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A Codimension-2 Bifurcation Controlling Endogenous Bursting Activity and Pulse-Triggered Responses of a Neuron Model

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Introduction

The dynamics of individual neurons are crucial for producing functional activity in neuronal networks. An open question is how temporal characteristics can be controlled in bursting activity and in transient neuronal responses to synaptic input. Bifurcation theory provides a framework to discover generic mechanisms addressing this question. We present a family of mechanisms organized around a global codimension-2 bifurcation. The cornerstone bifurcation is located at the intersection of the border between bursting and spiking and the border between bursting and silence. These borders correspond to the blue sky catastrophe bifurcation and the saddle-node bifurcation on an invariant circle (SNIC) curves, respectively. The cornerstone bifurcation satisfies the conditions for both the blue sky catastrophe and SNIC. The burst duration and interburst interval increase as the inverse of the square root of the difference between the corresponding bifurcation parameter and its bifurcation value. For a given set of burst duration and interburst interval, one can find the parameter values supporting these temporal characteristics. The cornerstone bifurcation also determines the responses of silent and spiking neurons. In a silent neuron with parameters close to the SNIC, a pulse of current triggers a single burst. In a spiking neuron with parameters close to the blue sky catastrophe, a pulse of current temporarily silences the neuron. These responses are stereotypical: the durations of the transient intervals—the duration of the burst and the duration of latency to spiking—are governed by the inverse-square-root laws. The mechanisms described here could be used to coordinate neuromuscular control in central pattern generators. As proof of principle, we construct small networks that control metachronal-wave motor pattern exhibited in locomotion. This pattern is determined by the phase relations of bursting neurons in a simple central pattern generator modeled by a chain of oscillators.

Abstract

The dynamics of individual neurons are crucial for producing functional activity in neuronal networks. An open question is how temporal characteristics can be controlled in bursting activity and in transient neuronal responses to synaptic input. Bifurcation theory provides a framework to discover generic mechanisms addressing this question. We present a family of mechanisms organized around a global codimension-2 bifurcation. The cornerstone bifurcation is located at the intersection of the border between bursting and spiking and the border between bursting and silence. These borders correspond to the blue sky catastrophe bifurcation and the saddle-node bifurcation on an invariant circle (SNIC) curves, respectively. The cornerstone bifurcation satisfies the conditions for both the blue sky catastrophe and SNIC. The burst duration and interburst interval increase as the inverse of the square root of the difference between the corresponding bifurcation parameter and its bifurcation value. For a given set of burst duration and interburst interval, one can find the parameter values supporting these temporal characteristics. The cornerstone bifurcation also determines the responses of silent and spiking neurons. In a silent neuron with parameters close to the SNIC, a pulse of current triggers a single burst. In a spiking neuron with parameters close to the blue sky catastrophe, a pulse of current temporarily silences the neuron. These responses are stereotypical: the durations of the transient intervals—the duration of the burst and the duration of latency to spiking—are governed by the inverse-square-root laws. The mechanisms described here could be used to coordinate neuromuscular control in central pattern generators. As proof of principle, we construct small networks that control metachronal-wave motor pattern exhibited in locomotion. This pattern is determined by the phase relations of bursting neurons in a simple central pattern generator modeled by a chain of oscillators.
Bifurcation theory explains how dynamical systems like neurons precisely change their dynamics in response to the variation of a controlling parameter like neuromodulatory tone or descending drive [1–3,6,11,12,30–51]. A variety of network mechanisms producing specific temporal patterns of activity have been previously studied [1,10,18,31,34,41,52–60].

Our study emphasizes the dynamics of single neurons as an organizing principle underlying pattern formation. We present a global codimension-2 bifurcation and assert that this so-called cornerstone bifurcation can precisely control the regulation of temporal characteristics in periodic and transient neuronal activity. We suggest that this bifurcation generates a family of cellular mechanisms which can answer the aforementioned questions. These mechanisms also explain the operation of conditional oscillators in parameter space near a functional bursting regime.

In neurons, control over regimes of activity is ubiquitously executed through neuromodulation. Potassium currents and hyperpolarization-activated currents are common targets for neuromodulation [61–68]. Among the usual targets of neuromodulation are the kinetic parameters controlling the voltage dependence of conductances of ionic currents such as a non-inactivating potassium current, \( I_{K2} \), and a hyperpolarization-activated current, \( I_h \). \( I_{K2} \) is an outward current that is activated during the burst and creates a mechanism controlling termination of the burst. To serve this role, the activation variable of this current must be significantly slower than the duration of a single spike. On the other hand, \( I_h \) is an inward current that is activated during the silent phase of bursting activity, and it creates a mechanism controlling the duration of the silent phase. The voltage of half-activation of \( I_{K2} \), \( \theta_{K2} \), could control burst duration, and the voltage of half-activation of \( I_h \), \( \theta_h \), could control the interburst interval. Here, we present a model in which the dependence of burst duration and interburst interval on \( \theta_{K2} \) and \( \theta_h \) can be quantitatively described by inverse-square-root laws.

Biophysically accurate neuronal models allow the utilization of well developed techniques from bifurcation theory. In a model, bifurcations can predict the dependence of temporal characteristics of oscillatory regimes near the bifurcation [11,12,42,43,48]. In type I spiking neuronal dynamics, a saddle-node bifurcation on an invariant circle (SNIC) describes a transition from tonic spiking into silence; the interspike interval obeys the inverse-square-root law imposed by the bifurcation. The period of spiking grows proportionally to \( 1/\sqrt{\pi-2 \pi^2} \) where \( \pi^2 \) is the bifurcation value of the parameter \( \pi \) and \( \pi \gg \pi^2 \). In our case, the bifurcation parameters are either \( \theta_{K2} \) or \( \theta_h \). The blue sky catastrophe, a special case of the saddle-node bifurcation for orbits, imposes the same inverse-square-root law on burst duration [45,69]. A saddle-node bifurcation for periodic orbits can control the transition from bursting into tonic spiking [11,12,43,70]. When the criteria for the SNIC and the blue sky catastrophe are simultaneously satisfied, a global bifurcation of codimension-2 occurs [69,70]. By coregulating these currents, we reveal the cornerstone bifurcation and how the bifurcation parameters control burst duration and interburst interval, latency to spiking in the response of a spiking neuron to inhibition, duration of a burst elicited by stimulation of a silent neuron, and multistability of spiking and silence. We apply these mechanisms to construct simple proof of principle models of CPGs. One of these mechanisms describes the coregulation of two biophysical parameters in the vicinity of the cornerstone bifurcation such that the duty cycle of bursting in a single cell is maintained across a wide range of periods. Connected into a chain, such oscillators self-organize their activity into a metachronal wave pattern, which is preserved against variation in period.

We present this model in regard to crawling behavior in *Drosophila* larvae.

**Results**

**Regimes of Activity**

We developed a generic low-dimensional Hodgkin-Huxley type neuronal model stemming from a model of the leech heart interneuron under certain pharmacological conditions [44,45]. In order to reduce the mathematical complexity of the system, we simulated the activity of the interneuron in bath with Co²⁺ and 4-aminoypyridine (4-AP). Application of Co²⁺ blocks Ca²⁺ currents and the synaptic current. Application of 4-AP blocks most of the K⁺ currents. The remaining currents in this model are the leak current, the non-inactivating potassium current, \( I_{K2} \), the fast sodium current, \( I_{Na} \), and a constant polarizing current. Our model also includes the hyperpolarization-activated current, \( I_h \), which is present in this pharmacological scenario but has not been previously represented in our reduced models. Intracellular recordings in these conditions show slow seizure-like oscillations with periods that are tens of seconds long [71,72].

The slow variable in this system is the activation of \( I_{K2} \), \( m_{K2} \); its time constant was 2 s. As such, the temporal characteristics of these variables are instrumental in the dynamics of bursting activity. By systematically manipulating parameters that determine the dynamics of the slow variables, we observed the range of bursting activity and bifurcations between qualitatively distinct regimes of neuronal activity. We investigated the effects of changes to the kinetics of \( I_{K2} \) and \( I_h \).

The parameters \(-\theta_{K2} \) and \(-\theta_h \) represent the voltages of half-activation of the variables \( m_{K2} \) and \( m_h \). We conducted an empirical investigation of the model by systematically varying \( \theta_{K2} \) and \( \theta_h \). The model exhibited a variety of regimes. Silence and tonic spiking corresponded to equilibria and periodic orbits. Bursting activity was either periodic or weakly chaotic. For low values of \( \theta_{K2} \) and \( \theta_h \) (region I), the model exhibited tonic spiking (Fig. 1A). For high values of \( \theta_{K2} \) and low values of \( \theta_h \) (region II), the model exhibited bursting activity (Fig. 1A). For high values of \( \theta_{K2} \) and \( \theta_h \) (region III), the model was silent (Fig. 1A). For low values of \( \theta_{K2} \) and high values of \( \theta_h \) (region IV), the model exhibited bistability of tonic spiking and silence (Fig. 1A).

We determined the boundaries between bursting and tonic spiking and between bursting and silence by integrating the system over a range of values for \( \theta_h \) and \( \theta_{K2} \). Samples varied from one another in the burst duration and the interburst interval (Table 1). The transition from bursting to tonic spiking occurred near \( \theta_{K2} = -0.0105 \) \( V \) almost independently of \( \theta_h \) (Fig. 1A). The transition from bursting to silence occurred near \( \theta_h = 0.0413 \) \( V \) with weak dependence on \( \theta_{K2} \) (Fig. 1A). As the parameters were changed to approach the transition from bursting to tonic spiking or from bursting to silence, the burst duration or interburst interval increased, respectively. We described the changes in the bursting wave form by duty cycle: the ratio of burst duration to cycle period. The duty cycle generally increased as \( \theta_{K2} \) or \( \theta_h \) decreased (Fig. 1A).

**The Inverse-Square-Root Laws Control Bursting Activity**

We associated saddle-node bifurcations of equilibria and periodic orbits with the transitions between different regimes. The transitions between I and IV and between II and III coincided with the saddle-node bifurcation for equilibria (Fig. 1A, black curve). The borders between regions I and II and between regions III and IV coincided with the saddle-node bifurcation for periodic orbits (Fig. 1A, grey curve). The location of the
The codimension-2 bifurcation was interpolated as the intersection of the curves representing the saddle-node bifurcation for equilibria and the saddle-node bifurcation for periodic orbits at $(\theta_{K2}, \theta_h)$: $(\theta_{K2}, \theta_h) = (-0.0105 V, 0.0413564925 V)$, $(\theta_{K2}, \theta_h) = (-0.0075 V, 0.041326 V)$, $(\theta_{K2}, \theta_h) = (-0.0075 V, 0.038 V)$, and $(\theta_{K2}, \theta_h) = (-0.0105 V, 0.038 V)$. The point at (B) is near but not on the codimension-2 bifurcation point.

We show that the saddle-node bifurcations along the border between I and II and between II and III are a blue sky catastrophe and a SNIC, respectively, through a series of curve fits and analysis of slow motion (Text S1; Text S2). We investigated temporal characteristics of bursting near these bifurcation curves (Text S1; Fig. S1). The system was directly integrated for a series of parameter values approaching the bifurcation values. As $\theta_h$ approached the saddle-node bifurcation for equilibria, the interburst interval grew in a fashion asymptotic to the parameter value of bifurcation (Fig. S1A–B). The parameter $\theta_{K2}$ was similarly varied so as to approach its critical value for the saddle-node bifurcation for periodic orbits. Samples of bursting activity for values of $\theta_{K2}$ successively closer to the bifurcation value showed an asymptotic increase in the burst duration (Fig. S1C–D). The curve...
fits of the burst duration and interburst interval conformed to the $1/\sqrt{x-x^2}$ temporal law at multiple locations along the bifurcation curves (Text S1). We also confirmed these results for the value of the time constant of activation of the potassium current 0.9 s used in the [45].

The Inverse-Square-Root Laws of Pulse Triggered Responses

For parameter values in regions I and III (Fig. 1), tonic spiking and silence were attracting regimes. However, a perturbation from the tonic spiking regime in I or the rest state in III triggered transient activity that shared the inverse-square-root laws with the corresponding characteristics of periodic bursting in region II (Fig. 2). For example, activity at parameter values in region III was quiescent, but a brief hyperpolarizing pulse of appropriate duration and amplitude triggered a single burst (Fig. 2 A-C). These individual bursts closely resembled the characteristic waveforms of bursting activity observed in Region II. We investigated this parameter space by observing pulse-triggered bursts at parameter values such that $\theta_0$ was fixed at 0.0415 $V$ and $\theta_{K^2}$ approached the value for the saddle-node bifurcation for periodic orbits at the border between regions III and IV. As $\theta_{K^2}$ approached the value of the bifurcation, the burst duration of pulse triggered bursts grew (Table 2). Even though there was no closed periodic orbit in region III, the phase point slowed down as it passed near the ghost of the saddle-node periodic orbit. We performed a curve fit of this data to the function $f_{K^2}(\theta_{K^2})$ (Eq. 1 in Text S1) (Fig. 2D).

We carried out a similar analysis for the border of the transition from regions I to IV. When tonic spiking activity was perturbed with a hyperpolarizing pulse, the system spent some time in a transient hyperpolarized silent state before returning to spiking activity. The trajectory during this time interval resembled the trajectory during the interburst interval of an endogenously bursting cell. We fixed $\theta_{K^2}$ at $-0.0107$ $V$, and we sampled region I for values of $\theta_0$ close to the saddle-node bifurcation for equilibria (Fig. 2E-G). As $\theta_0$ grew, the latency to spiking grew (Table 2). As the parameter $\theta_0$ took values close to the saddle-node bifurcation for equilibria, the phase point spent more and more time near the ghost of the saddle-node equilibrium. We quantified the dependence of the latency to spiking on $\theta_0$ by performing a curve fit to the function $f_{K^2}(\theta_0)$ (Eq. 2 in Text S1) (Fig. 2H).

The Manifolds of Slow Motion

This model exhibit bursting of square-wave type. The bursting phase is controlled by the activation of $I_{K^2}$. Over the course of the burst, $m_{K^2}$ incrementally grows with each spike. Vice versa, $m_{K^2}$ controls the shape of each spike. The shape of each spike depends on the value of $m_{K^2}$, and the next increment of $m_{K^2}$ depends on the shape of the spike. To account for this mutual interaction and to formalize the progression of the burst towards termination, we used slow-fast decomposition. This technique allows us to describe this process in terms of the dynamics of one variable: the slow variable.

We considered the model as a slow-fast system. The variable $m_{K^2}$ was the slow subsystem, and $\{V$, $\theta_{Na}$, $m_{K^2}\}$ composed the fast subsystem. The magnitude of $m_{K^2}$ controls rest states (equilibria) and spiking (periodic orbits) of the fast subsystem. These equilibria and periodic orbits are located on manifolds that could be determined by decoupling the slow-fast system and reintroducing the slow variable as a parameter [35,49]. Trajectories of the full system stay near these manifolds of slow motion [35,49,73]. The slow motion manifold that determines periodic orbits describes the breadth of specifications of spike shapes in the fast subsystem.

The slow motion manifold that determines the silent phase describes the characteristics of equilibria of the fast subsystem. We used a more accurate approach, whereby we manipulated a parameter of the slow variable ($\theta_{K^2}$) to compute the manifolds of slow motion in the full system [45–47,74]. We performed slow-fast decomposition and determined the slow motion manifolds to study the structure of the codimension-2 bifurcation. We applied the Pontryagin & Rodygin averaging method [45–47,74]. The stability of the regimes in the full system was determined by bifurcation analysis (Text S2).

The parameter space that we analyzed is partitioned by two bifurcation curves: one horizontal and one vertical (Fig. 3). The horizontal curve is a saddle-node bifurcation for equilibria ($\text{SNI}_E$ in Fig. S2). It divides the 2-D parameter space into two subspaces (red curve in Fig. 3A). At this bifurcation, a stable equilibrium disappears; the parameter space above this curve supports the stable equilibrium. The vertical solid blue curve represents a saddle-node bifurcation for periodic orbits (Fig. 3A; $\text{SNI}_O$ Fig. S2). At this bifurcation, a stable periodic orbit—representing tonic spiking—disappears; the parameter space to the left of this curve supports tonic spiking.

The cornerstone bifurcation, which is located at the intersection of the two bifurcation curves (red and blue solid curves), organizes bursting, spiking, and silent regimes (Fig. 3A). The bifurcations defining the borders of region II (SNI and blue sky catastrophe) lead to qualitative changes in the vector field which eliminate periodic bursting by obstructing the passage of the phase point over one or both manifolds of slow motion (Fig. 3A).

At the codimension-2 bifurcation point, the slow motion manifold determined by the equilibria of the fast subsystem is obstructed by a saddle-node equilibrium, and the slow manifold for spiking is obstructed by a saddle-node bifurcation for periodic orbits (Fig. 3B).

Perturbations in the ($\theta_0$, $\theta_{K^2}$) parameter space from the codimension-2 point lead to the disappearance of the saddle-node equilibrium or simple periodic orbit or cause the saddle-node orbit or equilibrium to split in two (Fig. 3C–F).

In region I, a stable orbit and a saddle orbit were found on the slow manifold for spiking (Fig. 3C). The stable periodic orbit corresponds to the tonic spiking regime, and is the only attracting regime in this region. Perturbations can reveal characteristics of the slow manifold associated with the equilibrium of the fast subsystem. The phase point can spend a significant amount of time in a quiescent state before firing resumes after the cessation of an inhibitory perturbation (Fig. 2E-G). This perturbation causes a fast transition of the phase point from a stable orbit onto the slow-motion manifold of equilibria for the fast subsystem. The phase point followed the slow-motion manifold, and the neuron exhibited a transient quiescence. For parameter values near the border between regions I and IV, the phase point could spend a long time near the ghost of the saddle-node equilibrium before the onset of firing.

---

Table 1. Temporal characteristics of bursting for different parameter values in region II.

<table>
<thead>
<tr>
<th>Figure 1: (B)</th>
<th>(C)</th>
<th>(D)</th>
<th>(E)</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\theta_{K^2}$ ($V$)</td>
<td>$-0.0105$</td>
<td>$-0.0075$</td>
<td>$-0.0075$</td>
</tr>
<tr>
<td>$\theta_0$ ($V$)</td>
<td>$0.0413564925$</td>
<td>$0.041326$</td>
<td>$0.038$</td>
</tr>
<tr>
<td>burst duration (s)</td>
<td>$412.0$</td>
<td>$9.8$</td>
<td>$5.4$</td>
</tr>
<tr>
<td>interburst interval (s)</td>
<td>$281.6$</td>
<td>$217.5$</td>
<td>$2.0$</td>
</tr>
</tbody>
</table>

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In region II, both the manifolds of slow motion are unobstructed. This parameter region corresponds to the endogenous bursting regime (Fig. 3D). During a burst, the phase point winds around the slow motion manifold for oscillations; during the interburst interval, the phase point follows the slow motion manifold for equilibria in the fast subsystem.

In region III, a stable equilibrium obstructs the slow manifold for equilibria of the fast subsystem (Fig. 3E). The stable equilibrium is the only attracting regime in this region. A perturbation can trigger a train of action potentials (Fig. 2A–C). During a transient burst, a fast transition occurred as the phase point moved from the stable equilibrium onto the slow manifold that controls oscillating activity in the fast subsystem. The model neuron executed a stereotyped burst as the phase point evolved across this manifold. Near the border between regions III and IV, the phase point could spend a long time near the ghost of the saddle-node orbit, producing a very long transient burst.

In region IV, there were stable and unstable orbits as well as stable and unstable equilibria. As a result, tonic spiking co-existed with a silent regime, and perturbations could elicit switches from one regime to another (Fig. 3F).

Table 2. Measure of burst duration and latency to spiking in response to inhibition from Figure 2 (A–E–G).

<table>
<thead>
<tr>
<th>Figure 2</th>
<th>$\theta_h$ (V)</th>
<th>$\theta_{k2}$ (V)</th>
<th>burst duration (s)</th>
<th>latency to spiking (s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>(A)</td>
<td>0.0415</td>
<td>$-0.0077$</td>
<td>10.327403</td>
<td>NA</td>
</tr>
<tr>
<td>(B)</td>
<td>0.0415</td>
<td>$-0.01043$</td>
<td>103.48097</td>
<td>NA</td>
</tr>
<tr>
<td>(C)</td>
<td>0.0415</td>
<td>$-0.010496$</td>
<td>309.27622</td>
<td>NA</td>
</tr>
<tr>
<td>(E)</td>
<td>0.04134</td>
<td>$-0.0107$</td>
<td>10.287</td>
<td>NA</td>
</tr>
<tr>
<td>(F)</td>
<td>0.041358041</td>
<td>$-0.0107$</td>
<td>103.378</td>
<td>NA</td>
</tr>
<tr>
<td>(G)</td>
<td>0.0413580468</td>
<td>$-0.0107$</td>
<td>317.679</td>
<td>NA</td>
</tr>
</tbody>
</table>

Doi:10.1371/journal.pone.0085451.g002
The mechanisms controlling burst duration and interburst interval in region II determine a scheme of two-parameter coregulation that supports bursting with a given duty cycle across a wide range of cycle periods. This scheme could explain the maintenance of phase relations in a network of coupled oscillators.

**Pattern-scaling in a Chain of Oscillators**

We applied a mechanism maintaining the duty cycle of endogenously bursting activity to show scaling in a metachronal-wave locomotor pattern. We developed a basic model of a central pattern generator for crawling of larval *Drosophila*. *Drosophila* larvae crawl by means of peristaltic contractions of the body. Posterior-to-anterior waves of peristaltic contraction produce forward motion. The phase lag of motor activity in neighboring segments scales proportionally to the period so that the phase relations between activity in neighboring segments is maintained. This scaling occurs over a two-fold range of cycle periods [24]. Activity in the nerve cord in isolated preparations occurs on the time scale of tens of seconds [20,21]. The phase delay from segment-to-segment of segmental nerve cord activity is roughly 10% of the cycle period [20]. We suggest that the scheme of coregulation controlling the duty cycle of participating neurons could present a cellular mechanism for the metachronal-wave pattern scaling. We modeled the metachronal-wave central pattern generator (CPG) as a chain of coupled oscillatory neurons.

The network was assembled from five endogenously bursting neurons from region II (Fig. 1A). If these neurons receive an inhibitory pulse during the interburst interval, they respond with a...
burst, and the activity is reset. This observation suggests that if these neurons are coupled in a chain, they will trigger burst responses one after another as though in a domino effect. We expect that this effect would be robust if the neurons were connected in a chain such that the strength of inhibitory coupling is stronger in one direction than in the other. This sequential propagation of excitation along a chain is referred to as a metachronal wave which would travel in the direction of stronger coupling. With this organization of activity, the phase delay is determined by burst duration. Thus by scaling the duty cycle, one can scale the pattern of the metachronal wave. We manipulated the network by changing the values of $\theta_{k2}$ and $\theta_b$ to control the burst duration and interburst interval. We coordinated these changes to maintain a constraint of 10% duty cycle within 0.1% tolerance for individual oscillators over a wide range of periods from 15 s to 85 s (Table 3). We predicted that a chain of coupled oscillators each with an intrinsic duty cycle of 10% would produce a metachronal wave with phase delay 10% at the period of the individual oscillators. With this duty cycle, in order to produce rhythms faster than 68 s, we obtained parameter sets with values of $\theta_{k2}$ beyond the range depicted in Figure 1A. They are located so close to the SNIC that they appear to sit on the curve (Fig. 1A). We did not investigate whether the SNIC extends beyond this range, yet we were able to obtain parameter sets that supported activity with a duty cycle of 10%.

The model is represented by a chain of identical bursting neurons with duty cycle 10% as described in Methods. To show pattern scaling, we investigated this model with parameters of individual cells varied according to our coregulation scheme defined in Table 3. Network activity self-organized into a metachronal-wave pattern following an interval of transient activity in all ten instantiations (Fig. 4). After the interval of transient activity, waves of bursts propagated along the chain of oscillators from the most posterior cell (segment 7) in the anterior direction. We quantified the time it took a network to reach a periodic state by counting the number of cycles of the posterior cell before the transient activity subsided and the propagating wave fully formed. In each instantiation of the network, the formation of the metachronal wave occurred within two cycles of the bursting neuron in segment 7.

We characterized metachronal waves in the activity of each network instantiation by the period and duty cycle of activity as well as the relative phase shift of bursting from one segment to its nearest neighbor. We compared the temporal characteristics of network bursting activity to those of activity produced by a single cell with the same values of $\theta_{k2}$ and $\theta_b$ (Table 3). The period and duty cycle of bursting activity of a single cell and the network matched well; these periods differed by less than 0.1% and 3%, respectively. The disparity between the period of single-cell and network activity was greater for network instantiations with larger period. In each network instantiation, the phase shift of the metachronal wave exhibited some dependence on cycle period (Fig. 5). For parameter values that produced activity with lower periods, the phase shift tended to be greater than that predicted by the duty cycle. For example, in network instantiations with

### Table 3. Cycle period, duty cycle, and values for $\theta_{k2}$ and $\theta_b$. Labeled parameter sets were used to produce activity found in Figure 4.

<table>
<thead>
<tr>
<th>$\theta_{k2}$ ($V$)</th>
<th>$\theta_b$ ($V$)</th>
<th>Period (s):</th>
<th>Cell</th>
<th>Network</th>
<th>Duty Cycle (%):</th>
<th>Cell</th>
<th>Network</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.0059</td>
<td>0.040736</td>
<td>15.1</td>
<td>15.1</td>
<td>10.0</td>
<td>9.9</td>
<td>Fig. 4(C)</td>
<td></td>
</tr>
<tr>
<td>0.0019</td>
<td>0.041048</td>
<td>21.0</td>
<td>21.0</td>
<td>10.0</td>
<td>11.0</td>
<td></td>
<td></td>
</tr>
<tr>
<td>-0.0001</td>
<td>0.041145</td>
<td>27.4</td>
<td>27.4</td>
<td>9.9</td>
<td>9.9</td>
<td></td>
<td></td>
</tr>
<tr>
<td>-0.0011</td>
<td>0.041183</td>
<td>31.8</td>
<td>31.8</td>
<td>9.9</td>
<td>9.8</td>
<td></td>
<td></td>
</tr>
<tr>
<td>-0.0031</td>
<td>0.041244</td>
<td>39.5</td>
<td>39.5</td>
<td>9.9</td>
<td>10.4</td>
<td></td>
<td></td>
</tr>
<tr>
<td>-0.0041</td>
<td>0.041268</td>
<td>48.0</td>
<td>48.0</td>
<td>10.0</td>
<td>9.9</td>
<td>Fig. 4(B)</td>
<td></td>
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<tr>
<td>-0.0057</td>
<td>0.041298</td>
<td>61.8</td>
<td>61.8</td>
<td>10.0</td>
<td>10.3</td>
<td></td>
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</tr>
<tr>
<td>-0.0061</td>
<td>0.041306</td>
<td>68.2</td>
<td>68.1</td>
<td>10.0</td>
<td>10.3</td>
<td></td>
<td></td>
</tr>
<tr>
<td>-0.0069</td>
<td>0.041318</td>
<td>82.3</td>
<td>82.1</td>
<td>10.0</td>
<td>10.0</td>
<td></td>
<td></td>
</tr>
<tr>
<td>-0.0070</td>
<td>0.041319</td>
<td>85.3</td>
<td>85.1</td>
<td>9.9</td>
<td>9.9</td>
<td>Fig. 4(A)</td>
<td></td>
</tr>
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All parameter sets were used to produce activity used in the analysis found in Figure 5. Average network cycle period and average network duty cycle each had coefficients of variation less than $2 \times 10^{-4}$ % and 4%, respectively. 

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![Figure 4. Scaling of the metachronal-wave pattern in chains of coupled endogenously bursting neurons.](image)
parameter values that produced activity with period 15.1 s and 21.0 s, the average relative phase between segments was 13.0% and 13.2%, respectively. For parameter values that produced activity with higher periods, the phase shift tended to be close to 10%. The instantiation of the network that produced activity with period 85.1 s had an average relative phase shift of 10.9%.

The cornerstone bifurcation controls periodicity through both burst duration and interburst interval. By coregulating \( \theta_{Kz} \) and \( \theta_b \), we vary the period of bursting while keeping the duty cycle fixed. As instantiations of the network become closer to the critical parameter values for the cornerstone bifurcation, the period of activity increases according to the inverse-square laws governing burst duration and interburst interval. Since duty cycle is fixed over variation in period, the burst duration in this model increases via spike addition. As such, the cycle period varies linearly with spike number (Fig. 5).

The period of a metachronal wave was well approximated by the intrinsic period of the individual oscillator on which that network instantiation was based. For parameter sets that produced activity with higher periods of bursting, the temporal characteristics of individual oscillators better predicted the temporal characteristics of network activity. The metachronal-wave pattern was maintained over a five-fold change in the period of activity.

**Discussion**

The flexibility of the central nervous system relies on the ability of neurons to meet a breadth of temporal specifications. Oscillatory neuronal networks, such as central pattern generators, maintain functional output over a wide range of cycle periods in order to produce appropriate behavior [8–10, 13–24, 75]. Neuro-modulation provides control over regimes of activity according to motor tasks by coordinating adjustments to the biophysical parameters of ionic currents [28, 29]. To understand this process, we must answer a key question in neuroscience. How do biophysical characteristics govern functional activity? This question is particularly important for understanding motor control of rhythmic movements. Among the biophysical characteristics that affect excitability, the maximal conductances and voltages of half-activation of ionic currents are the most prominent targets for neuromodulation [25, 26, 61–68, 76–81].

The maximal conductances of ionic currents are coregulated to produce functional activity [63, 64, 78–80]. Homeostasis of a functional pattern of activity, either activity-dependent or activity-independent, has been well documented in terms of covariation of maximal conductances. One mechanism for activity-independent homeostasis of functional activity is determined by the pattern of gene expression of ionic channels. The activity of an identified neuron is specified by the patterns of gene expression of a set of ionic channels [63, 78–80]. Correlations have been shown in the quantities of mRNA that code for various voltage gated ion channels in crab [78, 79]. Modeling studies have explored the role of correlation in biophysical parameters in the maintenance of functional activity [60, 82, 83]. The biophysical parameters of hyperpolarization-activated currents and potassium currents are often correlated [63, 78, 79]. For example, coregulation has been revealed by the injection of the mRNA coding A-type potassium current which induced a corresponding increase in \( I_b \) [63]. In most cells considered, the mRNA quantities for these currents are strongly correlated [78, 79]. The mRNA correlations map to correlations in expression of channel protein, which in turn determines the maximal conductance of ionic currents. Correlations in the maximal conductance that support functional activity has been shown in models [60, 82, 83]. Maximal conductances are also correlated in mechanisms of activity-dependent homeostasis [84, 85].

The voltages of half-activation of ionic conductances are also subject to neuromodulation [61, 62, 65–68]. These parameters are shifted in neuromodulation of potassium currents and hyperpolarization-activated currents [61, 62, 65–68]. For \( \text{Kv}2.1 \) expressed in HEK293, the voltage dependencies of activation and inactivation are shown to be shifted negatively by 30 mV and 22 mV in response to phosphorylation by AMPK [67]. In the pyloric network of the spiny lobster, depending on the time of exposure and concentration of dopamine, the magnitude and direction of shifts varies but in all cases the shifts appear to be small [63, 65]. Hour-long application of dopamine induces dose-dependent shifts in the voltage dependence of the conductance of an A-type potassium current [66]. In LP neurons, nM and \( \mu \)M concentrations positively shift the activation by 4.6 mV and 1.7 mV, respectively, while the shifts of the inactivation curve have different directions: 3.3 mV and −1.2 mV, respectively. In PD neurons, both nM and \( \mu \)M concentrations negatively shift the activation and inactivation curves by 1.3 mV or less [66]; moreover, a ten minute application of a larger concentration of dopamine (100 \( \mu \)M) induces a larger negative shift of 7.6 mV to the activation curve [65]. In ferret thalamocortical neurons, repeated negative pulses depolarize the activation of \( I_b \) by 3.7 mV, and application of cAMP depolarizes the activation of \( I_b \) by 12 mV [61, 62].

Neuromodulators can coregulate currents that have complementary effects on membrane dynamics—such as hyperpolarization-activated currents and potassium currents—to tune aspects of excitability [68]. These shifts appear to be small and negligible. However, our results show that precise control of these biophysical parameters in these reported ranges could be sufficient for effective

**Figure 5. Temporal characteristics of metachronal wave pattern across a range of cycle periods.** The phase relations of a metachronal-wave pattern are determined by the duty cycle of each element in a chain of inhibitory coupled bursting neurons. The parameters are changed in a coordinated fashion to support different cycle periods while the duty cycle is kept constant. (A) The phases of oscillators are portrayed relative to the oscillator in the seventh segment. (B) The average number of spikes per burst varies linearly with cycle period as \( \theta_{Kz} \) and \( \theta_b \) are changed. The black markers indicated average spike number with error bars. The grey line was the linear function for spike number fitted to our data: 0.49(cycle period) + 0.93.

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neuromodulation. Amendola et al. [68] show independence of neuromodulation of half-activation of h current and half-inactivation of A-type potassium current. Such ability to independently control these parameters supports the biological feasibility of the mechanisms of independent control of burst duration and interburst interval by variation of half-activations of $I_{K2}$ and $I_h$, presented in this report.

We have demonstrated a family of cellular mechanisms that provides three different kinds of control for temporal characteristics in neuronal activity by coregulating the kinetics of activation of a non-inactivating potassium current and a hyperpolarization-activated current. These controlling mechanisms are organized by a global codimension-2 bifurcation: the cornerstone bifurcation. The cornerstone bifurcation occurs at the intersection of two codimension-1 bifurcation curves—the saddle-node bifurcation for periodic orbits and the saddle-node bifurcation for equilibria—in the two-dimensional parameter space defined by the voltages of half-activation of $I_{K2}$ and $I_h$.

In our model, the kinetics of activation of $I_{K2}$ and $I_h$ have distinctly separate, complementary roles in rhythmogenesis. $I_{K2}$, with its half-activation between 0.006 and 0.011 V (Fig. 1), activates incrementally during each spike in the burst. For this mechanism to function, the activation variable must be significantly slower than the time scale of spiking. Over the course of the burst, this activation accumulates until the current is sufficiently activated to terminate oscillations in the fast subsystem. As the parameter $\theta_{K2}$ is changed such as to approach the critical value for bifurcation, the steady-state activation curve of $I_{K2}$ is shifted so that the current is less activated by each spike. Finally, at the critical value for bifurcation, the activation of this current no longer accumulates sufficiently over the course of spiking activity to terminate a burst. $I_h$ plays a similar role during the interburst interval. Its voltage of half-activation lies roughly between −0.038 and −0.042 V (Fig. 1), and it is responsible for the slow depolarization leading to the first spike in the burst. As the parameter $\theta_h$ approaches its critical value for bifurcation, the steady state activation curve of $I_h$ is shifted so that the current is less involved during the interburst interval. At the critical value for bifurcation, this current does not sufficiently activate to initiate the depolarization of the burst.

Bifurcation Theory and Control of Neuronal Activity

Bifurcation theory considers transitions (bifurcations)—either continuous and smooth or discontinuous and catastrophic—in response to a smooth change in a physical parameter. Applications of bifurcation theory in physics, engineering, and neuroscience have established a new interdisciplinary field: bifurcation control [1,86,87]. Bifurcation control offers techniques to control regimes of activity using knowledge of bifurcation locations in parameter space and techniques to control the type of a bifurcation. These techniques prescribe how to precisely control the target characteristics of dynamical regimes by changing bifurcation parameters. Common targets of control include steady state characteristics of equilibria, temporal characteristics and magnitude of oscillatory regimes and transient activity, the borders of basins of attraction, and properties of and the routes to turbulence in deterministic systems [2,3,6,11,12,31–40,42–50,87]. For example in the control of turbulent activity, different types of intermittency have been associated with specific bifurcation types [86,88]. Intermittency of type I and type II are related to the saddle-node bifurcation and the Andronov-Hopf bifurcation, respectively [86,88].

Bifurcation theory has been applied to describe the general laws of neuronal dynamics that govern characteristics of spiking and bursting activity as a controlling bifurcation parameter approaches a transition between qualitatively different regimes of activity [2,3,6,11,12,31–40,42–50,87]. Excitability in neuronal systems has been characterized by type of bifurcation. A saddle-node bifurcation on an invariant circle generates class I excitability [31,32,43,87]. At this bifurcation, a neuron makes a smooth transition from silence into tonic spiking as the bifurcation parameter smoothly changes. This bifurcation controls the interspike interval according to the inverse-square-root law. At the bifurcation point the neuron is silent and, thus, the frequency is zero. The spiking orbit appears at the bifurcation with a large, full-scale amplitude. It is well described by a canonical model: the $\theta$ neuron [31,32,89]. A supercritical Andronov-Hopf bifurcation generates class II excitability [31,32,39,43]. At this bifurcation, the rest state loses stability, and a spiking orbit with zero amplitude and non-zero frequency is born. The amplitude of spiking grows according to the square root of the bifurcation parameter. Depending on the class of excitability, the response of the neuron to a stimulus, in type 2, may advance or delay the next spike depending on phase or, in type 1, will advance in response to depolarizing pulses and delay in response to hyperpolarizing pulses [31,32]. The type of phase response describes synchronization in neuronal networks [31,32,43]. The quantitative laws described by bifurcation theory are generic and are not uniquely associated with specific biophysical properties of ionic currents. For example, similar dynamical mechanisms may appear in neuronal systems relying on distinct ionic currents [90], and qualitatively distinct dynamics such as type 1 versus type 2 can be realized in a neuronal system by variation of a few biophysical parameters [87]. Dynamical mechanisms have also been used to explain the generation of bursting in terms of slow-fast systems [35,49,87]. Bursting is usually based on bistability of spiking activity and a subthreshold regime in the fast subsystem and dynamics of a slow variable governing switches between these regimes [35,49,89]. The bursting activity in our model is a square-wave burster [45,87].

Similarly to spiking activity, temporal laws have been described for bursting activity where the length of the burst duration or interburst interval is subject to control by a bifurcation parameter [11,12,42,45,48]. The control of a specific temporal characteristic has been shown by the manipulation of single bifurcation parameter. In models of the leech heart interneuron under various experimental conditions, we have extensively studied how bifurcations determine the mechanisms that support bursting activity. Where the blue sky catastrophe controls the transition from bursting to tonic spiking, it imposes the inverse-square-root law on burst duration [45]. Where homoclinic bifurcations control the transition from bursting to the spiking regime—such as the Luykanov-Shilnikov scenario—a logarithmic law is imposed on burst duration [47]. In this report, we describe independent control of burst duration and interburst interval with two parameters.

Examples of control over burst duration and interburst interval have been shown in a model of the electrosensory lateral line lobe pyramidal cell [11]. In [11], the model shows a unique type of bursting called ghostbursting. In slow-fast decomposition, the fast subsystem of the ghostbursting model does not exhibit bistability [11]; such bistability is a common feature of bursting models [35]. This lack of bistability is described as unique to the ghostbursting mechanism, and as such, it makes a key difference between ghostbursting and our model in the topology of the manifolds that govern slow motion. In our model, bursting activity is based on bistability of spiking and a stable equilibrium in the fast subsystem, thus classified as square-wave bursting; the slow manifolds of spiking activity and equilibria are separated in the phase space.
This bistability of regimes in the fast subsystem allows for the bifurcation curves obtained in the full system—the saddle-node bifurcation for periodic orbits and saddle-node bifurcation for equilibria—to cross in the two parameter space. This feature of the bifurcation curves allows the cornerstone bifurcation to generate a family of mechanisms that independently control the temporal characteristics of bursting and transient responses to external stimuli. The curves for saddle-node bifurcation of periodic orbits and saddle-node bifurcation for equilibria cross, dividing parameter space into four regions: spiking, bursting, silence, and bistability of spiking and silence. Similarly to our model, the scenario presented in [11] involves a saddle-node bifurcation for fixed points (SNFP) and a saddle-node bifurcation for periodic orbits (SNLC). In contrast to our model, on the two-dimensional bifurcation diagram in [11], SNLC terminates at SNFP. It is important to emphasize that the saddle-node bifurcation for periodic orbits and the saddle-node bifurcation for equilibria are local bifurcations. In our model, the control of bursting activity is governed by specific global bifurcations: the blue sky catastrophe and the saddle-node bifurcation on an invariant circle are global bifurcations, which include the saddle-node bifurcation for periodic orbits and the saddle-node bifurcation for equilibria, locally. In our model, the two mechanisms, by which burst duration and interburst interval of periodic bursting activity are controlled, are uncoupled.

Another key feature that distinguishes ghostbursting from our scenario, is that in the ghostbursting model the SNLC describes the transition from tonic spiking to chaotic bursting, and SNFP describes the transition of chaotic bursting into silence. Doiron et al. show that these transitions to chaos are intermittency type I [11,86,88]. This bursting activity is governed by inverse-square-root laws for burst duration and interburst interval on average. In contrast, our model shows periodic bursting: the blue-sky catastrophe describes the transition between periodic bursting and tonic spiking [45], and SNIC describes the transition between periodic bursting and silence. The ability to control temporal characteristics of periodic bursting activity is critical for the control of precise rhythmic movements. For the first time, a biophysically realistic neuronal model is described which combines two inverse-square-root laws and realizes independent control over burst duration and interburst interval. Moreover, this is the first model of a physical system showing the cornerstone bifurcation. By coordinating burst duration and interburst interval, the duty cycle could be preserved over a wide range of cycle periods under neuromodulatory control.

A Cellular Mechanism of Pattern Scaling

We suggest that these cellular mechanisms could contribute to control of various types of motor activities. For example, phase shifts of a metachronal-wave pattern could be determined by the duty cycle of each element in a chain of inhibitory coupled bursting neurons. Metachronal-wave patterns are ubiquitous in animal locomotion such as that of the leech, the crayfish, Drosophila larvae, and the lamprey [15,16,19–24]. In each case, the motor pattern is maintained across a range of periods. The lamprey can swim with one body wave cycle with period in the range from 0.13 s to 0.66 s; the leech can swim with one body wave cycle with period from 0.39 to 1.1 s; or Drosophila larvae can crawl with body contractions of cycle period from 0.6 s to 1.3 s [13,15,16,23,24]. Phase lags between spinal segments are scaled such that the lag between neighboring segments is approximately 1% of the cycle period in lamprey, 5% of the cycle period in leech, or 10% of the cycle in Drosophila [13,15–17,19–21,23,24]. Coupled oscillators have been commonly used in the study of central pattern generators. Phase delay in oscillating patterns have been explained with various network mechanisms such as frequency gradient, coupling gradient, and coding phase delays through adjustment of coupling [18,34,52–54,56,58,39,91]. For example, by tuning the coupling strengths and synaptic delays in neuronal networks, complex activity patterns can be generated, stored, and retrieved [58,59].

Drosophila larvae crawl by means of peristaltic contractions of body wall muscles. Posterior-to-anterior waves of contraction propel the animal forward at varying speeds. The segmental phase delay of the propagation of these waves is proportional to the cycle period of oscillations [24]. Fictive motor patterns persist in the isolated nerve cord [19,20]. The period of this fictive pattern occurs on a time scale of tens of seconds [20,21]. The phase delay measured from segment to segment in segmental nerve cords is 10% of the period of the motor pattern [20].

We modeled this phenomenon in a chain of coupled bursting neurons. The model corresponded to segments three through seven of the larval Drosophila (see Methods). By preserving duty cycle and varying cycle period we achieved maintenance of delay in phase between neighboring neurons in the chain. Moreover, the duty cycle of a model of a single cell predicts the phase delay in the network. Our model demonstrated a five-fold scaling of the metachronal-wave pattern, which is comparable to the two-fold scaling of the motor pattern in Drosophila [24]. This is an example of a cellular mechanism which translates to network activity. Although this mechanism does not require extensive synaptic tuning, it could work in conjunction with synaptic mechanisms to produce more a more sophisticated phase delay-cycle period relation.

Transient Response to Stimulus

Dynamical mechanisms govern rhythmic neuronal activity in the vicinity of bifurcations. These types of mechanisms are often discussed within the context of steady state activity. Stereotyped transient responses and activity could process or represent sensory stimulation [1,4–6,11,12,92]. Such transient responses can be represented by heteroclinic connections in the phase space of a neuronal model. Heteroclinic connections can reliably dominate or control transient neuronal activity by drawing trajectories from a large basin of attraction into a specific and functional response to perturbation before relaxing into steady state activity. Strong attraction to slow manifolds makes evoked responses reproducible. This control manifests as a temporally precise response to inhibition.

These parameters provide two types of control over temporal characteristics for two types of transient activity. First, we demonstrate control of the duration of pulse-triggered bursts in a silent neuron. As the voltage of half-activation of I\textsubscript{K}\textsubscript{2} approaches the critical value for the saddle-node bifurcation for periodic orbits, the duration of individual bursts increases. Second, we demonstrated control of the duration of latency to spiking in response to a stimulus in an a periodically spiking neuron. As the voltage of half-activation of I\textsubscript{K} approaches the critical value for the saddle-node bifurcation for equilibria, there is an increase in the latency to spiking after inhibition in periodically spiking neurons. The dynamical mechanisms underlying transient responses described here feature the inverse-square-root law shared with type-I intermittency [11,12,86,88]. In [12], pulses of injected current induced individual transient bursts from a stable periodic spiking regime, called type I burst excitability.

Inverse-square-root Law for Pulse-Triggered Burst. In this article, we showed that the silent model could respond to a hyperpolarizing pulse by a stereotypical burst of spikes. We report
that the burst duration of the pulse-triggered bursts scaled as $\theta_{K2}$ approached the critical value for the saddle node bifurcation for periodic orbits (transition from III to IV). Similarly, Roa et al. have shown the inverse-square-root law in transient spiking activity for type II excitability neuronal models with parameter values near a saddle-node bifurcation for periodic orbits [2]. In Roa et al., a constant applied current is used as the controlling parameter, and an additional brief pulse of current is used to move the phase point away from a stable equilibrium [2]. These models then spend significant time spiking as the phase point passes near the ghost of the saddle-node periodic orbit before relaxing back onto the stable equilibrium. The time neurons spend spiking is proportional to the inverse of the square root of the value of the constant applied current. This mechanism is similar to the inverse-square-root law we reported here for pulse-triggered burst duration with values of $\theta_{K2}$ and $\theta_b$ near the border between regions III and IV. The main difference between underlying mechanisms is that in our model, the phase point moves along a slow motion manifold for oscillations and slows down near the ghost of a saddle-node orbit on that manifold. In our model, the inverse-square-root law in transient activity occurred as the phase point wound around the slow motion manifold.

**Latency to Spiking After Inhibition.** Our model produces a stereotypical latency to spiking after a brief hyperpolarizing pulse in a spiking neuron. By coordinating $\theta_{K2}$ and $\theta_b$, we place the system in a region of the parameter space with a stable tonic spiking regime such that it is close to the saddle-node bifurcation for equilibria (in region I close to the border with region IV). In this region after inhibition, the phase point traveled along the manifold of slow motion and slowed down as it passed the ghost of a saddle-node equilibrium. Control over latency to spiking could explain phenomena crucial for pattern coding and motor control.

In Steuber et al., pauses in Purkinje cell firing are implicated in pattern coding [93]. In their model, a potassium current plays a key role in the mechanism supporting delay to spiking. It is activated by the calcium influx which is induced by sufficient activation of parallel fiber synapses. Pauses in the spontaneous firing of the Purkinje cell occur after parallel fiber-evoked bursts, and long-term depression of parallel fiber synapses shorten pause duration. These pauses code the pattern of synchronous activation of multiple parallel fibers.

Potassium and hyperpolarization-activated currents have been implicated in the dynamics of delay to spiking. In Meng et al., the authors show control over delay to firing by manipulating the ratio of transient potassium conductances [6]. In the brainstem superior paraolivary nucleus, $I_h$ controls the timing of firing in rebound from hyperpolarization [5]. The kinetics of slow conductances may contribute to duration sensitivity after inhibition from a sensory stimulus [4].

**Bistability of Spiking and Silence**

Another important feature of the dynamics of our model is that it shows bistability of spiking and silence (region IV, Fig. 3). Perturbations can trigger a switch from one regime to the other (not shown). This phenomenon has been described in the squid giant axon model [39]. In the Hodgkin-Huxley model of the squid giant axon, the stable equilibrium and the oscillating regime are separated by the stable manifold of an unstable orbit [39]. This mechanism was also described in a simplified model of the leech heart interneuron [38]. This model supports a number of types of multistability including a case where a stable equilibrium and a spiking regime co-exist and are separated by the manifold of a saddle equilibrium [38]. Similar mechanisms are described for co-existence of spiking and silence and of bursting and silence in the canonical model of the leech heart interneuron [36,37]. In the example presented here, the stable periodic orbit sits on the strongly attracting slow manifold for oscillations which is separated from the stable equilibrium by the manifolds of a saddle equilibrium and saddle orbit.

**Designing Artificial Neurons**

In the example presented here, a codimension-2 bifurcation that satisfies the criteria for the SNIC and the blue sky catastrophe controls burst duration, interburst interval, pulse triggered bursting, and latency to spiking. In conclusion, we suggest that the control over the temporal characteristics of neuronal activity presented here is critical for designing functional artificial neurons in biomedical and neuroengineering fields [94]. The parameter $\theta_{K2}$ asserts control of burst duration in an endogenously bursting neuron and of duration of pulse-triggered bursts in a silent neuron. The parameter $\theta_b$ asserts control of interburst interval in an endogenously bursting neuron and latency to spiking in a spiking neuron. By coordinating these parameters, one could tune an artificial neuron to produce burst activity with any burst duration and interburst interval over a large range of values for the cycle period. Moreover, the transitions between different regimes of activity are smooth and safe over the range of $\theta_{K2}$ and $\theta_b$ addressed here, so parameters can be tuned without fear of the onset of multistability or catastrophe.

**Methods**

We developed a Hodgkin-Huxley style neuronal model. It contains three voltage gated currents: a fast $Na^+$ current, $I_{Na}$; a non-inactivating $K^+$ current, $I_{K2}$; and a hyperpolarization-activated current, $I_h$. From our previous models, it inherited the blue sky catastrophe [44–47]. The model is as follows:

$$\frac{dV}{dt} = -g_{Na}m_{Na,x}(V)h_{Na}[V-E_{Na}] + g_{K2}m_{K2}^{2}[V-E_{K}] + g_Ks_m^2[V-E_{K}] + g_{leak}[V-E_{leak} + 0.006],$$

$$\frac{dh_{Na}}{dt} = \left[\frac{1}{1 + \exp(500[V+0.0325])} - h_{Na}\right]/0.0405,$$

$$\frac{dh_b}{dt} = \left[\frac{1}{1 + 2\exp(180[V+0.0325]) + \exp(500[V+0.0325])} - m_b\right]/0.1,$$

$$\frac{dm_{K2}}{dt} = \left[m_{K2,x}(V, \theta_{K2}) - m_{K2}\right]/2,$$

where

$$m_{Na,x}(V) = \frac{1}{1 + \exp(-150[V+0.0305])},$$

and

$$m_{K2,x}(V, \theta_{K2}) = \frac{1}{1 + \exp(-83[V+\theta_{K2}])}.$$
The activation of $I_{Na}$ is instantaneous and is denoted as $m_{Na}(V)$. The inactivation of $I_{Na}$ and the activations of $I_K$ and $I_h$ are $h_{Na}$, $m_{K2}$, and $m_h$. The maximal conductances of ionic currents are $g_{Na} = 105 \, nS$, $g_{K2} = 30 \, nS$, $g_h = 4 \, nS$, and $g_{ leak} = 6 \, nS$. The reversal potentials are $E_{Na} = 0.045 \, V$, $E_K = -0.07 \, V$, $E_h = -0.021 \, V$, and $E_{ leak} = -0.046 \, V$. The regulating parameters $\theta_{K2}$ and $\theta_h$ represent the voltages of half-activation of the variables $m_{K2}$ and $m_h$. The function $m_{K2,\sigma}(V, \theta_{K2})$ is the steady state activation function of $I_{K2}$. The capacitance, $C$, is 2 $nF$. Similar to [45–47], $m_{K2}$ is the slow variable.

The majority of the parameters of this model were pulled directly from our previous model [45]. These models stem from the model explaining the dynamics of slow plateau-like bursting activity in the leech heart interneurons under the conditions when $Ca^{2+}$ currents are blocked with divalent ions and most of the $K^+$ currents are blocked with 4-aminopyridine [44,71,72]. Here, we added a hyperpolarization activated current as was described in Hill et al. [55]. We adjusted $g_{Na}$ and the time constant of $I_h$ during the search for the cornerstone bifurcation. We also adjusted the time constant of $m_{K2}$, the slow variable, to make the variable slower in order to emphasize the separation of time scales.

We built a model representing the CPG that produces a metachronal-wave pattern of locomotion in Drosophila larvae. It follows the generic motif of a chain of coupled oscillators [16,34,52–54,56,91]. This network was composed of a sequence of endogenously bursting model neurons (parameters taken from region II in Fig. 1). We connected these cells into a chain of coupled oscillators, where each node in the chain was an endogenously bursting neuron. We numbered these cells from anterior to posterior according to the segments in the Drosophila, such that the most anterior neuron corresponded to the third segment, and the most posterior neuron corresponded to the seventh segment. Coupling was accomplished through inhibitory synapses. The nearest of the posterior cells provided the strongest synaptic input. The synaptic input of the anterior cell was weaker than any of those from posterior cells. The total synaptic current onto cell $j$ was aggregated into the term $I_{syn,j}$, and the current balance equation and expression for synaptic activation became:

$$\frac{dV_j}{dt} = -[I_{Na,j} + I_{K2,j} + I_{h,j} + I_{leak,j} + 0.006 + I_{syn,j}]$$

$$\frac{ds_{i,j}}{dt} = \left[1 + \exp(-5000|V_i - 0.02|)\right]^{-1}/0.01$$

We defined $I_{syn,j}$ as the sum of the synaptic currents from each presynaptic cell $i$ [3–7],

$$I_{syn,j} = \sum_{i=3}^{7} g_{syn,i,j} s_{i,j} [V_j - E_{syn}]$$

where $g_{syn,i,j} = 0.03 ~ i-j = -1$

$$3 ~ i-j = 1$$

$$0.3 ~ i-j = 2$$

$$0.3 ~ i-j = 3$$

$$0 ~ \text{otherwise}.$$
Figure S2 The dependence of equilibria and periodic orbits on the parameter $\theta_{K2}$. For each orbit, we plot the maximum, minimum, and average voltage. The green curves represent the evolution of a stable orbit as $\theta_{K2}$ is varied. This stable orbit coalesced with a saddle orbit at a saddle-node bifurcation for periodic orbits ($SN_{01}$). We back-traced this saddle orbit (dashed light blue curves) between $SN_{01}$ at $\theta_{K2} = -0.010500 V$ and a second saddle-node bifurcation for periodic orbits ($SN_{02}$) at $\theta_{K2} = -0.013027 V$ where it coalesced with a stable orbit (solid orange curves). This orbit lost stability in a period doubling bifurcation ($PD$) at $\theta_{K2} = -0.011948 V$. The saddle orbit (dashed dark blue) terminated in a homoclinic bifurcation (Hom). The purple curve represents the equilibria states of the system. The solid purple component indicates a stable equilibrium. The stable equilibria coalesced with the saddle equilibrium ($SN_{01}$) in a saddle-node bifurcation at $\theta_{K2} = -0.010506 V$, and this saddle equilibrium coalesces with another saddle equilibrium in a saddle-saddle bifurcation at the point labeled $SN_{02}$ at $\theta_{K2} = 0.029936 V$.

Figure S3 Structure of the manifolds of slow motion. (A) The slow motion manifolds for parameter values of both the SNIC and the blue sky catastrophe calculated at $\theta_{K2}$. The stable and unstable portions of the slow motion manifold for oscillations are represented by $M^s$ and $M^u$, respectively in green and blue. The manifold is composed of many orbits calculated for different values of $\theta_{K2}$ (see Fig. S2). The average voltage is plotted against the average slow variable for each orbit in dark green ($\langle V \rangle$). The average nullcline of the slow variable is plotted in orange ($\langle m_{K2} \rangle = 0$). The nullcline for the slow variable is represented by the grey curve $m_{K2} = 0$, and the equilibrium state for the fast subsystem is the purple curve $M_{eq}$. The saddle-node orbit is the closed orange curve labeled as $SN_{01}$. The saddle-node equilibrium is the green dot labeled as $SN_{eq}$. (B) $M^s_{eq}$ and $M^u_{eq}$ are calculated at $\theta_{K2} = 0.038 V$. The closed orange curve is a sample periodic burst computed at $\theta_{K2} = -0.0105 V$ and $\theta_{K2} = 0.038 V$. The trajectory of bursting closely follows the manifolds of slow motion.

Text S1 Inverse-Square-Root Curve Fits Confirm SNIC and Blue Sky Catastrophe Bifurcation Curves. (PDF)

Text S2 Computing the Manifolds of Slow Motion. (PDF)

Author Contributions
Conceived and designed the experiments: WHB GSC. Performed the experiments: WHB. Analyzed the data: WHB GSC. Contributed reagents/materials/analysis tools: WHB GSC. Wrote the paper: WHB GSC.

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