GABAergic Excitatory Synapses and Electrical Coupling Sustain Prolonged Discharges in the Prey Capture Neural Network of Clione limacina

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Neuronal afterdischarges represent an essential and widespread phenomenon in brain functioning. The main important implication of afterdischarges during signal processing in the brain is a transformation of a brief sensory input from a prey into a lasting prey capture response. The present study, which focuses on the neuronal mechanisms of afterdischarges, demonstrates that a single pair of interneurons [cerebral A interneuron (Cr-Aint)] is responsible for afterdischarge generation in the network. Cr-Aint neurons are electrically coupled to all other neurons in the network and produce slow excitatory synaptic inputs to them. This excitatory transmission is found to be GABAergic, which is demonstrated by the use of GABA antagonists, uptake inhibitors, and double-labeling experiments showing that Cr-Aint neurons are GABA-immunoreactive. The Cr-Aint neurons organize three different pathways in the prey capture network, which provide positive feedback necessary for sustaining prolonged spike activity. The first pathway includes electrical coupling and slow chemical transmission from the Cr-Aint neurons to all other neurons in the network. The second feedback is based on excitatory reciprocal connections between contralateral interneurons. Recurrent excitation via the contralateral cell can sustain prolonged interneuron firing, which then drives the activity of all other cells in the network. The third positive feedback is represented by prominent afterdepolarizing potentials after individual spikes in the Cr-Aint neurons. Afterdepolarizations apparently represent recurrent GABAergic excitatory inputs. It is suggested here that these afterdepolarizing potentials are produced by GABAergic excitatory autapses.

Key words: recurrent inputs; autapses; afterdepolarization; afterdischarge; positive feedback; neural network; mollusc; feeding; GABAergic interneuron; GABA immunoreactivity

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a pathway for recurrent excitation via the contralateral cell. The third possible feedback suggested by the present investigation is the existence of recurrent synapses produced by Cr-Aint neurons on themselves (“autapses”; Van der Loos and Glaser, 1972).

**MATERIALS AND METHODS**

**Preparation.** Adult specimens of *Clione limacina*, body length of 1–3 cm, were collected from the breakwater at Friday Harbor Laboratories, University of Washington (Friday Harbor, WA) in the Spring–Summer season and were held in 1-gallon jars in a refrigerator at 5–7°C. Animals were anaesthetized in a 1:1 mixture of seawater and isotonic MgCl₂, and dissected in a Sylgard (Dow Corning, Midland, MI)-coated Petri dish. Electrophysiological experiments were performed on reduced preparations consisting of the CNS, head, and wings. All nerves running from the central ganglia to the head and wings remained intact. Before electrophysiological recordings, the sheaths of the central ganglia were softened by bathing the preparation in a 1 mg/ml solution of protease (type XIV; Sigma, St. Louis, MO) for ~5 min, followed by a 30 min wash.

**Electrophysiological recordings.** Intracellular recordings were made with glass microelectrodes filled with 2 M potassium acetate. Microelectrodes had resistances of 10–30 MΩ. Electrophysiological signals were amplified, displayed, and recorded by using standard electrophysiological techniques. Intracellular stimulation was provided via amplifier bridge circuit, which was balanced before each experiment. Changes in the membrane potential during measurements of electrical coupling or reversal potentials were provided by using two electrodes in one cell: one for current injections and the other for voltage recording. Changes in the membrane conductance were estimated by measuring changes in the amplitude of the membrane potential hyperpolarization steps, which were induced by injecting constant negative current pulses via a second intracellular electrode. Individual neurons were identified based on their electrophysiological properties, morphology, and motor effects. To test for monosynaptic connections, a high-Mg²⁺/high-Ca²⁺ solution (2.5× normal; 110 mM MgCl₂ and 25 mM CaCl₂) was used. High divalent solutions work appropriately in sensitive to GABA, neurons were chemically isolated by a high-Mg²⁺ from 10 to 20 Hz. To test whether a particular neuron was directly affected by given drugs, the effects of the drugs on the postsynaptic responses were studied (duration of the drug treatment was usually 0.5 sec; the firing frequency varied in different experiments). Artificial seawater composition was (in mM): NaCl 420, KCl 110, CaCl₂ 10, and NaHCO₃ 2, pH adjusted to 7.4. Na⁻⁻ replaced saline was made by substituting Na⁺ with equimolar choline (Sigma) on an equimolar basis. Ten normality NaOH was used to adjust pH to 7.4; therefore, the final concentration of Na⁺ was ~25%. Four hundred micromolar t-arginine hydrochloride in normal seawater did not have an essential effect on the Cr-A neuron activity, only slightly depolarizing their membrane potential (not more than 5 mV; n = 4). It also did not interfere with the effect of GABA on the Cr-A neurons (n = 4). Replacement of Na⁺ in zero-Ca⁺⁺ saline was done by substituting Mg²⁺.

**Reagents and drugs.** GABA, its antagonists 5-aminovaleric acid, saxitoxin, bicuculline, t-dihydrolipoic acid (dihydrolipoic acid) from Research Biochemicals, Natick, MA), bicuculline methochloride, picrotoxin (both from Sigma), and uptake inhibitor nipecotic acid (Research Biochemicals) were applied with the use of a graduated 1 ml pipette. The final concentration was estimated from the known volume of injected solution and the known volume of saline in the recording dish. The duration of the effects was estimated from those of the induced bursts of spikes in interneurons were constant in the controls and during drug application, when effects of the drugs on the postsynaptic responses were studied (duration was usually 0.5 sec; the firing frequency varied in different experiments from 10 to 20 Hz.) To test whether a particular neuron was directly sensitive to GABA, neurons were chemically isolated by a high-Mg²⁺ solution consisting of a 1:3 mix of 0.33 M MgCl₂ in seawater superfused into the preparation dish.

**GABA-immunocytochemistry and double-labeling experiments.** Because *Clione limacina* is a relatively small mollusc, entire reduced preparations were used for the whole-mount immunocytochemical procedure. The preparations were fixed for 3 hr in 4% paraformaldehyde and 0.1% glutaraldehyde in PBS (pH 6.5–7.0) at room temperature (15–20°C). To reduce high nonspecific fluorescence caused by glutaraldehyde fixation, the tissues were incubated overnight in 4% sodium borohydride in PBS (Kosaka et al., 1986). The preparations were then transferred to 3 ml vials, where they were washed for 12 hr in PBS and preincubated in PBS containing 0.1% Triton X-100 to increase tissue permeability. The tissues were then exposed to 6% goat serum in PBS and 0.1% Triton X-100 for 6 hr to reduce nonspecific staining and incubated 36 hr at 5°C in GABA antibody (polyonal GABA antiserum raised in guinea pig; Eugene Tech International, Inc., Ridgefield Park, NJ). The dilution of primary antibody was 1:500. After a 12 hr wash in PBS, the tissues were placed for 24 hr in fluorescein-labeled secondary antibody (working concentration of 40 μg/ml; anti-guinea pig IgG produced in goat; Vector Laboratories). The secondary antibody was removed with several PBS exchanges, and preparations were washed overnight. The tissues were then cleared in xylene, mounted in DPX, and examined in whole mount with a Nikon (Tokyo, Japan) fluorescence microscope and a Bio-Rad (Hercules, CA) MRC 600 laser scanning confocal microscope. The first set of the control experiments included preadsorption of the primary antibody with 50 μM GABA–BSA conjugate for 6 hr before processing the tissue. In the second set of controls, primary antibody was omitted from the procedure. No staining resulted in both sets of control experiments.

**RESULTS**

**Cerebral A interneuron is necessary and sufficient for afterdischarge generation**

A bilaterally symmetrical Cr-Aint neuron was found to be necessary and sufficient for generating afterdischarges in the Cr-A neuron network. All neurons in the network were normally silent, with membrane potentials of Cr-A motoneurons varying between –60 and –66 mV and Cr-Aint neurons varying between –58 and –63 mV. The level of excitability of the Cr-A neuron network, including its ability to generate afterdischarges, significantly varied in different preparations, which apparently reflected the level of feeding arousal (Norekian, 1993). For the first group of experiments, preparations with a high level of excitability were chosen in which each induced Cr-A neuron burst of spikes always triggered afterdischarges. Synchronous discharges were always produced in a simultaneously recorded Cr-Aint neuron (Fig. 1A). When the Cr-Aint neuron was removed from the network activity by injecting hyperpolarizing current, afterdischarges disappeared (n = 5 preparations with 5–10 presentations each) (Fig. 1A).

After release of the Cr-Aint neuron from hyperpolarization, induced Cr-A neuron bursts of spikes again triggered discharges in the Cr-Aint neuron and prolonged afterdischarges in the Cr-A neuron. In addition, when Cr-Aint neurons were mechanically removed from the ganglia, afterdischarges in the Cr-A neuron network disappeared and were never observed in these preparations (n = 3). For the second group of experiments, preparations with a low level of Cr-A neuron network excitability were chosen in which induced Cr-A neuron bursts of spikes did not trigger afterdischarges (Fig. 1B). In all of these preparations, only subthreshold depolarizing potentials were recorded in the Cr-Aint neuron, produced by Cr-A neuron bursts of spikes via electrical connections. When the Cr-Aint neuron was depolarized close to the spiking threshold and Cr-A neuron bursts became efficient in...
triggering Cr-Aint neuron firing, afterdischarges appeared in Cr-A neurons (n = 9 preparations with 5–10 presentations in each) (Fig. 1B). All simultaneous recordings of Cr-A and Cr-Aint neuron activities demonstrated that afterdischarges appeared only if the Cr-Aint neuron was spiking and were never observed when the Cr-Aint neuron did not produce action potentials (n = 32 preparations). Thus, the Cr-Aint neuron appears to be a key element for afterdischarge generation in the Cr-A neuron network.

**Synaptic connections between cerebral A interneurons and A motoneurons**

When a single Cr-A neuron was stimulated to trigger afterdischarge activity, it activated the Cr-Aint neuron via electrotonic transmission. Electrical coupling between a Cr-Aint neuron and Cr-A motoneurons was nonrectifying and relatively weak (considering high thresholds for spike generation in normal conditions). Coupling coefficients ranged from 0.1 to 0.25 (n = 7; measured at resting membrane potentials). These measurements explain why even strong bursts of spikes induced in a Cr-A neuron sometimes were not capable of producing afterdischarges; they could not activate Cr-Aint neurons via electrotonic transmission. On the other hand, induced Cr-Aint neuron bursts always generated Cr-Aint neuron afterdischarges and always produced simultaneous discharges in Cr-A neurons (n = 32 preparations, several presentations in each) (Fig. 2A). The important characteristic of the discharges in both Cr-Aint and Cr-A neurons was a prominent underlying depolarization. These facts suggested the existence of additional excitatory synaptic transmission from Cr-Aint to Cr-A neurons.

In high divalent solution, the Cr-Aint neuron bursts of spikes produced, in addition to the fast electrical membrane changes, slow depolarizations in the Cr-A neurons (n = 28 preparations) (Fig. 2B). These slow depolarizations had amplitudes of 7–15 mV and lasted 1–2 sec, depending on the strength of a Cr-Aint neuron burst of spikes. Each spike in a Cr-Aint neuron produced a single EPSP in a Cr-A neuron (Fig. 3A). This stable one spike/one EPSP ratio persisted in high-Mg<sup>2+</sup>/high-Ca<sup>2+</sup> saline, suggesting monosynaptic connections (n = 12 preparations). Each EPSP had an amplitude between 2 and 6 mV and had a slow decline lasting ~0.5 sec. All tested Cr-A neurons received those EPSPs from the Cr-Aint neurons, including cells on the dorsal and ventral sides and from the ipsilateral and contralateral cerebral ganglia (n = 18 different Cr-A neurons in five preparations) (Fig. 3A, B).

The Cr-Aint neuron is a bilaterally symmetrical cell (Arshavsky et al., 1993; Norekian and Satterlie, 1993a). Contralateral Cr-Aint neurons produced strong excitatory inputs to each other (n = 8) (Fig. 4A). Interneurons were found to be electrically coupled, with coupling coefficients between 0.1 and 0.2 (n = 5) (Fig. 4B). In addition to electrical transmission, each Cr-Aint neuron received from its contralateral compatriot slow depolarizing inputs similar to those produced in Cr-A neurons. These slow depolarizing potentials persisted in high-Mg<sup>2+</sup>/high-Ca<sup>2+</sup> saline, suggesting monosynaptic connections (n = 6) (Fig. 4C). Thus, each Cr-Aint neuron could activate its contralateral compatriot and, in turn, receive recurrent excitatory inputs from it. Positive feedback organized via these excitatory reciprocal connections apparently participates in the afterdischarge generation in Cr-Aint neurons and subsequently Cr-A neurons. However, it
was not the only positive feedback, as the following experiments demonstrated.

**Afterdepolarizing potentials in A interneurons**

In high-Mg\(^{2+}\)/high-Ca\(^{2+}\) saline, induced bursts of spikes in a Cr-Aint neuron produced prominent afterdepolarizations in the stimulated interneuron. Afterdepolarizations were very similar in shape to slow EPSPs in Cr-A neurons, had amplitudes of 7–15 mV, and lasted 1–2 sec, depending on the strength of the Cr-Aint neuron burst (n = 32) (Fig. 2B). Subthreshold depolarizations never induced Cr-Aint neuron afterdepolarizations; only spikes were effective (Fig. 3A). Each spike in a Cr-Aint neuron produced an afterdepolarizing potential that had an amplitude between 1 and 4 mV and had a slow decline lasting ~0.5 sec (n = 12) (Fig. 3A,C). The appearance of afterdepolarizing potentials after individual spikes in Cr-Aint neurons persisted in high-Mg\(^{2+}\)/high-Ca\(^{2+}\) saline (n = 16 preparations, several presentations in each) (Fig. 3A,C). Such stable occurrence of afterdepolarizations in high divalent solution suggested that these afterdepolarizations represented a property of the Cr-Aint neuron rather than polysynaptic recurrent inputs. Positive feedback provided by excitatory synaptic connections between contralateral Cr-Aint neurons was certainly not involved, because Cr-Aint neuron afterdepolarizations still persisted when the contralateral neuron was not spiking in high-Mg\(^{2+}\)/high-Ca\(^{2+}\) saline (Fig. 4C). In addition, when one Cr-Aint neuron was mechanically removed from the
ganglia (cell body with extended part of an axon and secondary branches), afterdischarges still persisted in the network \((n = 5)\). Afterdepolarizing potentials triggered by each single Cr-Aint neuron spike could have two theoretically possible mechanisms. First, spikes could activate slow voltage-sensitive inward current, which would depolarize the Cr-Aint neuron membrane potential. Slow inward currents play a major role in the generation of endogenous bursting activity in different pacemaker neurons (Thompson and Smith, 1976; Adams et al., 1980; Bulloch and Willows, 1981). Second, afterdepolarizations could be a result of the synaptic self-excitatory inputs. The fact that Cr-Aint neuron afterdepolarizations were very similar in shape to the simultaneously induced Cr-A neuron EPSPs suggested that they could be excitatory synaptic inputs produced by interneurons to themselves. Moreover, repetitive Cr-Aint neuron stimulation produced a prominent decrement of afterdepolarizations, with the same degree and time course as the parallel decrement of Cr-A neuron EPSPs \((n = 12)\). One general characteristic of any synaptic transmission is that it is a \(Ca^{2+}\)-dependent process. The goal of the first group of experiments was to test whether the Cr-Aint neuron afterdepolarizations were \(Ca^{2+}\)-dependent. When normal seawater in the recording dish was replaced with zero-\(Ca^{2+}\) seawater, Cr-Aint neuron afterdepolarizations disappeared, along with slow EPSPs in Cr-A motoneurons \((n = 4)\) (Fig. 5A). Cr-Aint neuron afterdepolarizations and Cr-A neuron EPSPs were also completely and reversibly blocked by \(Ca^{2+}\) channel antagonists: 120 mM Mg\(^{2+}\) \((n = 6)\) (Fig. 5B) and 0.2 mM Cd\(^{2+}\) \((n = 3)\). In addition, the Cr-Aint neuron afterdepolarizations, as well as Cr-A neuron EPSPs, were enhanced in high-\(Ca^{2+}\) saline (30 mM; \(n = 3)\). These experiments concluded that Cr-Aint neuron afterdepolarizing potentials were \(Ca^{2+}\)-dependent and that they could be synaptic inputs. However, there are too many \(Ca^{2+}\)-dependent processes in neural networks, and the only way to unequivocally establish synaptic transmission as a mechanism of Cr-Aint neuron afterdepolarizations would be to demonstrate the effects of the specific pharmacological agents.

**Depolarizations produced by A interneurons are GABAergic excitatory synaptic inputs**

In addition to Cr-A motoneurons, Cr-Aint neurons also targeted buccal cone retractor Cr-B motoneurons (Norekian and Satterlie, 1993a). Cr-Aint neurons produced strong inhibitory inputs to Cr-B neurons, which temporally interrupted their spontaneous spike activities \((n = 5)\) (Fig. 6). These inhibitory hyperpolarizing inputs persisted in high-Mg\(^{2+}\)/high-\(Ca^{2+}\) seawater, suggesting direct connection between neurons. The only neurotransmitter,
which mimicked all effects produced by Cr-Aint neurons, was GABA. GABA has been shown previously to exert a strong excitatory effect on Cr-A neurons (Arshavsky et al., 1993; Norekian and Satterlie, 1993b). In high-Mg$^{2+}$ seawater, 10 $\mu$M GABA produced prominent depolarization of Cr-A motoneurons and hyperpolarization of Cr-B motoneurons ($n = 4$) (Fig. 7) (Norekian and Satterlie, 1993b). Exogenous 10 $\mu$M GABA also produced a prominent depolarization of a Cr-Aint neuron itself ($n = 5$) (Fig. 7). Thus, GABA mimicked the effects induced by a Cr-Aint neuron and was suggested as its possible neurotransmitter.

The amplitude of 10 $\mu$M GABA-induced depolarizations in the Cr-A and Cr-Aint neurons was $\sim 20$ mV when measured from the resting membrane potentials ($n = 12$). When Cr-A motoneuron or Cr-Aint neuron membrane potentials were depolarized to $+20$ mV by injecting a positive current via a second electrode, GABA-induced depolarizations decreased in amplitude to 5 mV but still did not reverse ($n = 5$). Similarly, Cr-Aint neuron-induced EPSPs in the Cr-A motoneurons did not reach a reversal potential when the motoneuron membrane potential was depolarized to $+20$ mV ($n = 7$). GABA-induced depolarizations, as well as Cr-A motoneuron EPSPs and Cr-Aint neuron afterdepolarizations, were associated with increases in membrane conductance ($n = 10$). When seawater in the recording chamber was replaced with 2% Na$^+$ solution, GABA-induced depolarizations in the Cr-A and Cr-Aint neurons completely disappeared (GABA concentration of 50 $\mu$M; $n = 5$) (Fig. 8). After a 5 min wash in seawater, GABA again produced prominent depolarizing responses. Replacement of seawater with zero Ca$^{2+}$ and 1 mM Co$^{2+}$ solution did not influence GABA-induced depolarizations (50 $\mu$M GABA; $n = 3$). These observations suggest that GABA-induced depolarizations are Na$^+$-dependent.

The GABA antagonists bicuculline, phaclofen, saclofen, and picrotoxin, in concentrations of 50 $\mu$M and 1 mM, did not block 1 $\mu$M GABA-induced responses ($n = 9$). They also did not block Cr-A motoneuron EPSPs and Cr-Aint neuron afterdepolarizations ($n = 12$). The GABA antagonist 5-aminovaleric acid (50
produced a prominent depolarization and activation of the Cr-A and Cr-Aint neurons \((n = 4)\). Ten micromolar 5-aminovaleric acid depolarized neuron membrane potential 5–9 mV above the resting potential without initiating spikes and demonstrated a reversible blocking effect on the 1 \(\mu M\) GABA-induced responses \((n = 3)\). It also reversibly blocked chemical transmission from a Cr-Aint neuron to Cr-A motoneurons and Cr-Aint neuron afterdepolarizations \((n = 7)\). However, the best results were obtained with piperidine-4-S, which acts as a GABA antagonist in some systems (Woodward et al., 1993). Fifty micromolar piperidine-4-S did not significantly alter Cr-A and Cr-Aint neuron membrane potentials. Only a slight depolarization of <2 mV amplitude was observed in some experiments. Responses of Cr-A and Cr-Aint neurons to 1 \(\mu M\) GABA were completely and reversibly blocked by 50 \(\mu M\) piperidine-4-S, indicating that it works as a GABA antagonist in this system \((n = 3)\) (Fig. 9). Fifty micromolar piperidine-4-S reversibly blocked chemical transmission from Cr-Aint neurons to Cr-A motoneurons \((n = 8)\) (Fig. 10A). In these experiments, slow EPSPs in the Cr-A motoneurons completely disappeared, and the remaining fast responses were very similar in amplitude and shape to the Cr-A neuron responses in zero-Ca\(^{2+}\) or high-Mg\(^{2+}\) experiments (Fig. 5). Apparently, only fast electrical transmission remained functional between neurons. At the same time, 50 \(\mu M\) piperidine-4-S completely and reversibly blocked afterdepolarizations in the Cr-Aint neurons \((n = 8)\) (Fig. 10A). Nipecotic acid, a GABA uptake inhibitor, significantly enhanced both slow Cr-A neuron EPSPs and Cr-Aint neuron afterdepolarizations, increasing their amplitudes but most dramatically their durations \((n = 8)\) (Fig. 10B). Fifty micromolar nipecotic acid produced a twofold to threefold increase in the duration of both Cr-A neuron EPSPs and Cr-Aint neuron afterdepolarizations compared with controls. Application of the nipecotic acid always resulted in the 3–5 mV depolarization of Cr-A and Cr-Aint neurons and required the adjustment of the stimulating current to keep the firing frequency of the induced bursts of spikes in the Cr-Aint neurons constant. These experiments demonstrated that synaptic transmission from interneurons to Cr-A motoneurons was GABAergic. They also provided evidence that afterdepolarizations in Cr-Aint neurons are recurrent synaptic inputs, with GABA being a neurotransmitter.
Because synaptic transmission from a Cr-Aint neuron to its target cells was sensitive to GABA pharmacology, Cr-Aint neurons were expected to contain GABA. Validation of this expectation was important, because it would provide additional confirmation of its GABAergic nature and the direct nature of the synapses studied. One technique, which could demonstrate that Cr-Aint neurons contained GABA, was immunocytochemistry. GABA immunoreactivity was studied in 16 whole-mount preparations. General distribution of GABA-immunoreactive neurons in the CNS was similar to that described by Arshavsky et al. (1993). There were small groups of immunoreactive neurons in the buccal, pedal, and cerebral ganglia, and no stained neurons in the pleural and intestinal ganglia.

**GABA immunocytochemistry and double-labeling experiments**

Because synaptic transmission from a Cr-Aint neuron to its target cells was sensitive to GABA pharmacology, Cr-Aint neurons were expected to contain GABA. Validation of this expectation was important, because it would provide additional confirmation of its GABAergic nature and the direct nature of the synapses studied. One technique, which could demonstrate that Cr-Aint neurons contained GABA, was immunocytochemistry. GABA immunoreactivity was studied in 16 whole-mount preparations. General distribution of GABA-immunoreactive neurons in the CNS was similar to that described by Arshavsky et al. (1993). There were small groups of immunoreactive neurons in the buccal, pedal, and cerebral ganglia, and no stained neurons in the pleural and intestinal ganglia.

Immunoreactive cells in the buccal ganglia included three pairs of small neurons, which sent their axons into the cerebrobuccal connectives (Fig. 11). The pedal ganglia contained a cluster of five to six small neurons in the lateral region, one cell in the anterior region on the dorsal surface, and two cells in the medial region on

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**Figure 10.** A, The GABA antagonist piperidine-4-S completely and reversibly blocked Cr-Aint neuron afterdepolarizations and slow EPSPs in the Cr-A11 neuron. Apparently, only fast electrical transmission remained functional. B, Nipeptic acid, a GABA uptake inhibitor, increased the duration of both Cr-Aint neuron afterdepolarizations and slow EPSPs in the Cr-A11 neuron. All experiments were conducted in the seawater with the increased concentration of Mg$^{2+}$ and Ca$^{2+}$ (1.75× normal or half of the high-Mg$^{2+}$/high-Ca$^{2+}$ solution) to prevent prolonged firing and to unmask slow depolarizing responses in neurons. Calibration: 10 mV, 1 sec.

**Figure 11.** GABA immunoreactivity in the buccal ganglia: a composite confocal microscope reconstruction from several optical sections. Note the stained processes exiting the ganglia into the cerebrobuccal connectives (C-B). Scale bar, 100 μm.
the ventral surface (Fig. 12A). Although the pleural and intestinal ganglia did not contain any immunoreactive cell bodies, their neuropiles were innervated by immunoreactive processes (Fig. 12B). It appeared that only two symmetrical pairs of immunoreactive axon branches innervated the neuropile of the intestinal ganglia, and two pairs innervated the neuropile of the pleural ganglia. Approximately 10 pairs of immunoreactive neurons were found in the cerebral ganglia (Fig. 13B). One small brightly stained bilaterally symmetrical neuron was located near the head nerves. A group of small neurons was spread in the center of the dorsal surface of each cerebral ganglion. One bilaterally symmetrical neuron of middle size showed strong immunoreactivity and had a location similar to that of a Cr-Ainter neuron, suggesting that it might be the same cell (Fig. 13B).

Double-labeling experiments undoubtedly demonstrated that the Cr-Ainter neuron is GABA-immunoreactive (n = 10). One Cr-Ainter neuron in each preparation, from the left or right cerebral ganglion, was injected with neurobiotin and visualized with Texas Red (Fig. 13A). The same preparation was then processed for GABA immunoreactivity with the fluorescein-labeled secondary antibody (Fig. 13B). By switching filters in the same preparation, the Cr-Ainter neuron was identified as a GABA-immunoreactive neuron. Texas Red and fluorescein had very distinct emission wavelengths and were visible only in their own set of filters, thus allowing complete separation of images and clear interpretation of data.

One unusual observation, which resulted from the double-labeling experiments, was the lack of dye coupling in this system of electrically coupled neurons after neurobiotin injections (Fig. 13A). Neurobiotin is known to produce dye coupling, even when other junction-permeable dyes, such as Lucifer yellow, are not efficient (Vaney, 1991). The lack of neurobiotin coupling in the Cr-A neuron network could be explained by the specificity of the gap junctions, such as narrow junctional channels, and/or by the.

Figure 12. GABA immunoreactivity in the central ganglia. A. Left pedal ganglion. Note several stained processes in the pedal commissure (com). B. Pleural (pl) and intestinal (int) ganglia. Note that there are no stained cell bodies in these ganglia. All images represent a composite confocal microscope reconstruction. Scale bar, 200 μm.
the lack of a prolonged incubation period before fixation, which was always under 30 min.

**DISCUSSION**

**Mechanisms of afterdischarge generation in the prey capture network**

Afterdischarge activity in the *Clione* prey capture network has an important functional implication for the feeding behavior. It is responsible for transformation of a brief sensory input from a prey into a long-lasting motor output—prolonged protraction of the prey capture appendages (Norekian, 1993). The focus of the current investigation is how these afterdischarges are generated. It has been demonstrated previously that afterdischarge activity is not attributable to intrinsic properties of individual Cr-A motoneurons (Norekian, 1993). The present research revealed that Cr-A interneurons and their synaptic connections are necessary for afterdischarge generation.

The emerging picture of synaptic connections between Cr-A interneurons and Cr-A motoneurons is shown in Figure 14. Cr-A interneurons and Cr-A motoneurons are electrically coupled. There is also a slow excitatory synaptic transmission from Cr-A interneurons to Cr-A motoneurons. The excitatory synaptic inputs produced by Cr-A interneurons were found to be GABAergic, as demonstrated by the use of GABA antagonists, uptake inhibitors, and double-labeling experiments, which showed that Cr-A interneurons are GABA-immunoreactive. GABA was originally known as an inhibitory neurotransmitter. However, GABA-induced depolarizing effects were found in several neural systems (Gallagher et al., 1978; Ogata, 1987; Pfeiffer-Linn and Glantz, 1989; El-Beheiry and Puil, 1990; Mercuri et al., 1991; Michelson and Wong, 1991). In *Clione*, it appears that GABA works primarily as an excitatory neurotransmitter in the feeding system (Arshavsky et al., 1993; Norekian and Satterlie, 1993b). GABA-induced depolarizations in the Cr-A interneurons and Cr-A motoneurons have very high

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**Figure 13.** Double-labeling experiment. *A.* Image of the cerebral ganglia obtained via Texas Red filters. A single Cr-A interneuron was filled with neurobiotin and visualized by Texas Red. The cell body is indicated by an arrow. *B.* Image of the same cerebral ganglia obtained via fluorescein filters. This image shows GABA immunoreactivity in the cerebral ganglia (GABA antisera were labeled with fluorescein). The cell body of a Cr-A interneuron is indicated by an arrow. All images represent a composite confocal microscope reconstruction. Scale bars, 200 µm.
amplitudes, easily reach spike thresholds, and strongly activate the target cells. What makes these responses unusual compared with the previously known GABA-produced depolarizations is their Na$^+$ dependence and atypical pharmacology. Further investigation of these apparently novel GABAergic receptors will be an important direction for future experiments.

The Cr-Aint neurons organize three different pathways in the prey capture network, which provide the positive feedback necessary for sustaining prolonged spike activity. The first pathway includes electrical coupling and slow chemical transmission from the Cr-Aint neurons to Cr-A motoneurons. Cr-Aint neurons produce slow excitatory inputs to all Cr-A neurons in the network, and Cr-A motoneurons in turn activate the Cr-Aint neurons via electrotonic transmission. Electrical coupling between Cr-Aint neurons and Cr-A motoneurons, which all have high thresholds for spike generation, is relatively weak and does not efficiently propagate the impulses at the resting membrane potentials. However, when slow depolarizing inputs produced by Cr-Aint neurons bring all neurons close to the spiking thresholds, even weak electrical coupling could turn into a very efficient pathway for impulse transmission between neurons. The fact that neuron firing is very synchronized in the network during afterdischarges confirms the active involvement of electrical coupling (Norekian, 1993).

The second positive feedback mechanism, which apparently participates in sustaining neuron firing, is provided by reciprocal connections between contralateral Cr-Aint neurons (Fig. 14). Contralateral Cr-Aint neurons are electrically coupled. In addition, each interneuron produces slow excitatory synaptic inputs to its compatriot, similar to the synaptic inputs in the Cr-A motoneurons. Thus, when a brief sensory input induces a short burst of activity in the Cr-Aint neurons, each interneuron produces excitatory inputs to the contralateral interneuron, which returns this excitation. This recurrent excitation via the contralateral interneuron can sustain prolonged firing in the Cr-Aint neurons. Cr-Aint neuron firing in turn activates all Cr-A motoneurons in the network via excitatory synaptic transmission, and motoneurons fire as long as Cr-Aint neurons are active.

The third positive feedback mechanism is restricted to each individual Cr-Aint neuron. Single spikes in the Cr-Aint neurons produce afterdepolarizing potentials, which persist in high divalent solution. Induced bursts of spikes in a Cr-Aint neuron produce afterdepolarizations that are prominent enough to initiate the following set of spike activity, thus creating a positive feedback mechanism. Afterdepolarizations are sensitive to GABA pharmacology and apparently represent recurrent GABAergic excitatory inputs. How do Cr-Aint neurons activate themselves? Autoexcitatory effects of released neurotransmitters have been shown in previous studies. One well studied example is the group of bag cells in the mollusc Aplysia (Brown and Mayeri, 1989; Loechner and Kaczmarek, 1994). After stimulation, bag cells produce prolonged afterdischarges that typically last 15–30 min. Positive feedback responsible for this sustained firing activity is organized by autoexcitatory transmission of the several bag cell neuropeptides, which are nonsynaptically released during neuron firing. It appears highly unlikely that Cr-Aint neurons nonsynaptically release GABA. Cr-Aint neurons and Cr-A motoneurons do not represent a compact cluster of neurons as the bag cells do; their cell bodies and axons are spread around the entire cerebral ganglia. Nevertheless, they are the only neurons involved in the afterdischarge activity, although many other cerebral neurons are also sensitive to GABA (my unpublished observations). Cr-Aint neurons target only specific neurons, suggesting the existence of the specific synapses. Thus, afterdepolarizing potentials in the Cr-Aint neurons apparently represent GABAergic excitatory autaptic inputs (Fig. 14).

Self-excitatory synapses and afterdischarge generation

The existence of autapses in the brain has been documented by many morphological studies, starting from the work in rabbit neocortex by Van der Loos and Glaser (1972), who proposed the term autapse to describe a synapse between a neuron and a branch of its own axon. Autaptic connections were later found in various brain regions, such as monkey neostriatum (DiFiglia et al., 1976), cat substantia nigra (Karabelas and Purpura, 1980), rat striatum (Preston et al., 1980), and cat visual cortex (Tamás et al., 1995). Detailed light microscopic observations, combined with electron microscopy study, were recently accomplished in rat neocortex (Lubke et al., 1996) and cat visual cortex (Tamás et al., 1997). Thus, there are substantial morphological data obtained from several brain structures in different species, which demonstrate that autapses may be fairly common in the brain. Surprisingly, a physiological description of autapses and investigation of their possible significance in brain functioning are almost absent. Both excitatory and inhibitory autapses have been reported and extensively studied in cell culture (Crain, 1971; Bekkers and Stevens, 1991; Segal, 1991, 1994; Shi and Rayport, 1994). However, their formation in culture could be argued as an aberration in the absence of appropriate postsynaptic targets. Physiological studies in vivo are limited to only three reports describing inhibitory autapses. The first physiological study, which was performed on the cholinergic buccal neurons of the mollusc Aplysia, identified inhibitory autaptic connections as a mechanism of hyperpolarizing potentials that followed action potentials in these neurons and slowed their firing (Gardner, 1977; White and Gardner, 1981). The second report demonstrated that axon collaterals of the rat neostriatal spiny neurons mediate recurrent inhibition, a portion of which involves autaptic synapses (Park et al., 1980). A third, very recent report described autaptic inhibitory currents in interneurons from rat cerebellar slices and also suggested that their functional role is inhibition of interneuron firing during high-frequency discharges (Pouzat and Marty, 1998).
The present study suggests the existence of GABAergic excitatory autaptic connections in two Cr-Aint neurons, which are the key elements of the Clione prey capture neural network. Recurrent excitation organized by these autaptic connections apparently serves an important functional role, providing an efficient positive feedback for sustaining spike activity in the Cr-Aint neurons and generating afterdischarges in the entire network. A pair of interneurons [cerebral prey capture interneuron (Cr-PC)] has been previously identified in the cerebral ganglia that also produced electrical and excitatory connections to Cr-A motoneurons but without self-excitatory inputs (Norekian and Satterlie, 1995). These interneurons did not generate afterdischarge activity and triggered discharges in the Cr-A neural network only occasionally, when Cr-Aint neurons were apparently activated (Norekian and Satterlie, 1995). This comparison suggests that self-excitatory synapses in a single pair of key interneurons (Cr-Aint neurons) can influence dramatically the properties of the entire network and are very important for afterdischarge generation.

Although the physiological evidence presented here suggests that afterdepolarizing potentials in the Cr-Aint neurons represent the excitatory autaptic inputs, one alternative hypothesis exists that can explain the same data. According to this hypothesis, Cr-Aint neuron-induced excitatory synaptic inputs could produce depolarization of numerous Cr-A motoneurons, which in turn could feed back via electrical connections to maintain a depolarization in the Cr-Aint neurons. Electrical coupling measured between pairs of interneurons and motoneurons was found to be too weak to have such an effect. However, simultaneous depolarization of ~26 motoneurons could theoretically provide a substantial convergent current source back to Cr-Aint neurons. Unfortunately, there are no known blockers of electrical coupling in the Clione system, which would allow obtaining a direct physiological solution. One indirect consideration that favors the “excitatory autapses” hypothesis is the comparison with Cr-PC interneurons, which also have electrical connections with Cr-A motoneurons and produce strong excitatory synaptic inputs to all of them (Norekian and Satterlie, 1995). Although their synaptic transmission produces strong depolarization of all Cr-A motoneurons, Cr-PC interneurons do not have afterdepolarizing potentials and do not generate afterdischarge activity. However, the best and the final solution to the dilemma can be provided only by further computational and electron microscopic analysis. What this study unequivocally demonstrated is that a pair of GABAergic Cr-Aint neurons with their excitatory synaptic and electrical connections is responsible for sustaining prolonged discharges in a prey capture neural network of Clione.

REFERENCES

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