 1304 LP models are connected to one another, at least provisionally. (Note, however, that it does not imply that the set is convex, and, in fact, we found many pairs of models that had non-LP models on the line between them but that were connected via a path involving one or more intercalated models.) This provides strong evidence that our population of LP cells are all part of a single island.

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Figure 3. Activity of six LP models, chosen at random. A. Spontaneous activity in six LP models, chosen at random. This is the model activity in the absence of synaptic inputs. Somatic potential is shown. B. Activity in response to pyloric-like inputs. Models are the same as in A, with corresponding colors. C. Model parameters for each model, scaled to the ranges used for sampling (see Table 2). Note that the models have very different patterns of maximal conductances and other parameters, despite having grossly similar activity. max, Maximum; min, minimum.
connected volume of LP cells in parameter space, rather than being in a number of islands of LP models separated by one or more “oceans” of non-LP models.

Polynomial fits of the property functions

It is highly desirable to have a quantitative description of how each parameter in a neuronal model influences each electrophysiological property. The computational model provides such a description only implicitly. A more direct description would be provided by a single equation that computes the magnitude of the electrophysiological property given the parameters. To provide such a description, we performed a multiple regression on each electrophysiological property, using the model parameters as predictors. This approach is similar to approaches used previously (Rabitz and Alis, 1999; Feng et al., 2004, 2006; Feng and Rabitz, 2004; Feng, 2005), but those approaches are not well suited to our LP population because our LP model population was not generated via independent variation of parameters (because of the filtering step). We first tried linear fits and quadratic fits to each electrophysiological property, but neither provided acceptable fits. A linear fit has the property that a given change in parameters will have the same effect on output, independent of the starting parameters; for instance, increasing $g_A$ by 3 $\mu$S/NF and $g_{syn}$ by 1 $\mu$S/NF will always increase the slow-wave amplitude by 2 mV, regardless of the starting parameters. This will obviously not be true for many parameter-to-property maps arising in complicated biological systems, and allowing quadratic and higher-order terms in the polynomial fit allows a fit to avoid this constraint.

We then tried cubic fits to each electrophysiological property and found that these performed well: we obtained $R^2$ values $>0.85$ for six of the eight properties we examined (Table 4). This indicates that these six properties, as functions of the parameters, were well approximated by cubic polynomials.

As an example, the coefficients of the fit to one of these, the slow-wave amplitude, are shown in Figure 6. The quality of the fit is shown graphically in Figure 6A, which plots the true slow-wave amplitude in each model versus the slow-wave amplitude “predicted” by the fit. The points are near the identity line, indicating that these six properties, as functions of the parameters, were well approximated by cubic polynomials.

![Figure 4. Comparison of pyloric activity in experiment and model. A, Histograms of onset phase of firing in experiment (exp) and models. B–E, Histograms of other properties of LP activity in the pyloric rhythm (exp) and in response to a pyloric-like pattern of synaptic inputs (model).](image-url)

The quadratic coefficients of the cubic fit reveal nonlinear effects of changing model parameters (Fig. 6C). Two different kinds of quadratic terms in the cubic fit are possible: one kind involves the square of a single parameter (along the diagonal in Fig. 6C), and the other involves the product of two different parameters (off the diagonal in Fig. 6C). In the case of the slow-wave amplitude fit, the largest quadratic coefficient was for the square of $v_{1/2,pr,}$ the half-activation of the proctolin-activated current. The coefficient for this term was negative, meaning that either increasing or decreasing $v_{1/2,pr}$ relative to its mean value will tend to decrease the slow-wave amplitude, counteracting somewhat the linear effect of decreasing $g_{synPD}$.

Cubic terms in the polynomial also describe nonlinear effects of changing model parameters, but involving products of three parameters, rather than two. These three parameters can all be the same, resulting in terms containing single parameters to the third power, or combinations of different parameters. In the slow-wave amplitude fit, the largest-magnitude cubic coefficient is for the cube of $g_{synPD}$, reflecting a nonlinear influence of $g_{synPD}$ on slow-wave amplitude that somewhat counteracts the linear $g_{synPD}$ term (because the cubic coefficient is negative whereas the linear coefficient is positive and has a larger magnitude). These sorts of insights into what parameters affect other properties can also be obtained by examining the coefficients of those fits, at least in cases where the fit is good.

Using polynomial fits to quantify the effect of each parameter on each electrophysiological property

We wanted to quantify the overall contribution of each model parameter to determining each of the neuron’s electrophysiological properties. We did this by starting with the coefficients of the final fit to a property, and setting all of the model parameters to their mean values. We then set them back to their actual values one parameter at a time and noted how much of the variance in the property was explained with the addition of each parameter,
as a fraction of the total variance explained by the complete fit. Typically, the amount of variance accounted for by each parameter varies depending on the order in which parameters are added. Therefore, we repeated this process 3000 times, with a random order each time, and averaged the variance added over all repeats. This yielded a measure of the influence exerted by each parameter on the final fit such that the influence of all parameters summed to 1. These “influence factors” are shown in Figure 7 for each property we examined. Interestingly, we found that essentially all properties are affected by many parameters and that the relative importance of parameters varied from property to property.

Comparing polynomial fitting to traditional sensitivity analysis

Fitting a polynomial to a model property can be viewed as an alternative to more traditional sensitivity analysis. In sensitivity analysis, one typically starts with a single best model, and then each model parameter is varied, one at a time, and the change in the model property is plotted versus each parameter (Butera et al., 1999; Hill et al., 2001, 2002; Jezzini et al., 2004). In theory, fitting a polynomial gives the same information but also tells one how the model responds to combinations of parameter changes. Nevertheless, we were curious to see how the polynomial fits compared with sensitivity analysis. For individual models in the

Figure 5. Distributions of parameters and pairs of parameters in a population of 1304 model LP cells. A, Histograms of individual parameter values. Each x-axis spans the sampled range. Each y-axis is different. B, Scatterplots of pairs of parameter values. Each scatterplot contains 1304 points, corresponding to each of the LP models. Because each pair of parameters can be plotted either as x versus y or y versus x, the scatterplot matrix is symmetric about the diagonal. Scatterplots with blue/red background displayed a statistically significant positive/negative correlation between the two parameters. Histograms highlighted in yellow are those that arguably correspond to channels that Schulz et al. (2007) found to be positively correlated at the level of mRNA.
indicated by low or even negative cases, the cubic fits did not predict the sensitivity curves well, as sensitivity analysis fairly well in most cases. However, in some indicates that the cubic fits were able to predict the results of property for which the cubic fit performed worst overall). This models was/extracted the fit property value at the central model from the fit in the neighborhood of each test model, using data from the square root of the variance of that property, calculated over all 1304 LP models. 

In many cases, the polynomial fit lines matched the sensitivity analysis curves fairly well (Fig. 8A). To quantify the quality of these fits, we calculated $R^2$ values for the cubic fit to each property in the neighborhood of each test model, using data from the sensitivity analysis around each test model. For this analysis, we were mainly interested in how well the cubic fits predicted changes from the property value at the test model, so we subtracted the fit property value at the central model from the fit values and similarly subtracted the simulation property value at the central model from the simulation values. (Failure to perform this step yielded $R^2$ values that were generally much lower.) This analysis yielded a range of $R^2$ values, but the median across test models was >0.5 for all but one property (phase of burst end, the property for which the cubic fit performed worst overall). This indicates that the cubic fits were able to predict the results of sensitivity analysis fairly well in most cases. However, in some cases, the cubic fits did not predict the sensitivity curves well, as indicated by low or even negative $R^2$ values (a negative $R^2$ value indicates a fit worse than would be achieved by simply fitting the data by a constant). Although the cubic fits are not a substitute for traditional sensitivity analysis, on the whole the ability of the cubic fits to predict the results of sensitivity analysis were striking, especially because the test models were not used in constructing the fits.

### Discussion

How does the combination of voltage-gated conductances in a neuron determine its electrophysiological properties? In this study, we analyzed a population of 1304 model neurons, all of which meet stringent criteria that match those of a specific biological neuron. We found (1) that a population of neurons can have a well constrained electrophysiological phenotype but still display disparate underlying maximal conductances with only weak correlations between conductances and (2) that each property of an identified neuron with well defined behavior is affected by a different set of maximal conductances. These results have implications for the interpretation of experimental results about the effect of neuronal parameters on electrophysiological properties.

### Table 4. Polynomial fits to model properties

<table>
<thead>
<tr>
<th>Property</th>
<th>RMSE for constant fit$^a$</th>
<th>Number of coefficients$^b$</th>
<th>Test RMSE$^c$</th>
<th>Test $R^2$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Input conductance (nS)</td>
<td>8.69</td>
<td>112</td>
<td>2.88</td>
<td>0.89</td>
</tr>
<tr>
<td>Resting membrane potential (mV)</td>
<td>2.12</td>
<td>107</td>
<td>0.48</td>
<td>0.94</td>
</tr>
<tr>
<td>Resting spike rate (Hz)</td>
<td>3.44</td>
<td>135</td>
<td>0.82</td>
<td>0.95</td>
</tr>
<tr>
<td>Phase of burst onset (%)</td>
<td>3.20</td>
<td>158</td>
<td>1.85</td>
<td>0.70</td>
</tr>
<tr>
<td>Phase of burst offset (%)</td>
<td>2.64</td>
<td>30</td>
<td>1.95</td>
<td>0.43</td>
</tr>
<tr>
<td>Spike rate in burst (spikes/cycle)</td>
<td>3.40</td>
<td>140</td>
<td>0.92</td>
<td>0.93</td>
</tr>
<tr>
<td>Slow-wave amplitude (mV)</td>
<td>3.54</td>
<td>151</td>
<td>0.65</td>
<td>0.96</td>
</tr>
<tr>
<td>Peak slow-wave potential (mV)</td>
<td>2.51</td>
<td>122</td>
<td>0.54</td>
<td>0.95</td>
</tr>
</tbody>
</table>

$^a$The root mean-squared error (RMSE) for the constant that best fits the model property. This is also equal to the square root of the variance of that property, calculated over all 1304 LP models.

$^b$The number of polynomial terms used in the best fit, as determined using cross-validation (see Materials and Methods).

$^c$The RMSE for the final polynomial fit, as evaluated on the test set.

$^d$The $R^2$ for the final polynomial fit, as evaluated on the test set.

Are correlations between maximal conductances required for a given electrophysiological phenotype? The population of LP models did not show strong correlations between pairs of model parameters, including maximal conductance parameters. This is interesting, because experimental work found strong correlations between mRNA copy numbers for voltage-gated channels in LP and other STG cells (Schulz et al., 2006, 2007). Although in some cases the mRNA levels correlate strongly with conductance magnitudes measured in voltage clamp (Schulz et al., 2006), it is not known whether mRNA copy numbers are well correlated with maximal conductance for all kinds of channels in all neuron classes.

There are two possible explanations for the lack of strong correlations in the model population: either the model fails to capture something important about the neuron’s behavior or the neuron’s electrical firing properties do not require the correlations seen in the molecular studies. Consistent with the latter explanation, it is possible that the correlations observed experimentally do not arise from electrophysiological constraints, since it is possible to construct LP neurons with conductance ratios other than those observed experimentally and still “act” like LP neurons electrophysiologically. If this were the case, the ratios observed experimentally might be a consequence of the transcriptional regulation of channel genes (Latchman, 1997; Fan et al., 2000; Brivian Lou and Darnell, 2002; Rosati and McKinnon, 2004). For instance, having subunits of multiple channel types under the control of a common transcription factor might be simpler than independent transcriptional control of each channel type. This could result in different subunits being expressed in fixed ratios. Thus, the presence of correlations between conductances does not automatically imply that these correlations are required for proper electrophysiological behavior.

### Using polynomial fits to capture the relative roles of multiple conductances on neuronal activity

The cubic fits allowed the quantification of the influence of each parameter on each electrophysiological property (Fig. 7). Each property was affected, at least to some extent, by many parameters in the model, but different properties were affected by various parameters to different extents. Recent work on a model of globus pallidus neurons yielded a similar result (Gunay et al., 2008), suggesting that this is a feature common to many neurons. 

The large influence of the axonal conductances, and particularly the axonal leak and $I_A$ conductance ($g_{IA}$), on the input conductance was somewhat unexpected. This suggests that, at least in the model, the axon is not as electrically irrelevant to the current–voltage relationship at the soma as might be supposed given the extreme attenuation of spikes in the soma. Although the contributions of multiple parameters on different intrinsic properties were generally different, those for the resting spike rate and the spike rate in the burst were similar. This reflects the fact that these two properties were strongly correlated in the LP model population (data not shown).

Small influence factors should be interpreted with caution; the synaptic conductances ($g_{synABB}$, $g_{synPDF}$, and $g_{synPY}$) had small, but nonzero, influence factors for the input conductance, resting membrane potential, and resting spike rate, despite the fact that all of these conductances were set to zero when these properties are measured in the model. This is possible because of high-order correlations between these parameters and parameters that do affect these properties. These correlations are exploited by the fitting routine to give significantly better fits than if the synaptic conductance parameters are not used (data not shown). These
potentially misleading influence factors are uniformly small and do not detract from the utility of the influence factors overall.

The finding that each electrophysiological property is determined by a combination of several different conductances (Fig. 7) has important implications for how one interprets experiments involving pharmacological blockers, genetic deletions, and neuromodulation. When the levels of conductances vary from animal to animal, and when an electrophysiological property depends on multiple conductances, blocking one of these conductances may have highly variable effects on that property, because in some animals the conductance may already be low and be compensated for by the other conductances to keep the property at a normal level. A similar argument holds for genetic deletions, but with the added complication that if the deletions are nonacute, the animal may compensate for the knock-out by regulating levels of other conductances (Guo et al., 2000; Zhang et al., 2002; Xu et al., 2003; Zhou et al., 2003). Similarly, neuromodulator effects may vary from preparation to preparation because neuromodulators act on a particular set of conductances, which may be high or low depending on the individual animal.

Approaches to understanding the effect of parameters on electrophysiological properties

In a model, the simplest approach to understanding the effect of parameters on electrophysiological properties is traditional sensitivity analysis: identifying one “best” set of parameters, varying one parameter at a time, and plotting the electrophysiological property of interest against the varied parameter. This approach has been fruitfully applied (Butera et al., 1999; Hill et al., 2001, 2002; Jezzini et al., 2004) but has two limitations: it is based on examining departures from a single model, and it is based on varying one parameter at a time, and therefore yields no information about simultaneously changing multiple parameters.

Another approach is to generate a population of models, ei-
ther by picking random values for parameters or by systematically scanning the parameter space (Foster et al., 1993; Goldman et al., 2001; Prinz et al., 2003, 2004b; Achard and De Schutter, 2006; Tobin et al., 2006; Hobbs and Hooper, 2008; Weaver and Wearne, 2008). One can then use visualization techniques to determine how electrophysiological properties change as parameters are varied (Ward et al., 1994; Keim, 2000; Taylor et al., 2006), although these techniques are often qualitative and choosing a good method of visualization can be difficult.

In some cases, one can derive closed-form expressions for model properties as functions of parameters (Abbott, 1990), and this is arguably the best solution, when it is possible.

In this work, we used cubic functions to fit electrophysiological properties as a function of model parameters. This allowed a relatively simple description of model behavior over a population of models (not just in the vicinity of a single model), and it described how these models respond to multiple simultaneous parameter changes. Similar methods have been used in the chemistry literature (Rabitz and Alis, 1999; Feng et al., 2004, 2006; Feng and Rabitz, 2004; Feng, 2005) but have not been commonly used to understand neuronal dynamics. We focused here on electrophysiological properties that were real functions of the parameters. For example, the slow-wave amplitude is described by a single real number for each possible choice of parameters. Other cases of interest include properties that are integer (e.g., number of spikes per burst), category (e.g., activity type), or boolean (e.g., spiking or not) functions of the parameters. Fitting polynomials might be useful for integer properties if followed by a rounding operation but would not be ideal for category properties. For these sort of problems, an approach analogous to the one used here would be to use statistical classification (as opposed to regression) techniques (Hastie et al., 2001).

A related problem is finding the set of points in parameter space that all result in the same value for an electrophysiological property (the level set). Olypher and Calabrese (2007) used a numerical approximation to find the level sets of several electrophysiological properties. The method used here offers a more general answer to how parameters affect electrophysiological behavior over a population of models (not just in the vicinity of a single model), and it described how these models respond to multiple simultaneous parameter changes. Similar methods have been used in the chemistry literature (Rabitz and Alis, 1999; Feng et al., 2004, 2006; Feng and Rabitz, 2004; Feng, 2005) but have not been commonly used to understand neuronal dynamics. We focused here on electrophysiological properties that were real functions of the parameters. For example, the slow-wave amplitude is described by a single real number for each possible choice of parameters. Other cases of interest include properties that are integer (e.g., number of spikes per burst), category (e.g., activity type), or boolean (e.g., spiking or not) functions of the parameters. Fitting polynomials might be useful for integer properties if followed by a rounding operation but would not be ideal for category properties. For these sort of problems, an approach analogous to the one used here would be to use statistical classification (as opposed to regression) techniques (Hastie et al., 2001).

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**Relation to previous models of STG neurons**

These LP models capture many important features of the biological LP neuron in C. borealis, including the response of LP to pyloric inputs, the attenuation of spikes in the soma, and LP’s input impedance. Previous models of LP have captured some of these features but not all of them (Buchholtz et al., 1992; Nowotny et al., 2008). One difference between the behavior of the model and that of the biological LP neuron (Golowasch and Marder, 1992a) is that the model fires more rapidly than the biological neuron at rest, but without this we were not able to obtain the appropriate firing rates in response to inhibitory inputs.

Previous multicompartement models of STG neurons have had only two compartments, one to model the generation of spikes and another that contains the currents typically measured in voltage-clamp experiments (Soto-Trevino et al., 2005; Nowotny et al., 2008). Additional compartments were needed to accurately model both the input impedance of LP (data not shown) and the attenuation of spikes when measured in the soma.

**Conclusions**

In this work, we found that a population of neurons can have a well constrained electrophysiological phenotype but still display disparate underlying maximal conductances with only weak correlations between conductances, and that each property of an identified neuron with well defined behavior is affected by a different subset of several maximal conductances. These results have important implications for how one interprets experiments involving pharmacological blockers, genetic deletions, and neuromodulation.

**References**

**Figure 8.** Sensitivity analysis and cubic fits in the neighborhood of test models. \(A\). Each panel shows the result of varying one parameter at a time. Note the close agreement between simulations and the cubic fit in most cases. Missing points in the \(g_{Na}\) and \(g_{K}\) panels indicate parameter values for which the models no longer spiked. The gray backdrop indicates axonal parameters. \(B\), Summary of comparisons between sensitivity analysis and cubic fits. Box-and-whisker plots show the distribution of \(R^2\) values found when comparing the sensitivity data and cubic fits for each property. (The \(R^2\) values were calculated using only the changes relative to the respective central values; see Results.) Each box-and-whisker plot summarizes the distribution of \(R^2\) values for all 130 test models. The upper and lower limits of each box give the 75th and 25th percentiles of each distribution, and the line in the middle of each box gives the median. The dashed “whiskers” give the extent of all nonoutlier data points. A data point was classified as an outlier if it was more than 1.5 box heights away from the box. Outliers are shown as diamond symbols. Some properties had a few points with \(R^2\) values below the lower limit of the plot. The number of such points for the different properties was 5, 17, 0, 58, 8, 2, and 7, respectively. The large black circle indicates the \(R^2\) value for the example shown in \(A\). Properties not in bold had poor overall \(R^2\) values (see Table 4).

<table>
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<th>Parameter</th>
<th>(g_{Na})</th>
<th>(g_{K})</th>
<th>(\Delta E_{\text{leak}}) (mV)</th>
<th>(\Delta A)</th>
<th>(\Delta F_{\text{Ca}})</th>
<th>(\Delta F_{\text{synP}})</th>
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Eisen JS, Marder E (1982) Mechanisms underlying pattern generation in lobster stomatogastric ganglion as determined by selective inacti-
crustacean stomatogastric nervous system. In: Dynamic biological networks: the stomatogastric nervous system (Harris-Warrick RM, Marder E, Selverston AI, Moulines M, eds), pp 31–86. Cambridge, MA: MIT.


