Serotonin Enhances the Resistance Reflex of the Locomotor Network of the Crayfish through Multiple Modulatory Effects that Act Cooperatively

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Serotonin (5HT) is an endogenous amine that modifies posture in crustacea. Here, we examined the mechanisms of action of 5HT on the resistance reflex in crayfish legs. This reflex, which counteracts movements imposed on a limb, is based on a negative feedback system formed by proprioceptors that sense joint angle movements and activate opposing motoneurons. We performed intracellular recordings from depressor motoneurons while repetitively stretching and releasing a leg joint proprioceptor in a resting in vitro preparation (i.e., a preparation that lacks spontaneous rhythmic activity). 5HT increased the amplitude of the depolarization during the release phase of the proprioceptor (corresponding to an upward movement of the leg) and the discharge frequency of the motoneurons. The 5HT-induced increase in the resistance reflex is caused, to a large extent, by polysynaptic pathways because it was very attenuated in the presence of high divalent cation solution. In addition to this activation of the polysynaptic pathways, 5HT also has postsynaptic effects that enhance the resistance reflex. 5HT causes a tonic depolarization, as well as an increase in the time constant and input resistance of motoneurons. We developed a simple mathematical model to describe the integrative properties of the motoneurons. The conclusion of this study is that the input frequency and the decay time constant of the EPSPs interact in such a way that small simultaneous changes in these parameters can cause a large effect on summation. Therefore, the conjuncture of presynaptic and postsynaptic changes produces a strong cooperative effect on the resistance reflex response.

Key words: crustacea; stretch reflex; neuromodulation; posture; serotonin; crayfish

Behavioral/Systems/Cognitive

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Introduction

The operation of neural networks controlling behaviors is based on both membrane properties and connectivity of the component neurons. These two levels of a network have been shown to be controlled by neuromodulators in vertebrates and invertebrates (Hultborn and Kiehn, 1992; Katz, 1998; Sillar et al., 1998; Cantrell and Catterall, 2001; Nusbaum et al., 2001; Marder and Thirumalai, 2002). In crustacea, serotonin (5HT) was shown to modify synaptic transmission (Fischer and Florey, 1983; Delaney et al., 1991; Pearlstein et al., 1998; Wang and Zucker, 1998; Southard et al., 2000) and endogenous properties of neurons (Rossi-Durand, 1993; Alvarado-Alvarez et al., 2000; Nagayama, 2002). At the behavioral level 5HT can influence posture and walking (Teshiba et al., 2001; Tierney and Mangiamele, 2001), aggressiveness (Huber et al., 1997a,b), and social status in crayfish (Huber et al., 1997b, 2001). For example, when injected into freely moving lobsters, 5HT and octopamine generate stable and stereotypical postures that resemble those seen in dominant (5HT) and subordinate (octopamine) lobsters (Livingstone et al., 1980; Kravitz, 1988). The analysis of the mechanisms of 5HT-induced neuromodulation is complex because these effects are themselves modulated by the pattern of exposure of the cell to serotonin and the social status of the animal (Edwards et al., 2003).

In this paper, we have studied the effect of 5HT on postural control in the crayfish Procambarus clarkii. The control of posture involves proprioceptive feedback. In crayfish, the resistance reflex consists of the monosynaptic and polysynaptic excitation of the motoneurons (MNs) that oppose any imposed joint movement (El Manira et al., 1991a), and of the polysynaptic inhibition of the antagonistic MNs (Le Bon-Jego and Cattaert, 2002).

In the in vitro preparation of the crayfish walking system (El Manira et al., 1991a), the perfusion of the CNS in a resting preparation (i.e., one that lacks rhythmical activity) with 10 μM 5HT induces a dramatic increase in the amplitude of the resistance reflex of MNs in most cases. Here, we demonstrate that polysynaptic excitatory pathways of the resistance reflex are activated by 5HT, and we show that 5HT modifies the intrinsic properties of

Received June 20, 2003; revised Oct. 21, 2003; accepted Oct. 21, 2003.

This work was supported by The Centre National de la Recherche Scientifique and by a Groupement d’Intérêt Scientifique (sciences de la cognition) contract (CIAM 10-2). M.L.B.-J. received a fellowship from the Ministère de l’Enseignement Supérieur et de la Recherche. We thank Drs. F. Clarac and P. Meyrand for valuable discussions during the course of this work. We are also grateful to Fahi Issa and Andrew Hill for comments and corrections on this manuscript and for improving the English.

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DOI:10.1523/JNEUROSCI.4032-03.2004

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MNs. 5HT causes a depolarization of the resting potential of motoneurons and increases their input resistance and time constant. These multiple effects induced by serotonin act cooperatively to enhance the resistance reflex.

Materials and Methods

Experimental animals
Experiments were performed on male and female crayfish (Procambarus clarkii; n = 48) weighing 25–30 gm. The animals were purchased from a commercial supplier (Château Garreau, La Bastide d’Armagnac, France), maintained in indoor aquaria containing 80–100 animals at 15–18°C, and fed once a week with mussels.

In vitro preparation. An in vitro preparation of the thoracic nervous system was used (Sillar and Skorupski, 1986; El Manira et al., 1991a). It consists of the last three thoracic (T3–T5) and the first abdominal (A1) ganglia dissected out with all the nerves of the two proximal segments of the left fifth leg (Fig. 1A). The chordotonal organ (CBCO), which monitors the movements of the second joint (coxoharaposipodite), was also dissected out and kept intact, and the distal end of its elastic strand was attached to an electromagnetic puller VT101 (Ling Dynamic Systems, Meudon-la-Forêt, France) controlled by a homemade function generator allowing the application of sinewave movements to the CBCO strand that mimic upward (during stretch) and downward (during release) movements of the leg. The preparation was pinned dorsal side up on a Sylgard-lined Petri dish (Dow Corning, Wiesbaden, Germany). The nervous system was superfused continuously with oxygenated control saline composed of (in mm): 195 NaCl, 5 KCl, 13 CaCl2, 2 MgCl2 and 3 HEPES (Sigma, St Louis, MO) with a pH of 7.65. The fourth and fifth ganglia were desheathed to improve the superfusion of the central neurons and to allow for intracellular recordings (Fig. 1). The CNS was insulated from the CBCO by a Vaseline wall (Fig. 1A) to restrict the superfusion of 5HT (10 μM) to the ganglia. At the same time the CBCO was superfused with control saline. In some experiments, a high divalent cation solution was used to raise the spiking threshold of the interneurons containing (in mm): 34 CaCl2, 6.4 MgCl2 with sodium concentration reduced accordingly to preserve the osmolarity of the solution (157 NaCl).

Recordings. Extracellular recordings from the motor nerves innervating the depressor (Dep) and levator (Lev) muscles from the sensory nerve of the CBCO were made using stainless steel pin electrodes contacting the nerves and insulated with Vaseline. Intracellular recordings from depressor MNs (Fig. 1B, Dep MN) were performed with glass micropipettes (Clark Electromedical Instruments, Reading, UK) filled with 3 M KCl (resistance 10–20 MΩ) connected to an Axoclamp 2B amplifier (Axon Instruments, Foster City, CA) used in the current-clamp mode. In some experiments, the injection of fluorescent dye (5% dextran rhodamine in 0.2M potassium acetate) was used to stain and analyze the geometry of the recorded MN after the experiment (Fig. 1B). In crustacean MNs, the soma do not participate in the electrical activity of the neuron. MN soma lie outside of the neuropile (the region in which neurons form their synaptic contacts) (Fig. 1B) and are linked to the arbor of the neuron by a thin neurite. For these reasons, intracellular recordings were made from the main neurite, where EPSPs could be recorded (Fig. 1B). The resting membrane potential of MNs was usually in the range of −78 to −65 mV. Stability of resting membrane potential over long period of time (>4 hr) was used as a criterion for evaluation of cell health in recordings. In crustacean MNs, the soma and neurites do not actively convey spikes. Therefore, spike amplitude was generally small (<20 mV) at the recording site. Data were digitized and stored onto a computer hard disk through an appropriate interface (1401plus) and software (Spike2) from Cambridge Electronic Design Ltd. (Cambridge, UK).

Data analysis. Data were analyzed using the Spike2 analysis software. Spikes recorded from the CBCO nerve were discriminated according to their waveform on the basis of a template matching protocol (wavenmark). Templates were built automatically and corresponded to the mean duration of sensory spikes (1.5 msec in duration). The sampling rate for CBCO nerve recording was set to 15 kHz, which resulted in templates containing 20–22 points. The procedure used two criteria to identify a spike: (1) >90% of the points should be in the confidence limits of the template; (2) the maximum amplitude change for a match was <5%. This procedure was applied off-line. After the completion of this protocol, each identified CBCO unit (spike shape) was assigned an arbitrary number. Subsequently, a spike-triggered average was performed for each CBCO unit, allowing us to observe in a given MN the occurrence of any postsynaptic events related to this unit. Statistical analyses were done with the Prism program (Graphpad Software, San Diego, CA). The results are given as mean values ± SEM.

Results

5HT enhances resistance reflex
Application of stretch and release movements to the CBCO strand in the in vitro preparation resulted in reflex activation of MNs (El Manira et al., 1991a,b; Le Ray et al., 1997a,b). During sinewave movements applied to the CBCO strand, Dep and Lev
MNMs were cyclically activated: Dep MNMs during release (mimicking upward movement of the leg in intact animals) and Lev MNMs during stretch (mimicking downward movements of the leg in intact animals). These responses corresponded, therefore, to resistance reflex (Fig. 2). All the experiments reported here (intact animals). These responses corresponded, therefore, to resistance reflex during release (mimicking upward movement in the intact animals), Lev MNMs were cyclically activated: Dep MNs during stretch (mimicking upward movements of the leg in intact animals) and Lev MNs were cyclically activated: Dep MNs during stretch (mimicking upward movements of the leg in intact animals) and Lev MNs were cyclically activated: Dep MNs during release (mimicking downward movements of the leg in intact animals) and Lev MNs were cyclically activated: Dep MNs during release (mimicking upward movement in the intact animals), the Dep MNMs were activated. The mean frequency of the discharge of the motor activity recorded is presented above each recording (Dep n Mean Freq, Lev n Mean Freq). The mean frequency was calculated by a floating average with a time window of 100 msec. When 5HT was superfused on the ganglia, the resistance reflex response of Dep and Lev MNs was increased.

Figure 2. Effect of 5HT on the resistance reflex in a resting preparation. Sinusoidal movements imposed on the CBCO strand induced resistance reflex responses recorded extracellularly from the levator nerve (Lev n) and the depressor nerve (Dep n). When the CBCO strand was released (R; mimicking upward movement in the intact animals), the Dep MNMs were activated. In contrast, during stretch movements (S), the Lev MNs were activated. The mean frequency of the discharge of the motor activity recorded is presented above each recording (Dep n Mean Freq, Lev n Mean Freq). The mean frequency was calculated by a floating average with a time window of 100 msec. When 5HT was superfused on the ganglia, the resistance reflex response of Dep and Lev MNs was increased.

Figure 3. Effect of 5HT on intracellularly recorded Dep MNs. A, After application of 5HT (10 \(\mu M\)), an intracellularly recorded Dep MN displays a slight tonic depolarization. During this recording, sinusoidal movements were applied continuously to the CBCO strand and evoked a resistance reflex in the Dep MN (see enlarged view in B). The amplitude of this reflex response was increased after 25 min in the presence of 5HT. B, Enlarged view of recordings presented in A. The numbers 1 and 2 refer to the control situation (A, control; \(t = 0\)) and to 25 min after onset of 5HT perfusion (4, 5HT; \(t = 25\) min). This increase in the amplitude of the resistance reflex response was not caused by any change in the sensory activity from the CBCO nerve as demonstrated by the mean frequency of discharge of the sensory nerve of the CBCO (CBCO n Mean Freq). The resting potential in control situation is indicated by a dashed line (48 mV). This tonic depolarization was in-
confirmation of the absence of leakage, and (2) the mean frequency of the global sensory discharge recorded from the CBCO sensory nerve was monitored to ensure that it did not change during an experiment (Fig. 3B).

The enhancement of the MN resistance reflex by 5HT is long lasting
The two effects produced by 5HT application on Dep MN (slight tonic depolarization and an increase in the amplitude of the resistance reflex response) occurred with different time courses. In the presence of 5HT, although the small tonic depolarization started after ~1 min (Fig. 3A), the increase in the amplitude of the reflex response was not observed before 5–10 min in the presence of 5HT (Fig. 4A) at a time when the tonic depolarization was maximum. In Figure 4A, the intracellular responses to sinewave movements imposed on the CBCO strand were averaged by groups of 10 cycles (Fig. 4A,B), and the error bars represent the SEM. In the control situation, the amplitude of the reflex response was 1.94 ± 0.32 mV (n = 10 cycles). The amplitude of the response increased to 3.95 ± 0.33 mV (n = 10 cycles) after 10 min in the presence of 5HT, and the maximum response was 4.88 ± 0.32 mV (n = 10 cycles) after 25 min in the presence of 5HT. Therefore, 5HT application caused the response to CBCO movement to increase by 151%. This effect persisted for >1 hr (Fig. 4A,B). When an effect of 5HT on resistance reflex was observed (n = 40; see below), its time course was very consistent and more or less similar to the experiment presented in Figure 4A, regardless of the 5HT application protocol used (long application >1 hr to short 10 min applications). Note that in this series of experiments, the rate of the perfusion remained unchanged (1 ml/min). In all cases, the reflex response increased progressively after a delay of 5–10 min, and reached a plateau 25 min after the beginning of 5HT application, and persisted for >1 hr after washing. Therefore, in all subsequent experiments, we applied 5HT for 10 min and measured its effects during 1 hr (as in Fig. 4). Subsequently, in all of the figures and in all the following text, the expression “after 5HT application” will mean that the presented effect of 5HT was measured after 25 min in the presence of 5HT, at a time during which the effect was maximum.

In the experimental conditions used, generally all Dep MNs were silent in the Dep motor nerve. In this paper, only Dep MNs with naturally hyperpolarized resting membrane potentials are presented (Figs. 4, 6A). The spike threshold (~55 to ~50 mV) was never reached in these MNs in response to imposed CBCO movements even after 5HT application. This allowed us (1) to measure continuously the change in response to movement and the input resistance of MNs without contamination by sodium and potassium conductances supporting spikes and (2) to analyze the changes in EPSP amplitude (see below). Dep MNs that responded to CBCO-imposed movements by bursts of spikes in the experimental conditions were generally rare (fewer than two per preparation).

Diversity of MN response to 5HT
Although 5HT increased the resistance reflex response in most experiments, in 16.6% of the 48 experiments, 5HT did not produce any effect on the recorded MN. The absence of effect was not correlated with the amplitude of the control reflex response: among the experiments in which 5HT had no effect on the intracellularly recorded MNs, we found MNs with either small or large reflex responses in control situations. Figure 5 presents two experiments. In the control situation, a sinewave movement imposed on the CBCO strand elicited a cyclic resistance reflex response in the two Dep MNs (Fig. 5A,B, left). The mean amplitude of the response was 1.81 ± 0.04 mV (n = 16 cycles) in Dep MN1 and 0.87 ± 0.04 mV (n = 11 cycles) in Dep MN2. At t = 0, the preparation was bathed in 10 μM 5HT. At t = 10 min, the preparation was washed in normal Ringer’s solution. The
effect of 5HT on Dep MN1 was maximum at \( t = 25 \) min: its response increased two-fold (2.59 ± 0.06 mV; \( n = 16 \)) (Fig. 5A, right). By contrast, in the other experiment, Dep MN2 did not show any significant change (\( t \) test, \( p = 0.39 \)) in its response (0.82 ± 0.05 mV; \( n = 11 \)) (Fig. 5B, right). Such a discrepancy between recorded MNs could indicate either a difference between MNs or a difference between the preparations.

To clarify this point, we performed simultaneous intracellular recordings from up to four Dep MNs in a given experiment. In Figure 6, four Dep MNs were intracellularly recorded simultaneously. All of them displayed a cyclic resistance reflex response during sinewave movement applied to the CBCO strand. As in Figure 3, after 1 min in the presence of 5HT, these four Dep MNs showed a small tonic depolarization (Fig. 6A), and the amplitude of the movement response increased to reach a maximum after 25 min (Fig. 6A, right). Superimposition of control and 5HT (25 min) responses indicates that the four MNs displayed an increased response under 5HT condition; however, the magnitude of the effects was different (8–98%) among the different recorded MNs (Fig. 6A, right). In addition to the intracellularly recorded Dep MNs, other Dep MNs showed an increase of their reflex response as monitored by the progressive increase of their reflex discharge recorded from the depressor nerve. During the experiment, application of 5HT did not affect the activity of CBCO sensory neurons as attested by the enlarged view of the CBCO nerve recording (Fig. 6B). Similar results were obtained in five other experiments.

By contrast, in other experiments, 5HT did not induce any changes in all of the simultaneous recordings from four Dep MNs (Fig. 7). It is noteworthy that in such experiments (\( n = 4 \)), neither the small tonic depolarization nor the slower increase in the movement response was observed.

Remarkably, in no experiment in which four simultaneous intracellular recordings from Dep MNs involved in resistance reflex were performed was a combination of 5HT-sensitive and 5HT-insensitive Dep MNs observed. Therefore, it seems that the preparations could be classified into two categories: the ones in which 5HT induced a general increase in the reflex response in all Dep MNs and the ones in which 5HT did not induce any change in any of the Dep MNs. Hereafter, we have focused on experiments in which 5HT induced an increase in the resistance reflex response.

### 5HT increases input resistance in Dep MNs

We have tested the idea that the increase in the response could be caused by a direct effect of 5HT on the MNs. Therefore, during subsequent experiments, we made continuous measurements of the input resistance of intracellularly recorded Dep MNs. Two examples of such experiments are shown in Figure 8. In one of the experiments (Fig. 8A, C, Dep MN1), the resistance reflex response was 1.25 ± 0.26 mV in the control situation and reached 1.65 ± 0.34 mV after 5HT application. In the same Dep MN1, the input resistance that was 2.05 ± 0.15 Ω in control situation increased to 2.46 ± 0.22 Ω after 5HT application. In another experiment, the amplitude of the response of a Dep MN (Fig. 8B, Dep MN2) was 1.96 ± 0.15 mV in the control situation and reached 4.12 ± 0.34 mV after 5HT application. Accordingly, the input resistance of Dep MN2 increased from 2.16 ± 0.12 Ω in the control situation to 5.83 ± 0.16 Ω after 5HT application (Fig. 8D).

The increase in input resistance was generally in accordance with the increase in the amplitude of the resistance reflex response. For example a moderate increase in movement response (32% in the case of Dep MN1) corresponded with a moderate increase (20%) in input resistance. Similarly, a large increase in movement response (110% in the case of Dep MN2) corresponded with a large increase in input resistance (169%). These
observations were consistent over the eight Dep MNs tested in this series of experiments.

This result was confirmed further by measuring the repolarization time constant in control and 5HT conditions (Fig. 8E,F). In each of the eight tested Dep MNs, the time constant was significantly increased after 5HT application. This increase was highly correlated with the increase in the movement response (Fig. 8G). Similarly, the increase in input resistance was also correlated with the increase in movement response (Fig. 8G). Therefore, it was very likely that the increase in the movement response was directly related to the increase in input resistance of the recorded Dep MN. To confirm this hypothesis, however, we analyzed the effect of 5HT on the amplitude of unitary EPSPs evoked by identified CBCO sensory neurons.

Effect of 5HT on unitary EPSPs from CBCO sensory neurons
The extracellular recording of the CBCO nerve was analyzed by Spike2 software to separate the different sensory units according to their shape. Then each spike shape was used to perform spike-triggered averaging of the intracellular recording of the Dep MN. In Figure 9, only the spikes that were associated with an EPSP in the Dep MN are shown. Each of the spike shapes was monitored continuously during an experiment. In this experiment, after 5HT application, the movement response increased from 0.78 ± 0.08 to 1.52 ± 0.09 mV (95.3% increase) (Fig. 9A). At the same time, eight unitary EPSPs triggered by identified sensory units were analyzed (Fig. 9B,C). Each unitary EPSP is shown in control (gray trace) and 5HT (black trace) conditions. With each unitary EPSP, the shape of the corresponding sensory spike (Unit) is presented, together with the histogram of its firing activity during the sinusoidal movement imposed on the CBCO strand. As can be seen, these sensory units fired during release movement of the CBCO strand (i.e., upward movement of the leg in intact animals), and because they evoked an EPSP in a Dep MN, they therefore were involved in a resistance reflex. Note that some sensory units were also slightly activated during the stretch movement imposed on the CBCO strand (corresponding to downward movement of the leg). During the downward movement, Dep MNs receive IPSPs via a polysynaptic pathway (Le Bon-Jego and Cattaert, 2002). These IPSPs reduce the amplitude of the unitary EPSPs. Therefore the analysis of unitary EPSP amplitude was performed exclusively during the upward movements. The most remarkable result of this analysis was that the amplitude of unitary EPSPs was only slightly affected if at all in the presence of 5HT (Table 1). For six of eight resistance reflex units, the unitary EPSP was either unchanged (units 13, 15 and 21; t test; p > 0.2) or even slightly but significantly reduced (units 12, 17 and 27; t test; p < 0.0001, p = 0.0008, p = 0.026, respectively) in the presence of 5HT (Fig. 9B, Table 1). This result was very surprising because in the presence of 5HT the input resistance was increased almost fourfold, leading us to expect a proportional increase in the amplitude of individual EPSPs. Indeed, only two resistance reflex units (the two with the largest spike amplitudes) presented a net increase of EPSP amplitude under 5HT application (Fig. 9C).

These two EPSPs were characterized by complex shapes: an initial rapid EPSP followed by a second slower depolarizing wave. The detailed analysis of one of these EPSPs (triggered by Unit28) is presented in Figure 9D. In the control condition, when several EPSPs triggered by this unit were superimposed, only the first depolarizing event was reliably present (Fig. 9D, Control) and therefore was likely attributable to a monosynaptic EPSP. This hypothesis is supported by the observation that the delay between the extracellularly recorded sensory spike and the onset of this first EPSP was constant. The second wave in the averaged EPSP was caused by polysynaptic events, which are characterized by nonconstant delays (Fig. 9D). On the averaged traces (Fig. 9C, Unit28), the rapid likely monosynaptic EPSP was increased from 0.832 ± 0.022 mV (n = 70) in control to 0.991 ± 0.051 mV (n = 62) in the 5HT condition. This increase was significant (t test; p = 0.039) (Table 1). In addition, the amplitude of the late compound EPSP (measured 6 msec after the peak of the monosynaptic EPSP) (Fig. 9C, arrow) was significantly increased from 0.428 ± 0.075 mV in control to 0.980 ± 0.130 mV in the 5HT condition (t test; p = 0.0002). This increase was a result of the recruitment of large and late polysynaptic EPSPs after 5HT application (Fig. 9D, 5HT). Similar results were obtained in 12 preparations (Table 2).

To verify that EPSP amplitude did not depend on the phase of
the sinewave movement imposed on the CBCO strand, which could mask some of the effects of 5HT on unitary EPSPs, in some experiments we have systematically averaged the amplitude of unitary EPSPs with respect to the phase of the imposed movement. The result of such an analysis is shown in Figure 10. In the example presented, the global resistance reflex response of the Dep MN was increased by 94.8% in the 5HT condition. One unitary EPSP is presented (Fig. 10A). The amplitudes of the averaged EPSPs of this unit in control (0.590 ± 0.016 mV; n = 526) and 5HT (0.580 ± 0.017 mV; n = 489) conditions were not significantly different (t test; p = 0.663) (Fig. 10A). Moreover, the plot of unitary EPSP amplitude against phase of the movement cycle demonstrates stationarity of EPSPs along movement cycle
The increase of resistance reflex response involves polysynaptic pathways

To estimate the role of polysynaptic pathways in the 5HT-induced increase of the resistance reflex response, we performed a series of experiments in which the CNS was bathed successively in normal saline and in a high divalent cation solution, both in control and in 5HT conditions. The response of an intracellularly recorded Dep MN to CBCC-imposed sinewave movement is presented in Figure 11A. In high divalent cation solution (Fig. 11A2), the amplitude of the control response (Fig. 11A1) was reduced from 0.96 ± 0.04 to 0.67 ± 0.02 mV, indicating that even in the absence of exogenous 5HT, polysynaptic pathways are involved in the global resistance reflex response. When back in control saline (Fig. 11A3), the amplitude of the resistance reflex response was not totally restored to its initial value (0.78 ± 0.02 mV). Application of 5HT (10 μM in normal saline) induced a large increase of

Table 1. Statistical analysis of amplitude of unitary EPSPs presented in Figure 9

<table>
<thead>
<tr>
<th>CBCO unit</th>
<th>Control unitary EPSP mean amplitude ± SEM</th>
<th>5HT unitary EPSP mean amplitude ± SEM</th>
<th>t test p value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Unit12</td>
<td>0.657 ± 0.0013</td>
<td>0.564 ± 0.019</td>
<td>p &lt; 0.001****</td>
</tr>
<tr>
<td>n = 567</td>
<td></td>
<td>n = 483</td>
<td></td>
</tr>
<tr>
<td>Unit13</td>
<td>0.153 ± 0.015</td>
<td>0.128 ± 0.015</td>
<td>p = 0.244</td>
</tr>
<tr>
<td>n = 737</td>
<td></td>
<td>n = 856</td>
<td>NS</td>
</tr>
<tr>
<td>Unit15</td>
<td>0.137 ± 0.016</td>
<td>0.115 ± 0.017</td>
<td>p = 0.335</td>
</tr>
<tr>
<td>n = 496</td>
<td></td>
<td>n = 424</td>
<td>NS</td>
</tr>
<tr>
<td>Unit16</td>
<td>0.759 ± 0.020</td>
<td>0.788 ± 0.021</td>
<td>p = 0.332</td>
</tr>
<tr>
<td>n = 240</td>
<td></td>
<td>n = 271</td>
<td>NS</td>
</tr>
<tr>
<td>Unit17</td>
<td>0.478 ± 0.010</td>
<td>0.419 ± 0.014</td>
<td>p = 0.008***</td>
</tr>
<tr>
<td>n = 575</td>
<td></td>
<td>n = 606</td>
<td></td>
</tr>
<tr>
<td>Unit21</td>
<td>0.368 ± 0.012</td>
<td>0.348 ± 0.014</td>
<td>p = 0.301</td>
</tr>
<tr>
<td>n = 737</td>
<td></td>
<td>n = 677</td>
<td>NS</td>
</tr>
<tr>
<td>Unit27</td>
<td>0.496 ± 0.012</td>
<td>0.453 ± 0.015</td>
<td>p = 0.026*</td>
</tr>
<tr>
<td>n = 607</td>
<td></td>
<td>n = 607</td>
<td></td>
</tr>
<tr>
<td>Unit28</td>
<td>0.832 ± 0.022</td>
<td>0.991 ± 0.051</td>
<td>p = 0.0039**</td>
</tr>
<tr>
<td>n = 70</td>
<td></td>
<td>n = 62</td>
<td></td>
</tr>
</tbody>
</table>

*p < 0.05; **p < 0.01; ***p < 0.001; ****p < 0.0001.

For each experiment, the number of EPSPs presenting a type of response to 5HT (no effect, increase, decrease) is given. We have distinguished the early EPSPs (occurred during the release movement of the CBCO: the EPSPs counted in filled bars) and late compound EPSPs (occurred during the movement of the leg in the intact animal) and therefore occurred in conjunction with IPSPs of the inhibitory component of the resistance reflex (Le Bon-Jego and Cattaert, 2002). Their amplitude therefore was generally smaller, and they were not included in the analysis. Similar results were obtained with all of the identified unitary EPSPs analyzed (n = 37).

Table 2. Effect of 5HT on the amplitude of unitary EPSPs in 12 experiments

<table>
<thead>
<tr>
<th>Experiment</th>
<th>Number of CBCO units evoking an EPSP in the MN</th>
<th>Effect of 5HT on early EPSP</th>
<th>Effect of 5HT on late EPSP</th>
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<td>No change</td>
<td>Increase</td>
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For each experiment, the number of EPSPs presenting a type of response to 5HT (no effect, increase, decrease) is given. We have distinguished the early EPSPs from the late compound EPSPs. The significance of effects was assessed by t test as in Table 1, with p < 0.05.

*Only a fraction of the analyzed EPSPs presented a clearly visible late compound EPSP.
A noticeable increase of peak amplitude or recovery time constant, this was a result of polysynaptic events as in Figure 11C.

5HT and EPSP amplitude
In the presence of 5HT the input resistance of the MNs was generally increased by a factor 1.2–2.9 (Fig. 8G). However, during the application of movements to the CBCO strand, many EPSPs did not increase their amplitude in the presence of 5HT (Fig. 9, Table 1). This intriguing result could be caused by three phenomena. (1) In the presence of 5HT, the membrane potential of the MN is more depolarized than in the control condition (Figs. 3B, 4, 6). Therefore, the electrotonic driving force supporting the EPSPs is smaller in 5HT condition, and the amplitude of EPSPs is expected to become smaller. (2) Because of massive synaptic inputs, the input resistance may be locally reduced (shunting excitation), causing the decrease of EPSP amplitude. (3) During a volley of EPSPs, each EPSP summates with the falling phase of preceding EPSPs, resulting in a smaller EPSP amplitude as frequency increases. We have analyzed these hypotheses. A simulation study demonstrated that the small change in driving force caused by the slightly more depolarized membrane potential in the 5HT condition did not contribute significantly to the change in EPSP amplitude observed (data not shown). Similarly, the effect of shunting excitation was tested in simulation studies, and it was concluded that it did not contribute significantly to the change in...
EPSP amplitude observed (data not shown). The decrease in EPSP amplitude was caused mainly by an increase of EPSP frequency in the volley in the presence of 5HT. Indeed, during a volley, the amplitude of each EPSP summates with the falling phases of preceding EPSPs. This phenomenon is illustrated in Figure 13A, in which the amplitude \( V_1 \) of the second EPSP is smaller than the amplitude \( V_0 \) of the first EPSP, because the second EPSP summates with the falling phase of the first EPSP. The amplitude of subsequent EPSPs is decreased further because of accumulation of falling phases (Fig. 13B). Because of this phenomenon, increasing the frequency of the volley results in a further decrease in the amplitude of EPSPs.

This hypothesis was tested in EPSPs intracellularly recorded from Dep MNs. When amplitude measurements were performed on an isolated identified unitary EPSP in the absence of movement stimulation (in this condition most of the sensory neurons were silent or, when active, their discharge frequency was less than a few hertz), the amplitude of such an “isolated” EPSP was always increased in preparation in response to 5HT application (Fig. 12). In the experiment reported here, EPSPs from a sensory neuron have been recorded in four conditions: in control situation and in the presence of 5HT both during and in the absence of sinewave movement imposed on the CBCO. In the control situation, in the absence of movement, this sensory neuron produced a volley of EPSPs (Fig. 12A1, Isolated EPSPs). During a series of sinewave movements imposed on the CBCO, this same sensory neuron presented a burst of sensory spikes that evoked a volley of EPSPs in the Dep MN (data not shown). These volley EPSPs are shown in Figure 12A2. Isolated EPSPs were significantly larger \((t \text{ test}; p < 0.0001)\) in the 5HT condition \((0.458 \pm 0.013 \text{ mV})\) than in control \((0.372 \pm 0.011 \text{ mV})\) (Fig. 12B1,C1), and the likely monosynaptic part of the volley EPSPs was significantly smaller \((t \text{ test}; p = 0.0176)\) in the 5HT condition \((0.311 \pm 0.016 \text{ mV})\) than in control \((0.365 \pm 0.015 \text{ mV})\) (Fig. 12B2,C2). Such measurements were not possible on most of the CBCO units because they were activated only during movement application. Only 12 units projecting on Dep MNs and presenting a slow rate of tonic activity in the absence of sinewave movement were analyzed. For all of these units, isolated EPSPs were significantly larger in the 5HT condition than in the control condition. For six of these units the volley EPSP was significantly smaller in 5HT condition than in control. For four of these units there was no significant difference in volley EPSP between control and 5HT conditions. Two units displayed larger volley EPSPs in 5HT condition.

**Integrative properties of MNs**

The integrative properties of a neuron and its input–output relationship depend on several parameters such as input resistance, EPSP frequency, time constant, and membrane potential. In all of the experiments in which 5HT induced an increase in the resistance reflex, an increase in the input resistance of the Dep MN was observed (Fig. 8). To understand the effect of this parameter on the integrative properties of the MN, we will examine first the relationship between the shape of an EPSP and the summation...
properties of the MN. Let us consider a volley of \( n \) EPSPs at a frequency \( N \). Each EPSP has a decay time constant \( \tau \). For simplicity, suppose that each EPSP can be modeled as an instantaneous rising phase followed by an exponential decay. The last EPSP will summate with the exponentially decreasing trace of the previous EPSP. In the case of a volley containing only two EPSPs (Fig. 13A), the maximum amplitude of depolarization (\( V \)) reached during the volley by the second EPSP is:

\[
V = V_0 + V_0 \times e^{-\frac{1}{\tau}}
\]

with

\[
t = \frac{1}{N},
\]

where \( V_0 \) is the amplitude of a single EPSP.

In a volley containing \( n \) EPSPs (Fig. 13B), we summate all preceding EPSPs:

\[
V = V_0 + V_0 \times e^{-\frac{1}{\tau N}} + V_0 \times e^{-\frac{2}{\tau N}} + \cdots + V_0 \times e^{-\frac{n-1}{\tau N}}
\]

\[
V = V_0 \times \left(1 + e^{-\frac{1}{\tau N}} + e^{-\frac{2}{\tau N}} + e^{-\frac{3}{\tau N}} + \cdots + e^{-\frac{n-1}{\tau N}}\right).
\]

In an infinite series of EPSPs:

\[
V = V_0 \times \left(\frac{1}{1 - e^{-\frac{1}{\tau N}}} \right).
\]

The value of \( V \) depends on the value of \( e^{-\frac{1}{\tau N}} \). If this value is close to 1, \( V \) will be very large, but generally this value is small because of the small time constant \( \tau \). In most of our recordings \( \tau = \approx 10 \) msec (10\(^{-2}\) sec). Therefore, with a frequency of 100 Hz (which is on the order of magnitude of EPSP frequency received by a given Dep MN), \( N\tau = 1 \), and \( e^{-\frac{1}{\tau N}} = 0.37 \), then \( V = V_0 \times 1.58 \).

This model is very simple and underestimates temporal summation because we considered each EPSP to have an instantaneous rising phase followed by a simple exponential decay that begins immediately (Fig. 13C). In fact, a real EPSP has a discrete rising phase that causes a delay in the onset of the exponential falling phase. To model this delay using EPSPs with single exponential falling phases, we artificially increased by a factor \( K \) the amplitude of all EPSPs preceding the final EPSP that was used in the measurement of \( V \) (Fig. 13D, E); then the estimation of \( V \) can be written for \( n \) EPSPs as:

\[
V = V_0 + K \times V_0 \times e^{-\frac{1}{\tau N}} + K \times V_0 \times e^{-\frac{2}{\tau N}} + K \times V_0 \times e^{-\frac{3}{\tau N}} + \cdots + K \times V_0 \times e^{-\frac{n-1}{\tau N}}.
\]

and with an infinite number of EPSPs:

\[
V = V_0 + K \times V_0 \times X(1 + X + X^2 + X^3 + \cdots + X^n) = V_0 + K \times V_0 \times \left(\frac{X}{1 - X}\right).
\]

In this new expression, with the same parameters (\( \tau = 10 \) msec; \( n = 100 \) Hz), we get a depolarization \( V = V_0 \times 2.16 \) during the 100 Hz volley. The maximum amplitude is directly proportional to the amplitude of EPSPs, whereas the two other parameters (\( \tau \) and \( n \)) are equally important and induce a nonlinear effect on summation.

Let us now examine the effect of 5HT on these parameters. We can hypothesize that any change in input resistance of the MN will result in a change of the same order of magnitude in the amplitude of EPSPs. If this is true, then in the presence of 5HT, the amplitude of EPSP should be increased by a factor of 1.2–2.9 (Fig. 8G). This first effect on the amplitude of EPSPs will directly produce an increase in the global response to movement of the same order of magnitude. In addition, 5HT will also increase the decay time constant \( \tau \) of EPSPs, which will increase the global response to movement during the volley of EPSPs. Finally, in the 5HT condition, polysynaptic EPSPs are added to the monosynaptic EPSPs. This addition is equivalent to increasing the frequency \( n \). For example, if the decay time constant \( \tau \) of EPSPs is doubled (from \( \tau = 7 \) msec to 14 msec, with \( n = 100 \) Hz) then \( X = e^{-\frac{1}{\tau}} \) is increased from 0.24 to 0.49, and the global response is increased from 0.81 to 1.46 mV (80% increase). If we add now the polysynaptic EPSPs (\( n = 200 \) Hz), \( X \) increases to 0.70 and the global response increases to 2.83 mV (247% increase). It is notable that these values are compatible with the experimental observations of the global response in the presence of 5HT (Fig. 8G). Thus realistic changes in the parameters (\( n \) and \( \tau \)) can result in considerable increase in summation.

On the other hand, the magnitude of the effects of increasing \( n \) or \( \tau \) will depend on the values of these parameters. If \( n \) is too small, summation will not be very effective because of the exponential form of \( X (X = e^{-\frac{1}{\tau N}}) \), and in this case, increasing \( \tau \) may not lead to a noticeable increase in the global response. For example, when the polysynaptic excitatory pathways are blocked, the frequency of synaptic input (\( n \)) is lower, leading to less temporal summation. This phenomenon was likely responsible for the small, statistically insignificant increase in the global response to 5HT in high divalent cation solution shown in Figure 11B2 (0.67 ± 0.02 mV in control to 0.78 ± 0.02 mV in 5HT condition), although \( \tau \) was increased by 5HT.

**Discussion**

We demonstrated that in the CNS 5HT modulates proprioceptive sensorimotor integration in crayfish walking legs. Indeed, 5HT enhances the resistance reflex involved in posture control via two main effects: (1) increasing input resistance of the Dep MN and (2) activation of the polysynaptic pathway (Figs. 9D, 11).

**Effect of 5HT on MNs**

The two effects of 5HT on Dep MNs (slow tonic depolarization and increase in input resistance) occur with different time scales. Depolarization started after 1 min in the presence of 5HT and reached a maximum after 5 min. In a series of experiments in which 5HT was pressure applied close to an intracellularly recorded Dep MN, each puff of 5HT elicited a rapid (a few tens of milliseconds) and transient depolarization (D. Cattaert, unpublished observations). Such 5HT-induced depolarization was observed in other preparations (Katz and Harris-Warrick, 1989; McCormick and Pape, 1990; Kiehn and Harris-Warrick, 1992). The second effect of 5HT on Dep MNs
consisted of an increase in the input resistance. In our experiments, this effect was detectable after 10 min in the presence of 5HT and reached a maximum after 25 min. Because of the long latency of this effect, the input resistance increase is likely mediated through metabotropic 5HT receptors and second messengers. The multiplicity of effects of 5HT on a given target was shown in the stomatogastric ganglion of the crab, in which 5HT evoked a depolarization in dorsal gastric (DG) neurons that was accompanied by a decrease in input conductance (Kiehn and Harris-Warrick, 1992). The authors demonstrated that 5HT modulated more than one target conductance in DG neurons and that the effect depended on the membrane potential of DG. In addition, in the escape circuit of the crayfish, 5HT induces dual opposite effects depending on the mode of application (Teshiba et al., 2001). In the present work, 5HT was applied with a slow perfusion system, and the level of 5HT increased gradually to reach 10 μM in the bath. Therefore, our findings are in accordance with the effects of low doses of 5HT on the escape circuit of the crayfish.

Effect of 5HT on the polysynaptic excitatory pathway
The polysynaptic pathways activated in the presence of 5HT involved spiking interneurons because they were blocked in the presence of high divalent cation solution. This result indicates that in the presence of 5HT, the involved spiking interneurons increased their excitability, allowing their membrane potential to reach the threshold for spike generation, and thereby evoked an EPSP in postsynaptic MNs. Such a 5HT-induced increase in excitability was demonstrated in rat facial MNs (Rasmussen and Aghajanian, 1990), pyramidal neurons from rat piriform cortex (Sheldon and Aghajanian, 1991), and neurons of the lamprey spinal cord (Van Dongen et al., 1986).

On the other hand, the increase in the polysynaptic response recorded from the MNs (Fig. 9) may result from an increase in the amplitude of the polysynaptic EPSPs. This later effect may result from the postsynaptic effect of 5HT on input resistance as analyzed in the mathematical model presented in this paper. However, the increase of EPSP amplitude in Dep MNs could also involve presynaptic mechanisms as was shown in rat dorsolateral septal nucleus via 5HT2A receptors (Hasuo et al., 2002).

Multiplicity of 5HT effects
It is notable that each of the 5HT-induced changes reported here would lead to minor effects if occurring alone, but that it is their conjunction that produces a dramatic enhancement of the resistance reflex. The small but consistent depolarization of the MNs will bring their membrane potential closer to spike threshold, thereby causing an increase in firing rate. The increase in the time constant of the MNs increases temporal summation of EPSPs and thereby allows the MNs to reach more depolarized potentials during a volley of EPSPs. The likely increase in excitability of the interneurons causes these neurons to fire more readily in response to a given sensory input and thereby reinforces the polysynaptic component of the compound EPSPs triggered by sensory spikes.

In addition to the central effects analyzed here, 5HT also induces peripheral effects on the sensory neurons and on muscles. It is striking that these peripheral effects act in the same direction as the central effects demonstrated here. When 5HT is bath applied on the CBCO proprioceptor, the excitability of the sensory neurons is increased (Rossi-Durand, 1993); however, the maximal effect was obtained with a concentration of 1 μM. Higher concentrations seemed to be less effective, and for 20% of the recorded cells, 100 μM 5HT induced a decrease of the sensory discharge. The duality of the effects on sensory neurons for 5HT concentrations of 10 μM are in accordance with the present report on MNs. The significance of this dual effect is discussed below.

The other peripheral target of 5HT is the neuromuscular transmission. In the crayfish opener muscle, EPSP amplitudes increase up to 18-fold in the presence of 5-HT (Fischer and Florey, 1983). This effect was caused predominantly by a serotonin-induced increase in the release of quanta of excitatory transmitter with nerve stimulation (Glusman and Kravitz, 1982). This increased synaptic transmission at the neuromuscular junction is use dependent (Qian and Delaney, 1997). In the locust, 5HT was also shown to increase muscle contraction. For example, bath application of 5HT on the mandibular closer muscles of the locust while the muscles are active produces marked changes in the contractions, increasing their amplitude, rate of contraction, and rate of relaxation (Baines et al., 1990). These authors have shown that 5HT elevates the levels of cAMP in isolated muscle fibers.

All of these effects of 5HT seem to contribute to the same goal: increasing the resistance reflex activity. In the presence of 5HT, sensory neurons increase their sensitivity to movement, providing the CNS target neurons with higher rates of spike discharge. In the presence of 5HT, interneurons of polysynaptic pathways of the resistance reflex are more excitable and elicit large polysynaptic EPSPs in the MNs. In the presence of 5HT, these MNs increase their responsiveness in at least two ways: a moderate tonic depolarization and an increased input resistance. As a consequence, their input–output relationship is steeper, and because of the increased sensory inflow relayed and amplified by interneurons of polysynaptic pathways, they are recruited more readily in response to any movement of the leg. As a consequence, previously silent MNs are activated. Finally, the two last stages (neuromuscular synaptic transmission and muscle contraction) are also enhanced in the presence of 5HT (Fischer and Florey, 1983; Baines et al., 1990). All of these combined effects may be caused by the activity of the 5HT neurons that exist by pairs in thoracic and abdominal ganglia (Beltz and Kravitz, 1983). It is not known, however, whether these 5HT-containing neurons project on the various targets mentioned here, and the modality of their discharge in the living animal is not known.

Variability of 5HT-induced responses
In experiments using simultaneous intracellular recordings from Dep MNs involved in resistance reflex, either all recorded Dep MNs displayed an increase of the resistance reflex or none of them was affected. Therefore, one source of variability of the response to 5HT is inter-individual. On the other hand, in a given experiment, the magnitude of the effect of 5HT was different among the different intracellularly recorded Dep MNs (Fig. 5), suggesting a variability between MNs. This inter–MN variability could have several nonexclusive origins. First, the increase in the resistance reflex response can depend on the distance between the recording site and synapses of the polysynaptic pathway. Dep MNs indeed have a complex morphology with many branches (Cattaert et al., 1994b; Clarac and Cattaert, 1996; Watson et al., 2000), and there are few data concerning the location of monosynaptic (Cattaert and El Manira, 1999) and no data about polysynaptic input synapses. Second, the inter–MN variability
could suggest an heterogeneity in the MN population. Such a heterogeneity was shown for the response of Dep MNs to acetylcholine (ACh). Some Dep MN are conditional oscillators and present rhythmic oscillations of their membrane potential in the presence of ACh, whereas others never do (Cattaert et al., 1994a,b).

One possible source of inter-individual variability could be related to the social status of the animal. In crayfish, the effect of 5HT on the neural circuit for tail-flip escape behavior was found to depend also on the animals’ social experience. 5HT reversibly enhanced the response to sensory stimuli of the lateral giant tail-flip command neuron in socially dominant crayfish and reversibly inhibited it in subordinate animals (Yeh et al., 1996). All of the animals used in our experiments were maintained in large tanks containing many animals. Therefore, we did not know the social status of each experimental animal. Given the fact that dominant animals generally present a more elevated posture than subordinate animals, we hypothesize that 5HT can enhance the resistance reflex response of the Dep MN in dominant animals and have no effect (or a negative effect) on the resistance reflex in subordinate animals. However, because of the fact that 5HT increases reflex activation in both Dep and Lev MNs, it is likely that if this hypothesis is true, 5HT modulation would lead to an overall increase in joint stiffness rather than toward an asymmetric change in joint posture in dominant animals. These effects are also dependent on which MNs participate to maintaining posture, and therefore the effect of 5HT on posture in vivo cannot be drawn directly from the present results.

References


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