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A role for compromise: synaptic inhibition and electrical coupling interact to control phasing in the leech heartbeat CPG

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How can flexible phasing be generated by a central pattern generator (CPG)? To address this question, we have extended an existing model of the leech heartbeat CPG’s timing network to construct a model of the CPG core and explore how appropriate phasing is set up by parameter variation. Within the CPG, the phasing among premotor interneurons switches regularly between two well defined states – synchronous and peristaltic. To reproduce experimentally observed phasing, we varied the strength of inhibitory synaptic and excitatory electrical input from the timing network to follower premotor interneurons. Neither inhibitory nor electrical input alone was sufficient to produce proper phasing on both sides, but instead a balance was required. Our model suggests that the different phasing of the two sides arises because the inhibitory synapses and electrical coupling oppose one another on one side (peristaltic) and reinforce one another on the other (synchronous). Our search of parameter space defined by the strength of inhibitory synaptic and excitatory electrical input strength led to a CPG model that well approximates the experimentally observed phase relations. The strength values derived from this analysis constitute model predictions that we tested by measurements made in the living system. Further, variation of the intrinsic properties of follower interneurons showed that they too systematically influence phasing. We conclude that a combination of inhibitory synaptic and excitatory electrical input interacting with neuronal intrinsic properties can flexibly generate a variety of phase relations so that almost any phasing is possible.

Keywords: neuronal networks, entrainment, neuronal oscillators

INTRODUCTION

Underlying many rhythmic activities like breathing or walking are rhythmically active neuronal networks that produce motor patterns in the absence of sensory input with the same rudimentary timing and coordination as in vivo (Marder and Calabrese, 1996; Marder and Bucher, 2007). Analysis of these central pattern generators (CPGs) has helped not only to elucidate how motor patterns are controlled by nervous systems but the general mechanisms of network function that carry over into all neuronal networks, both sensory and motor. Modeling has been essential to this analysis (De Schutter et al., 2005; Marder et al., 2005; Grillner et al., 2007).

CPGs are also remarkably plastic and through neuromodulation they can be reconfigured so that different forms of the motor pattern are produced (Hooper and DiCaprio, 2004; Marder et al., 2005). Moreover, CPGs can produce motor variants that reflect changes in coordination between motor elements necessary for opposing functions, as for example egostive versus ingestive biting behavior in the mollusk Aplysia (Cropper et al., 2004) or different forms of scratching in turtles (Stein, 2005). In both these instances, the relative phasing of pattern generating elements changes with resultant changes to motor outflow. Understanding how phasing is established and how it may be modified is key to understanding CPG function. Thus the control of phasing in CPGs is a subject of active investigation using both physiological and modeling approaches (Bose et al., 2004; Mamiya and Nadim, 2004; Mouser et al., 2008; Hooper et al., 2009).

How can flexible phasing be generated in a CPG? We used a model of a core part of the leech heartbeat CPG that we constructed by extending an existing model CPG’s timing network (Hill et al., 2002; Jezzini et al., 2004). In the heartbeat CPG, premotor interneurons are coordinated differently on the two sides in distinct peristaltic and synchronous coordination modes. Phase and duty cycle of the activity of all the interneurons of the modeled CPG core have been rigorously quantified and animal-to-animal variability determined (Norris et al., 2006). Moreover, synaptic interactions in the CPG timing network have been extensively characterized (see Kristan et al., 2005 for a review). Thus we are in a strong position to constrain both the parameters and the output of our CPG model and to explore how parameters and output are related.

BACKGROUND TO THE CURRENT MODEL

The heartbeat central pattern generator (CPG) of medicinal leeches has been studied intensively for over two decades (for a recent review see Kristan et al., 2005) and has been characterized and modeled extensively. Medicinal leeches have two tubular hearts that run the length of the body and move blood through the closed circulatory system (Thompson and Stent, 1976; Krahl and Zerbst-Boroffka, 1983; Wenning et al., 2004a). The beating pattern (beat period 4–10 s) is asymmetric with one heart generating high systolic pressure through a front-directed peristaltic wave (peristaltic coordination mode) along its length, and the other generating low systolic pressure through near synchronous constriction (synchronous coordination mode) along its length. The fictive motor pattern
for heartbeat is correspondingly bilaterally asymmetric (Wenning et al., 2004b). Heart motor neurons, which occur as bilateral pairs in midbody segmental ganglia 3–18 fire in a rear-to-front progression (peristaltic) on one side, while those on the other fire in near synchrony (synchronous) but with strict side-to-side coordination (Wenning et al., 2004b). The asymmetry is not permanent, but rather the motor neurons of the two sides change roles (patterns) every 20–40 heartbeat cycles.

The leech heartbeat CPG consists of seven identified and well-characterized bilateral pairs of heart interneurons that occur in the first seven segmental ganglia: heart interneuron HN(1)–HN(7), indexed by midbody ganglion number (Figure 1). Two additional pairs of premotor interneurons (HN(15) and HN(16), termed rear premotor interneurons), which do not feedback onto the rest, have recently been identified (Wenning et al., 2008). An unidentified HN(X) pair has only been indirectly characterized (Norris et al., 2006). We focused on the first seven pairs which generate the beat timing and provide the only inputs to motor neurons in midbody segments 7–14 (Norris et al., 2007a). In this CPG core, interneurons can be subdivided into overlapping functional groups. The HN(1)–HN(4) interneurons constitute a timing network (Figure 1A), in which activity phase relations are fixed, albeit subject to modulation (Masino and Calabrese, 2002a,b). The timing network does not receive feedback from the other identified heart interneurons and imposes the regular beat rhythm on the entire CPG through its synaptic contacts (Figure 1). In the timing network, each of the HN(3) and HN(4) interneuron pairs form reciprocal inhibitory connections with their respective contralateral homologs, forming half-center oscillators that pace activity in the circuit. These oscillator interneurons (Figure 1A) and their reciprocal synaptic interactions have been biophysically characterized and a working model has been produced through several cycles of experimental testing and revision (Olsen and Calabrese, 1996; Hill et al., 2001). The HN(1) and HN(2) interneurons coordinate, via mutual ipsilateral inhibitory connections, the two half-center oscillators so that they assume a stable phase relationship; on average the HN(4) oscillator leads the HN(3) oscillator by 0.04 in phase (Masino and Calabrese, 2002b). The synaptic connections and functional interaction in this circuit have been extensively characterized, and a detailed model of this eight neuron circuit has been produced through two cycles of experimental testing and revision (Hill et al., 2002; Jezzini et al., 2004).

The oscillator interneurons (HN(3) and HN(4) pairs) are also premotor, making specific inhibitory synaptic connections with ipsilateral heart motor neurons (Norris et al., 2007a). The phase relations of these front premotor interneurons (Figure 1A) are fixed but the middle premotor interneurons [HN(6) and HN(7)] pairs (Figure 1A) on the two sides are phased differently with respect to the front premotor interneurons by intervening switch interneurons [HN(5)] (Norris et al., 2006). These switch interneurons receive ipsilateral inhibitory input from the front premotor interneurons (oscillator interneurons) and make bilateral inhibitory connections to the middle premotor interneurons. The middle premotor interneurons also receive electrical input (thought to be rectifying) from ipsilateral front premotor interneurons (Calabrese, 1977). Only one of the two switch interneurons is active in bursts at any time; the other is silent (Figure 1). The result is that on the side of the active (bursting) switch interneuron, the premotor interneurons fire in near synchrony, while on the side of the silent switch interneuron they fire in a distinct rear-to-front progression, leading to two different coordination modes of the two lateral heart tubes, left synchronous/right peristaltic and left peristaltic/right synchronous, respectively (Figure 1). It is convenient to speak of peristaltic and synchronous coordination modes, but it is important to realize that saying one side is peristaltic at any given time necessarily means that the other side is synchronous at the same time. Moreover, periodic changes (20–40 times the heartbeat period) in the activity pattern (silent vs. bursting) of the switch interneurons lead to periodic side-to-side changes in the coordination mode within the CPG by shifting the phase of middle premotor with respect to front premotor interneurons (Figure 1) (Norris et al., 2006). We infer from these regular switches that there are no permanent asymmetries in the heartbeat CPG, i.e., in its synaptic connections or in the intrinsic properties of its components neurons. The synaptic connections in the CPG core have been extensively characterized (Calabrese, 1977; Peterson, 1983a,b; Ivanov and Calabrese, 2000, 2003, 2006a,b). Most importantly, the activity and phase relations of the interneurons in the CPG core have been exhaustively quantified and animal-to-animal variability determined (Norris et al., 2006). Therefore, we have target values for phase and duty cycle of all the component neurons to constrain our model of the CPG.

**OVERVIEW OF PRINCIPAL EXPERIMENTS AND FINDINGS**

To reproduce experimentally observed phasing in our CPG model, we varied the strength of inhibitory synaptic and excitatory electrical input from the timing network to follower premotor interneurons. Neither inhibitory nor electrical input alone was sufficient to produce proper phasing on both sides, but instead a balance was required. Our model suggests that the different phasing of the two sides arises because the inhibitory synaptic and excitatory electrical inputs oppose one another on one side (peristaltic) and reinforce one another on the other (synchronous) (see below). Our search of parameter space defined by the strength of inhibitory synaptic and excitatory electrical input strength led to a CPG model that well approximates the experimentally observed phase relations. The strength values derived from this analysis constitute model predictions that we tested by measurements made in the living system. Further, variation of the intrinsic properties of follower interneurons showed that they too systematically influence phasing. We conclude that a combination of inhibitory synaptic and excitatory electrical input interacting with neuronal intrinsic properties can flexibly generate a variety of phase relations within a rhythmically active neuronal network.

**MATERIALS AND METHODS**

We modeled a core part of the heartbeat CPG in medicinal leeches. This CPG core consists of heart (HN) interneurons from HN(R,1)–HN(L,7), which are indexed by body side (L or R) and midbody ganglion number (1–7). Recently we have identified two more pairs of heart interneurons in midbody ganglia 15 and 16; these neurons switch in time with the ipsilateral HN(6) and HN(7) interneurons but their connections from within the CPG are not currently well defined (Seaman and Calabrese, 2008; Wenning...
et al., 2008). Because they do not feedback (rear-directed axons) to the CPG core they were not considered. One unidentified heart interneuron pair called HN(R/L,X) because its ganglionic origin is unknown was also not considered since they have no known synaptic connections onto the HN(5), HN(6) and HN(7) interneurons that were the main subject of this study. Unless otherwise noted (e.g., Figure 8), model interneurons on the left side were always in the synchronous coordination mode while those on the
right were in the peristaltic mode, so we dispensed with body side indexing and simply labeled the interneuron or motor neuron as synchronous or peristaltic.

**GENERAL MODELING STRATEGY**

The heart interneuron CPG model was implemented using GENESIS (GENeral NERual SLimulation System) software (Bower and Beeman, 1998). The 14 heart interneurons (7 bilateral pairs) (Figure 1) in our model CPG were outfitted with intrinsic conductances and inhibitory synaptic conductances largely derived from biophysical studies (See Kristan et al., 2005 for a review.), however conductances for the rectifying electrical junctions linking ipsilateral premotor interneurons – the HN(3), HN(4), HN(6), and HN(7) interneurons – were only estimated from voltage recordings (Calabrese, 1977).

The network connectivity diagram of the heartbeat CPG model is given in Figure 1, illustrating the connections among the 14 heart interneurons. The timing network of the CPG (Figure 1A), consisting of the HN(1)–HN(4) interneurons, has been modeled in considerable detail (Hill et al., 2001, 2002; Jezzini et al., 2004), and we implemented the model of this timing network by Jezzini et al. (2004). Because of their similar connectivity and interaction with the oscillator interneurons of the third and fourth ganglia, coordinating heart interneurons of the first and second ganglia were combined and modeled as a single bilateral pair of intersegmental cables (multi-compartmental fiber models) for computational efficiency. For each coordinating heart interneuron model, we implemented a “two-site model” (Δf = 2.1; see Jezzini et al., 2004 for details) that includes inhibitory synaptic conductances and intrinsic conductances tuned for the primary spike initiation site to be located in the fourth ganglion. The oscillator heart interneurons of the third and fourth ganglia were modeled as single-compartment neurons with the appropriate intrinsic conductances, inhibitory synaptic conductances as originally described in Hill et al. (2001). Within the timing network the intersegmental conduction delays (20 ms per segment), the strengths of each synaptic input, and the estimated time course of synaptic plasticity, were obtained from averaged voltage-clamp recordings and modeled as before (Hill et al., 2001, 2002; Jezzini et al., 2004). The h-current maximum conductance (g_h) used was 4.0 nS, corresponding to a free-run timing network cycle period of 9.3 s (Jezzini et al., 2004).

Each of the remaining six heart interneurons (three bilateral pairs) were modeled as single-compartment neurons with the appropriate intrinsic conductances, inhibitory synaptic conductances, and a conductance for the rectifying electrical junctions linking ipsilateral oscillator interneurons (also front premotor heart interneurons) – the HN(3) and HN(4) – with the middle premotor heart interneurons – HN(6) and HN(7) (Calabrese, 1977) (Figure 1A). The inhibitory synaptic input onto the middle premotor heart interneurons arises from the HN(5) switch heart interneurons that connect to them bilaterally.

**Switch interneurons**

The primary focus of this study is to investigate the mechanisms of phasing of the middle premotor interneurons. It was thus important to develop model HN(5) switch interneurons that would faithfully reproduce activity pattern of the living switch interneurons so that the model middle premotor interneurons would receive the proper pattern of inhibitory synaptic input. The switch interneurons are very different in their electrical properties from the other heart interneurons but are not favorable for voltage-clamp analysis of voltage-gated currents (Lu et al., 1999). We therefore developed a reduced activity-based single-compartment model of the switch interneurons (HN(5)) using four voltage-gated conductances that were tuned via a genetic algorithm (Houck et al., 1997; Tobin and Calabrese, 2006) to fit experimentally recorded burst, phase, and spike frequency characteristics (Gramoll et al., 1994; Lu et al., 1999). The four conductances include three from the model oscillator interneurons – \( I_{la} \) (Fast Na⁺), \( I_p \) (Persistent Na⁺), and \( I_{in} \) (Delayed Rectifier) – and a boot-strapped fast activating, very slowly inactivating (τ = 21.3 s) outward current. In contrast to oscillator (front premotor) interneurons, switch interneurons fire only on the synchronous side and in bursts with an accelerating spike frequency. The tuned switch [HN(5)] model interneurons were able to reproduce these accelerating bursts (see HN(5) in Figures 2, 5 and 8). The synaptic inhibition onto the switch interneurons arising from the ipsilateral HN(3) and HN(4) interneurons was modeled with both spike-mediated and graded conductances with a constant delay of 1.6 s. This delay was implemented to align artificially the model switch interneuron’s phase with experimental recordings of its phase with respect to the timing network (Norris et al., 2006). We used the same model equations for these synapses as for the synapses between oscillator interneurons (Hill et al., 2001) but the maximal conductances (\( g_{syn} \) and \( g_{vps} \)) were set to 60 nS and 30 nS respectively.

In addition to the synaptic inputs arising from the timing network, the peristaltic-side switch interneuron (silent) is tonically inhibited by a persistent leak current (reversal potential of −60 mV) that arises from unknown origins outside of the CPG (Gramoll et al., 1994; Lu et al., 1999). This current was modeled as an additional tonic leak conductance with a reversal potential of −60 mV and a maximum conductance (\( g_{switch} \)) of 15 nS. For Figure 8, this conductance was alternated across the two sides every 20–40 cycles as observed in the experimental preparation.

**Middle premotor interneurons**

The middle [HN(6) and HN(7)] premotor interneurons show similar activity to the [HN(3) and HN(4)] oscillator (front premotor) interneurons and were modeled similarly; they share the same complement of intrinsic conductances (Hill et al., 2001). All conductance parameters of the front and middle premotor interneurons were the same, except the leak-current maximum conductance and reversal potential of the middle premotor interneurons were chosen so that they were brought into an endogenous bursting regime (\( g_{h} \): 9.9 nS, \( E_h \): −63.5 mV; (Cymbalyuk et al., 2002), and the h-current maximum conductance (\( g_h \): 2.0 nS) was tuned to give them a free-run cycle period of 8.1 s (87% of the timing network period). The current through the rectifying electrical junction from the front premotor interneurons was calculated as a constant multiplied by the difference between low-pass filtered voltage waveforms of the coupled interneurons (τ: 0.2 s) and restricted to pass only depolarizing current onto the middle premotor interneurons (the equations for such junctions are presented in Garcia et al., 2008). The synaptic inhibition onto the middle premotor interneurons arising from the HN(5) switch interneurons was modeled as being spike-mediated and showing short-term synaptic.
Due to the substantial computational time of modeling the entire CPG (especially the coordinating interneurons), all neurons within the heartbeat timing network were initially modeled fully and their relevant synaptic parameters (e.g., $V_{m}$, $I_{CaS}$, and $I_{CaF}$) for determining their inhibitory synaptic currents and junctional currents onto the interneurons of the fifth through seventh ganglia were recorded as a time series for 81.2 s of model time, corresponding to 10 full cycles of network activity. During the parameter searches, the timing network was no longer computed and this previously recorded time series plasticity identical to the spike-mediated synaptic component of the synapses between oscillator interneurons (Hill et al., 2001) but the maximal conductance ($g_{SynS}$) was varied as described below.

**RUNNING SIMULATIONS AND PARAMETER SEARCHES**

The model equations were integrated with an exponential Euler method using a time step of 0.0001 s for a sufficient period (≥10 s) to allow the model to settle and then data were recorded for 50–200 s. All complete bursts of the interneurons from the middle of the recording period were used in the analysis.

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In Figure 3. For all other model studies, all heart interneurons (including the entire timing network) were fully modeled with no play-back.

In the parameter searches, we systematically varied the strength of synaptic inhibition (\(g_{\text{SynS}}\)) from the switch interneurons and the electrical coupling from the front premotor interneurons onto the middle premotor interneurons (Figures 2 and 3) to investigate data was “played-back” in a recurring loop of 10 cycles to the appropriate synaptic conductances to investigate the effects of these parameters on the middle heart interneurons. The ends of the loop were carefully spliced in order not to cause any abrupt changes in these parameters. This play-back saved an enormous amount of computational time and thus allowed us to pursue a more fine-grained analysis of the parameter space as evidenced in Figure 3.
the role of synaptic coupling on the relative phasing and duty cycle of the middle premotor interneurons. The values of synaptic inhibition maximum conductance were varied from \( (g_{\text{max}}) \) 0.6 to 84 nS in 0.6 nS increments. Values of electrical coupling strength \( (g_{\text{coupl}}) \) were varied exponentially from 1 to 301 nS. In plotting the effects of these variations on phasing of the middle premotor interneurons, we plotted the electrical coupling parameter values on a logarithmic scale to enhance data separation at low conductances.

**Model data analysis**

Simulation data were analyzed for period, intraburst spike frequency, duty cycle and phase using the same methods for the corresponding data from the living system in Norris et al. (2007b). Briefly, custom analysis programs were written in MATLAB, and ≥10 full bursts of simulation data from a given model interneuron were analyzed for each data point. Data points reported are mean ± SD. Our burst marker for measuring period and phase was the middle spike of each burst. Our burst detection paradigm recognized a burst as groups of at least four spikes separated from other spikes by a minimum inter-burst interval of 300 ms.

We calculated bilateral (absolute) phase of the model heart motor neurons using the middle spike of the HN(4) premotor interneuron input pattern in the peristaltic coordination mode as our phase reference (assigned 0.0 phase with no standard deviation) in accordance with our convention for the living system (Norris et al., 2007b), thus facilitating comparisons between the model and the living system. We also calculated first and last spike phase for each burst and constructed bilateral phase diagrams as described in Norris et al. (2007b).

**EXPERIMENT VERIFICATION OF MODEL PREDICTIONS**

Leeches (*Hirudo sp*) (Siddall et al., 2007) were obtained from commercial suppliers (Leeches USA, Westbury, NY, USA and Biopharm, Charleston, NC, USA) and maintained in artificial pond water at 15°C. After the animals were anesthetized in cold saline, chains of ganglia were dissected consisting of midbody ganglion 2 to at least midbody ganglion 8 (G2–G8) for recording the strength of the HN(5)-mediated IPSCs in the HN(6) and HN(7) middle premotor interneurons. The preparations were pinned (ventral surface up) in 60-mm Petri dishes lined with Sylgard™ 184 (Dow Corning, Midland, MI, USA). Ganglia in which heart interneurons were to be recorded were desheathed using fine scissors or microscalps. The preparation was superfused continuously with normal leech saline containing (in mM): 115 NaCl, 4 KCl, 1.8 CaCl₂, 10 glucose, 10 HEPES buffer, adjusted to pH 7.4 with NaOH, at 1–2 ml/min (bath volume 6–8 ml).

**Extracellular and intracellular recording techniques**

We used conventional electrophysiological procedures for leech neurons described in Norris et al. (2007b). For extracellular recordings from heart interneurons, we used suction electrodes filled with normal saline. Electrodes were pulled on a Flaming/Brown micropipette puller (P-97, Sutter Instruments, Novato, CA, USA) from borosilicate glass (1 mm o.d., 0.75 mm i.d., A.M. Systems) and placed in a suction electrode holder (E series, Warner Instruments Corp., Hamden, CT, USA). To ensure a tight fit between the cell and electrode, the electrode tips had a final inner diameter of ~20 μM, approximately the diameter of a heart interneuron’s soma. The electrode tip was brought in contact with the cell body and light suction was applied using a syringe until the entire cell body was inside the electrode. Extracellular signals were monitored with a differential A.C. amplifier (model 1700, A-M Systems, Carlsborg, WA, USA) at a gain of 1000 with the low and high frequency cutoff set at 100 and 1000 Hz, respectively. Noise was reduced with a 60-Hz notch filter and a second amplifier (model 410, Brownlee Precision, Santa Clara, CA, USA) amplified the signal appropriately for digitization. Heart interneurons were identified based on soma size, soma location in the ganglion, and ultimately identified by their characteristic bursting activity (e.g., Figure 1B). The HN(5) switch interneurons are very difficult to identify and record extracellularly because their somatic spikes are small (~5 mV recorded intracellularly). To aid our search, we always monitored an easily identified and recorded front premotor interneuron. Signal to noise ratios were often poor for the switch interneuron recordings, necessitating off-line filtering so that the spikes could be easily discerned and detected.

For intracellular recordings from middle premotor interneurons, we used sharp intracellular electrodes (~20–30 MΩ filled with 4 M KAc, 20 mM KCl) and an Axoclamp-2B amplifier (Molecular Devices, Sunnyvale, CA, USA) operating in discontinuous current-clamp or discontinuous single electrode voltage-clamp mode with a sample rate of 2.5–2.8 kHz. The electrode potential was monitored to ensure that it settled during each sample cycle. Output bandwidth was 0.3 kHz. Voltage-clamp gain was 0.8 to 2.0 nA/mV. The voltage-clamp holding potential for recording spontaneous IPSCs in interneurons was ~45 mV and for recording spontaneous spike-mediated coupling currents was ~55 mV. At the end of each experiment the electrode was withdrawn from the neuron and only data in which the electrode potential was within ±5 mV of ground were included. Thus holding potentials were accurate within ±5 mV.

Data were digitized (5-kHz sampling rate) using a digitizing board (Digi-Data 1200 Series Interface, Axon Instruments, Foster City, CA, USA) and acquired using pCLAMP software (Axon Instruments) on a personal computer (PC).

**Determining the strength of HN(5)-mediated IPSCs and HN(3) and HN(4)-spike-mediated coupling currents in the HN(6) and HN(7) middle premotor interneurons**

To determine the strength of each inhibitory synaptic connection from a switch interneuron to a middle premotor interneuron, we recorded extracellularly from one (a minority of cases, \( n = 8 \)) or both (\( n = 17 \)) switch interneurons. We then voltage clamped as many of the middle premotor interneurons as possible in that preparation (~45-mV holding potential) one after another, recording spontaneous IPSCs for several interneuron burst cycles in each coordination mode, spanning several switches in the activity state of the switch interneuron (switches in coordination mode). \( N = 25 \) total preparations were used in these experiments. We then used off-line spike-triggered averaging during periods when the switch interneuron was in its active state. These spike-triggered averages gave us a direct measure of synaptic strength. When we recorded only one HN(5) switch interneuron we inferred the synaptic strength of its bilateral homolog in the same premotor
interneuron, during the recorded switch interneuron’s silent state, by manually measuring and averaging the spontaneous rhythmic IPSCs phased with the activity of the monitored front premotor interneuron. For both spike-triggered and manually averaged IPSCs, we averaged over at least 10 spike bursts, and we ignored the first 5 and last 5 spikes in a burst. When both HN(5) switch interneurons were recorded, direct comparisons of spike-triggered averaged IPSCs were made.

To determine the strength of each electrical connection from a front premotor interneuron to a middle premotor interneuron, we recorded extracellularly from one \( (n = 23) \) or both \( (n = 5) \) front premotor interneurons. We then voltage clamped the ipsilateral middle premotor interneurons (−55 mV holding potential), recording spontaneous spike-mediated coupling currents for a minimum of 15 interneuron burst cycles. \( N = 28 \) preparations were used in these experiments. For averaging spike–mediated coupling currents, we averaged over at least 10 spike bursts, and we ignored the first 5 and last 5 spikes in a burst. To assess the impact of switches in coordination mode on the spike-mediated coupling currents, continuous voltage-clamp measurements were made across a minimum of two switches \( (n = 4) \). Synchronous and peristaltic coordination modes were compared with a pairwise, 2-tailed \( t \)-test.

Spike detection and IPSC/spike-mediated coupling current averaging were performed off-line using custom-made MATLAB software (Mathworks, Natick, MA, USA); see Norris et al. (2006, 2007a,b) for more details. The average strength of a connection was defined as the amplitude (measured from the preceding baseline current) of the largest peak of the spike-triggered average IPSC or spike-mediated coupling current.

**Statistics**

Mean values are presented ± standard deviation (SD) and in some cases the coefficient of variation (CV) expressed as a decimal fraction of the mean. Conductance and current measurements were subjected to single factor ANOVA to determine significant differences between effects. \( F \) statistic, \( df \), and \( p \) are reported. Where appropriate, post hoc testing was done with Tukey’s HSD test. In cases where ANOVA was not appropriate, we performed paired \( t \)-tests (two-tailed). For all tests \( p < 0.05 \) was the criterion for significant difference.

**RESULTS**

**MODEL STRATEGY**

We extended our existing model of the timing network (Jezzini et al., 2004) to construct a model of the heartbeat CPG core (Figure 1A). In our CPG model, we implemented known synaptic and neuronal properties. An activity-based model of the switch interneurons was constructed that was tuned to fit experimentally recorded burst and phase characteristics. The middle premotor interneurons show similar activity to and were modeled in the same manner as the front premotor (oscillator) interneurons.

To reproduce experimentally observed CPG phasing, we systematically varied the strength of inputs onto the middle premotor interneurons, i.e., inhibitory synapses from the switch interneurons and excitatory electrical coupling from the front premotor interneurons (Figure 1A). Once suitable strengths were determined we then varied the intrinsic properties of the middle premotor interneurons to determine their effects on activity characteristics.

**THE EFFECT OF INHIBITORY SYNAPTON INPUT AND EXCITATORY ELECTRICAL COUPLING ON PHASING OF THE MIDDLE PREMOTOR INTERNEURONS**

First we determined whether inhibitory synaptic input from the switch interneurons, \( \bar{g}_{\text{Syn}} \) or excitatory electrical coupling from the front premotor interneurons, \( g_{\text{Coup}} \) alone could establish entrainment and appropriate phasing of the middle premotor interneurons (Figures 2A,B). HN(6) and HN(7) middle premotor interneurons were modeled with identical intrinsic properties. With a minimum of 6 nS for inhibitory synaptic maximal conductance (\( \bar{g}_{\text{Syn}} \)) or 1 nS for electrical coupling conductance (\( g_{\text{Coup}} \)) stable entrainment of the middle premotor interneurons was established (Figures 2A,B). Beyond these threshold values, varying either over an eight-fold range in the absence of the other had no discernable effect on middle premotor interneuron phasing or duty cycle on either the peristaltic or synchronous side. With electrical input alone (\( g_{\text{Coup}} \)) both sides assume highly synchronous phasing between front and middle premotor interneurons. With inhibitory synaptic input (\( \bar{g}_{\text{Syn}} \)) alone, the basic structure of the asymmetric peristaltic-synchronous pattern seen in the living system is established. The middle premotor interneurons lead the front premotor interneurons by a large phase difference on the peristaltic side, and the middle premotor interneurons slightly lag the front premotor interneurons in phase on the synchronous side very similar to the living system (c.f. Figure 5C).

This observation highlights the primacy of the asymmetric activity in the switch interneuron pair in establishing the asymmetric coordination in the heartbeat CPG. To attain the smooth phase progression of middle and front premotor interneurons observed in the living system on the peristaltic side (c.f. Figure 5C), however, it is necessary to be able to generate phase differences intermediate between the large phase lead seen with only inhibition and the synchrony seen with only electrical coupling. We concluded that to establish appropriate peristaltic phasing of the middle premotor interneurons neither inhibitory synaptic nor excitatory electrical input alone would suffice.

Next we determined whether inhibitory synaptic input from the switch interneurons, \( \bar{g}_{\text{Syn}} \), in conjunction with excitatory electrical coupling from the front premotor interneurons, \( g_{\text{Coup}} \), could establish entrainment and appropriate phasing of the middle premotor interneurons (Figures 2C,D). We independently varied \( \bar{g}_{\text{Syn}} \) or \( g_{\text{Coup}} \) in the presence of a fixed suprathreshold amount (established above for entrainment) of the other (Figures 2C,D; \( g_{\text{Coup}} = 2 \) nS and \( \bar{g}_{\text{Syn}} = 24 \) nS respectively). Beyond the threshold values, varying either over an eight-fold range in the presence of the other had a monotonic effect on middle premotor interneuron phasing on both the peristaltic and synchronous side. Increasing \( \bar{g}_{\text{Syn}} \) caused the middle premotor interneurons to fire slightly later in phase on the synchronous side and earlier in phase on the peristaltic side. Increasing \( g_{\text{Coup}} \) caused the middle premotor interneurons to fire slightly earlier in phase on the synchronous side and later in phase on the peristaltic side. Moreover, duty cycle also varied monotonically with variations of \( \bar{g}_{\text{Syn}} \) or \( g_{\text{Coup}} \) decreasing with increasing \( \bar{g}_{\text{Syn}} \) on both sides and increasing with increasing \( g_{\text{Coup}} \).
on both sides, but more prominently on the peristaltic side. We concluded that it should be possible to obtain appropriate phasing and duty cycle for each of the middle premotor interneurons in either coordination mode by adjusting the balance of $g_{\text{SynS}}$ and $g_{\text{Coup}}$ for each premotor interneuron in the two coordination modes.

We next systematically co-varied $g_{\text{SynS}}$ and $g_{\text{Coup}}$ and determined the phasing and duty cycle of the middle premotor interneurons on the two sides. We used 120 values in the range of 1–302 nS for $g_{\text{SynS}}$ and 140 values in the range 0.1–14 nS for $g_{\text{Coup}}$ and constructed 3-dimensional contour plots for phase and duty cycle (data not shown). We compared phase and duty cycle from these simulations to the living system for each of the middle premotor interneurons on the two sides (Figure 3). Then, the simulation data that matched well with both phase and duty cycle in the living system were combined into a single point. These combined plots show where in the $g_{\text{SynS}}$ vs. $g_{\text{Coup}}$ plane both phase and duty cycle were appropriate to the living system – within the experimentally observed range and/or within $±1.5 \times SD$ of the mean of the living system (corresponding to 86.6% of expected normal data) for each middle premotor interneuron on the two sides (Norris et al., 2006). For each of the four plots, there is a broad area of the $g_{\text{SynS}}$ vs. $g_{\text{Coup}}$ plane where both criteria are met.

**Choosing inhibitory synaptic input, $g_{\text{SynS}}$, and excitatory electrical coupling, $g_{\text{Coup}}$, to obtain appropriate phasing and duty cycle of the middle premotor interneurons**

We looked for parameter pairs ($g_{\text{SynS}}$ and $g_{\text{Coup}}$) for each of the four premotor interneurons (HN(6) and HN(7) synchronous and HN(6) and HN(7) peristaltic) that would match well the living system (Figures 2C, D). Our main criteria for this match were the observed phasing and duty cycle of each of the middle premotor interneurons. Due to the regular switching between coordination modes in the living system, we also required that the parameters meet a symmetry constraint; connections made by left and right neurons of the same type are equal in strength. We then simplified the parameter choice further by making the electrical input arrive only when this summed error was a minimum for each of the four premotor interneurons and plotted this point as an asterisk on each of the corresponding panels of Figure 3. These four points constitute a clear model prediction. To obtain appropriate phasing and duty cycle for the four middle premotor interneurons, the ipsilateral inhibitory connections of the switch interneurons must be stronger than their corresponding contralateral connections and the connections of the switch interneurons onto the HN(7) premotor interneurons must be stronger than their corresponding ipsilateral connections onto the HN(6) interneurons.

**How do inhibitory synaptic and excitatory electrical input interact to produce appropriate phasing and duty cycle of the middle premotor interneurons?**

Using the values of $g_{\text{SynS}}$ and $g_{\text{Coup}}$ for each of the model middle premotor interneurons obtained in the analysis of Figure 3 (hereafter referred to as canonical parameters), we explored how inhibitory synaptic and electrical coupling currents interact in the intrinsically bursting middle premotor interneurons to produce appropriate phasing and duty cycle. The middle premotor interneurons were tuned as intrinsically bursting neurons with an intrinsic free-run cycle period of 8.1 s, which was 87% of the canonical timing network period of 9.34 s. Consequently, for stable entrainment of the middle premotor interneurons by the timing network to be effectuated, these premotor interneurons must be slowed during each cycle. Analysis of the synaptic and coupling currents that occur in these neurons during entrainment by the canonical timing network illuminate how phase and duty cycle are controlled (Figure 4A). It is important to note when viewing these records that the coupling conductance is always present but that currents flow only when there is a difference in potential between the two coupled neurons as when the HN(3) or HN(4) neurons produce spikes. In contrast, the synaptic conductance occurs only when the presynaptic HN(5) neuron produces a spike, and the postsynaptic current will depend on the conductance amplitude and the driving force ($V_m-E_{\text{syn}}$).

On the synchronous side, in the middle premotor interneurons, the strong inhibitory synaptic input from the switch interneurons arrives earlier than the electrical input from the front premotor interneurons and quickly terminates the ongoing burst in the HN(6) interneuron and delays the onset of the next burst (Figure 4A). When the electrical input does arrive, it generates sufficient initial inward current, however, to bring on the next HN(6) burst before the termination of the inhibitory input causing the initial part of the burst to seem ragged. Entrainment is achieved mainly through the delaying effect of strong inhibition. The situation is similar in the synchronous HN(7) interneuron – $g_{\text{SynS}}$ from the ipsilateral active switch interneuron in the HN(7) interneuron (43.2 nS) is comparable to $g_{\text{SynS}}$ in the HN(6) interneuron (34.8 nS) (Figure 3) – and so there is not much phase difference between these middle premotor interneurons.

On the peristaltic side, in the middle premotor interneurons, inhibitory synaptic and electrical input arrives nearly simultaneously and the corresponding currents that flow are balanced but with a steady inward bias, particularly in the HN(6) interneuron (Figure 4A) where $g_{\text{SynS}}$ from the contralateral active switch interneuron is relatively small (6.0 nS) compared to $g_{\text{SynS}}$ in the peristaltic HN(7) interneuron (21.0 nS) (Figure 3). During entrainment, the middle premotor interneurons begin their burst in the absence of input due to their intrinsic bursting ability, and the mixed synaptic/coupling current prolongs their burst and delays the next burst onset. Because the peristaltic HN(7) interneuron receives more inhibitory current but the same excitatory coupling current, its bursts terminate earlier and begin earlier, i.e., it leads the peristaltic HN(6) interneuron in phase.

Using the canonical values of $g_{\text{SynS}}$ and $g_{\text{Coup}}$, the activity pattern of our CPG model and average data from the living system (Norris et al., 2006) are remarkably similar (Figure 5). The only
Figure 4B gives a conceptual framework for understanding how this realistic phasing of the middle premotor interneurons is achieved in the CPG model. In the middle premotor interneurons on detail that appears off in the model is that the bursts of the HN(6) and HN(7) interneurons are a bit prolonged with a low spike frequency tail at their end.
it would be with only the electrical excitation, because the excitation determines the time of burst onset (Figure 4B). Entrainment is established, however, mainly by the delaying effect on burst onset of the synchronous side, inhibitory synaptic and excitatory electrical input reinforce one another. The strong inhibitory synaptic input arrives earlier than electrical input and phasing is nearly the same as it would be with only the electrical excitation, because the excitation determines the time of burst onset (Figure 4B). Entrainment is established, however, mainly by the delaying effect on burst onset of the synchronous side, inhibitory synaptic and excitatory electrical input reinforce one another. The strong inhibitory synaptic input arrives earlier than electrical input and phasing is nearly the same as it would be with only the electrical excitation, because the excitation determines the time of burst onset (Figure 4B). Entrainment is established, however, mainly by the delaying effect on burst onset of
the strong inhibitory current. In the middle premotor interneurons on the peristaltic side, the relatively weak inhibitory synaptic input and electrical input occur nearly simultaneously and thus oppose one another (Figure 4B). The weak inhibition cannot force the termination of the burst, which is thus extended by the excitation, and this extension of the burst establishes entrainment by delaying the next burst. On both sides, phasing is a compromise between the phase of entrainment with only the synaptic inhibition (orange dashed lines) or only the electrical excitation (green dashed lines) (Figure 4B). Specifically, in both coordination modes, the total burst activity phase (duty cycle) of the middle premotor interneurons completely overlaps the burst activity expected with electrical coupling alone, but on the peristaltic side the burst beginning (first spike phase) is advanced (as is middle spike phase) and the total activity phase (duty cycle) is correspondingly expanded.

HOW DO THE INTRINSIC MEMBRANE PROPERTIES AFFECT PHASING IN THE CPG MODEL?
In a model with canonical values of $g_{\text{syn}}$ and $g_{\text{Coup}}$, we explored how intrinsic membrane properties, particularly those underlying the period of the timing network and of the intrinsic bursting of the middle premotor interneurons, affected entrainment and phasing of the middle premotor interneurons in the CPG model. Figure 6A shows how we varied these periods by varying $h$-current ($g_h$) in either the front (period of the timing network) or middle (intrinsic period of the middle premotor interneurons) premotor interneurons. Varying timing network period caused a monotonic change in phasing of middle premotor interneurons on both sides, and there was a limited range of timing network period where entrainment was established with middle premotor interneuron phasing appropriate to the living system (Figure 6B). Stable entrainment on the peristaltic side could not be achieved with timing network periods exceeding $\sim 13.5$ s, and the timing network cannot be driven to periods shorter than $\sim 6$ s. Appropriate middle premotor interneuron phasing for the two sides was limited to timing network periods in the range of $\sim 7$ to $\sim 10$ s, which is relatively close to the intrinsic burst period of the middle premotor interneurons of $8.1$ s. The analysis shows that robust appropriate middle premotor interneuron phasing is possible either when the timing network has a longer or a shorter period than the intrinsic period of the middle premotor interneurons.

Varying $g_h$ in the middle premotor interneurons also caused a monotonic change in phasing of middle premotor interneurons on both sides that saturated at high as well as low values of $g_h$, on both sides, though the effect was more pronounced on the peristaltic side (Figure 6C1). When $g_h$ in the middle premotor interneurons is set below $\sim 1.7$ nS, then the model interneurons no longer burst intrinsically but become silent (Figure 6A) and coordinated bursting is achieved solely by post-inhibitory rebound. When $g_h$ in the middle premotor interneurons is set above $\sim 8.7$ nS, then the model interneurons no longer burst intrinsically but fire tonically (Figure 6A) and coordinated bursting is achieved by inhibitory sculpting. In Figure 6C2, the variation of $g_h$ in the middle premotor interneurons of Figure 6C1 is transformed into a variation in middle premotor interneuron intrinsic period. Figure 6C2 reveals that appropriate phasing is achieved on the peristaltic side only over a limited range of intrinsic burst periods, whereas appropriate phasing is achieved on the synchronous side over the entire range of intrinsic burst periods. We conclude that intrinsic properties of both the timing network and the middle premotor interneurons are important for establishing appropriate phase relations for the middle premotor interneurons. These properties should be matched appropriately with the strength of inhibitory synaptic and electrical excitatory input onto the middle premotor interneurons.

TESTING MODEL PREDICTIONS
Given how important the values of $g_{\text{syn}}$ and $g_{\text{Coup}}$ are in obtaining appropriate phasing of the model middle premotor interneurons (Figure 3), it seems appropriate to consider the canonical values of these parameters as model predictions. We tested these predictions by measuring in the living system the strength of the inhibitory synaptic currents evoked in middle premotor interneurons by ipsilateral and contralateral HN(5) interneurons and of spike-mediated coupling currents evoked in middle premotor interneurons by ipsilateral HN(3) and HN(4) interneurons (Figure 7). Specifically, the model predicts that (1) ipsilateral connections of the switch interneurons onto the middle premotor interneurons should be stronger than their corresponding contralateral connections and (2) connections of the switch interneurons onto the HN(7) middle premotor neuron should be stronger than corresponding connections onto the HN(6) middle premotor interneurons (Figure 3).

During switches in coordination mode recorded in voltage clamp in middle premotor interneurons, the size of the envelope of inhibitory synaptic currents was dramatically larger when the ipsilateral HN(5) interneuron was active than when the contralateral HN(5) interneuron was active (Figure 7A). This gross observation was corroborated by spike-triggered (or hand) averaged inhibitory synaptic currents, which in every case of a middle premotor interneuron thus recorded showed the inhibitory synaptic current from the ipsilateral HN(5) interneuron to be larger than that from the contralateral HN(5) interneuron (Figure 7B). Data combined across $N = 25$ preparations also showed for both the HN(6) and the HN(7) middle interneurons that the average inhibitory synaptic current from the ipsilateral HN(5) interneuron was significantly larger than from the contralateral HN(5) interneuron (Figure 8C) ($\text{HN}(6): n = 11, p = 0.014; \text{HN}(7): n = 14, p = 0.011$ – paired t tests). Moreover, the inhibitory synaptic currents evoked by the HN(5) interneurons in HN(6) interneurons were significantly smaller than in side-corresponding HN(7) interneurons (Single Factor ANOVA followed by post hoc testing using Tukey’s HSD – $F = 11.99, df = 3, p = 0.00001$. All post hoc pairwise comparisons were significant to $p < 0.01$ with Tukey’s HSD). These size relationships correspond well to the model “predictions” of Figure 3.

Moreover, during switches in coordination mode recorded in voltage clamp in middle premotor interneurons, the size of the envelope of spike-mediated coupling currents remained similar when the ipsilateral HN(5) interneuron was active or when the contralateral HN(5) interneuron was active (Figure 7A). This gross observation is expected given that there is no change in the sources or phase of this input during switches. Spike-triggered (or hand) averaged coupling currents in middle premotor interneuron thus recorded showed coupling currents from both the ipsilateral HN(3) and HN(4) interneurons to be similar (i.e., not significantly different) in both ipsilateral HN(6) and HN(7) middle
premotor interneurons (data not shown). Combined data across \( N = 28 \) preparations also showed for both the HN(6) and the HN(7) middle interneurons that the average coupling currents from the ipsilateral HN(3) and HN(4) interneurons were similar (Figure 7C) (Single Factor ANOVA \( F = 1.42; df = 3, \ p = 0.259 \)). These size relationships correspond well to the model “predictions” of Figure 3.

As can be seen from the standard deviation bars in Figure 7C, there was considerable animal-to-animal variation in the size of inhibitory synaptic and coupling currents measured in middle premotor interneurons. We observed that the strength of the inhibitory synapses varied over a nearly six-fold range from animal-to-animal and that the strength of electrical coupling varied over a nearly three-fold range from animal-to-animal. Ranges for inhibitory synapses were 28.5–131 pA for HN(5) to the ipsilateral HN(6), 21–73 pA for HN(5) to the contralateral HN(6), 47–278 pA for HN(5) to the ipsilateral HN(7), and 27–139 pA for HN(5) to the contralateral HN(7). Ranges for electrical coupling were 4.5–117 pA for HN(3) to HN(6), 38–84 pA for HN(4) to HN(6), 34–92.5 pA for HN(3) to HN(7), and 13–101.5 pA for HN(4) to HN(7). These ranges indicate that the living network arrives at different solutions to appropriate middle interneuron phasing, possibly within a framework like that of Figure 3.

In the living system, often a switch interneuron in the “silent” state fires weakly or sporadically (e.g., Figures 1 and 7). We used spike-triggered averaging to determine if these spikes in the silent state cause detectable IPSCs. In six different middle premotor interneurons (three HN(6) – one contralateral and two ipsilateral to the switch interneuron – and three HN(7) – one contralateral and two ipsilateral to the switch interneuron), we found
weaver et al. phasing in a model CPG

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no detectable IPSCs by spike-triggered averaging even when the silent switch interneuron was firing 10–20 spikes per burst at a frequency of 2 Hz. In other heart interneurons [e.g., HN(3) and HN(4)], spikes arising off a relatively hyperpolarized baseline are ineffective at causing transmitter release (Ivanov and Calabrese, 2003); presumably the same is true for switch interneurons in the silent, relatively hyperpolarized, state, and we need not consider such firing in our model.

SWITCHES IN COORDINATION MODE

Figure 8 shows an induced switch in coordination mode in our CPG model. Prior experimental work has shown that the inactive switch interneuron is silenced by the input of a tonic leak current of unknown origin (Gramoll et al., 1994). Halfway through the switch interneuron is silenced by the input of a tonic leak current. Prior experimental work has shown that the inactive switch interneuron was firing 10–20 spikes per burst at a frequency of 2 Hz. In other heart interneurons [e.g., HN(3) and HN(4)], spikes arising off a relatively hyperpolarized baseline are ineffective at causing transmitter release (Ivanov and Calabrese, 2003); presumably the same is true for switch interneurons in the silent, relatively hyperpolarized, state, and we need not consider such firing in our model.

**DISCUSSION**

Within natural motor patterns and within the CPGs that produce them activity does not just assume simple in phase (0.0) and antiphase (0.5) coordinations. All possible phase relationships may be appropriate for producing the proper sequences of motor neuron discharge; indeed in metachronal behaviors like undulatory swimming in lampreys (Grillner et al., 2007) and leeches (Kristan et al., 2005) and swimmeret beating in crayfish (Jones et al., 2003; Smarandache et al., 2009) a smooth intersegmental variation in phase leads to motor discharge in a multiplicity of phases. How can a CPG generate phases of activity intermediate between in phase, typically promoted by electrical coupling, and antiphase, typically promoted by moderate to strong inhibition? Our results suggest that a core network displaying canonical in phase and antiphase activity can produce any phasing required by balancing electrical coupling and synaptic inhibition as inputs to inherently bursting follower neurons.

We constructed a model of the core of the leech heartbeat CPG by extending an existing model of the CPG’s timing network (Hill et al., 2002; Jezzini et al., 2004). In the heartbeat CPG, premotor interneurons are coordinated differently on the two sides in peristaltic and synchronous modes that regularly and reciprocally switch sides (Figure 1A). Our model suggests that the different coordination modes (phasing) of the two sides arise because the inhibitory synaptic and excitatory electrical inputs onto middle premotor interneurons oppose one another on one side (peristaltic)
and reinforce one another on the other (synchronous). This asymmetry results from asymmetry in the activity of a bilateral pair of switch interneurons with bilateral inhibitory connections onto the middle premotor interneurons. One of the pair is active leading to synchronous coordination and the other is silent leading to peristaltic coordination (Figure 1B). In our model, to reproduce the details of the experimentally observed phasing on each side, we varied the strength of inhibitory synaptic input arising from the switch interneurons and excitatory electrical input arising from front premotor interneurons onto the middle premotor interneurons. Neither inhibitory nor electrical input alone was sufficient to produce proper phasing on both sides, but instead a balance was required. Our search of parameter space defined by the strength of inhibitory synaptic and excitatory electrical input strength led to a CPG model that well approximates the experimentally observed phase relations. The strength values derived from this analysis constitute model predictions that we tested and confirmed by measurements made in the living system. Further, variation of the intrinsic properties of middle premotor interneurons in the model showed that they too systematically influence phasing. We conclude that a combination of inhibitory synaptic and excitatory electrical input interacting with neuronal intrinsic properties can flexibly generate a variety of phase relations within a rhythmically active neuronal network.

ENDOGENOUS BURSTING GIVES MORE FLEXIBILITY IN ACHIEVING A MULTITUDE OF PHASE RELATIONS IN FOLLOWER NEURONS

It is interesting to note that we were not able to achieve experimentally observed phasing by the balancing mechanism we explored here if the follower interneurons were not inherently bursting. This was particularly true on the peristaltic side where inhibition and excitation are nearly in phase and thus opposed. Tonic spiking neurons tend to fire in response to electrical excitation and turn off whenever inhibited but bursting neurons (and we suspect neurons with strong rebound plateau properties) tend to have more fixed burst durations and defined inter-burst intervals that bring them into entrainment in a variety of phases dependent on the excitatory/inhibitory balance. The middle premotor interneurons do possess intrinsic bursting properties when isolated from inhibitory synaptic input with bicuculline (Cymbalyuk et al., 2002). The neuron models

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**FIGURE 8** | The phasing of the middle premotor neurons responds rapidly to a switch in coordination mode from left synchronous/right peristaltic to left peristaltic/right synchronous in the CPG model. This switch is illustrated in the top panel: the left active switch interneuron becomes silent as the right silent switch interneuron becomes active. The bottom panel shows recordings from one pair of front premotor interneurons, both pairs of middle premotor interneurons and the switch interneurons in the CPG model during this switch. Note the rapid readjustment of the phase of the middle premotor interneurons after the switch. The dark/light green dashed line indicates the firing phase of the peristaltic/synchronous HN(4) interneuron.
used for the middle premotor interneurons have been extensively studied for their inherent bursting properties and are dominated by the inactivation and de-inactivation time constants of a slowly inactivating Ca current that supports bursting (Hill et al., 2001, 2002; Cymbalyuk et al., 2002; Olypher et al., 2006). These time constants restrict the burst duration and inter-burst interval and fit well the requirements for entrainment at a variety of phases given the period of the entraining timing network. As Figure 6B shows, the inherent period of the entraining timing network can be either faster or slower than the inherent burst period of the follower middle premotor interneurons and still achieve appropriate phasing. Nevertheless, there are period limits for appropriate phasing and one-to-one entrainment.

With endogenously bursting follower neurons, like the model middle premotor interneurons we used, the excitatory drive associated with electrical coupling appears more effective than even strong inhibition at establishing the phasing of the follower neuron. Consider Figure 4B; very strong inhibition as on the synchronous side barely moves the phase from what would be established by the electrical coupling alone. Even when inhibition and electrical coupling are opposed as on the peristaltic side, phasing is still closer to what would be established by electrical coupling alone despite the inhibition being stronger in terms of current (Figure 4A). This observation suggests that electrical coupling is more effective at synchronizing bursting neurons than inhibition.

**IMPLICATIONS FOR ANIMAL-TO-ANIMAL VARIABILITY OBSERVED IN THE LEECH HEARTBEAT CPG**

The subject of animal-to-animal variability in intrinsic membrane and synaptic parameters and their implication for network activity has received considerable attention, particularly in the crustacean stomatogastric ganglion CPGs (Prinz et al., 2004; Bucher et al., 2005; Marder et al., 2007; Goaillard et al., 2009). The leech heartbeat CPG shows similar variability in its synaptic parameters (Figure 7) (Marder et al., 2007; Norris et al., 2007a; Seaman and Calabrese, 2008). The phasing of the middle premotor interneurons is quite variable between animals, particularly the HN(7) interneuron (peristaltic HN(6) phase = 0.893 ± 0.043; synchronous HN(6) phase = 0.578 ± 0.035; peristaltic HN(7) phase = 0.802 ± 0.094; synchronous HN(7) phase = 0.581 ± 0.072, Norris et al., 2006); correspondingly the motor pattern shows considerable phase variability as does the constriction pattern of the hearts (Norris et al., 2007b). Our parameter variations of Figure 3 show that the observed phase variability of the middle premotor interneurons is easily accounted for by the observed variability in inhibitory synaptic strength and electrical coupling (Figure 7). Moreover, simple changes in parameters that determine the intrinsic membrane properties of model neurons can traverse the entire criterion range of middle premotor interneuron phases as shown in Figure 6C1. These comparisons point out the importance of building neuronal network models that can accommodate observed animal-to-animal variability in output. Our model of the leech heartbeat CPG was able to reproduce such variability in output through variation of model parameters that correspond to experimentally observed variability in synaptic strengths; in this sense our model is a resounding success.

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Conflict of Interest Statement: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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