Bistable nerve conduction

Zhaoyang Zhang1 and Zhilin Qu1,2,*
1Department of Medicine, David Geffen School of Medicine, University of California, Los Angeles, Los Angeles, California and 2Department of Computational Medicine, David Geffen School of Medicine, University of California, Los Angeles, Los Angeles, California

ABSTRACT It has been demonstrated experimentally that slow and fast conduction waves with distinct conduction velocities can occur in the same nerve system depending on the strength or the form of the stimulus, which give rise to two modes of nerve functions. However, the mechanisms remain to be elucidated. In this study, we use computer simulations of the cable equation with modified Hodgkin-Huxley kinetics and analytical solutions of a simplified model to show that stimulus-dependent slow and fast waves recapitulating the experimental observations can occur in the cable, which are the two stable conduction states of a bistable conduction behavior. The bistable conduction is caused by a positive feedback loop of the wavefront upstroke speed, mediated by the sodium channel inactivation properties. Although the occurrence of bistable conduction only requires the presence of the sodium current, adding a calcium current to the model further promotes bistable conduction by potentiating the slow wave. We also show that the bistable conduction is robust, occurring for sodium and calcium activation thresholds well within the experimentally determined ones of the known sodium and calcium channel families. Since bistable conduction can occur in the cable equation of Hodgkin-Huxley kinetics with a single inward current, i.e., the sodium current, it can be a generic mechanism applicable to stimulus-dependent fast and slow conduction not only in the nerve systems but also in other electrically excitable systems, such as cardiac muscles.

INTRODUCTION

The major function of the nerve system is to transmit information via electrical excitation and conduction for the body or parts of the body to respond to environmental changes. The response time is determined by the conduction velocity (CV) of the nerve cable. It is well established that CV is determined by the membrane conductance of the sodium (Na+) current (I_{Na}) and calcium (Ca^{2+}) current (I_{Ca}), the radius of the cross section of the nerve cable, temperature, and myelination, etc. (1–4). Namely, a larger Na+ or Ca^{2+} current conductance gives rise to a steeper upstroke of the action potential (or a faster depolarization) to result in a larger CV. A larger nerve cable or myelination effectively enhances the diffusive coupling to give rise to a larger CV. Temperature affects both the ionic currents and the coupling to affect conduction. CV in different nerve fibers or species spans over a wide range, differing in orders of magnitude (1). These conduction properties can be well described by the cable equation with the Hodgkin-Huxley (HH) model (5–10).

Besides the regular conduction properties, an interesting conduction behavior was observed experimentally, i.e., stimulus-dependent fast and slow conduction waves occur in the same nerve fiber. For example, Aglantha digitale, a species of jellyfish, has two modes of swimming—a slow swimming for fishing and a fast swimming for escaping away from predators (11,12). In an experimental study (13), Mackie and Meech showed that the slow and fast swimming modes were caused by a slow conduction (~0.3 m/s) and a fast conduction (~1.4 m/s) in the same motor giant axon, respectively. The slow conduction is a

SIGNIFICANCE Stimulus-dependent fast and slow conduction waves in the same nerve system have been shown experimentally, which give rise to two modes of nerve functions. We use computer simulations to show that the fast and slow waves are a bistable conduction caused by a positive feedback loop of the wavefront upstroke speed, mediated by the sodium channel inactivation properties, which is further potentiated by the presence of the calcium current. The mechanism is robust with respect to the experimentally determined activation thresholds of the known sodium and calcium channel families. The theoretical insights provide a generic mechanism for stimulus-dependent fast and slow conduction in the nerve systems, which may be also applicable to other electrically excitable tissues, such as cardiac muscles.
low-amplitude and long-duration wave mediated by $I_{Ca}$ and the fast conduction is a high-amplitude and short-duration wave mediated by $I_{Na}$ (Fig. S1). The fast wave was induced by an external stimulus and the slow wave was spontaneous and endogenous. This same conduction dynamics was also shown in experiments of conduction in cockroach giant axons by Hochner and Spira (14), who demonstrated that two distinct conduction waves (0.1–0.6 and 3–6 m/s) occurred in the same axon treated with ethanol. Differing from *Aglantha digitale*, the slow conduction in the cockroach giant axon was not mediated by $I_{Ca}$ but still by $I_{Na}$. Besides conduction in the giant axons of jellyfish and cockroaches, evidence of stimulus-dependent fast and slow conduction has also been shown in experiments of rat visual cortex (15,16), i.e., the CV of the spontaneous waves differ from those of the evoked waves in the same cortex. These experimental observations imply that, besides the regular conduction behavior, a new conduction dynamics can occur in the nerve systems, i.e., the same nerve fiber or tissue can exhibit two stable conduction states depending on the initial conditions. As in *Aglantha digitale*, the two conduction behaviors of the same fiber accomplish two distinct biological functions. However, the underlying mechanism remains to be elucidated.

In this study, we investigate the mechanisms of the stimulus-dependent fast and slow conduction using analytical methods and computer simulations in a cable equation of the HH model (17) with modifications (see Methods). An $I_{Ca}$ formulation is added to the HH model to investigate the role of $I_{Ca}$. We show that stimulus-dependent fast and slow conduction capturing the experimental observations can occur in the cable with the modified HH model. This is a bistable behavior, namely bistable conduction, emerging during conduction in the cable, caused by a positive feedback loop of the wavefront upstroke speed mediated by the $Na^+$ channel inactivation properties. The addition of $I_{Ca}$ can further potentiate bistable conduction. Using simulations of randomly selected parameter sets, we show that the bistable conduction mechanism is robust, i.e., the activation thresholds of $I_{Na}$ and $I_{Ca}$ for bistable conduction detected in a wide range of parameters are well within the experimentally determined activation thresholds of the known $Na^+$ and $Ca^{2+}$ channel families. Since the bistable conduction is mediated by the $I_{Na}$ alone in the HH model, it is likely a generic mechanism applicable to conduction not only in the nerve systems, but also in other electrically excitable media, such as cardiac muscles.

**METHODS**

**Mathematical model**

The HH model (17) with modifications is used to simulate action potential conduction in a one-dimensional cable with the following partial differential equation for voltage ($V$):

$$\frac{\partial V}{\partial t} = - \frac{I_{ion} + I_{stim}}{C_m} + D \frac{\partial^2 V}{\partial x^2},$$

(1)

where $C_m = 1 \text{ mF/cm}^2$ is the membrane capacitance, and $D = 0.0045 \text{ cm}^2/\text{ms}$ is the diffusion constant. $I_{stim}$ is the stimulus current density and $I_{ion}$ is the total ionic current density consisting of different types of ionic currents, i.e.,

$$I_{ion} = I_{Na} + I_{Ca} + I_K + I_L.$$

(2)

In Eq. 2, $I_{Na}$ is the $Na^+$ current density described by $I_{Na} = G_{Na}m^3h(V - E_{Na})$. $I_K$ is the potassium ($K^+$) current density described by $I_K = G_Kn^4(V - E_K)$. $I_L$ is the leak current density described by $I_L = G_L(V - E_L)$. $I_{Ca}$ is the $Ca^{2+}$ current density described by $I_{Ca} = G_{Ca}d^2f(V - E_{Ca})$. This formulation is adopted from Medlock et al. (18) with the addition of an inactivation gate $f$, $m$, $h$, $n$, $d$, and $f$ are gating variables which are described by the following type of differential equations:

$$\frac{dy}{dt} = (y_m - y)/\tau_y.$$

(3)

In Eq. 3, $y_m = \frac{m^4}{n^2}$, and $\tau_y = \frac{1}{a \cdot \tau_y}$ in which $\alpha$ and $\beta$ are rate constants and functions of $V$.

To observe bistable conduction, we make modifications to the original HH kinetics. We shift the kinetics (see Table 1) to give rise to a resting potential of around $-65 \text{ mV}$ (the measured resting potential in *Aglantha digitale* is close to this value (13)). Some of the shifts are required for facilitating bistable conduction. Besides the voltage shifts, we also alter the magnitudes of the time constants $\tau_y$ by multiplying a prefactor $\gamma_{y}$, i.e.,

$$\tau_y(V) \rightarrow \gamma_{y} \cdot \tau_y(V).$$

(4)

**TABLE 1 Changes of parameters from the original HH model**

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Original HH model</th>
<th>Modified HH model</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\alpha_{Na}$</td>
<td>0.1 $25 - V$</td>
<td>0.1 $-35 - V$</td>
</tr>
<tr>
<td>$\beta_{Na}$</td>
<td>$4 \exp \left( \frac{-V}{10} \right)$</td>
<td>$4 \exp \left( \frac{-35 - V}{10} \right)$</td>
</tr>
<tr>
<td>$\alpha_{k}$</td>
<td>0.07 $20 - V$</td>
<td>0.07 $-75 - V$</td>
</tr>
<tr>
<td>$\beta_{k}$</td>
<td>$1 \exp \left( \frac{30 - V}{10} \right)$</td>
<td>$1 \exp \left( \frac{-45 - V}{10} \right)$</td>
</tr>
<tr>
<td>$\alpha_{l}$</td>
<td>0.01 $10 - V$</td>
<td>0.01 $-15 - V$</td>
</tr>
<tr>
<td>$\beta_{l}$</td>
<td>$0.125 \exp \left( \frac{-V}{80} \right)$</td>
<td>$0.125 \exp \left( \frac{-25 - V}{80} \right)$</td>
</tr>
<tr>
<td>$E_{Na}$</td>
<td>120 mV</td>
<td>55 mV</td>
</tr>
<tr>
<td>$E_{K}$</td>
<td>$-12 \text{ mV}$</td>
<td>$-77 \text{ mV}$</td>
</tr>
<tr>
<td>$E_{L}$</td>
<td>$10.6 \text{ mV}$</td>
<td>$-65 \text{ mV}$</td>
</tr>
</tbody>
</table>

$\alpha_{Na}$ and $\beta_{Na}$ are shifted 60 mV toward more negative voltages. $\alpha_{k}$ and $\beta_{k}$ are shifted 75 mV toward more negative voltages. $\alpha_{l}$ and $\beta_{l}$ are shifted 25 mV toward more negative voltages. $E_{Na}$ and $E_{K}$ are shifted 65 mV toward more negative voltages, and $E_{L}$ is shifted 75.6 mV. The shifts are to change the resting potential from around zero in the original HH model to $-65 \text{ mV}$ and to facilitate bistable conduction in the cable.
For \( d_u \) and \( f_u \), we use the following formulations:

\[
d_u = \frac{1}{1+e^{(-V_{tu})}} \quad \text{and} \quad f_u = \frac{1}{1+e^{(-V_{fu})}}
\]

\( \tau_d \) and \( \tau_f \) are set as constants independent of \( V \).

The default parameters are set as: \( G_{Na} = 36 \text{ mS/cm}^2 \), \( G_{Ca} = 0.3 \text{ mS/cm}^2 \), \( \gamma_{Na} = 0.2 \), \( \gamma_{Ca} = 0.35 \), \( \gamma_{I} = 3 \), \( \tau_d = 3 \) ms, \( \tau_f = 20 \) ms, \( k_d = 5.8 \), and \( k_f = 6 \). The values of \( G_{Na} \) and \( G_{Ca} \) are stated in the figure legends.

We also simulate a nonspatial single-point (or single-cell) model by dropping the diffusion term in Eq. 1, which becomes an ordinary differential equation, i.e.,

\[
\frac{dV}{dt} = - \frac{I_{stim} + I_{Na}}{C_m}.
\]

Note that after the modifications, the model is still in the excitable regime with a single stable fixed point, i.e., the resting potential. Traveling waves are initiated by \( I_{stim} \) which is applied at the beginning of the cable. Details of \( I_{stim} \) are stated in the figure legends. Bistable fixed points and oscillatory solutions of Eq. 5 can occur when we use random parameter sets and we exclude these sets from our data shown in Fig. 4.

### Numerical methods

A forward Euler method is used for numerical simulation of Eq. 1 with the following discretization:

\[
V_i(t + \Delta t) = V_i(t) - \frac{I_{Na}}{C_m} \Delta t
\]

\[
+ D \left[ \delta \frac{V_{i+1}(t) + V_{i-1}(t)}{\delta x^2} - 2V_i(t) \right] \Delta t,
\]

with \( \Delta x = 0.045 \) cm and \( \Delta t = 0.005 \) ms. No-flux boundary condition is used. The numerical method is stable when \( \Delta t < \frac{\Delta x}{D} = 0.225 \) ms. The \( \Delta t \) we use is much smaller than this value, ensuring the numerical stability. Eq. 5 is also numerically solved using the Euler method with \( \Delta t = 0.005 \) ms. The gating variables (Eq. 3) are integrated using the method by Rush and Larsen (19), i.e.,

\[
y(t + \Delta t) = y_t - [y_t - y(t)]e^{-\Delta t/\tau_t}
\]

### RESULTS

#### Bistable conduction in the cable equation

To observe bistable conduction in the cable equation with the HH model (17), we modify the model as described in Methods. To examine the role of \( I_{Ca} \), we add an \( I_{Ca} \) formulation to the model. Fig. 1, A and B show a fast wave (1.4 m/s) induced by a strong stimulus and a slow wave (0.21 m/s) induced by a weak stimulus, respectively. The action potential of the fast wave (inset in Fig. 1 A) exhibits a steep upstroke, a high amplitude (≈95 mV, from −65 to 30 mV), and a short duration (≈10 ms), while that of the slow wave (inset in Fig. 1 B) exhibits a shallow upstroke, a low amplitude (≈45 mV, from −65 to −20 mV), and a long duration (≈40 ms). These features recapitulate well the experimental observations in the jellyfish (see Fig. S1) and cockroach experiments.

To show how the bistable conduction in the cable is affected by \( I_{Na} \) and \( I_{Ca} \), we scan the maximum conductance of \( I_{Na} (G_{Na}) \) and that of \( I_{Ca} (G_{Ca}) \), and plot the CV in color in the two-parameter plane (Fig. 1 C). For each parameter set, a strong stimulus and a weak stimulus, as in Fig. 1, A and B, are applied to elicit conduction waves. The gray region in Fig. 1 C is where bistable conduction occurs. The white region is where both stimuli fail to elicit conduction (conduction failure). The colored regions (I, II, and III) are where both stimuli give rise to a single stable conduction with CV color coded. Regions I and II exhibit stable conduction mediated by \( I_{Na} \), and region III exhibits stable slow conduction mediated by \( I_{Ca} \). The presence of \( I_{Ca} \) promotes bistable conduction; however, \( I_{Ca} \) is not required since the bistable conduction occurs in the absence of \( I_{Ca} \) (\( G_{Ca} = 0 \)). In the absence of \( I_{Ca} (G_{Ca} = 0) \), bistable conduction occurs in an intermediate range of \( G_{Na} \), and the fast conduction occurs when \( G_{Na} \) is either above or below this range. Therefore, bistable conduction can occur in the HH model mediated by \( I_{Na} \) alone without requiring the presence of \( I_{Ca} \) or another inward current. In Fig. 1, D and E, we plot CV versus \( G_{Na} \) for \( G_{Ca} = 3 \) μA/cm² and \( G_{Ca} = 0 \), respectively, showing typical hysteresis, a hallmark of bistability.

#### Mechanism of bistable conduction

Since bistable conduction can occur in the absence of \( I_{Ca} \), we investigate how it occurs in the cable without \( I_{Ca} \) focusing on the role of \( I_{Na} \). Fig. 2 A shows a fast wave induced by a stimulus slightly above the critical strength and a slow wave by a stimulus slightly below the critical strength in the absence of \( I_{Ca} \). In the first 10 ms, the action potentials are almost identical for the two cases, which then bifurcate (as indicated by the arrow) into a stable fast wave and a stable slow wave. Since the stimuli are very close to the critical strength, the two waves in the first 40 ms are close to the solution of the unstable conduction. The open circles in Fig. 1 D are calculated from the unstable conduction. In the three conduction behaviors, besides the difference in the amplitude and duration of the action potentials, another important difference is the upstroke speed of the wavefront. For the degeneration from the unstable conduction to the stable fast wave, the upstroke speed of the wavefront becomes faster and faster until reaching the steady-state conduction. For the degeneration from the unstable conduction to the stable slow wave, the upstroke speed of the wavefront becomes slower and slower until reaching the steady-state conduction. Because of the different upstroke speeds, the inactivation of \( I_{Na} \) is different. In Fig. 2 B, we plot the steady-state inactivation curve of \( I_{Na} \) (\( h_{Na}, \text{ thick green} \)) and the trajectories of the three types of conduction in the V-h plane. The trajectory of the
steady-state slow wave \((\text{thick red})\) undergoes a path that is very close to the steady-state inactivation curve, while the trajectory of the steady-state fast wave \((\text{thick blue})\) undergoes a path that is far away from the steady-state curve. The thin red and blue trajectories are the transient ones degenerating from the steady-state unstable wave \((\text{dashed})\) to the two stable waves, respectively. In other words, the steady-state unstable trajectory \((\text{dashed})\) is the separatrix of the two stable waves.

As indicated by the two cases above, a slower upstroke speed causes more \(I_{Na}\) inactivation, which in turn results in an even slower upstroke in the next conduction site, or vice versa, forming a positive feedback loop. Therefore, the formation of the stable fast and slow conduction in the cable is a result of the positive feedback in upstroke speed of the wavefront mediated by \(I_{Na}\) inactivation, result in two modes of \(I_{Na}\) activation/inactivation. This causes bistable conduction to occur in the intermediate range of \(G_{Na}\). When \(G_{Na}\) is too large, the high \(I_{Na}\) mode is always activated and thus only the fast wave occurs. When \(G_{Na}\) is too small, the low \(I_{Na}\) is not large enough to support the slow wave and thus only the high \(I_{Na}\)-mediated fast wave is observed. However, \(I_{Ca}\) can rescue the slow wave and thus extends the bistable conduction in a wider parameter space (Fig. 1 C).

To further understand the mechanism of bistable conduction, we obtain an analytical solution using a two-variable model, which includes only \(I_{Na}\) and \(I_L\), i.e.,

\[
\frac{\partial V}{\partial t} = \frac{\partial^2 V}{\partial x^2} + \sigma_{Na} m(V) h - G_{Ca}(V - E_L)
\]

\[(8)\]

\[
\frac{\partial h}{\partial t} = \frac{h(V) - h}{\tau_h},
\]

\[(9)\]

where \(V\) is voltage and \(h\) the inactivation gating variable of \(I_{Na}\). \(\sigma_{Na}\) is a parameter proportional to \(G_{Na}\). \(m(V)\) and \(h(V)\) are Heaviside functions:

\[
m(V) = \begin{cases} 
0, & V < V_m \\
1, & V \geq V_m 
\end{cases}
\]

\[(10)\]

and

\[
h(V) = \begin{cases} 
1, & V < V_h \\
h_0, & V \geq V_h 
\end{cases}
\]

\[(11)\]

By using the moving coordinate system and continuation conditions, one can obtain that the CV \((c)\) satisfies the...
properties during the fast and slow waves. The thick green line is the steady-state slow wave and the thin red curves are the ones recorded from the transition period from the unstable conduction in the absence of I_{Ca}. The fast wave is induced by a stimulus (I_{stim} = 22.32155910275 \mu A/cm^2 with 20 ms in duration) slightly above the critical strength, and the slow wave by a stimulus (I_{stim} = 22.32155910270 \mu A/cm^2 with 20 ms in duration) slightly below the critical strength. Voltage traces are recorded at sites 0.225 cm apart along the cable. The blue and red traces are overlapped until a distance away (marked by the arrow) from the stimulation site. G_{Na} = 92 mS/cm^2 and G_{Ca} = 0. (A) Na^+ channel activation properties during the fast and slow waves. The thick green line is the steady-state inactivation curve of the Na^+ channel (h). The thick red curve with arrows is the trajectory of the steady-state slow wave and the thin red curves are the ones recorded from the transition period from the unstable conduction to the stable slow conduction. The thick blue curve with arrows is the trajectory of the steady-state fast wave and the thin blue curves are the ones recorded from the transition period from the unstable conduction to the fast fast conduction. The black dashed curve is the separatrix of the two types of waves, which is the trajectory taken from the unstable conduction marked in (A) before the arrow. To see this figure in color, go online.

following equation (see supporting material for detailed derivation):

\[ - (V_m - E_L) \sqrt{c^2 + 4G_L} + \sigma_{Na} \left( \beta + \frac{h_0}{G_L} \right) - \frac{\beta}{\tau_h c} = 0, \]

where \( \delta = \frac{c + \sqrt{c^2 + 4G_L}}{2}, \beta = \frac{(1 - h_0) e^{\beta \delta}}{\tau_h (c + \sqrt{c^2 + 4G_L})}, \) and \( \varepsilon = \frac{\ln \left( \frac{V_m - E_L}{c - \delta} \right)}{\varepsilon - \beta}. \) One can rewrite Eq. 12 in the form as \( \sigma_{Na} = f(c, \tau_h, V_m, h_0, G_L, E_L), \) i.e., \( \sigma_{Na} \) is expressed as a nonlinear function of \( c. \) Fig. 3 A plots \( c \) versus \( \sigma_{Na} \) calculated using this formulation with other parameters fixed, showing that \( c \) is bistable in a certain range of \( \sigma_{Na}. \) One can also numerically solve Eq. 12 to obtain \( c \) when the parameters are given. Fig. 3 B shows the conduction behaviors versus \( \sigma_{Na} \) and \( \tau_h, \) showing that the bistable region decreases as \( \tau_h \) increases. We perform additional simulations of the one-dimensional cable with the HH model to investigate the effects of I_{Na} kinetics on the conduction behaviors. Fig. 3 C shows the conduction behaviors versus \( \tau_m \) and \( \tau_h \) (activation time constant of I_{Na}), and Fig. 3 D shows the conduction behaviors versus \( \sigma_{Na} \) and \( \tau_h. \) Note that the phase diagram in Fig. 3 D is similar to that in Fig. 3 B, indicating that theoretical predictions agree well with the numerical simulation results of the HH model.

**Robustness of bistable conduction**

In the case of Aglantha digitale (13), based on their observation that the fast wave is mediated by I_{Na} and the slow wave by I_{Ca}, the authors hypothesized that to facilitate the fast and slow conduction in the same axon, the Na^+ channel activation threshold is much higher than the Ca^2+ channel activation threshold so that during I_{Ca}-mediated conduction (the slow wave), the peak voltage remains low enough to avoid activation of I_{Na}. On the other hand, in the case of cockroach experiments (14), since the slow wave was still mediated by I_{Na}, the authors then hypothesized that there might exist another type of I_{Na} with a low activation threshold in the same axon to explain their observations. In other words, these authors hypothesized that the fast and slow conduction are mediated separately by two inward currents with a certain required difference in activation thresholds (we call it dual-threshold hypothesis). In the original HH model, since there is only one inward current (i.e., I_{Na}), it can only exhibit a single-threshold response (Fig. 4 A). On the other hand, after adding I_{Ca} to the HH model, it can exhibit a dual-threshold response in which two types of action potentials occur depending on the stimulus strength (Fig. 4 B). Since there are two distinct types of action potentials, one would expect that each type will give rise to a conduction in the cable, resulting in stimulus-dependent fast and slow conduction. This raises a question: which of the two mechanisms is robust or more likely to occur in the real systems?

To address this question, we perform the following investigations. We first carry out a large number of simulations of the cable equation with randomly selected parameters in assigned intervals (see details in supporting material). We detect the parameter sets giving rise to both fast and slow conduction. We then use these parameter sets to perform simulations using the nonspatial single-point model (Eq. 5) to filter out the single-threshold and dual-threshold responses by examining the stimulus-response relationship as shown in Fig. 4, A and B. The parameter sets (open dots) exhibiting single-threshold responses are plotted in Fig. 4 C and those exhibiting dual-threshold responses in Fig. 4 D. The data points are plotted on the plane of the half-activation voltages for Na^+ channels.
(V_{Na,1/2}) and Ca^{2+} channels (V_{Ca,1/2}). The definitions and ranges of V_{Na,1/2} and V_{Ca,1/2} in the model are detailed in supporting material. We also plot the experimentally measured V_{Na,1/2} (red squares) and V_{Ca,1/2} (green circles) surveyed from literature by Catterall et al. (20,21). V_{Na,1/2} for the known 10 members of the Na^+ channel family ranges from −56 to −8.8 mV (20), and V_{Ca,1/2} for the 10 members of the Ca^{2+} channel family ranges from −46 to 14 mV (21). V_{Na,1/2} and V_{Ca,1/2} are plotted in a way so that all possible combinations of the 10 members of the Na^+ channel family and the 10 members of the Ca^{2+} channel family fall inside the dashed red box. Therefore, we argue that the parameter sets fall inside the box are the ones that can occur in the real system.

The data sets exhibiting single-threshold response fall both inside and outside the dashed box (Fig. 4 C). We plot the distributions of CV for the fast and slow conduction in Fig. 4 E, and the ratios of the fast-to-slow CV in Fig. 3 F. The average slow CV is 0.23 ± 0.08 m/s and the average fast CV is 1.10 ± 0.32 m/s. The average of the fast-to-slow CV ratio is 7.0 ± 8.84. The fast-to-slow CV ratios were 5–10 in the experimentally measured CVs in the giant axons of Aglantha digitale (13) and cockroaches (14), which are in the same range as obtained in the simulations. Therefore, the bistable conduction mechanism agrees well with experimental data and is robust.

The data sets exhibiting dual-threshold response fall into two groups (Fig. 4 D). One group is completely outside the box (lower right), which occurs for V_{Na,1/2} is 40 mV higher than V_{Ca,1/2}. The other group can still fall into the box (upper left), which occurs for V_{Ca,1/2} is 30 mV higher than V_{Na,1/2}. However, some caveats of this group of data are worth noting. The slow wave is mediated by I_{Na} and the fast wave by I_{Ca}, which is not what occurs in either the jellyfish (13) or the cockroaches (14). Although the data sets were filtered through the dual-threshold criterion using the nonspatial single-point model (Eq. 5), when we check the behaviors in the cable, the Na^+ channel exhibit similar response as in Fig. 2, indicating that the mechanism may still be bistable conduction as in the single-threshold case. For example, when we remove I_{Ca} from the data sets in Fig. 4 D, we can still observe bistable conduction. To show this, we perform the same simulations using the parameter sets in Fig. 4 D without I_{Ca} (see Fig. S2 A), the parameter sets outside the box disappear, but about 50% of the parameter sets inside the box are retained. Another evidence to support this is that if we slow the Na^+ channel inactivation, this group of parameter sets will disappear. We perform the same simulations as in Fig. 4 except that the inactivation is slowed by increasing \( \gamma_{h} = 0.35 \) to \( \gamma_{h} = 1 \) (see Fig. S2 B). The upper group disappears, but the lower group is retained and still outside the dashed box. The results shown in Figs. 4 and S2 indicate that the dual-threshold mechanism is difficult to be satisfied in the real systems.

In the model, the resting potential is set at −65 mV. If we lower the resting potential, e.g., to −75 mV, we observe almost the same results (Fig. S3) as those shown in Fig. 4 and S2 except that the minimum \( I_{Na} \) or \( I_{Ca} \) activation threshold for bistable conduction has a roughly 10 mV shift due to the lowering of the resting potential.
DISCUSSION

Nerve conduction can be well described by the cable equation with the HH model (5–9). It is well known that the cable equation with FitzHug-Nagumo (FHN) model or the HH model exhibits monostable conduction (10). In this study, we show that the cable equation with the HH model can exhibit bistable conduction in which a fast and a slow stable conduction occur in the same cable depending on the stimulus strength. The bistable behavior is a result of the positive feedback of the wavefront upstroke speed mediated by the Na$^+$ channel inactivation properties. In other words, this positive feedback cause two stable modes of $I_{Na}$ activation, which occurs when the Na$^+$ channel inactivation is relatively fast (see also Fig. 3). Unlike the fast conduction that occurs as long as $G_{Na}$ is greater than a critical value, bistable conduction can only occur in an intermediate range of $G_{Na}$. This is because when $G_{Na}$ is too large, the high $I_{Na}$ mode is always activated and thus only the fast wave can occur. When $G_{Na}$ is too small, the low $I_{Na}$ cannot support a stable slow conduction, and thus only the fast conduction can occur. However, the failed slow conduction can be rescued by the addition of $I_{Ca}$, which can substantially extend the bistable conduction regime (see Fig. 1 C). We use an analytical treatment of the simplified cable model to demonstrate that bistable conduction can be mediated by a single inward current, namely $I_{Na}$. Using simulations with randomly drawing parameter sets, we also show that the bistable conduction is robust, which can occur well within the experimentally determined ranges of the activation thresholds of the known Na$^+$ and Ca$^{2+}$ channel families.

Our computer simulation results agree well with the experimental observations of electrical conduction in the giant axons of jellyfish and cockroaches. For example, the action potential profiles in the fast and slow waves and the ratios of the fast-to-slow CV in the simulations agree with those shown in experiments in both jellyfish and cockroaches. More importantly, our theoretical study unifies the seemingly different experimental observations in jellyfish and cockroaches to the same general mechanism. In the jellyfish experiments, Mackie and Meech (13) showed that the slow wave was blocked by Ca$^{2+}$ channel blockers, and thus concluded that the slow wave was mediated by $I_{Ca}$. This led them to hypothesize that the activation threshold of $I_{Na}$ has to be much higher than that of $I_{Ca}$ to allow the two waves to occur. As shown in our simulations (Fig. 1 C), although $I_{Ca}$ is not required for bistable conduction, it potentiates bistable conduction by rescuing the slow conduction. Therefore, blocking it will suppress the slow conduction, agreeing with the experimental observation. On the other hand, in the cockroach experiments, Hochner and Spira (14) found that the slow wave was blocked not by Ca$^{2+}$ channel blockers but by Na$^+$ channel blockers,
differing from the jellyfish experiments. They then hypothesized that a low-threshold $I_{Na}$ must be responsible for the slow wave. Our simulations showed that $I_{Ca}$ or another low-threshold $I_{Na}$ is not required since a single $I_{Na}$ could produce both the slow and fast conduction via the bistable conduction dynamics. In other words, the fast and slow conduction observed in cockroaches may originate from the same $I_{Ca}$. A moderate $I_{Na}$ reduction may block the slow wave but not the fast wave while a strong $I_{Na}$ reduction can block both waves (see Fig. 1, C and E). Therefore, the experimental observations in both jellyfish and cockroaches can be explained by the same mechanism in different parameter regimes.

Besides the giant axons in jellyfish and cockroaches, bistable conduction may also occur in other nerve systems. For example, it was shown that the spontaneous waves and the evoked waves exhibit distinct CVs in the same rat visual cortex (15,16), which could be a result of bistable conduction. In addition, bimodal CV distributions were widely observed in sensory and motor nerves (22–28), which were traditionally attributed to the size difference of the fibers. However, bistable conduction can be a candidate mechanism for bimodal CV distributions, which needs to be verified in future experimental studies.

Note that, although the duration of action potential in the slow wave (~40 ms) shown in Fig. 1 B matches quantitatively well with the ones measured in experiments (Fig. S1 and Fig. 1 in Meech and Mackie (29)), but the action potential of the fast wave in Fig. 1 A has a duration of about 10 ms, which is much longer than the one recorded in the experiment (~2 ms, see Fig. S1). The main cause of action potential lengthening is due to the shift of the $K^+$ channel kinetics (see Table 1), which reduces activation of the $K^+$ current. The purpose of shifting the $K^+$ channel kinetics is to result in bistable conduction in the HH model. However, there are multiple $K^+$ currents in the giant axon of Aglantha digitale. In the experimental studies by Meech and Mackie (29,30), they showed that there are three kinetically distinct $K^+$ currents in the giant axon of Aglantha digitale, and these currents behave like the A-type current that is widely observed in neurons (31). If one adds an A-type current with a proper conductance into the cable model (Fig. S5), one can shorten the action potential to quantitatively match the ones recorded in experiments (compare the action potentials in Fig. S5 with the ones in Fig 1 in Meech and Mackie (29)), but still retains the bistable conduction behavior.

Different modes of conduction in the same cable have been shown in the HH model (3,32) as well as in the FHN model (10,33), i.e., for the same parameter set, there are a fast and a slow conduction. However, since the slow conduction is unstable and decremental, the two conduction modes are not a bistable behavior. A bistable behavior in the cable model with the FHN equations was demonstrated by Rinzel and Terman (34), in which two action potential repolarization behaviors were observed in the same cable during conduction. Namely, in one state, the action potential repolarizes normally but, in the other state, the action potential fails to repolarize. However, since in the two states, the upstrokes of wavefront are roughly the same, the conduction velocities do not exhibit two distinction values. In other words, it is a bistable behavior in repolarization not in depolarization, differing from the bistable conduction shown in this study. Note that the bistable conduction is mediated by fast Na$^+$ channel inactivation, which is absent in the FHN model, and thus the bistable conduction behavior cannot occur in the FHN-type models.

Finally, as shown in our simulations (Fig. 4), the dual-threshold mechanism is theoretically plausible; however, it requires a very large minimum difference in the activation thresholds of $I_{Na}$ and $I_{Ca}$. Meech and Mackie (29,35) showed that the slow conduction is mediated by the T-type Ca$^{2+}$ current and fast conduction by the Na$^+$ current, with the Na$^+$ current activation threshold being much higher than that of the Ca$^{2+}$ current. However, using the data of Na$^+$ and Ca$^{2+}$ channel activation threshold surveyed by Catterall et al. (20,21), we show that this mechanism is much less robust, i.e., the condition for the mechanism cannot be easily satisfied in the real systems. On the other hand, the bistable conduction mechanism can be easily achieved without requiring any gap or correlation between the activation thresholds of the two types of ionic currents. Moreover, the bistable conduction mechanism may also have an evolutionary advantage over the dual-threshold mechanism since a single rather than two inward currents can provide two survival functions as in the jellyfish. Although it is unclear what are the functional roles of the fast and slow conduction in other species or diseases, as it is well known that bistability is a ubiquitous phenomenon in biology and responsible for many biological functions (36–39), we believe that bistable conduction may also play important roles in nerve functions under healthy and diseased conditions. As shown in our study, bistable conduction occurs in the cable equation with the HH model with modifications, we believe that it is a generic mechanism that is applicable to not only the nerve systems but also other electrically excitable tissues, such as cardiac muscles.

**SUPPORTING MATERIAL**

Supporting material can be found online at [https://doi.org/10.1016/j.bpj.2022.08.006](https://doi.org/10.1016/j.bpj.2022.08.006).

**AUTHOR CONTRIBUTIONS**

Z.Q. conceived the project, supervised the research, provided funding, and wrote the manuscript. Z.Z. performed the simulations and mathematical analyses. Z.Q. and Z.Z. analyzed the results and edited the manuscript.
ACKNOWLEDGMENTS

This study was supported by National Institutes of Health grants R01 HL134709 and R01 HL139829.

DECLARATION OF INTERESTS

The authors declare no competing interests.

REFERENCES