Serotonin in the Developing Stomatogastric System of the Lobster, *Homarus americanus*

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ABSTRACT: We studied the development of the serotonergic modulation of the stomatogastric nervous system of the lobster, *Homarus americanus*. Although the stomatogastric ganglion (STG) is present early in embryonic development, serotonin immunoreactivity is not visible in the STG until the second larval stage. However, incubation of the STG with exogenous serotonin showed that a serotonin transporter is present in embryonic and early larval stages. Serotonin uptake was blocked by paroxetine and 0% Na+/H+ saline. The presence of a serotonin transporter in the embryonic STG suggests that hormonally liberated serotonin could be taken up by the STG, and potentially released as a “borrowed transmitter”. Consistent with a potential hormonal role, serotonin is found in the pericardial organs, a major neurosecretory structure, by midembryonic development. The rhythmic motor patterns produced by embryonic and larval STGs were decreased in frequency by serotonin.

INTRODUCTION

Serotonin is an important neuromodulator in lobsters, where it acts both as a circulating hormone and a neu-
considerably after the time at which the ganglion is first rhythmically active (Casasnovas and Meyrand, 1995; Richards et al., 1999). Specifically, although the STG is rhythmically active by midway through embryonic life (Casasnovas and Meyrand, 1995; Richards et al., 1999), serotonin immunoreactivity in the neuropil of the STG does not strongly appear until the second larval stage (LII) (Kilman et al., 1999), when the animal is already freely feeding (Factor, 1995).

Numerous studies have suggested that, in addition to its actions as a neuromodulator in adult nervous systems, serotonin plays important roles in the development of neuronal circuits. For example, serotonin influences process outgrowth and synapse formation (Haydon et al., 1984; Goldberg and Kater, 1989; Diefenbach et al., 1995; Mercer et al., 1996), and the manipulation of serotonin levels during development alters the structure and function of identified neurons and connections (Mooney et al., 1998; Persico et al., 2000; Sullivan et al., 2000; Benton and Beltz, 2001). This suggests that serotonin might play a developmental role in the stomatogastric nervous system at the time it first appears, around the second larval stage. As a first step in approaching this question, we were curious to determine if serotonin receptors and serotonin transporters are present in the stomatogastric nervous system early in development. In the process of asking these questions we discovered significant developmental alterations in the properties of one of the neuromuscular junctions in the lobster stomach, and in the effects of serotonin on this junction. Specifically, in young animals, serotonin appears to accentuate the intrinsic excitability of the muscle fibers, while in adults there is a strong modulation of the gain of the nerve-muscle synapse.

METHODS

Animals and Saline

Adult lobsters, *H. americanus*, of both sexes were purchased from local fishermen (Commercial Lobster, Boston, MA) and kept in aerated artificial seawater tanks. Embryos and larvae were either obtained from the New England Aquarium and held in artificial seawater tanks at Brandeis University, or obtained from the Massachusetts State Lobster Hatchery and Research Station at Martha’s Vineyard and held in chilled running sea water at the Marine Biological Laboratory, Woods Hole. The saline composition (in mM) was: 479.12 NaCl, 12.74 KCl, 13.67 CaCl2, 20 Mg2SO4, 3.91 Na2SO4, 5 HEPES, pH = 7.4–7.5.

Embryos were staged according to Helluy and Beltz (1991), and are referred to as their percentage of embryonic development. In the terminology we use in this article, E50 is 50% of embryonic development. Embryos hatch to become the first larval stage, LI.

Immunocytochemistry

Embryonic and LI stomachs were dissected in saline according to Kilman et al. (1999). Embryonic ventral nerve cords and pericardial organs (POs) were dissected by making a dorsal incision along the thoracic midline, and pinning down the thoracic body wall alongside the ventral nerve cord. The dissected preparations were incubated for 30 min to 1 h in indicated serotonin concentrations, washed thoroughly with saline, then fixed overnight in 4% paraformaldehyde in 0.1 M sodium phosphate buffer, pH 7.4. They were then washed four to five times in PTA (0.1 M sodium phosphate buffer, 0.3% Triton X-100, and 0.1% NaN3, pH 7.4) at approximately 1 h intervals. The preparations were then incubated overnight in a rat antiseroptotonin monoclonal antibody (Accurate Chemical and Scientific Corp.) at a final dilution of 1:100 with 10% goat normal serum (GNS). The next day, the tissues were washed again with PTA four to five times at 1 h intervals and then incubated in secondary antibody [FITC or Alexafluor 488 (Molecular Probes)] at 1:400 with 10% GNS. Finally, the preparations were washed as above in 0.1 M sodium phosphate buffer without Triton and mounted in 80% glycerol and 20% 0.02 M sodium phosphate buffer. Preparations were then imaged with the appropriate filters on either a BioRad MRC-600 or Leica TCS laser scanning confocal microscope. Optical sections were taken approximately every 1–2 μm and maximum projections of the z series were made using Confocal Assistant (BioRad). Photomicrographs were processed in Photoshop 6.0 and Canas 8.0 before printing on an Epson 1280 printer.

The serotonin uptake inhibitor paroxetine (Smith, Kline, and Beecham) was suspended in normal saline with vigorous shaking and then used immediately. Sodium free saline had the following composition (in mM): N-methyl d-Glutamate 479.12, KCl 12.74, CaCl2 13.67, MgSO4 20, K2SO4 3.91, and HEPES 5. For all recipes, the pH was adjusted to 7.4 using HCl and NaOH.

Recordings from the Neuromuscular Junctions

For adult recordings, the stomach was dissected out and the p1 muscle (Maynard and Dando, 1974) was isolated and pinned in a small Sylgard-lined Petri dish (Dow Corning, Midland, MI) along with the lateral ventricular nerve (lvn), which contains the axon of the Lateral Pyloric (LP) neuron that innervates p1 [Fig. 1(B)]. The end of the lvn was surrounded with a Vaseline™ well and stimulated to produce excitatory junction potentials (EJPs) with bipolar stainless steel pins using an A-M Systems (Carlsborg, WA) isolated pulse stimulator (Model 2100).

For the earlier developmental stages, the entire stomach was dissected from the animal and pinned in the Sylgard dish [Fig. 1(A)]. When necessary, the STG input to the
muscles was removed by cutting the descending lvn, and a suction electrode was used to stimulate the lvn with a Grass S88 stimulator. Intracellular recordings were then made from muscle bundles that are referred to as lpm [LP-innervated muscle] and lgm [Lateral Gastric (LG) innervated muscle]. We use this nomenclature because the defining boundaries of the adult muscles are not present at this early developmental stage. When these boundaries are defined in the adult, lpm fibers will make up muscles p1 and cpv6, and lgm fibers will become gm6 and gm8 (Maynard and Dando, 1974).

Preparations were continuously superfused with physiological saline at a rate of 8–10 mL/min, and cooled to 9–13°C with a Peltier cooling system. Serotonin (Sigma, St. Louis, MO) was dissolved in saline and added to the superfusion system via a switch at the inflow port. Adult EJPs were recorded in single electrode current clamp with an AxoClamp 2A amplifier and viewed in Clampex 8.1 (Axon Instruments, Foster City, CA). Recording electrodes for adult p1 fibers had resistances of 8–15 MΩ, filled with 0.6 M K$_2$SO$_4$ and 20 mM KCl. Electrodes used in the earlier developmental stages were 50–100 MΩ and were filled with 4 M K acetate and 20 mM KCl.

Reported EJP amplitudes are the average amplitude of 30 consecutive EJPs stimulated at 1 Hz (adult) or the first of two paired stimuli at 250 ms intervals (embryo and LI). The stimulus frequency was chosen to prevent facilitation of the postsynaptic response. The coefficient of variation was computed as the standard deviation of the amplitudes of the 30 events divided by the mean amplitude. The decay time was computed as the time from the peak of the EJP to one-third of the peak amplitude. Quantitative analysis of EJP kinetics was performed with Clampfit 8.1 (Axon Instruments) and statistical analysis was performed with SigmaStat 2.0 (Jandel Scientific Software, San Rafael, CA). Data are reported as means ± S.E.M (standard error of the mean).

**RESULTS**

The lobster, *H. americanus*, undergoes embryonic development over several months before hatching into its first larval stage (LI). It then molts several times, progressing through larval stages LII and LIII before metamorphosing into the postlarval LIV stage (Factor, 1995). Although the STG is present and active by the midembryonic stage (Casasnovas and Meyrand, 1995), serotonin (5-HT) immunoreactivity in the STG

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**Figure 1** Schematic diagram of the stomatogastric nervous system and recording configuration for various stages of development. In embryos, larvae, and adults, the paired commissural ganglia (CoGs) and esophageal ganglion (OG) project to the STG via the stomatogastric nerve (stn). Motor neurons in the STG project through the lateral ventricular nerve (lvn) and innervate the stomach muscles. (A) Embryo/LI recording configuration. Intracellular recordings were made from the Lateral Pyloric neuron innervated p1 muscle (lpm in embryos and larvae). A suction electrode was placed on the lvn to stimulate synaptic events. (B) Adult recording configuration. The p1 muscle and attached ossicle were dissected free of the stomach and the lvn stimulated with an extracellular electrode. In both (A) and (B), fibers were impaled with glass microelectrodes.
does not first appear until about the LII stage (Kilman et al., 1999).

5-HT Uptake in Embryos

Figure 2(A) shows an embryonic STG (E91) labeled with an antibody against 5-HT in control conditions. Note that there is no immunoreactivity present. The preparation in Figure 2(B) was incubated in $10^{-4} \text{M}$ 5-HT for 1 h, washed briefly, and then processed for 5-HT immunoreactivity. Note the dense 5-HT labeling in the processes throughout the neuropil. These processes are derived from the serotonergic Gastro-Pyloric Receptor (GPR) neurons (Katz et al., 1989; Katz and Harris-Warrick, 1989). Figure 3(A) shows 5-HT labeling in the posterior portion of an E91 preparation incubated in $10^{-4} \text{M}$ 5-HT for 30 min. Clearly visible are two groups of GPR cell bodies (asterisks and double asterisks) as well as 5-HT immunoreactive fibers in the lvn and dorsal ventricular nerve (dvn) projecting towards the STG (STG not shown). Figure 3(B) and (C) show single 2 μm optical sections of the GPR cell bodies shown in Figure 3(A), 5-HT labeling is present in both the cytoplasm and nuclei of the GPR cell bodies.

On the basis of data from 29 embryonic stomach preparations incubated in concentrations of 5-HT from $10^{-4} \text{M}$ to $10^{-9} \text{M}$, the threshold for significant 5-HT loading into the embryonic STG was between $10^{-8} \text{M}$ and $10^{-7} \text{M}$, and maximal intensity of staining was seen by about $10^{-5} \text{M}$ serotonin. Following uptake of 5-HT by the GPR neurons, an intriguing aspect of the immunocytochemical labeling for 5-HT is the apparent staining of the nuclei of GPR cells [Fig. 3(B,C)].

Paroxetine is a potent inhibitor of 5-HT transport (Bourin et al., 2001). To test the effects of paroxetine on 5-HT transport in the STG, we incubated dissected stomachs with $10^{-4} \text{M}$ 5-HT and varying concentrations of paroxetine. Figure 4(A) shows that there was no 5-HT labeling in the STG of an E86 animal treated with $10^{-4} \text{M}$ 5-HT and $10^{-7} \text{M}$ paroxetine ($n = 4$). When incubated with $10^{-4} \text{M}$ 5-HT with $10^{-7} \text{M}$ paroxetine [Fig. 4(B); $n = 3$], faint 5-HT labeling was visible in the STG neuropil. Figure 4(C) shows that $10^{-5} \text{M}$ paroxetine failed to block uptake of $10^{-4} \text{M}$ 5-HT ($n = 3$).

In other systems, 5-HT uptake depends on the presence of extracellular Na$^+$ (Humphreys et al., 1994). Figure 4(D) shows an example of an STG from
an E93 embryo that was incubated in $10^{-4} M$ serotonin in Na$^+$-free saline. Note that the staining in the STG was significantly attenuated. In three out of five preparations the STGs incubated in 5-HT and Na$^+$-free saline showed only faint staining in the main process, while the other two preparations showed slight staining in the finer processes as well. All preparations showed attenuated staining when compared with the controls.

**Serotonin Staining in Embryonic POs**

The precocious appearance of the 5-HT uptake system suggests the possibility that hormonally liberated 5-HT could be taken up by the STG, and possibly used as a “borrowed neurotransmitter”, growth factor, or transcriptional regulator (Lebrand et al., 1996, 1998; Cases et al., 1998; Vitalis et al., 1998) early in development. The POs are a major neurosecretory structure that liberates a variety of substances into the hemolymph (Cooke and Sullivan, 1982). Because the STG is in a major artery just anterior to the heart, it is ideally positioned to receive hormones liberated from the POs. The POs in embryos contain 5-HT (Pulver and Marder, 2002).

Figure 4 Serotonin uptake is blocked by paroxetine and low sodium. (A) An E86 STG incubated with $10^{-4} M$ 5-HT and $10^{-4} M$ paroxetine. (B) An E80 STG incubated with $10^{-4} M$ 5-HT and $10^{-7} M$ paroxetine. (C) An E83 STG incubated with $10^{-4} M$ 5-HT and $10^{-8} M$ paroxetine. (D) An E93 STG incubated with $10^{-4} M$ serotonin in 0% sodium saline. All images are maximum projections of 10–13 1.8-μm optical sections. Scale bar is 50 μm.

Figure 5 Serotonin in the ventral nerve cord and pericardial organs in an E52 embryo. Fibers project from the SEG and T1-5 to putative hormonal release sites in the POs. Fibers originating in the SEG also ramify in a second neurosecretory region, the AR. Image is a maximum projection of 23 3-μm optical sections. Scale bar is 50 μm.
densely labeled varicosities in the PO region as well as less extensive varicosities in the anterior region (AR), another presumed neurosecretory structure. The 5-HT stained segmental nerves that connect the POs to the ventral nerve cord are visible, showing that the 5-HT-containing PO terminals are derived from ventral nerve cord structures. Similar data were seen in seven complete preparations with the ventral nerve cord attached to the POs. These data show that much of the serotonin neurosecretory system is a potential source of hormonally delivered serotonin to the STG and its musculature in the embryo.

**Effects of Serotonin on Embryonic and Larval Motor Patterns**

Serotonin is an important modulator of the STG motor patterns in adult animals (Marder and Eisen, 1984; Flamm and Harris-Warrick, 1986a,b; Katz et al., 1989; Katz and Harris-Warrick, 1989, 1990a,b), including *H. americanus* (Beltz et al., 1984; Marder and Richards, 1999). The embryonic STG is already rhythmically active by the midembryonic stage (Casasnovas and Meyrand, 1995; Le Feuvre et al., 1999, 2001; Richards et al., 1999; Richards and Marder, 2000), and this activity is similar to that seen in LI animals (Casasnovas and Meyrand, 1995; Richards et al., 1999; Richards and Marder, 2000). Therefore, we were curious to determine if serotonin could influence STG motor patterns in embryos and LI animals, before the STG itself acquires its serotonin.

Figure 6(A) shows dual intracellular recordings from two muscles, the lpm (innervated by the LP motor neuron) and the lgm (innervated by the LG motor neuron) in a LI animal. Under control conditions the preparation was rhythmically active, and the discharges recorded in the two muscles occurred almost synchronously. In response to bath application of 10^{-5} M 5-HT the burst frequency decreased and the burst duration increased [Fig. 6(B)]. These effects were reversible upon washing the preparation in normal saline [Fig. 6(C)].

In these experiments the STG was removed and the lvn was stimulated. Figure 8(A) shows overlays of the averages of 30 consecutive EJPs recorded at each time period. Note that the amplitude in the adult is considerably smaller than what is seen in the embryo and LI. The EJP amplitude increased slightly from 12.9 ± 3.9 mV in embryos to 17.9 ± 2.8 mV in larvae. However, in adults, EJP amplitude decreased significantly from both embryonic and larval values to 3.3 ± 0.5 mV (one-way ANOVA on ranks, p < .05, n = 8 for embryos, n = 10 for larvae, n = 15 for adults).

In the process of studying the effects of serotonin on the motor patterns in the embryos we noticed that in some preparations serotonin had a marked effect on the duration and shape of the EJPs. The top panel of Figure 7 shows a recording from a lpm in an E91 preparation. The bottom trace shows the preparation in the presence of 10^{-5} M 5-HT. Note that in 5-HT some of the depolarizations are broader, and it appears that single EJPs are triggering action potentials (asterisks). These regenerative events are more common in the presence of 5-HT than in control conditions (top panel, single asterisk).

Of the experiments included in this study, four of nine embryonic preparations and three of six larval preparations had these altered events when serotonin was applied to spontaneously active preparations driven by the STG. These data suggested the possibility that serotonin could be acting directly on the muscle membrane, and made it important to study nerve-evoked EJPs under more controlled conditions to determine if serotonin was having direct actions at the neuromuscular junction.

**Developmental Regulation of LP-Evoked EJPs in Control Saline**

In this section we first compare nerve-evoked EJPs in embryos, LI, and adult muscles. In the analysis of EJP properties described below, only preparations that did not show either spontaneous action potentials or serotonin-evoked action potentials were used. This was done to study separately the properties of the EJPs and the action potentials. Thus, we have excluded for this analysis those preparations in which the amplitude and time course of EJPs were likely to be “contaminated” by muscle fiber regenerative currents. The criteria for inclusion in this data set was that successive EJPs had to be essentially identical in time course and wave-form.

In these experiments the STG was removed and the lvn was stimulated. Figure 8(A) shows overlays of the averages of 30 consecutive EJPs recorded at each time period. Note that the amplitude in the adult is considerably smaller than what is seen in the embryo and LI. The EJP amplitude increased slightly from 12.9 ± 3.9 mV in embryos to 17.9 ± 2.8 mV in larvae. However, in adults, EJP amplitude decreased significantly from both embryonic and larval values to 3.3 ± 0.5 mV (one-way ANOVA on ranks, p < .05, n = 8 for embryos, n = 10 for larvae, n = 15 for adults).

Figure 8(B) shows a comparison of the EJP decay at the three time periods. When the EJPs are scaled to the same amplitude, changes in decay times are readily seen. Figure 8(B) shows that the time to decay for adults was faster than for the embryos and larvae. Times to one-third amplitude for EJPs were 95 ± 24 ms in embryos, 98 ± 16 ms in larvae, and 37 ± 3 ms in adults. The decay times are highly variable, but the decrease in decay time from both embryos and larvae to adults is significant (one-way ANOVA, p < .05, n = 8 for embryos, n = 10 for larvae, n = 15 for adults).
Figure 8(C) compares the variability in the release of neurotransmitter at the three developmental time periods. In each preparation we recorded 30 consecutive EJPs stimulated at 1 Hz, and computed the coefficient of variation (CV) of the 30 EJP amplitudes. Figure 8(C) shows 10 consecutive EJPs overlaid in embryos, larvae, and adults to demonstrate the variability in amplitude. The CV of embryonic EJPs was 0.227 ± 0.038 while the CV in larvae was 0.311 ± 0.045 and the CV in adults was 0.135 ± 0.015. The decrease in CV from embryos and larvae to adults was significant (one-way ANOVA, $p < .05$, $n = 8$ for embryos, $n = 10$ for larvae, $n = 15$ for adults).

Age-Dependent Effects of Serotonin on EJPs

Figure 9 compares the effects of $10^{-5} \text{M 5-HT}$ on EJP amplitude, time to decay, and CV in embryos, LIs,
and adults. Figure 9(A,B) shows that there was no significant difference in EJP amplitude in embryos (control: 13.3 ± 4.4 mV, 5-HT: 14.0 ± 5.1 mV, p = .554, paired t test, n = 7) and larvae (control: 20.2 ± 2.7 mV, 5-HT: 24.5 ± 3.1 mV, p = .108, paired t test, n = 6), but EJP amplitude in adults increased significantly from 3.9 ± 0.8 mV in control to 9.6 ± 1.9 mV in 5-HT (p = .006, paired t test, n = 7).

Consistent with its failure to increase EJP amplitude in embryos and larvae, 5-HT did not significantly decrease the CV of EJP amplitude, as seen in Figure 8(B) (embryo: control, 0.215 ± 0.042, 5-HT, 0.208 ± 0.035, p = .781, paired t test, n = 7; larvae: control, 0.292 ± 0.056, 5-HT, 0.180 ± 0.032, p = .132, paired t test, n = 6). In adults, the CV significantly decreased in the presence of serotonin, indicative of an enhancement of transmitter release (from 0.122 ± 0.027 in control saline to 0.076 ± 0.017, p = .034 in 5-HT, paired t test, n = 7).

Serotonin did not significantly affect the decay time in embryos (control, 81 ± 23 ms, 5-HT, 182 ± 40 ms, p = .064, paired t test, n = 7), larvae (control, 82 ± 17 ms to 134 ± 51 ms, p = .315, paired t test, n = 5), or adults (from 35 ± 3 to 30 ± 3 ms, p = .241, paired t test, n = 7). Although there are no statistically significant differences between the

Figure 7 Bath application of serotonin can change the shape of EJPs in rhythmically active preparations. Intracellular recordings from lpm in an E84 animal. Top trace, control saline; bottom trace, 10−5 M 5-HT. Asterisks indicate onset of action potentials triggered by single EJPs. Note that one widened depolarization in 5-HT fails to trigger a muscle action potential (arrow). Resting potentials: −64 mV (control), −58 mV (5-HT).

Figure 8 Comparison of evoked EJPs in embryos, LIs, and adults. All recordings are from the lpm/p1 muscle. (A) Overlaid averages of 30 consecutive EJPs evoked at 1 Hz in an embryo, LI, and adult. (B) Average of 30 consecutive EJPs in an embryo, LI, and adult scaled to the same amplitude. (C) Thirty consecutive EJPs overlaid to show variability in amplitude.
Serotonin and Muscle Action Potentials

Spontaneous muscle action potentials were never seen in the adult muscle preparations. However, spontaneous muscle action potentials were sometimes seen in embryonic and larval preparations, and serotonin increased the likelihood that EJPs would trigger muscle spikes. Figure 10 shows recordings from a LI preparation in which nerve-evoked EJPs sometimes triggered muscle action potentials in control saline. The top panel shows recordings of 35 responses triggered on the stimulus. Most of the EJPs failed to trigger a muscle action potential, but seven of the EJPs were followed by a muscle action potential, with a variable latency after the stimulus. The bottom trace of Figure 10 shows the same preparation in the presence of $10^{-5}$ M 5-HT. Note that all of the EJPs triggered spikes, most of them immediately on the rising phase of the EJP, but others at a longer latency. Also note that the triggered action potentials were wider than in control, and that several stimuli elicited two spikes. The spike width measured at 20 mV went from an average of 20 ± 6 ms in control to 40 ± 4 ms in serotonin ($p = .022$). Similar data were obtained in three preparations.

DISCUSSION

In the stomatogastric nervous system of the lobster, *H. americanus*, serotonin is found in the peripherally located GPR neurons that project into the neuropil of the STG and from there to more anterior ganglia (Beltz et al., 1984; Katz et al., 1989; Katz and Harris-Warrick, 1989). In a previous study we found that serotonin does not first appear in the stomatogastric nervous system until about LII, the second larval stage of development (Kilman et al., 1999). The late appearance of serotonin in the stomatogastric nervous system until about LII, the second larval stage of development (Kilman et al., 1999). The late appearance of serotonin in the stomatogastric nervous system suggested that it might play an important developmental role in the maturation of the circuitry and motor patterns of the STG. This may, in fact, still be true, but the data in this article suggest that hormonally released serotonin could be acting physiolog-
ically much earlier in development, both on the STG itself, and on the stomach neuromuscular junctions.

The POs are a major source of neurohormones for crustaceans (Cooke and Sullivan, 1982; Keller, 1992). The POs acquire a large number of their neuromodulators, including serotonin (Fig. 5), early in embryonic development (Pulver and Marder, 2002). Hormones released from the POs will be rapidly delivered both to the STG itself and to the stomach musculature, which is bathed in hemolymph. Interestingly, in this article we demonstrate that the STG itself, as well as one of the neuromuscular junctions, might be early sites of action of hormonally delivered serotonin.

**What Role Does Serotonin Play in the Embryonic STG?**

We have shown that a serotonin transporter is present in embryos and LI animals (Figs. 2 and 3), and that the nuclei as well as cytoplasm of the GPR cells (Fig. 3) apparently become immunoreactive following serotonin uptake. This raises the possibility that serotonin acts as a borrowed transmitter and/or as an intracellular regulator at these early stages. The concept of serotonin acting as a borrowed transmitter suggests that the GPR cells take up and concentrate serotonin, possibly to release it at a later time. In this way hormonally liberated serotonin could temporarily alter the cotransmitter phenotype of the embryonic GPR neurons. Alternatively, or in addition to this function, serotonin may act intracellularly as a growth or transcriptional regulator within the GPR cells. The apparent staining of the GPR nuclei [Fig. 3(B,C)] suggests that after uptake serotonin may, indeed, be transported retrogradely to the nucleus.

There are precedents for these nontraditional actions of serotonin in both invertebrates and vertebrates. In the mouse brain, the serotonin transporter is expressed widely at embryonic stages before synapses have formed (Bruning et al., 1997). Thalamocortical relay neurons do not synthesize serotonin, but become immunoreactive for it as a result of taking up extracellular serotonin, probably released by Raphe neurons (Lebrand et al., 1996). The thalamocortical neurons also express the vesicular monoamine transporter, and are responsible for a transient serotonergic innervation of cortex thought to be developmentally important. Thus, it is unclear whether these relay neurons use serotonin as a growth regulatory molecule, as a transient borrowed transmitter, or both. Likewise, in the lobster brain, newborn olfactory interneurons show a transient uptake of serotonin within a few hours of their birth (Beltz et al., 2001). These neurons never show serotonin immunoreactiv-
EJPs to the range of the embryonic EJPs, these larger EJPs did not evoke action potentials, nor did depolarization with intracellular microelectrodes. Obviously, it is possible that the adult muscle fibers may still be able to fire action potentials under some modulatory conditions, but it appears that they are largely without this function in control saline or serotonin.

In other crustacean preparations, muscle action potentials are Ca2+ mediated. In many developing systems, early Ca2+ spikes are thought to be important for setting the stage for the development of excitability and synaptic structure (Spitzer, 1994; Moody, 1998a,b; Spitzer and Ribera, 1998). Similarly, here it is possible that when the EJPs trigger a spike, the increase in Ca2+ concentration is used to stabilize developing neuromuscular junctions.

As at other crustacean neuromuscular junction systems (Dudel, 1965; Glusman and Kravitz, 1982; Dixon and Atwood, 1985; Delaney et al., 1991), the serotonin-mediated enhancement of transmitter release at the adult LP-p1 neuromuscular synapse is seen as an increase in EJP amplitude and a corresponding decrease in the variability of the EJPs. The decrease in the CV of the EJPs is consistent with a presynaptic action of serotonin, and this is consistent with the finding that serotonin has no effect on the amplitude of the response to iontophoretically applied glutamate (data not shown). Because serotonin is known to activate multiple receptors and multiple signal transduction pathways in crustacean nerve muscle preparations (Dixon and Atwood, 1989a,b,c; Goy and Kravitz, 1989), it is possible that there is a developmental switch in the extent to which these multiple serotonin-activated pathways are expressed. Thus, our data are consistent with a predominantly postsynaptic action of serotonin on muscle excitability in the young animals, and a predominantly presynaptic action of serotonin on transmitter release in the adult.

CONCLUSIONS

Serotonin has physiological actions in both embryonic and adult stomatogastric preparations, but there appears to be a complex suite of developmental changes in serotonin’s actions over development. All neurally released serotonin in the adult is thought to derive from the GPR terminals (Katz et al., 1989; Katz and Harris-Warrick, 1989). The GPR neurons are stretch receptors and contain several cotransmitters, which in H. americanus include acetylcholine, serotonin, FLRFamide-like peptides, and allatostatin-like peptides (Kilman et al., 1999). In the crab, C. borealis, the GPR neurons produce both rapid cholinergic actions and slower modulatory actions that have been attributed to serotonin (Katz et al., 1989; Katz and Harris-Warrick, 1989, 1990a,b, 1991). We know that the GPR neurons are present early in development because they stain with their other cotransmitters (Kilman et al., 1999) and take-up serotonin. Nonetheless, if they are active early in development this means that they will function with their other cotransmitters, but not with serotonin unless hormonally released serotonin loads the GPR neurons sufficiently to allow them to release serotonin. This suggests that the cotransmitter complement of the GPR neurons early in development may be hormonally regulated, or hormonally gated, via the ability of the GPR neurons to take up serotonin. At present, we do not know the physiological circumstances under which the embryonic POs are likely to release serotonin, so we have no way of knowing whether the GPR neurons are likely to load and possibly release serotonin often or rarely.

Serotonin is active on the LP to p1 neuromuscular junction in both embryos and adults, but these actions change over development, as do the properties of the neuromuscular junctions themselves. Early in development the neurally evoked EJPs are large and variable, and serotonin appears to increase the likelihood that muscle action potentials or prolonged EJPs will be seen. In adults, the neurally evoked EJPs are smaller and more constant, and the effect of serotonin appears to predominantly regulate the amplitude of the EJP, but not muscle fiber membrane excitability. Together, all of the data in this article demonstrate that serotonin elicits a complex suite of actions during development and adult life. The dual roles of serotonin as a rapid modulator and a developmental regulator are inextricably intertwined, as together these mechanisms shape the motor patterns produced throughout the lifetime of the animal.

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