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# Activation of high and low affinity dopamine receptors generates a closed loop that maintains a conductance ratio and its activity correlate

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Neuromodulators alter network output and have the potential to destabilize a circuit. The mechanisms maintaining stability in the face of neuromodulation are not well described. Using the pyloric network in the crustacean stomatogastric nervous system, we show that dopamine (DA) does not simply alter circuit output, but activates a closed loop in which DA-induced alterations in circuit output consequently drive a change in an ionic conductance to preserve a conductance ratio and its activity correlate. DA acted at low affinity type 1 receptors (D1Rs) to induce an immediate modulatory decrease in the transient potassium current ( $I_A$ ) of a pyloric neuron. This, in turn, advanced the activity phase of that component neuron, which disrupted its network function and thereby destabilized the circuit. DA simultaneously acted at high affinity D1Rs on the same neuron to confer activity-dependence upon the hyperpolarization activated current ( $I_h$ ) such that the DA-induced changes in activity subsequently reduced  $I_h$ . This DA-enabled, activity-dependent, intrinsic plasticity exactly compensated for the modulatory decrease in  $I_A$  to restore the  $I_A:I_h$  ratio and neuronal activity phase, thereby closing an open loop created by the modulator. Activation of closed loops to preserve conductance ratios may represent a fundamental operating principle neuromodulatory systems use to ensure stability in their target networks.

**Keywords:** activity-dependent intrinsic plasticity, metaplasticity, metamodulation, HCN channel, stomatogastric, pyloric network

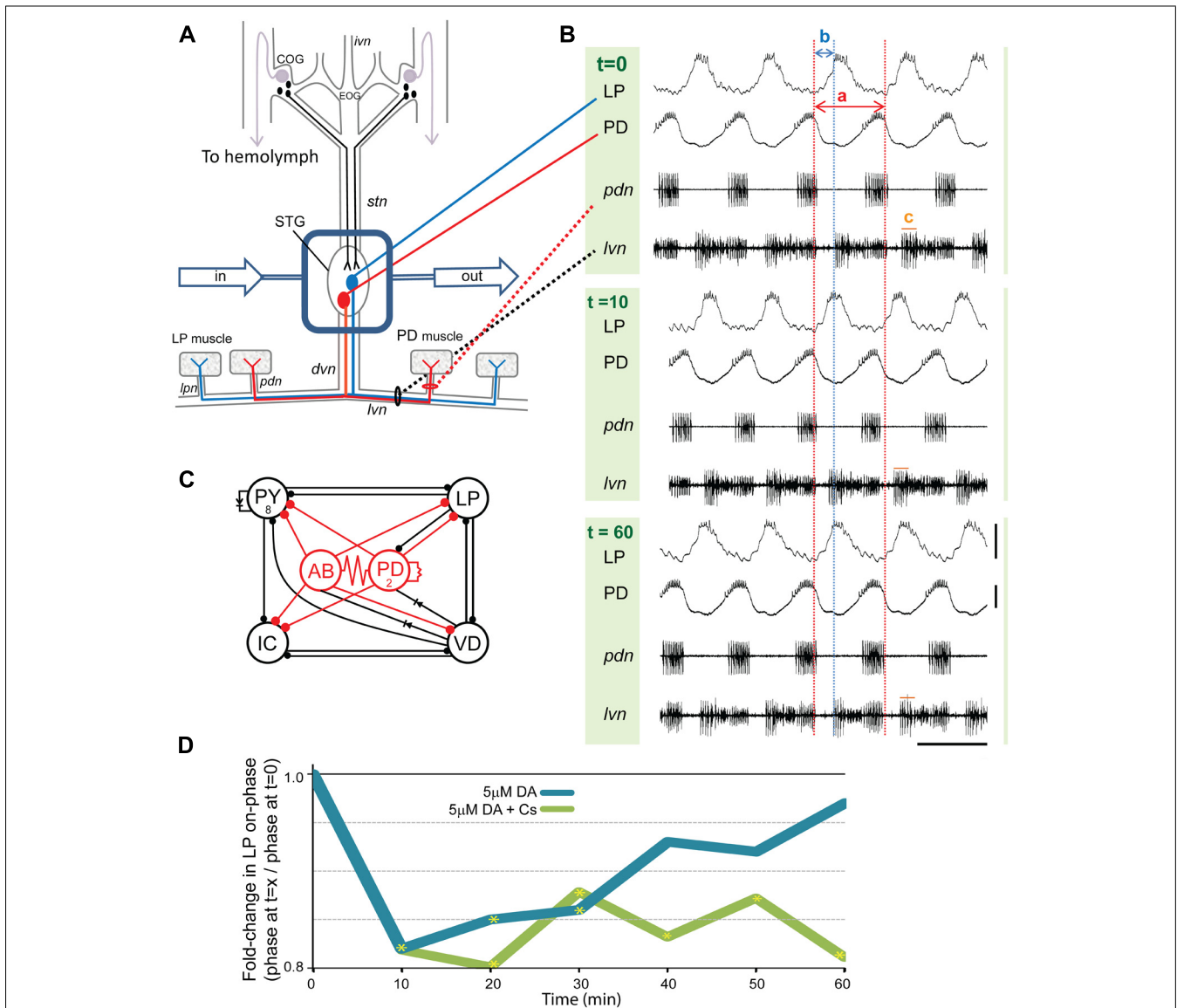
## INTRODUCTION

Neuromodulators reconfigure circuit output; but, they must confer stability as well as flexibility in order to maintain the functionality of a target network. Our knowledge of modulatory stabilizing mechanisms is limited. We suggest that modulators stabilize circuits by activating feedback loops that preserve conductance ratios and their activity correlates. Many cells maintain conductance ratios (Linsdell and Moody, 1994; MacLean et al., 2003; Schulz et al., 2006; Peng and Wu, 2007), and it is generally thought that a given conductance ratio sustains a specific activity parameter(s) (Marder and Goaillard, 2006; Hudson and Prinz, 2010; Soofi et al., 2012). A neuromodulator could establish a feedback loop if it modulated one of the conductances in the pair and conferred activity dependence on the other. In this case, modulation of the first current would contribute to changes in neuronal and circuit output that, in turn, would drive a change in the second current to restore the ratio and the activity feature. The work presented here establishes, for the first time, the existence of such a feedback loop.

The 14-neuron pyloric circuit in the spiny lobster, *Panulirus interruptus*, is a small central pattern generator (CPG) that drives the striated muscles surrounding the pylorus to produce an ordered series of contractions (Marder and Bucher, 2007). One cycle of contractions is continuously repeated to

produce constant filtering of the foregut contents. The repetitive cycle of muscle contractions is underpinned by the recurrent output of the pyloric CPG, which stems from a pace-maker kernel that rhythmically inhibits four follower neuron cell types. The follower neurons then display post-inhibitory rebound (PIR), and differences in their rates of PIR, together with the synaptic architecture, produce a tri-phasic motor pattern (Figure 1).

Follower neuron cell types have specific activity phases, meaning that a given cell type fires a burst of action potentials at the same point in each reiteration of the cyclic network output. The timing of neuronal activity phases is determined, in part, by their rate of PIR.  $I_A$  and  $I_h$  are opposing subthreshold conductances whose ratio regulates the rate of PIR (Harris-Warrick et al., 1995). Population studies on other species of crustaceans showed that pyloric neuron activity phases (Bucher et al., 2005; Goaillard et al., 2009) and their  $I_A:I_h$  ratios (Temporal et al., 2012) were invariant across individuals and lifetimes, suggesting compensatory mechanisms may exist to maintain the  $I_A:I_h$  ratio and its activity correlates. Such a compensatory mechanism(s) was revealed by overexpressing the Kv4 channels mediating  $I_A$  throughout days in organ culture. Overexpression of  $I_A$  in pyloric neurons resulted in compensatory increases in  $I_h$  that maintained the rates of PIR (MacLean et al., 2003, 2005).



**FIGURE 1 | Phase recovery in the pyloric network. (A)** *In situ* preparation: the stomatogastric nervous system (STNS) is dissected and pinned in a dish. The commissural ganglia (CoGs) contain DA neurons that project to the STG (black) and L-cells, which are the source of neurohormonal DA (purple). The well surrounding the STG (blue rectangle) is continuously superfused with saline (in/out arrows). There are ~30 neurons in the STG. The pyloric network comprises 14 STG neurons; two are drawn: pyloric dilator (PD, red), lateral pyloric (LP, blue). Network neurons interact locally within the STG and can project axons to striated muscles surrounding the foregut. The diagram shows that PD and LP neurons project their axons through identified nerves to innervate muscles (rectangles). **(B)** Spontaneous pyloric network output from one experiment during a 1 h 5  $\mu$ M DA application: one set of traces comprises two intra-cellular recordings (top) and two extra-cellular recordings (bottom) from the *in situ* preparation diagrammed in **(A)**. The three sets of traces represent recordings from the indicated time points, in minutes, directly before and after the start of DA application. Red and blue dashed lines reveal how cycle period and LP-on delay change with time. The two red lines demarcate one cycle. Cycle period (a) is defined as the time between the last spike in one PD burst and the last spike in the subsequent PD burst. Note that for each time point the last spike in the first PD burst is aligned with the first red line; however, the last spike in the second PD burst is not aligned with the second red line except at  $t = 0$ . This is because 5  $\mu$ M DA produces a

sustained average 10% reduction in cycle period. Thus, for  $t = 10$  and 60 min, the spike in the second PD burst occurs prior to the second red line. Within the indicated cycle, a blue line aligns with the first spike in LP at  $t = 0$ . The time between the last spike in PD and the first spike in LP (b) represents LP-on delay, and LP-on phase is: b/a. Note that for the  $t = 10$  min cycle, the first spike in LP occurs well before the blue line. This is because DA produces an average ~20% LP-on phase advance. LP-on phase recovery can be seen in the diagram because the first LP spike is again aligned with the blue line. Measures of pyloric output parameters can be obtained from either intra- or extra-cellular traces, and LP burst duration is indicated by (c) on the extracellular traces; scale bars: 20 mV and 500 ms. **(C)** The pyloric circuit: the diagram represents pyloric neuron interactions within the STG. Open circles represent the six cell types, numbers indicate more than one cell within a cell type: anterior burster (AB), inferior cardiac (IC), ventricular dilator (VD); filled circles, inhibitory chemical synapses; resistors and diodes, electrical coupling; red, pacemaker kernel and its output connections. **(D)** Phase recovery: the preparation shown in **(A)** was superfused with one of the two indicated treatments for 1 h and LP on-phase was measured every 10 min throughout the experiment ( $n \geq 6$ /treatment). Average fold-changes in LP on-phase are plotted for each group; yellow asterisks, significantly different from  $t = 0$ , data taken from Rodgers et al. (2011a). Note that phase recovery in 5  $\mu$ M DA was blocked by Cs.

We suggest that a mechanism to maintain the  $I_A:I_h$  ratio may also prevail during DA modulation of pyloric neurons (Rodgers et al., 2011a): there is a single LP follower neuron in the pyloric network, and it contributes to cycle frequency regulation (Weaver and Hooper, 2003). The timing of the LP activity phase is critical for this function (Johnson et al., 2011). LP expresses D1Rs but not D2Rs (Zhang et al., 2010), and a 10 min bath application of 5  $\mu$ M DA can disrupt the LP  $I_A:I_h$  ratio and induce an LP phase advance largely by decreasing LP  $I_A$  (Harris-Warrick et al., 1995). DA modulation also decreases LP burst duration and increases pyloric cycle frequency through intrinsic and network effects (Harris-Warrick et al., 1998; Rodgers et al., 2011a). During continuous DA application, the timing of LP activity phase is restored, while the DA-induced changes in burst duration and cycle frequency are maintained (Rodgers et al., 2011a); thus, a compensatory mechanism operates to restore neuronal activity phase during neuromodulation. Here we investigate this mechanism and show that LP phase recovery involves a DA- and activity-dependent (DAD) decrease in LP  $I_h$  that compensates for the modulatory decrease in LP  $I_A$  to restore the LP  $I_A:I_h$  ratio and LP activity phase.

## MATERIALS AND METHODS

### ANIMALS AND DRUGS

California spiny lobsters, *Panulirus interruptus*, were purchased from Catalina Offshore Products (San Diego, CA, USA) and Marinus Scientific (Long Beach, CA, USA) and housed at 16–18°C in saltwater aquaria at Georgia State University (Atlanta, GA, USA). Animals of both sexes were used in these experiments. TTX was from Tocris (Ellisville, MO, USA), all other reagents were from Sigma (St. Louis, MO, USA). Solutions containing DA were made fresh every 30 min in saline to prevent oxidation and reduced DA activity.

### PHYSIOLOGICAL RECORDINGS

Lobsters were anesthetized on ice for at least 30 min, followed by dissection of the stomatogastric nervous system (Figure 1), as previously described (Panchin et al., 1993). A Vaseline well was constructed around the stomatogastric ganglion (STG) which was continuously superfused for the remainder of the experiment with *Panulirus* (*P.*) saline (in mM: 479 NaCl, 12.8 KCl, 13.7 CaCl<sub>2</sub>, 39 Na<sub>2</sub>SO<sub>4</sub>, 10 MgSO<sub>4</sub>, 2 glucose, 4.99 HEPES, 5 TES; pH 7.4). Experiments were conducted at room temperature (19–21°C). Temperature was continuously monitored with a miniature probe inside the well. Temperatures changed by less than 1°C throughout the course of the day.

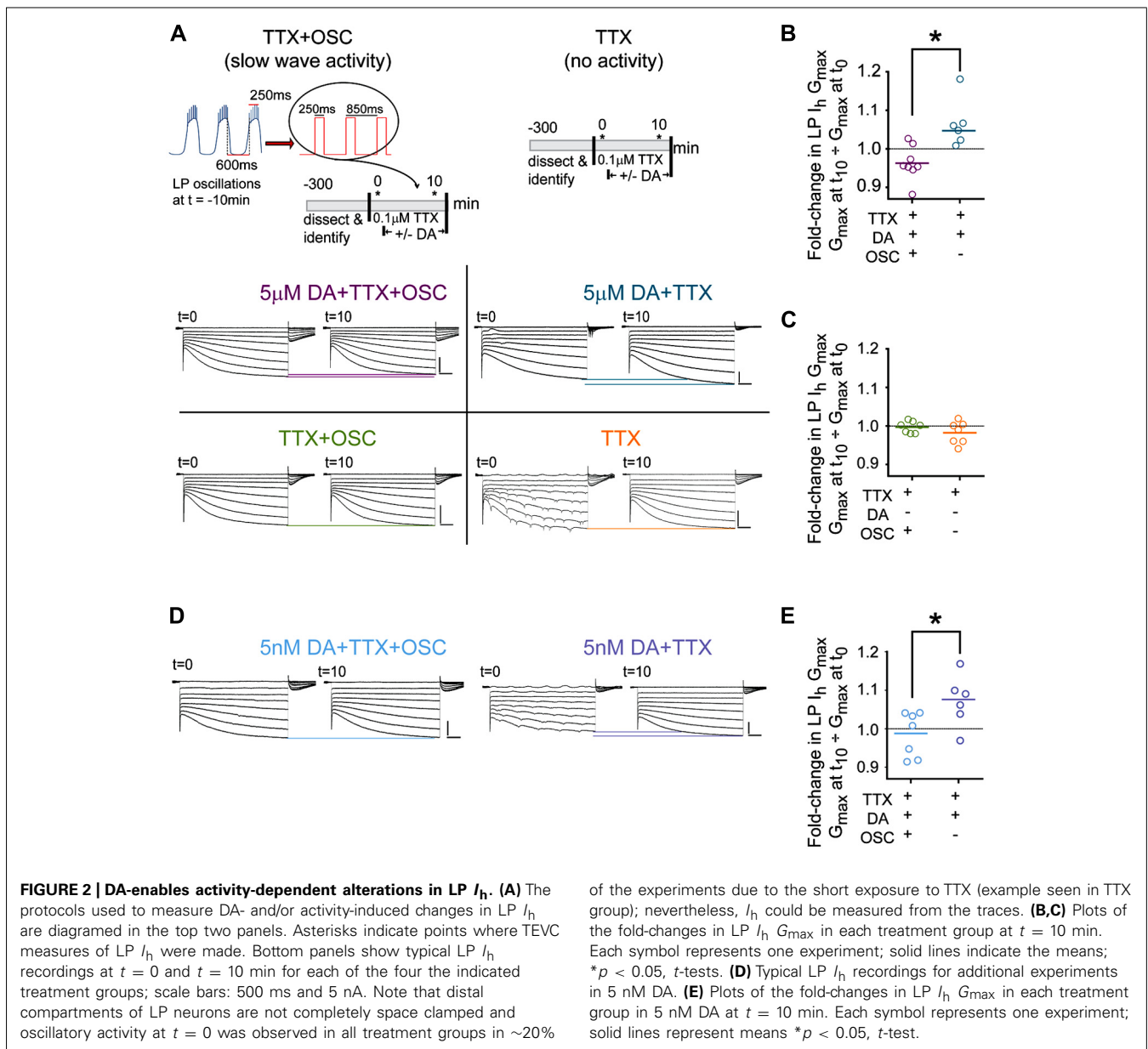
Cells were identified by combining standard intracellular and extracellular recording techniques. Lateral pyloric (LP) neurons were identified by their distinct waveforms, the timing of their voltage oscillations, and correlation of spikes on the extracellular and intracellular recordings. Intracellular somatic LP recordings were obtained using 20–40 M $\Omega$  glass microelectrodes filled with 3 M KCl connected to Axoclamp 2B or 900A amplifiers (Molecular Devices, Foster City, CA, USA). Extracellular recordings of identified motor neurons were obtained using a model 1700 differential AC amplifier (A-M Systems, Everett, WA, USA) and

stainless steel pin electrodes on the lateral ventricular nerve (*lvn*) and pyloric dilator nerve (*pdn*) and recorded with Axoscope v8.2 software (Molecular Devices, Foster City, CA, USA). Extracellular recordings were analyzed using DataView v6.3.2 (Heitler, 2009) to determine cycle period, spike frequency, burst duration, LP-on/off delays, and LP activity phase as previously described (Rodgers et al., 2011a). Reported values for all parameters represent a 10 cycle average.

Experiments in TTX blocked action potential firing and slow voltage oscillations in STG neurons. Under these conditions, the resting membrane potential of most pyloric neurons is between  $\sim -52$  and  $-62$  mV. Pyloric neuron input/output curves suggest that graded transmitter release will be minimal to non-existent at these voltages (Johnson et al., 1991, 1995). DA (100  $\mu$ M) can shift the curves (Johnson and Harris-Warrick, 1990), but a 10-fold lower concentration has a minimal effect on the strength of graded release (Kvarta et al., 2012). Pyloric neurons can oscillate in TTX if bathed in 100  $\mu$ M DA, but we do not observe pyloric oscillations in TTX at  $\leq 5$   $\mu$ M DA.

### TWO-ELECTRODE VOLTAGE CLAMP (TEVC)

For TEVC of LP  $I_h$ , the LP neuron was impaled with two micropipettes (8–10 M $\Omega$  when filled with 3 M KCl) connected to Axoclamp 2B or 900A amplifiers (Molecular Devices, Foster City, CA, USA). The well surrounding the STG was superfused with P. saline containing 100 nM TTX for  $\geq 5$  min. LP was clamped to a  $-50$  mV holding potential using pClamp software.  $I_h$  was elicited using a series of 4 s hyperpolarizing voltage steps, from  $-60$  to  $-120$  mV in 10 mV increments with 6 s between steps. Steady-state peak currents were measured by fitting the current trace back to the beginning of the hyperpolarizing voltage step using a single exponential equation. In some experiments small oscillations interrupted the current trace at  $t = 0$  (e.g., Figure 2) and prevented curve fitting. In those experiments, peak  $I_h$  was measured by subtracting the initial fast leak current from the slowly developing peak of  $I_h$  at the end of each negative voltage step. Currents were converted to conductance using ( $G = I_{\text{peak}}/(V_m - V_{\text{rev}})$ ) and fitted to a first-order Boltzmann equation.  $V_{\text{rev}} I_h = -35$  mV (Kiehn and Harris-Warrick, 1992). For TEVC measurement of peak  $I_A$  the command potential was stepped from  $-50$  to  $-90$  mV for 200 ms to remove resting inactivation. The deinactivating prepulse was immediately followed by an activation pulse to 60 mV for 400 ms to ensure that channels were maximally activated and observed changes could not be due to alterations in voltage dependence (Zhang et al., 2010). To subtract the leak current the hyperpolarizing prepulse was omitted and instead the prepulse was set to  $-40$  mV to remove  $I_A$  activation from the  $-50$  mV holding potential. For recordings to measure the LP  $I_A:I_h$  ratio in 5  $\mu$ M DA, the saline also contained 20  $\mu$ M TEA and 1  $\mu$ M PTX to block DA-induced modulatory changes in other conductances that could interfere with measures of peak currents. Recurring voltage steps to mimic slow wave oscillations and action potentials were constructed with pClamp software. When currents were not being measured, and recurring steps were not being implemented, LP was held at its initial resting membrane potential in TTX (on average,  $-59$  mV).

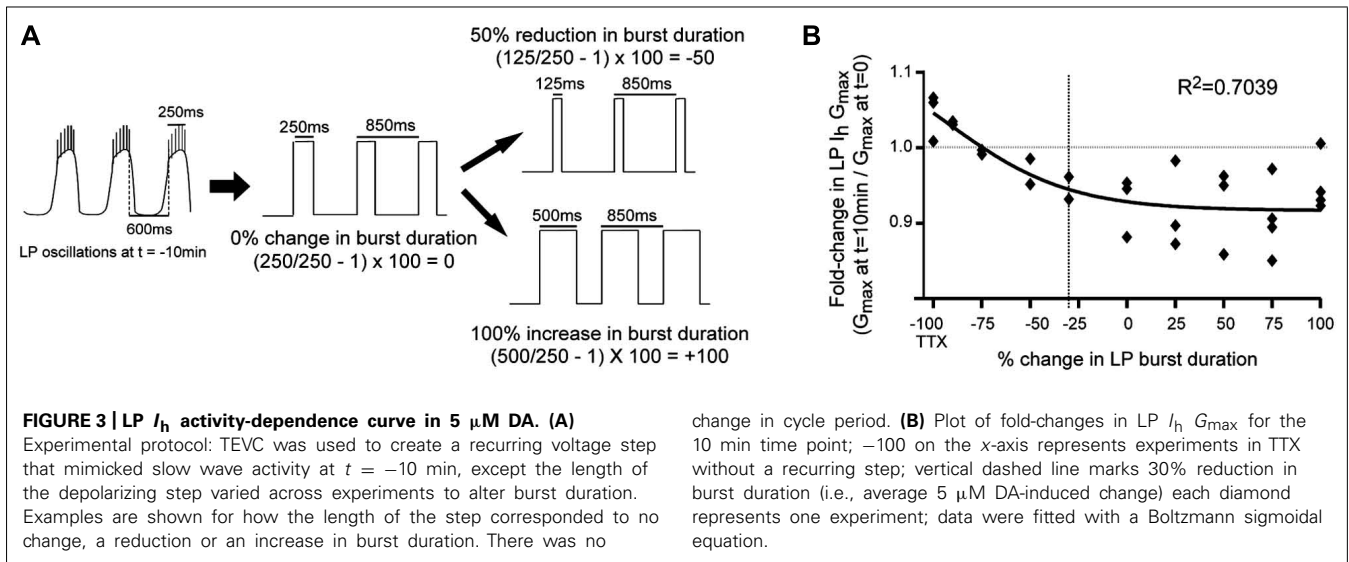


### DYNAMIC CLAMP

We used the dynamic clamp to introduce an artificial injection current ( $I_{inj}$ ) specified to counteract the metaplastic (DA modulation of activity dependent (AD) intrinsic plasticity) change in  $I_h$  in LP neurons during ongoing rhythmic pyloric activity following bath application of 5  $\mu$ M DA (Sharp et al., 1993a,b; Prinz et al., 2004a). The membrane potential of the LP soma was amplified, fed into a PCI-6052E DAQ board (National Instruments, Austin, TX, USA), and digitized at 20 kHz. The dynamic clamp program was written in the C programming language and designed to use the real time Linux dynamic controller (Dorval et al., 2001). This dynamic clamp software calculated the  $I_{inj}$  that would be active at the measured membrane potential ( $V_m$ ) given a set of model parameters as follows:

$$I_{inj} = G_{max} m (V_m - E_{rev}),$$

where  $m$  changed according to  $dm/dt = (m_{\infty} - m)/\tau_m$ , computed numerically using the first-order forward Euler method, and  $m_{\infty}$  was given by  $m_{\infty} = 1/(1 + \exp((V_m - V_{1/2})/V_{slope}))$ .  $E_{rev}$  was set to  $-35$  mV (Kiehn and Harris-Warrick, 1992). Values for  $I_h$   $\tau_m$  represent average TEVC measures from 12 experiments. LP  $I_h$  was measured in each LP neuron before the dynamic clamp experiment and  $G_{max}$ ,  $V_{1/2}$ , and  $V_{slope}$  were determined from a Boltzmann fit as described above. The predicted metaplastic change in LP  $I_h G_{max}$  was determined using the activity-dependence curve in **Figure 3** and the measured change in LP burst duration after a 10 min application of 5  $\mu$ M DA. The predicted metaplastic change in  $I_h$  conductance was subtracted with the dynamic clamp, which calculated and continuously injected current according to



the above model, where  $G_{\text{max}} = \text{measured LP } I_h G_{\text{max}} \times \text{predicted metaplastic change in LP } I_h G_{\text{max}}$ . Intracellular and extracellular recordings of LP activity throughout the experiment were obtained using a separate computer equipped with Axoscope and Clampex 9.2 software (Axon Instruments).

### STATISTICAL ANALYSIS

Data were checked for normal distribution and analyzed using parametric statistical tests with Prism software package v5.01 (GraphPad, La Jolla, CA, USA). Significance was set at  $p < 0.05$  in all cases. Individual samples that were more than 2 standard deviations from the mean were excluded from the analyses after determining the mean. This eliminated two experiments. ANOVAs are followed by *post hoc* tests that make all possible comparisons between columns (Tukey's) or that compare all columns to a single column, usually  $t = 0$  (Dunnett's). Means are followed by standard errors.

## RESULTS

### THE EXPERIMENTAL MODEL

The pyloric circuit is located in the crustacean STG (Figure 1A), and it produces a rhythmic motor output *in situ*. Each pyloric cell type displays repetitive oscillations in membrane potential with a burst of spikes riding on the depolarized plateau (Figure 1B). The circuit comprises six oscillatory cell types coupled by fast inhibitory synapses and/or gap junctions (Figure 1C). The pacemaker kernel (anterior burster (AB) + 2 PD neurons) rhythmically inhibits the four follower neuron cell types, which then display different rates of PIR. The different rates of PIR are due, in part, to differences in the expression of  $I_A$  in each follower neuron (Baro et al., 1997, 2000).  $I_A$  delays pyloric neuron PIR (Tierney and Harris-Warrick, 1992): the hyperpolarizing phase of the membrane potential oscillation removes resting inactivation from the Kv4 channels mediating  $I_A$  and activates the hyperpolarization activated cyclic nucleotide (HCN) gated channels mediating the depolarizing inward  $I_h$ . The subsequent depolarization activates Kv4 channels, and the resulting outward potassium current slows

the rate of depolarization. In this way, the ratio of  $I_A:I_h$  can influence when LP activity phase begins (termed LP-on phase). Figure 1B shows intra- and extra-cellular recordings from a typical experiment where the STG was superfused with 5  $\mu\text{M}$  DA for 1 h: DA was applied after the initial recording at  $t = 0$ . By 10 min, DA increased pyloric network cycle frequency by reducing the inherent period of the pacemaker AB neuron (Harris-Warrick et al., 1998; Rodgers et al., 2011a). DA application also reduced LP burst duration and advanced LP firing phase. The traces indicate that by 60 min in DA, network cycle frequency was still increased and LP burst duration was still decreased, but LP-on phase was restored. In previous experiments we clearly demonstrated that phase recovery was AD: if the experiment shown in Figure 1B was repeated with continuous injection of a depolarizing bias current into LP to block the DA-induced decrease in LP burst duration, then the LP phase advance occurred, but phase recovery did not (Rodgers et al., 2011a). We also showed that phase recovery in the presence of 5  $\mu\text{M}$  DA could be blocked by bath application of CsCl to reduce  $I_h$  (Figure 1D).

### DA- AND ACTIVITY-DEPENDENT (DAD) REGULATION OF LP $I_h$ IN 5 $\mu\text{M}$ DA

We first tested the idea that DA conferred activity-dependence upon LP  $I_h$  by measuring  $I_h$  in LP neurons that showed different activity patterns. In these experiments, LP neurons have one of two activity patterns: either LP activity is completely blocked (TTX), or LP displays normal slow wave but not spike activity (TTX + OSC). LP  $I_h$  is measured in each of these two groups in the presence and absence of DA resulting in four treatment groups. The experiment, which is diagrammed in Figure 2A, was as follows: after dissection and cell identification, the STG was superfused with TTX for 5 min to block spike and slow wave activity, and the TTX was present throughout the remainder of the experiment. Next, at  $t = 0$ , LP  $I_h$  was measured with somatic TEVC. After the first measure of LP  $I_h$ , DA was or was not added to the superfusate and LP  $I_h$  was re-measured after 10 min. The voltage of LP was continuously controlled with TEVC throughout the experiment.

Between measures of LP  $I_h$ , a recurrent step mimicking LP slow oscillatory activity at  $t = -10$  min was (TTX + OSC) or was not (TTX) implemented. Frequency, duration, and amplitude of the recurrent steps were chosen for each preparation individually depending upon measured activity at  $t = -10$  min: frequency and duration of the recurrent step corresponded to average cycle frequency and LP burst duration at  $t = -10$  min, respectively; the step and holding potentials corresponded to the average peak and nadir of the LP oscillation at  $t = -10$  min, respectively. In the absence of the recurring voltage step, LP was held at its initial resting membrane potential in TTX ( $-59$  mV on average). Typical LP  $I_h$  recordings for each treatment group are shown in **Figure 2A**.

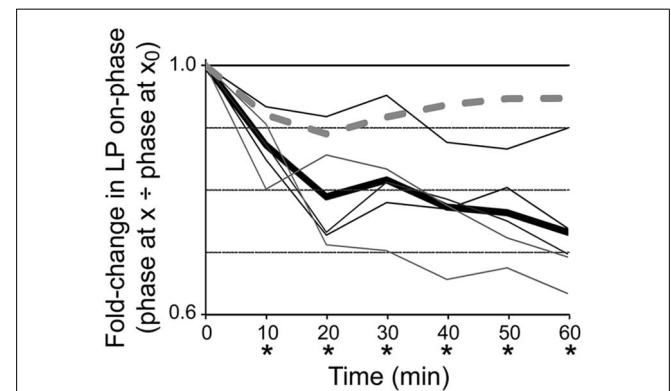
The results indicated that DA conferred activity dependence upon LP  $I_h$ : in the presence of DA, the fold-change in LP  $I_h G_{\max}$  varied according to LP activity (**Figure 2B**;  $t$ -test,  $p < 0.004$ ); by 10 min in 5  $\mu$ M DA average LP  $I_h G_{\max}$  was significantly decreased in preparations with the slow wave LP activity pattern (paired  $t$ -test,  $t = 0$  vs. 10 min,  $p = 0.0491$ ) and significantly increased in preparations showing no LP activity (paired  $t$ -test,  $p = 0.0285$ ). In the absence of DA the fold-change in LP  $I_h G_{\max}$  was not significantly different between treatment groups (**Figure 2C**,  $t$ -test,  $p = 0.256$ ) and there was no significant change in LP  $I_h G_{\max}$  by  $t = 10$  min relative to  $t = 0$  in preparations where slow wave activity was mimicked (paired  $t$ -test,  $p = 0.1166$ ) or activity was completely blocked (Wilcoxon matched pairs signed rank test,  $p = 0.2969$ ). We previously demonstrated that 5 nM DA acting at high affinity LP D1Rs permitted a decrease in LP burst duration to produce an increase in LP  $I_h G_{\max}$  that persisted well beyond DA washout (Rodgers et al., 2011a). This suggested that perhaps high affinity D1Rs receptors might also mediate the more rapid DAD regulation of LP  $I_h G_{\max}$  observed in **Figure 2B**. To test this hypothesis, we repeated the experiments diagrammed in **Figure 2A**, but applied 5 nM rather than 5  $\mu$ M DA (**Figure 2D**). The results were consistent with the hypothesis; in the presence of 5 nM DA, the fold-change in LP  $I_h G_{\max}$  at  $t = 10$  min varied according to activity (**Figure 2E**,  $t$ -test,  $p = 0.0321$ ). Interestingly, LP  $I_h G_{\max}$  did not change over time in 5 nM DA preparations where slow wave activity was mimicked (paired  $t$ -test,  $t = 0$  vs. 10 min,  $p = 0.5962$ ); however, a complete block of activity produced a clear trend toward an increase in LP  $I_h G_{\max}$  (paired  $t$ -test,  $p = 0.0596$ ), and the magnitude of the increase was similar to that observed in 5  $\mu$ M DA (compare **Figures 2B** vs. **2E**). The difference in the TTX + OSC treatment groups in 5 nM DA (no change in  $G_{\max}$ ) vs. 5  $\mu$ M DA (decrease in  $G_{\max}$ ) may be due to the fact that micromolar DA can regulate calcium dynamics during oscillations in membrane potential (Johnson et al., 2003; Kadiri et al., 2011). For all treatment groups the voltages of half activation changed by  $\leq 2.3$  mV on average, and LP  $I_h$  voltage dependence is not considered further here. In sum,  $\geq 5$  nM DA permitted activity to differentially regulate LP  $I_h G_{\max}$ ; but, neither 5 nM DA alone nor changes in activity alone significantly altered LP  $I_h G_{\max}$ ; i.e., DA did not modulate LP  $I_h$ , but conferred activity-dependence upon LP  $I_h$ .

#### DAD REGULATION OF LP $I_h$ IS NECESSARY FOR PHASE RECOVERY

Our previous study suggested that LP phase recovery during sustained DA modulation was triggered by a change in LP burst

duration (Rodgers et al., 2011a). In order to understand if and how DAD regulation of LP  $I_h$  restored the timing of the LP activity phase in 5  $\mu$ M DA, it was necessary to determine how LP  $I_h$  varied according to changes in LP burst duration. An LP  $I_h$  activity-dependence curve for changes in burst duration was constructed by repeating the previous experiments in 5  $\mu$ M DA for the TTX + OSC treatment group, except that the length of the depolarizing step varied across experiments to mimic a change in burst duration (**Figure 3A**). A plot of the fold-change in LP  $I_h G_{\max}$  vs. percent change in LP burst duration at  $t = 10$  min was best-fitted with a Boltzmann sigmoidal equation. DA (5  $\mu$ M) produced an average 30% decrease in LP burst duration (Rodgers et al., 2011a), and so, according to the activity-dependence curve, LP  $I_h G_{\max}$  should be reduced by  $\sim 6\%$  in 5  $\mu$ M DA during on-going activity (**Figure 3B**, dashed line). This decrease in LP  $I_h$  is consistent with our hypothesis that DAD regulation of LP  $I_h$  compensates for the DA-induced modulatory decrease in LP  $I_A$  to restore the  $I_A:I_h$  ratio and the timing of LP activity phase.

In order to determine if DAD regulation of LP  $I_h$  was necessary for phase restoration, we used the activity-dependence curve in conjunction with dynamic clamp experiments to abrogate DAD regulation of LP  $I_h$  (**Figure 4**). The experimental preparation was as shown in **Figure 1A**. After dissection and cell identification the STG was superfused with TTX for 5 min; LP  $I_h$  was measured with TEVC and values for  $G_{\max}$ ,  $V_{1/2}$  and  $V_{\text{slope}}$  were subsequently incorporated into the dynamic clamp model for  $I_{\text{inj}}$  (see Section "Materials and Methods"). TTX was immediately washed out with saline for 90 min. LP burst duration was measured at the end of the wash followed by application of 5  $\mu$ M DA from  $t = 0$ –60 min. The predicted fold-change in LP  $I_h G_{\max}$  due to DAD regulation was determined using the activity-dependence curve in **Figure 3**



**FIGURE 4 | DAD regulation of LP  $I_h$  is necessary for phase recovery in 5  $\mu$ M.**

Plots of fold-changes in LP-on phase over time for dynamic clamp (solid lines) and control (dashed line) experiments indicate that introduction of a dynamic clamp current to abrogate DAD regulation of LP  $I_h$  prevents phase recovery; thin lines, individual experiments with dynamic clamp ( $n = 5$ ); thick line, average for experiments with dynamic clamp; dashed line, control experiment that was exactly the same as the dynamic clamp experiments except that the dynamic clamp was turned off during the 1 h superfusion with 5  $\mu$ M DA. Repeated measures ANOVA with Dunnett's *post hoc* tests that compared all time points to  $t = 0$  showed that average LP-on phase did not recover in experimental preparations,  $F(6,4) = 16.04$ ,  $p < 0.0001$ ; \* $p < 0.05$ . Note that phase did recover in the control experiment.

and the measured change in LP burst duration from  $t = 0$  to  $t = 10$  min, and was subsequently incorporated into the dynamic clamp model for  $I_{inj}$  (see Section “Materials and Methods”). From  $\sim t = 10$  to 60 min, dynamic clamp was used to remove the predicted DAD regulation of LP  $I_h$ , i.e., to add back, in the form of dynamic clamp current, the same amount of  $I_h$  that was predicted to have been lost because of DAD regulation. LP-on phase was subsequently measured every 10 min from  $t = 0$ –60 min. Plots of the fold-change in LP-on phase over the course of the experiment demonstrated that 5  $\mu$ M DA-induced the usual phase advance, but removing DAD regulation of LP  $I_h$  prevented LP-on phase recovery (compare **Figures 4** vs. **1D**). It also prevented LP-off phase recovery [repeated measures ANOVA:  $F(6,4) = 3.119$ ,  $p = 0.0210$ ]. However, it should be noted that the recovery of LP-off phase may be complicated by the PY cell activity phase. The PY-LP synapse contributes to the timing of LP-off phase, especially in DA; thus, any change in LP-on phase that subsequently alters the timing of PY activity through the LP–PY synapse may also indirectly affect LP-off phase (Johnson et al., 2011). From these experiments we conclude that DAD regulation of LP  $I_h$   $G_{max}$  is necessary for LP-on phase restoration.

#### DAD REGULATION OF LP $I_h$ COMPENSATES FOR MODULATORY CHANGES IN LP $I_A$ TO RESTORE $I_A:I_h$

Thus far the data are consistent with our working model for how phase advance and recovery occur in 5  $\mu$ M DA: 5  $\mu$ M DA initially alters the LP  $I_A:I_h$  ratio by decreasing LP  $I_A$ , and this creates a phase advance (Harris-Warrick et al., 1995; Zhang et al., 2010). DA (5  $\mu$ M) also produces a 30% reduction in LP burst duration, and this subsequently initiates a process that generates a compensatory decrease in LP  $I_h$  to restore the LP  $I_A:I_h$  ratio and produce phase recovery. In order to further test this hypothesis, we repeatedly measured the LP  $I_A:I_h$  ratio during a 1 h 5  $\mu$ M DA application accompanied by a recurring step that mimicked a 30% reduction in LP burst duration. At  $t = 0$ , peak LP  $I_A$  was measured at +60 mV and peak LP  $I_h$  was measured at –120 mV. DA (5  $\mu$ M) was immediately applied for 1 h and peak currents were re-measured at  $t = 10, 30$ , and 60 min. During the DA application, whenever peak currents were not measured, LP received a recurring step. Plots of the average fold-changes in the peak  $I_A:I_h$  ratio (**Figure 5A**) and average peak  $I_A$  and  $I_h$  (**Figure 5B**) suggested that our hypothesis was incorrect or incomplete. The average  $I_A:I_h$  ratio significantly decreased over time (**Figure 5A**) because the decreases in peak LP  $I_h$  did not fully compensate for the decreases in peak LP  $I_A$  (**Figure 5B**).

It is noteworthy that DA-induced a change in both LP burst duration and cycle period (Rodgers et al., 2011a), but our step only mimicked the change in burst duration. We next asked if the DA-induced increase in cycle frequency contributed to DAD regulation of LP  $I_h$   $G_{max}$ , by repeating the experiments to measure the LP  $I_A:I_h$  ratio but using a recurring voltage step that mimicked both the average 30% decrease in LP burst duration and the 10% increase in cycle frequency. In this case, the average  $I_A:I_h$  ratio did not change significantly throughout the experiment [**Figure 5C**, repeated measures ANOVA,  $F(3,4) = 2.161$ ,  $p = 0.1457$ ], despite the fact that by 10 min, average peak LP  $I_A$  was significantly and stably reduced to  $81 \pm 4\%$  of its initial value [**Figure 5D**,

repeated measures ANOVA,  $F(3,4) = 16.91$ ,  $p = 0.0001$ ]. The ratio did not change because by 10 min in DA, average peak LP  $I_h$  was significantly and stably reduced to  $87 \pm 3\%$  of its original value [**Figure 5D**, repeated measures ANOVA,  $F(3,4) = 6.983$ ,  $p = 0.0057$ ]. We conclude that the AD mechanism that regulates LP  $I_h$   $G_{max}$  in the presence of DA integrates information on both neuronal burst duration and cycle period.

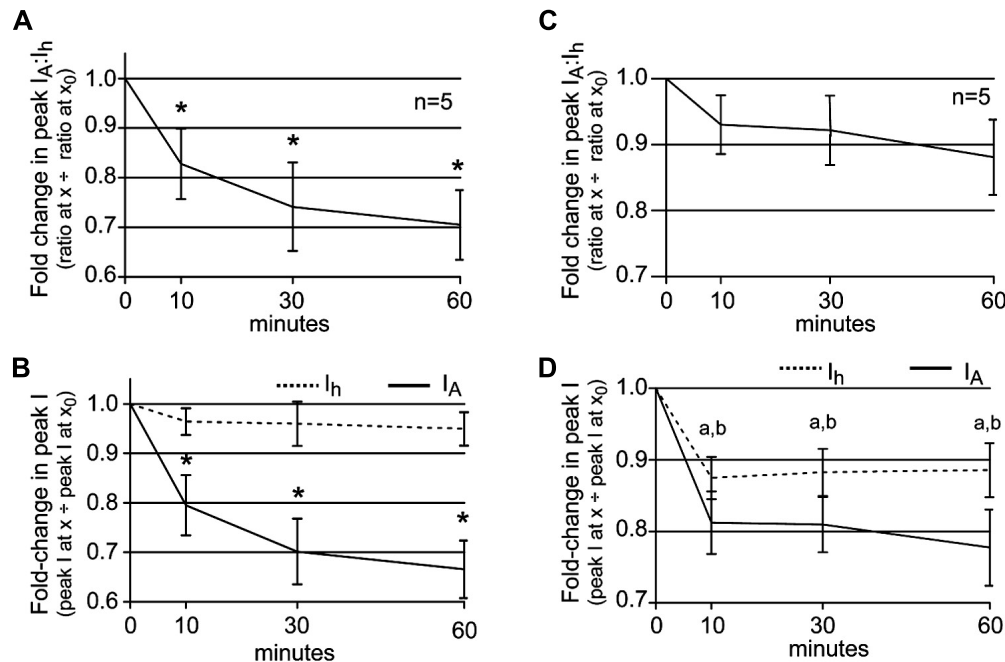
#### SPIKE ACTIVITY DELAYS THE EFFECT OF CHANGES IN SLOW WAVE ACTIVITY

Overall, the data supported our hypothesis: in the presence of 5  $\mu$ M DA and average DA-induced changes in LP slow wave activity, the DA-induced fold-change in LP  $I_A$  was compensated by a similar fold-change in LP  $I_h$ . However, one aspect of the data did not fit with our working model. The ratio could be restored by 10 min (**Figure 5**), but phase recovery required 60 min on average (**Figures 1B,D**). It is possible that restoration of the LP  $I_A:I_h$  ratio was necessary (**Figure 4**) but not sufficient for phase recovery, and that one or more unidentified slower processes were also involved. Alternatively, one major difference between the experiments shown in **Figures 1** vs. **5** was the presence vs. absence of spike activity. If a  $Ca^{2+}$  sensor participated in this homeostatic mechanism to maintain the LP  $I_A:I_h$  ratio (Gunay and Prinz, 2010), then spike activity and DA-induced changes in slow wave activity might have opposing effects on steady-state  $Ca^{2+}$ , and spike activity could delay the compensatory decrease in LP  $I_h$  by slowing the rate of change of steady-state  $Ca^{2+}$ . To investigate this idea, we repeated experiments to measure the LP  $I_A:I_h$  ratio using a recurring step that mimicked not only slow wave activity, but also, spike activity.

During normal LP activity, spikes passively spread to the soma and neuropil from a distal spike initiation zone (siz). We mimicked spike activity generated at the siz with depolarizing current injections into the soma. We reasoned that LP HCN channels, which are located in the neuropil (Goeritz et al., 2011), will experience a similar depolarization regardless of whether the spikes initiate at the soma or siz, because the two structures are roughly equidistant from the neuropil. This logic rests on the untested assumption that the electrotonic properties and protein composition of the entire primary neurite membrane between soma and spike initiation zone are homogeneous and that electrotonic potentials spread with similar efficiency in both directions. We also made untested assumptions about LP spike amplitude and duration. Peak voltage (+40 mV) and duration (2 ms) of PD spikes have been directly measured from intra-axonal recordings (Ballo et al., 2012). We assumed LP and PD spikes would be similar and used these values here.

Previous work suggested that activity-dependent regulation can be coded by the pattern of spike activity and not simply the total amount of depolarization (Gorbunova and Spitzer, 2002). We performed two series of experiments to determine if spike activity influenced the LP  $I_A:I_h$  ratio either by the total amount of depolarization produced or by the pattern of depolarization. The total amount of depolarization was mimicked with a step to +40 mV whose duration equaled the average number of spikes per burst multiplied by 2 ms. Patterned spike activity was mimicked by 2 ms depolarizations to +40 mV separated by the average interspike





**FIGURE 5 | The LP  $I_A:I_h$  ratio is maintained in 5  $\mu\text{M}$  DA when DA application is accompanied by DA-induced changes in slow wave activity. (A)** A plot of the fold-changes in the LP  $I_A:I_h$  ratio (mean  $\pm$  SEM) throughout a 1 h superfusion with 5  $\mu\text{M}$  DA and implementation of a recurring voltage step that mimicked the DA-induced 30% decrease in LP burst duration, but no change in cycle frequency. The ratio significantly decreased with time; repeated measures ANOVA with Dunnett's *post hoc* tests that compare all time points to  $t = 0$ ,  $F(3,4) = 7.322$ ,  $p = 0.0032$ . **(B)** Plots of the fold-changes in peak LP  $I_A$  and  $I_h$  (mean  $\pm$  SEM) from the same experiments as in **(A)**. Repeated measures ANOVAs with Dunnett's *post hoc* tests that compare all time points to  $t = 0$  indicate that only LP  $I_A$  was

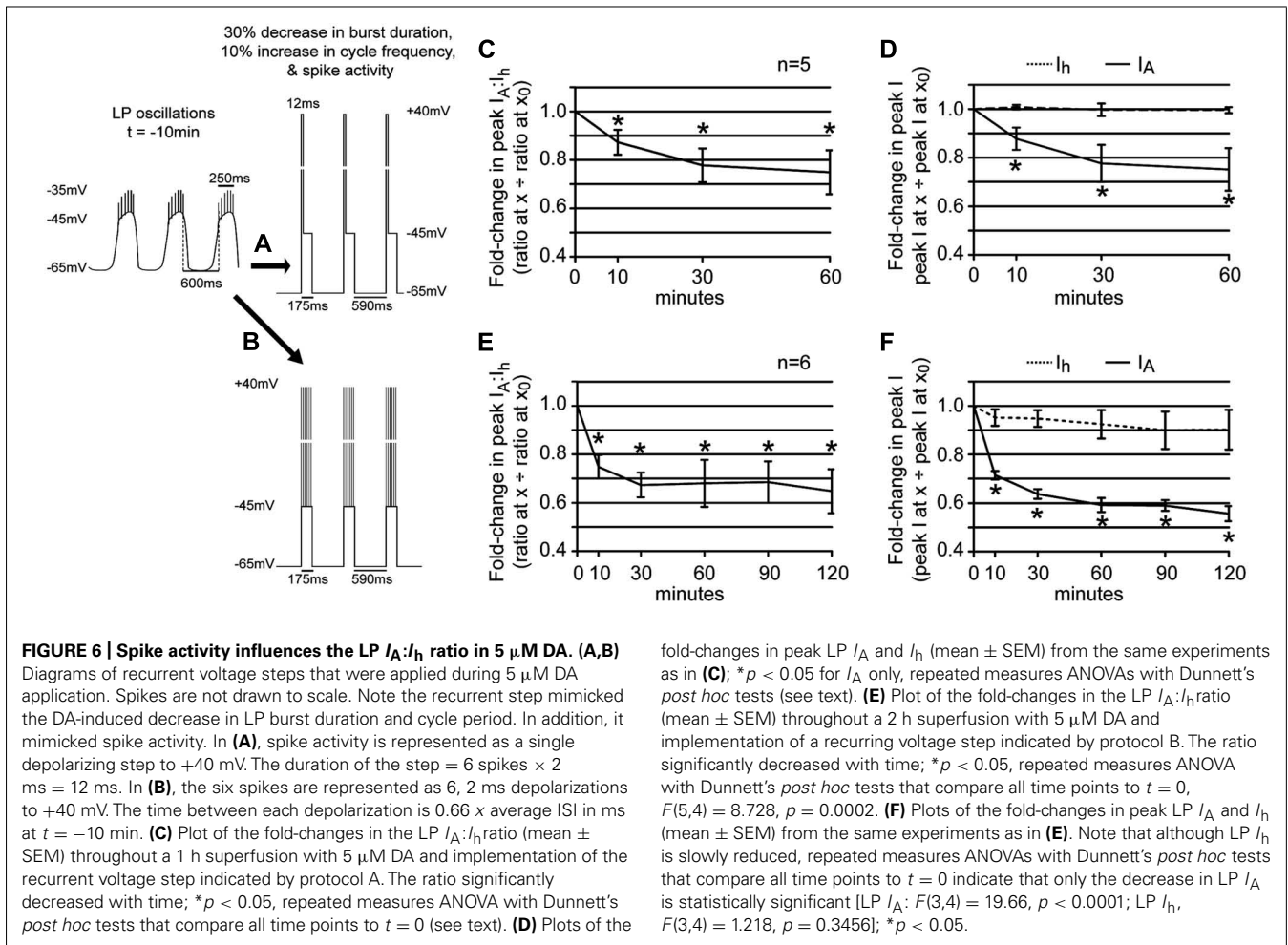
significantly decreased [LP  $I_A$ :  $F(3,4) = 19.66$ ,  $p < 0.0001$ ; LP  $I_h$ ,  $F(3,4) = 1.218$ ,  $p = 0.3456$ ]. \* $p < 0.05$ . **(C)** Plot of the fold-changes in the LP  $I_A:I_h$  ratio (mean  $\pm$  SEM) throughout a 1 h superfusion with 5  $\mu\text{M}$  DA and implementation of a recurring voltage step that mimicked the DA-induced 30% decrease in LP burst duration and a 10% increase in cycle frequency. The ratio did not change significantly over time (repeated measures ANOVA, see text). **(D)** Plots of the fold-changes in peak LP  $I_A$  and  $I_h$  (mean  $\pm$  SEM) from the same experiments as in **(C)** show that both currents are stably altered by 10 min; a and b indicate a significant change in LP  $I_A$  and  $I_h$ , respectively, based on repeated measures ANOVA with Dunnett's *post hoc* tests that compare all time points to  $t = 0$ ,  $p < 0.05$  (see text).

interval (ISI), and the number of depolarizations was equal to the average number of spikes per burst.

In the first set of experiments a depolarizing step to +40 mV was superimposed upon the recurrent voltage step that mimicked LP slow wave activity in 5  $\mu\text{M}$  DA (**Figure 6A**). The duration of the step to +40 mV corresponded to the average number of spikes per burst at  $t = -10$  min multiplied by 2 ms. Note that the average number of spikes per burst does not change significantly during a 1 h 5  $\mu\text{M}$  DA application [repeated measures ANOVA,  $F(6,8) = 0.8920$ ,  $p = 0.5083$ , data not shown]. Surprisingly, this short depolarization on top of the usual recurrent voltage step that mimicked a 30% decrease in LP burst duration and a 10% increase in cycle frequency completely abolished the effect of DA-induced changes in slow wave activity upon LP peak  $I_h$ . The LP  $I_A:I_h$  ratio significantly decreased under these conditions [**Figure 6C**; repeated measures ANOVA:  $F(3,3) = 6.114$ ,  $p = 0.0149$ ] because, there was no reduction in LP  $I_h$  (**Figure 6D**, mean  $\pm$  SEM fold-change in LP peak  $I_h$  at 10 min =  $1.008 \pm 0.010$ ). The insignificant change in LP  $I_h$  throughout the 1 h 5  $\mu\text{M}$  DA application could not compensate for the significant decrease in LP  $I_A$  [**Figure 6D**; repeated measures ANOVAs:  $I_h$ ,  $F(3,4) = 0.1801$ ,  $p = 0.9078$ ;  $I_A$ ,  $F(3,3) = 5.251$ ,  $p = 0.0228$ ]. Note that the change in LP  $I_A$  was not significantly different between experiments that did

(**Figure 6D**) vs. did not (**Figure 5D**) mimic spike activity along with DA-induced changes in slow wave activity [two-way ANOVA: treatment,  $F(1,28) = 0.08$ ,  $p = 0.7789$ ; time,  $F(3,28) = 6.83$ ,  $p = 0.0014$ ; interaction,  $F(3,28) = 0.33$ ,  $p = 0.8065$ ].

We next asked if we could delay, but not abolish the compensatory decrease in LP  $I_h G_{\text{max}}$  by better mimicking the spike pattern (**Figure 6B**). To do this, we included an ISI in between each 2 ms depolarization to +40 mV that was equal to the average ISI at  $t = -10$  min multiplied by 0.66, because a 1 h 5  $\mu\text{M}$  DA application reduced the mean ISI to 66% of its initial value [repeated measures ANOVA:  $F(6,4) = 4.002$ ,  $p = 0.0065$ , data not shown]. Including patterned spike activity in the recurrent voltage step delayed the compensatory reduction in LP  $I_h G_{\text{max}}$  (**Figure 6F**). By 10 min in 5  $\mu\text{M}$  DA, the compensatory reduction in LP peak  $I_h$  was significantly smaller for protocols that did (**Figure 6F**) vs. did not (**Figure 5D**) include patterned spike activity on top of the slow wave (Student's *t*-test,  $p = 0.0014$ ). Although a delayed and slowly developing compensatory reduction in LP  $I_h G_{\text{max}}$  was elicited with protocol B, it was not large enough to compensate for the decrease in LP  $I_A$ , even by 2 h (**Figure 6E**). This is because the patterned spike activity also unexpectedly regulated LP  $I_A$ : the reduction in peak LP  $I_A$  was significantly larger for protocols that did (**Figure 6F**) vs. did not



(Figure 5D) include patterned spike activity on top of the slow wave [two-way ANOVA, treatment,  $F(1,32) = 25.76$ ,  $p < 0.0001$ ; time,  $F(3,32) = 38.53$ ,  $p < 0.0001$ ; interaction,  $F(3,32) = 3.45$ ,  $p = 0.0278$ ]. This large decrease in LP  $I_A$  was most likely a technical artifact. Kv4 channels are located throughout the LP somatodendritic membrane (Baro et al., 2000). According to our untested assumption, Kv4 channels in the neuropil will experience typical changes in membrane potential with each spike mimic; however, this is not true for somatic Kv4 channels. The average LP membrane potential typically recorded at the soma at the peak of spike activity is  $-36 \pm 4$  mV because spikes are severely attenuated as they passively spread to the soma. Thus, a  $+40$  mV depolarization at the soma is unrealistic and most likely generates an artificially large decrease in the somatic LP  $I_A$ . Nonetheless, based on these experiments we can conclude that DAD regulation of LP  $I_h$   $G_{\text{max}}$  integrates information on burst duration, cycle period, and spike activity.

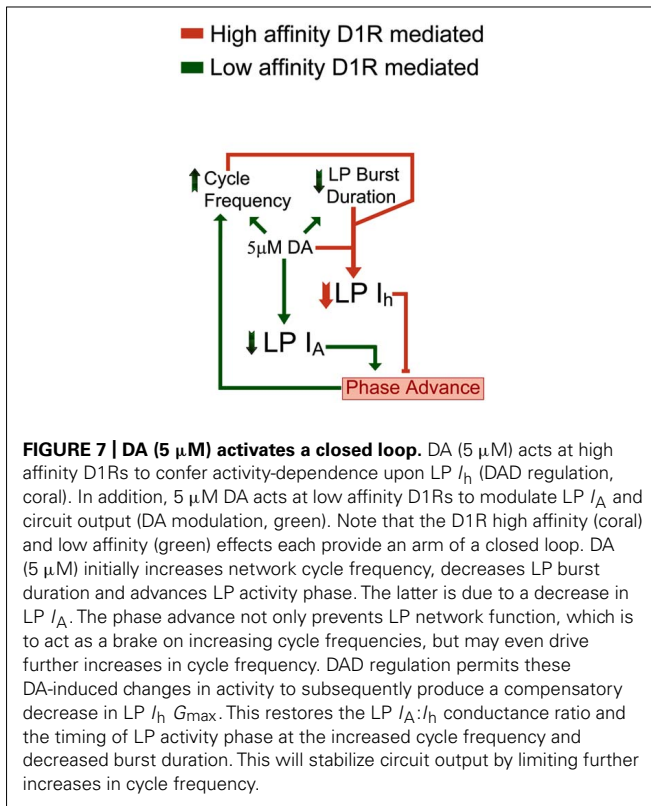
## DISCUSSION

The principal finding of our study is that  $5 \mu\text{M}$  DA simultaneously creates flexibility and stability in a rhythmically active neural network by activating a closed loop (Figure 7). DA acts at both low and high affinity D1Rs to alter activity and enable AD intrinsic

plasticity, respectively. The feedback loop re-established a conductance ratio that was modified by DA, and thereby restored a neuronal phase relationship during a sustained increase in cycle frequency. The generation of closed loops via modulator-enabled AD intrinsic plasticity may represent a fundamental organizing principle used by modulatory systems to preserve conductance ratios and their associated activity correlates, while at the same time altering other aspects of circuit output.

## DA SIMULTANEOUSLY GENERATES FLEXIBILITY AND STABILITY BY ACTIVATING HIGH AND LOW AFFINITY D1RS

Like most systems, DA transmission takes two forms in the stomatogastric nervous system, tonic, and phasic. DA neurons in the commissural ganglia project to the STG and release DA into open synapses; DA then diffuses to its sites of action before re-uptake (Oginsky et al., 2010). To the best of our knowledge, DA levels have not been measured in the STG, but in other systems that use volume transmission, DA is tonically present at  $\sim\text{nM}$  levels (range: 0.1–100 nM) and can transiently increase to  $\sim\mu\text{M}$  levels (range: 0.1–100  $\mu\text{M}$ ) near the release sites of bursting DA neurons (Zoli et al., 1998; Schultz, 2007; Fuxe et al., 2010). In addition, the STG is located in a blood vessel and is bathed by neurohormonal DA (Sullivan et al., 1977; Marder and Bucher, 2007). Generally



speaking, high affinity receptors respond to  $\sim\text{nM}$  DA (tonic) and low affinity receptors respond to  $\sim\mu\text{M}$  DA (phasic). We have previously shown that LP possesses both high and low affinity D1Rs that mediate different effects on  $I_A$ . High affinity receptors were activated by a tonic 1 h application of 0.5 nM but not 0.05 nM DA and produced a persistent (i.e., non-reversible) increase in LP  $I_A$  through a translation-dependent mechanism (Rodgers et al., 2011b, in press). On the other hand, low affinity D1Rs responded to bath application of  $\sim\mu\text{M}$  DA and immediately and reversibly decreased LP  $I_A$  by altering its biophysical properties (Zhang et al., 2010). In this study we showed that high affinity D1Rs do not simply act through slow mechanisms (hours) to produce persistent changes in ionic currents, but can also rapidly (seconds to minutes) confer activity-dependence upon an ionic conductance to generate a feedback loop.

Concomitant stimulation of both low and high affinity LP D1Rs activates a closed loop that maintains neuronal activity phase while other aspects of neuronal output are altered (Figure 7). A 5  $\mu\text{M}$  but not 5 nM DA application alters pyloric network activity (Rodgers et al., 2011a); therefore, DA acts at low affinity receptors to modulate circuit output. At least three key aspects of pyloric network output are modulated by DA (Rodgers et al., 2011a): on average, cycle frequency is increased by  $\sim 10\%$ , LP burst duration is decreased by 30% and LP firing phase is advanced by  $\sim 20\%$ . The LP phase advance is largely due to a DA-induced reduction in LP  $I_A$  (Harris-Warrick et al., 1995; Zhang et al., 2010). These alterations in network output disrupt LP network function (Johnson et al., 2011). Normally, LP acts through the LP-PD synapse to slow increasing cycle frequencies (Nadim et al., 1999; Weaver

and Hooper, 2003; Mamiya and Nadim, 2004, 2005; Johnson et al., 2011). The timing of LP activity phase is critical for this function because, LP inhibition has different effects according to when it occurs during the pacemaker oscillation, and a phase advance can even increase cycle frequency (Thirumalai et al., 2006; Johnson et al., 2011). This creates a potential for spiraling changes in network output that would destabilize the system. However, besides eliciting these alterations in network activity, DA acts at high affinity D1Rs to permit AD regulation of LP  $I_h$ . This allows the DA-induced changes in cycle frequency and LP burst duration to subsequently elicit a reduction in LP  $I_h$  that exactly compensates for the modulatory decrease in LP  $I_A$  to restore the timing of LP activity phase. Restoring LP firing phase re-establishes LP network function which is to slow increasing cycle frequency (Johnson et al., 2011). This could limit the DA-induced increase in cycle frequency driven by DA actions on the pacemaker and stabilize circuit performance at the increased network cycle frequency, decreased LP burst duration, and potentially altered LP input:output gain (Burdakov, 2005). It should also restore the initial phasing of rhythmic pyloric muscle contractions, but at an increased cycle frequency. Interestingly, burst duration and on/off-delays scale with cycle period in the natural population throughout development and over a wide range of temperatures (Bucher et al., 2005; Goallard et al., 2009; Tang et al., 2010). Thus, the closed loop uncovered here may be part of a more extensive control system that synchronizes these network characteristics over multiple time scales and through multiple mechanisms.

#### DOPAMINERGIC TONE MIGHT MAINTAIN THE $I_A:I_h$ RATIO DURING NON-DOPAMINERGIC PERTURBATIONS TO ACTIVITY

Landmark studies from the Marder group demonstrated equivalent neuronal and network firing patterns can arise from different sets of intrinsic and synaptic conductances (Golowasch et al., 1999a,b; Prinz et al., 2004b; Schulz et al., 2006, 2007). This work led to the idea that conductances co-vary over time in order to maintain a particular activity feature, an idea that was supported by existing ion channel overexpression studies (MacLean et al., 2003, 2005). These findings were unexpected and caused the Selverston group to ask: can the output of a network made up of disparate components be robust to perturbation (Szucs and Selverston, 2006)? Within a population, peak PD  $I_A$  and PD  $I_h$  each varies by  $>3$ -fold across individuals; but, all individuals maintain the same PD  $I_A:I_h$  ratio (Temporal et al., 2012). Selverston's group reasoned that if PD  $I_A$  were blocked with 4-AP in multiple preparations, then PD  $I_h$  would no longer be balanced in any preparation, and the variable amounts of PD  $I_h$  in each preparation would be revealed in distinct PD activity patterns (Szucs and Selverston, 2006; Nowotny et al., 2007). To their surprise, all blocked neurons produced similar activity patterns, suggesting either that the pyloric network is not made up of disparate components in each individual or that rapid compensatory mechanisms must exist to maintain activity. Our findings suggest the latter may be true: modulator-enabled, AD feedback loops could have produced compensatory changes in  $I_h$   $G_{\text{max}}$  that maintained activity in the Selverston group's experiments. Indeed, modulatory inputs were intact in the latter studies (Szucs and Selverston, 2006), and 4-AP significantly alters pyloric cycle period and neuronal burst

durations (Tierney and Harris-Warrick, 1992). Together, the data imply that modulatory tone could enable multiple AD feedback loops that maintain conductance ratios and their activity correlates under a variety of conditions.

### PHASE MAINTENANCE

Both intrinsic and synaptic mechanisms can operate over different time scales to maintain pyloric neuron phase relationships when cycle frequency varies. Synaptic depression rapidly promotes phase maintenance by proportionately delaying neuronal firing as synapses increasingly recover from depression with longer cycle periods (Nadim et al., 1999, 2003; Manor et al., 2003). DA can modulate synaptic dynamics to promote phase maintenance: 10  $\mu$ M DA decreased the time constants of short-term depression and its recovery at the PD–LP graded synapse, thus contributing to phase maintenance with changing network frequency (Kvarta et al., 2012). It is also worth noting that PY inhibition onto LP plays an important role in determining LP off-phase and this impact is enhanced in DA (Johnson et al., 1993, 1995), contributing to the shortening and stabilization of LP activity phase (Johnson et al., 2011). Fast intrinsic conductances, including  $I_A$ , can act in conjunction with synaptic mechanisms to promote phase maintenance in pyloric neurons (Bose et al., 2004; Greenberg and Manor, 2005; Rabbah and Nadim, 2005). Slower processes can also play a role in pyloric neuron phase maintenance. In a combined physiological and computational study on the spiny lobster, Hooper et al. (2009) demonstrated that a conductance with slow activating and inactivating kinetics (seconds to minutes) could explain adjustment of PIR and phase maintenance in PY neurons in the presence of altered cycle period. Goaillard et al. (2010) showed the crab LP neuron possessed a similar mechanism. Neither of these studies identified the slow conductance.  $I_h$  was considered, but blocking  $I_h$  did not terminate the mechanism. The authors suggested the conductance could be an unidentified slow potassium or calcium conductance, deinactivation of a fast sodium current, a pump current or a combination of opposing currents with fast kinetics. Our research extends these previous findings by revealing the existence of a DA-enabled mechanism(s) for phase maintenance that involves preserving the  $I_A:I_h$  ratio. DAD regulation of  $I_h$  may contribute to phase maintenance in other rhythmically active systems where phase relationships are maintained amidst changes in cycle frequency (Dicaprio et al., 1997; Jacobson et al., 2009).

### MECHANISM OF DAD REGULATION OF LP $I_h$

DAD regulation of LP  $I_h$   $G_{max}$  integrates information on multiple aspects of activity. The neurons under study exhibit slow membrane potential oscillations ( $\sim 20$  mV at 1–2 Hz) and action potentials riding on the depolarized plateau of each oscillation. DAD regulation integrated information on cycle period and burst duration, as well as spike activity. Integration may be an

epi-phenomenon created by voltage clamp measures of the entire population of HCN channels, and it is possible that distinct sub-cellular populations of HCN channels are differentially regulated by different types of activity.

It is not clear if DAD regulation of LP  $I_h$  represents a single integrator that is influenced by multiple types of activity; or, if multiple molecular integrators exist, each of which is sensitive to a distinct aspect of activity. AD mechanisms that regulate  $I_h$  density could rely on both  $Ca^{2+}$  release and  $Ca^{2+}$  entry. It is tempting to speculate that the mechanism(s) that is sensitive to burst duration and cycle frequency senses  $Ca^{2+}$  release from stores while the mechanism(s) that is sensitive to spiking senses  $Ca^{2+}$  entry through voltage-gated calcium channels. It was previously noted that  $Ca^{2+}$  release from stores can regulate  $I_h$  density in hippocampal neurons (Narayanan et al., 2010), and that in the pyloric AB neuron,  $Ca^{2+}$  release oscillates with oscillations in membrane potential (Kadiri et al., 2011). Thus, changes in cycle frequency and burst duration could alter steady-state  $Ca^{2+}$  contributed by store release. In addition,  $Ca^{2+}$  entry through glutamate receptors can regulate surface expression of HCN channels over minutes in cultured hippocampal neurons (Noam et al., 2010). Perhaps this mechanism may be generalized to  $Ca^{2+}$  entry through other types of channels, such as high threshold voltage-gated  $Ca^{2+}$  channels that open maximally during spike activity. In this case, spike frequency could also influence steady-state  $Ca^{2+}$ . Previous studies show that micromolar DA can enhance LP voltage-gated  $Ca^{2+}$  currents (Johnson et al., 2003; Kloppenburg et al., 2007), and in the AB neuron micromolar DA can act on  $IP_3$  receptors to increase release from stores (Kadiri et al., 2011). Since higher concentrations of DA can alter  $Ca^{2+}$  dynamics, these data suggest that DAD regulation of LP  $I_h$  may vary according to DA concentrations as well as activity patterns.

The mechanisms by which high affinity D1Rs permit AD regulation of LP  $I_h$   $G_{max}$  are not known. Traditionally, D1Rs are thought to act through  $G_{\alpha s}$  to regulate adenylyl cyclase activity and thereby cAMP levels, which in turn regulate PKA. D1R induced increases in PKA activity can regulate surface expression of cortical neuron glutamate receptors (Sun et al., 2005). Thus, in one scenario, a cAMP-PKA signaling pathway may modulate AD surface expression of HCN channels. Indeed such a pathway can regulate AD Kv4 channel trafficking in hippocampal neurons (Hammond et al., 2008). One of the invertebrate adenylyl cyclases, rutabaga, is a known coincidence detector that can be influenced by both  $G_{\alpha s}$  and  $Ca^{2+}$  (Tomchik and Davis, 2009; Gervasi et al., 2010), and rutabaga could underpin DA's permissive effect.

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