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The neural control of heartbeat in invertebrates

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Abstract

The neurogenic heartbeat of certain invertebrates has long been studied both as a way of understanding how automatic functions are regulated and for how neuronal networks generate the inherent rhythmic activity that controls and coordinates this vital function. This review focuses on the heartbeat of decapod crustaceans and hirudinid leeches, which remain important experimental systems for the exploration of central pattern generator networks, their properties, network and cellular mechanisms, modulation, and how animal-to-animal variation in neuronal and network properties are managed to produce functional output.

Invertebrates, owing to their small numbers of central neurons all of which are uniquely identifiable, have served as important experimental systems for the exploration of neuronal networks, their properties, network and cellular mechanisms, modulation, and how animal to animal variation in neuronal and network properties are managed to produce functional output. Automatic functions like neurogenic heartbeat have been particularly attractive for study because of their ongoing nature and the robustness of fictive neuronal activity in isolated nervous system preparations. Here we focus on two examples that are being actively pursued by experimental and computational approaches.

Crustacean heartbeat

In decapod crustaceans, the heart is neurogenic and receives innervation from a cardiac ganglion containing 9 neurons, five of which (Large Cells) are motor neurons. Heartbeat is driven by rhythmic bursts of impulses (burst period 1–5 s) in these motor neuron, which produce depolarizing EPSPs on heart muscle cells. The comparative anatomy and physiology of crustacean cardiac ganglia and their innervation of the heart have been comprehensively and lovingly reviewed in [1] and a small fraction only of this work is highlighted here. The cardiac ganglion is in essence a central pattern generator that produces fictive heartbeat in the Large Cells in isolation (Fig. 1). There are 4 Small Cells, interneurons
that are strongly electrically coupled, that pace activity in the ganglion by producing coherent bursts of action potentials that provide direct excitatory synaptic inputs to drive activity in the Large Cells (motor neurons). These motor neurons are themselves strongly electrically coupled and produce synchronous bursts to drive heart muscle. Both Small and Large Cells produce plateau-like regenerative potentials, called Driver Potentials, which support burst formation. Driver Potentials are triggered in the Large Cells by the rhythmic excitatory drive from the Small Cells; in Small Cells, Driver Potential are driven by a spontaneous slowly depolarizing pacemaker potential. Voltage-clamp analyses reveal that Driver Potentials result from the interplay of a Ca current ($I_{Ca}$) and a slow K current ($I_K$) and to a lesser extent a Ca activated K current ($I_{KCa}$); the source of the slowly depolarizing pacemaker potential remains controversial.

More recent work has occurred on two fronts: on the identification and role of the large number of neuropeptide modulators that are active both in the ganglion and at neuromuscular junctions to enhance cardiac output and on how homeostatic mechanisms regulate membrane conductances to ensure uniformity in ganglionic output across the life of an animal and across animals.

The neuropeptide work [2–4], in the American Lobster, illustrates the complex nature of neuromodulation when operating on multiple components of the multilayered neuromuscular system: interneurons (SCs) and motor neurons (LCs), the neuromuscular junctions (NMJs) and muscles, and importantly feedback systems. Both SCs and LCs have stretch sensitive dendrites on the heart, which feedback positively to decrease cycle period [5], and both SCs and LCs are modulated by NO released by heart muscle during contraction, which slows their burst period [6]. Two related peptides GYSDRNYLRFamide (GYS) and SGRNFLRFamide (SGRN) were studied in detail [3]. They act similarly at low concentration ($10^{-10}$ to $10^{-9}$ M) on both the ganglion and the neuromuscular system; they speed burst period, enhance stretch feedback, and increase contraction amplitude by acting either on the NMJs or the muscle, albeit by slightly different mechanisms. At higher concentration ($10^{-8}$ M) they continue to increase contraction amplitude, but GYS increases cycle period and LC burst duration, and SGRN has no effect on cycle period. These differences are attributable to how the two peptides influence stretch and NO mediated feedback; GYS enhances NO feedback while SGRN enhances stretch feedback.

The work on ion channel homeostasis [7–9] in the Jonah crab, Cancer borealis, is beginning to reveal homeostatic mechanisms and to uncover how ion channel expression levels that achieve similar activity still affect responses to perturbation. Using voltage-clamp measurements of ionic currents, molecular characterization and quantitative measurements of ionic channel mRNAs, various ionic currents have been identified in Large Cells [7] and shown to vary 2–4 fold across homologous neurons within the same ganglion as well as across animals [8,9]. These variable conductances make Large Cells susceptible to pharmacological perturbations so that these motor neurons become uncoordinated and show different firing patterns. Thus despite genetic identity within an animal and uniform and synchronous activity within the network, Large Cells show highly variable underlying maximal conductances that are tuned to achieve the same activity [9]. Remarkably, synaptically isolated Large Cells rapidly (30–90 minutes) compensate for changes in
excitability induced by $K^+$ channel blockers, by coordinately regulating two potassium currents ($I_A$ and $I_{KCa}$) presumably to homeostatically maintain a target level of excitability [8]. Thus the crab cardiac ganglion is rapidly becoming a test bed for sophisticated analyses of homeostatic regulatory mechanism at the cellular and molecular level. Given the strong electrical coupling among Large Cells and the resultant potential for intercellular communication, the relative cellular independence of homeostatic regulatory mechanism revealed thus far seems all the more remarkable.

**Leech heartbeat**

The leech (*Hirudo sp*) heartbeat system (see [10] for a past review) offers the opportunity to study and understand an on-going rhythmic behavior from neuron biophysics, to neuronal networks of interneurons and motor neurons, to circulatory movements. Four decades of research have made the CPG that drives heartbeat one of the best characterized and most comprehensively modeled.

Bilateral tubular hearts, running the length of the animal, undergo coordinated rhythmic (period ~6–10s at 22°–23° C) constrictions to propel blood flow [11,12]. At any one time the left and right hearts coordinate their constrictions differently, with one showing a rear-to-front peristalsis and the other near synchrony, switching roles every ~40 beats (~300–400s) (Fig. 2). The difference in the coordination mode of the two hearts corresponds to the pattern of blood flow. Peristalsis generates high systolic pressure and propulsive blood flow forward and into main dorsal and ventral longitudinal vessels which feed the main capillary beds and the CNS respectively. Synchrony generates low systolic pressure and drives local peripheral blood flow in each segment. Switching coordination modes presumably balances local segmental blood flow over the time course of minutes.

The tubular heart walls encompass a mesh of spiral muscle fibers that are innervated in each midbody segment (3–18) by a segmental motor neuron [13,14]. These heart (HE) motor neurons, one on each side, reside in their respective segmental ganglia of the leech ventral nerve cord and project ipsilateral axons to innervate their segment (with some segmental overlap) of the heart tube. This innervation has been described in detail, but simplifying for clarity, each motor neurons provides excitatory junctional potentials that sum to trigger plateau-like muscle potentials that produce constrictions (Fig. 2). Constrictions are timed by motor neuron bursts, such that maximal tension is achieved after the middle spike of each burst when motor neuron firing frequency has reached a plateau (usually >10 Hz) [15]. The middle spike of each motor neuron burst is a useful indicator of heart muscle contraction and roughly corresponds to the maximum rate of rise (MRR) of the tension. Peristalsis is thus characterized by a phase lead of heart motor neuron firing (middle spike) and of heart muscle tension (MRR) rear-to-front between segments and synchrony by near simultaneity of these characteristics between segments. Switches in coordination mode are accomplished within a beat cycle – as shown in Figure 2, we see a switch from right peristaltic (left synchronous) to left peristaltic (right synchronous) – and are always reciprocal. Thus the leech heartbeat motor pattern is one that has complex dynamics on two temporal scales - beat timing and switch timing - that serve clear functional roles and provide function-based...
metrics for assessing temporal characteristic: how they are generated and how they vary within and across animals.

**Leech fictive heartbeat and the heartbeat central pattern generator**

The isolated nerve cord of the leeches produces continuously a fictive motor program (Fig. 3a) that is startlingly similar to the motor program observed in nearly intact leeches, including the regular reciprocal switches in coordination mode [16,17]. Heart motor neurons fire in a rear-to-front progression (peristaltic coordination) on one side and a nearly synchronous pattern on the other (synchronous coordination) and coordination modes switch reciprocally abruptly within one beat cycle. Intersegmental phase differences (measured by the middle spikes of motor neuron bursts) in the fictive motor pattern characterize the peristaltic (substantial rear phase lead) and synchronous (near synchrony) coordination modes; these phase differences vary considerably in size across animals correspondingly as they do for the beat pattern of intact leeches measured with non-invasive video methods but remain characteristic of their coordination mode (Fig. 4b) [14].

A set of segmental (HN) heart interneurons located in midbody ganglia 1–7 form the core heartbeat CPG, and the heart motor neurons are rhythmically inhibited by the premotor heart interneurons of this CPG (Fig. 3a), which are located in segments 3, 4, 6 and 7 [16,17]. These premotor interneurons form a stereotypical pattern of ipsilateral synaptic contacts with heart motor neurons in more caudal segments (Fig. 3b). These connections vary in strength among the segments; each heart interneuron has its own segmental profile of synaptic strength with general interneurons HN(3) and HN(4) dominating in more rostral segments and interneurons HN(6) and HN(7) dominating in more caudal segments (Fig. 3c) [18,19]. These synaptic strength profiles have been described in great detail using voltage clamp and do not change across switches in coordination mode within any given animal. Nevertheless they vary significantly among individuals both in absolute and more importantly in relative terms (Fig. 4a) [19].

The heart interneuron network (heartbeat CPG) itself produces a complex rhythmic pattern that mirrors the fictive motor pattern [17,20]. Premotor interneurons on one side fire in a rear-to-front progression that is associated with peristaltic coordination of the motor neurons and on the other with near synchrony (or a small front-to-rear progression) that is associated with synchronous coordination of the motor neurons (Fig. 4b). These phase differences (measured from the middle spikes of their bursts) vary in size across animals correspondingly as they do for the fictive motor pattern but remain characteristic of their coordination mode (Fig. 4b). These characteristic firing patterns of the premotor interneurons switch to produce the switches in coordination mode of the motor pattern.

The heart interneurons form a network with a highly interconnected front kernel (HN(1)–HN(4) interneurons) characterized by reciprocal inhibition with feed forward inhibition and electrical excitation to the more reward interneurons (Fig. 5) [21]. The HN(5) interneurons serve to switch coordination modes within the network by interfacing between the front kernel with the HN(6) and HN(7) premotor interneurons. They are timed by rhythmic inhibition from the front kernel and make bilateral inhibitory connections to the HN(6) and HN(7) premotor interneurons. Repeated bilateral recording of HN(5) switch interneurons,
both extra cellular (Fig. 5) and intracellular, show that the switch interneurons on the peristaltic side of the network is quiescent while the one on the synchronous is robustly rhythmically active [21]. Switches in coordination mode are accomplished when the previously quiescent switch interneuron becomes suddenly active whereas the previously active switch interneuron becomes quiescent -within one beat cycle [22,23]. Modeling studies show that the balance of ipsilateral excitation from the HN(3) and HN(4) interneurons and inhibition from the active switch interneuron, which voltage clamp studies show is stronger ipsilaterally and in the HN(7) vs. the HN(6) interneuron, accounts for the characteristic peristaltic and synchronous phase differences among ipsilateral premotor interneurons [21,24].

Several modeling [25–27], and dynamic clamp [28] studies indicate that the coordination of the motor neurons by the premotor interneurons is realized by a combination of their characteristic peristaltic and synchronous firing patterns (intersegmental phase differences; rear-to-front vs. near synchrony) and the characteristic segmental synaptic profiles of the premotor interneurons onto the motor neurons. As a simplifying example consider that in peristaltic coordination the first firing HN(7) interneuron is strong in more rearward motor neurons and weaker up front, while the lagging HN(4) premotor interneuron is stronger up front and weaker rearward (Fig. 3c). This conjunction of temporal pattern in the interneurons and their segmental synaptic strength profiles causes the more rearward heart motor neuron to fire in advance of the more frontward. The temporal patterns determines the limits of what motor neuron phase difference can be achieved and the segmental synaptic strength profile determines what phase difference is realized within this limit. The animal-to-animal variability in the temporal pattern of premotor interneuron firing and in segmental synaptic profiles result in each animal arriving at a unique solution for peristaltic and synchronous coordination of heart motor neurons (Fig. 4) [19,28].

**Beat rhythmicity and switch rhythmicity within the heartbeat CPG**

The rhythmicity that underlies beat timing in the heart beat CPG arises in the HN(3) and HN(4) interneurons and percolates through the rest of the network. These interneurons (in addition to their premotor role detailed above) form strong reciprocal inhibitory connections with their bilateral homologs forming two classic half-center oscillators (HCOs) [21,29]. Numerous voltage clamp studies have detailed the ionic current that pace their activity, in addition to the fast inward and outward currents that produce spikes, they possess a non-inactivating K current [30], a persistent Na current that regulates general excitability [31] and most importantly a slowly-inactivating low-threshold Ca current ($I_{CaS}$) [32] and a hyperpolarization activated inward current ($I_h$) [33]. $I_{CaS}$ drives the burst phase slowly inactivating to release, as spike rate wanes, the opposite inhibited neuron of the half-center oscillator, while the $I_h$ activated by the hyperpolarization associated with inhibition, which also removes inactivation from $I_{CaS}$, drives the inhibited neuron of the HCO toward escape and burst formation. This mixed escape-release model for rhythmicity in a half-center microcircuit has been repeatedly corroborated by modeling [29,34–36] and dynamic-clamp studies [37,38]. The endogenous peptide myomodulin speeds the burst rhythm of the two HCOs by up-modulating $I_h$ and down modulating the current driven by the Na+/K+ pump.
Recent work indicates that the cycle-to-cycle dynamics of \( \text{Na}^+/\text{K}^+ \) pump current works in conjunction with \( I_h \) to regulate period of the HCOs [40].

The HN(1) and HN(2) interneurons serve as coordinating fibers that link the two HN(3) and HN(4) HCOs into a beat timing-network [21,41]. Inherent period differences between the two HCOs lead to the phase difference in activity observed ipsilaterally between them (Fig. 4c) [41–43]. Detailed modeling studies corroborate these findings [44,45].

The rhythmicity that underlies switch timing is not well understood. Switch interneurons do not interact synaptically, making a slow half-center oscillator comprising them highly unlikely [23]. Voltage-clamp studies show that switch interneurons in the quiescent state experience a tonic outward current that suppressing firing and that a switch between the quiescent and active state involves the termination of this current [22]. Artificially altering the state of a switch interneuron from quiescent to active or vice-versa with injected current or dynamic clamp does not influence the state of the opposite switch interneuron. Switch interneurons switch in isolated ganglia 5. Thus is seems likely that there exist a switch oscillator in ganglion 5 separate from the identified heartbeat CPG that engages the two switch interneuron with tonic inhibition alternately.

The leech heartbeat control system remains a vibrant test bed for ideas about the origins of rhythmicity in neuronal networks, intersegmental coordination, premotor control, the origins and importance of variability for neuronal network performance, and neuromodulation.

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**References and recommended reading**

Papers of particular interest, published within the period of review, have been highlighted as:

- of special interest
- • of outstanding interest


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Highlights

The cardiac ganglion reveals homeostatic mechanisms regulating electrical activity.

The leech heartbeat system can be studied at all levels from neurons to circulation.

In both systems individual variation is large, yet functional activity is maintained.
Figure 1.
Cardiac ganglion of the Jonah crab, *Cancer borealis*. Left: schematic of the ganglion showing the positions of the somata of the 5 Large Cells (LCs, motor neurons) and the 4 Small Cells (SCs, interneurons). Right: Typical electrical activity of an isolated cardiac ganglion. Top trace is an intracellular recording from two Large Cells (identified by color), showing their synchronous synaptic drive, driver potentials and spiking activity, and the bottom trace is an extracellular recording from a ganglionic trunk showing Small Cell activity (smaller spikes two different units clearly visible) and synchronous Large Cell activity (large spikes). Figure kindly provided by David J. Schulz.
Figure 2.
Heartbeat and heart motor neuron activity in a minimally dissected leech, *Hirudo sp.* Top: simultaneous extracellular (loose patch) recording from ipsilateral heart motor neurons in segments segments 8 and 12 (HE(R,8) and HE(R,12)) and tension recordings from the ipsilateral heart in the same segments. At the beginning of the record the right heart is in the peristaltic coordination mode; at the end of the record coordination mode of the heart has switched to synchronous. The switch occurs in the middle of the record and is accomplished in one beat cycle. Bottom: blow-ups of the shaded parts of the record at top to show how heart constriction is timed to heart motor neuron activity in each coordination mode. Diamonds mark the middle spike of each motor neuron burst and crosses the maximum rate of rise of heart tension (MRR). After [15].
Figure 3.
The heartbeat control system of the leeches: heart motor neurons and the heartbeat CPG. (a) Simultaneous ipsilateral extracellular recording of the four premotor interneurons of the core CPG and an intracellular recording of a heart motor neuron (HE(8)), which the interneurons all inhibit. The rhythmic inhibitory input sculpts out bursts in the heart motor neuron activity. (b) Bilateral circuit diagram including all the identified heart (HN) interneurons of the core CPG showing the inhibitory connections from the heart interneurons of the leech heartbeat CPG onto heart (HE) motor neurons in the first 12 midbody segmental ganglia. The ipsilateral HN(3) and HN(4) and the ipsilateral HN(6) and HN(7) premotor interneurons provide input to heart motor neurons [HE(3)–HE(12)]. (c) The segmental synaptic strength
profiles of heart interneurons for the HE(8) and HE(12) motor neurons. In the upper panel, an HE(12) motor neuron was recorded in voltage clamp (holding potential -45 mV) simultaneously with extracellular recording from four ipsilateral premotor heart interneurons, HN(3), HN(4), HN(6), and HN(7) (middle spikes indicated by symbols), that provide its input. HN spikes from 11 bursts from each interneuron were used to generate the spike-triggered averages of IPSCs in the HE(12) motor neuron, and subsequently similar recordings were used to generate the spike-triggered averages of IPSCs for the HE(8) motor neuron in the same animal. Note the difference in the premotor input synaptic strength profiles between the two motor neurons. Upward arrows indicate the time of the triggering HN spike and the downward arrows indicate the peak of the averaged IPSC used to measure amplitude. Spike-triggered average IPSCs, left to right arise from the HN(3)–HN(7) premotor interneurons respectively. All recordings from isolated nerve cord preparations. Cell identity coded with color consistently across figures. Adapted from [18,19].
Figure 4.
(a) The synaptic strength illustrated in Figure 3, shows considerable animal-to-animal variation both in absolute terms and in the relative strengths of the four inputs to a given motor neuron. Iconic unilateral circuit diagram (above) identifies the neurons recorded for each graph and standard colors and symbols are used to indicate the strength of each HN input in each graph. Absolute strength in nS (upper graphs) or relative strength (lower graphs) is plotted vs. preparation ordered by the absolute or relative strength of the HN(3) input, respectively. (b) Complete analysis of input and output temporal patterns and synaptic strength patterns (spatial pattern) for 2 different preparations from a sample of 12 (included in (a)) in which simultaneous extracellular recordings (loose patch) were made of ipsilateral HN(3), HN(4), HN(6), and HN(7) premotor interneurons (inputs) and HE(8) and HE (12) motor neurons (outputs) in both peristaltic (p) and synchronous (p) coordination modes and then subsequently synaptic strength of each interneuron in the two motor neurons was measured in voltage clamp as in Figure 3c. Summary phase diagrams of the premotor interneurons (standard color code) and the HE(8) and HE(12) motor neurons in both the peristaltic (pink-outlined boxes) and synchronous (light blue-outlined boxes) coordination modes are shown. For both motor neurons and interneurons, the average duty cycle is indicated by the length of the bar: the left edge of each bar indicates the average phase of the first spike of the burst and the right edge indicates the average phase of the last spike of the burst. Average phase is indicated by a vertical line within the bar. Error bars indicate SDs. The strength pattern of the inputs is shown to the right of each composite phase diagram. Note the characteristic peristaltic and synchronous firing patterns in both the premotor
interneurons and the motor neuron that yet demonstrate differences between the two preparations illustrated. Similarly the synaptic strength profiles are characteristic yet clearly different. Adapted from [19].
Figure 5.
The leech heartbeat CPG and the role of the switch interneurons. Top: Circuit diagram of the identified heart interneurons of the core CPG showing their synaptic interconnections, inhibitory and electrical. The two possible states of the heartbeat CPG are illustrated one with the left switch interneuron quiescent and the right switch interneuron active (corresponding to left synchronous), and the other with the left switch interneuron active and the right switch interneuron quiescent (corresponding to left peristaltic). Heart interneurons that have similar input and output connections are lumped together for ease of presentation. Bottom: Simultaneous extracellular recordings of two bilateral pairs of premotor heart interneurons [HN(R/L,3) and HN(R/L,7)] and the bilateral pair of switch interneurons [HN(R/L,5)] during a switch in coordination mode from left synchronous to left peristaltic as indicated in the circuit diagrams above. Note the invariant activity of the HN(3) (HCO) interneurons and the shift in activity of the HN(7) premotor interneurons during the switch, which is accomplished in one beat cycle as the HN(5) interneurons move reciprocally between quiescent and active states. Symbols mark the middle spike of each interneuron burst to illustrate phase relationships. Cell identity coded with color consistently across figures. Adapted from [20,24].