GABA INDUCED PRESYNAPTIC INHIBITION OF PERIPHERALLY LOCATED PARTS OF SPIDER CUTICULAR MECHANOSENSORY NEURONS

I. Panek and P.H. Torkkeli

Dept. of Physiology and Biophysics, Dalhousie University, Halifax, NS, B3H 4H7 Canada

INTRODUCTION

Presynaptic inhibition of mechanosensory afferents is an ubiquitous phenomenon throughout the animal kingdom. Inhibitory GABAergic synapses have been described in the axon terminals close to the output synapses of many types of primary afferent neurons in several verterbate and inverterbate species. This inhibition involves a depolarization of the afferent terminals called primary afferent neurons in several (PAD) that leads to blockade of action potential invasion into the terminals, or reduction of the amplitude of the propagate dation potentials and consequently to a reduction in the effectiveness of synaptic transmission to posts/mainer neurons. While several mechanisms have been suggested to be responsible for PAD, the most widely accepted model assumes that the depolarization is induced by efflux of CT from GABA-gated CT -channels which transiently drives the membrane potential loward the CT equilibrium potential. Because of the complicated morphology of afferent terminals it has not been possible to test this, or alternative models, using voltage-clamp experiments.

Recently, Fabian-Fine et al. (1999) described an extensive and complex efferent innervation of the cuticular mechanoreceptor neurons of the spider, *Cupiennius sulei* using electron microscopical and immunocytochemical approaches. One group of these, the VS-3 neurons innervating the lyriform silt sense organ on the pattel and of the spider, accessible to intracadinal their mechanically. (Höger et al. 1997) and voltage-Gekizawa et al. 1999; 2000; Torkkeli et al. 2001) activated currents have recently been investigated using the single-electrode voltage-champ (SEVC) method, creating a preparation that can be used not only to investigate the PAD, but also the currents that contribute to this phenomenon.

GABA EFFECTS ON VOLTAGE RESPONSE, MEMBRANE POTENTIAL AND MEMBRANE RESISTANCE



When 50 µM to 10 mM GABA or muscimol was applied to the bath solution the threshold for firing action potentials in the VS-3 neurons became higher, the action potential amplitude smaller (*right*) or the neurons ceased firing completely (*lcf*). This effect wars revensible. These GABA receptor agonists also induced a decrease of membrane resistance that can be seen when hyperpolarizing current pulses are given.



At -100 mV GABA application induced a membrane depolarization of about 40 mV (*left*). This recording was performed using short depolarizing voltage pulses (50 ms. 1 pulse 1.5 s) and it shows the change in the membrane resistance during the GABA nduced depolarization. The membrane resistance decreased about 74% after GABA applications ashown in the argraph (*right*) and this change was reversible

100µM GABA



Peak current (nA)

Voltage response of a VS-3 neuron to application of 100 µM GABA at four different holding voltages (*left*) and currents induced with application of 1 mM muscimol at four different holding voltages (*right*). Both GABA and muscimol induced a large membrane depolarization and ninward current at and below the resimg potential jobau -65 mV). This current reversed significantly above resting potential: Linear fit to the data from 22 different experiments with GABA(*gth below*) indicated that the reversal potential of GABA induced current was about -22 mV. The current voltage curve obtained from 6 experiments with muscimol indicated that the current eversel at about the same value, -20 mV.

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Schematic diagram of two sensory neurons in the spider patellar VS-3 slit sense organ Also shown are the fine efferent fibers that run parallel to the sensory neurons. Fabina-Fine et al. (1999) found three efferent fibers that show GABA like immunoreactivity. Synaptic connections are very complex-there are at least four different types of synaptic vesicle sensory neurons, glia and other efferent neurons. In addition to the simple undirectional synapses shore are reciprocal synapses, serial synapses, and convergent and divergent dyads.



The VS-3 neurons were detached from the spider cuticle and placed in a recording chamber. The preparation was continuously perfused by spider saline (in mM: 223 NaCl, 6.8 KCl, 8 CaCl₂, 5.1 MgCl₂, 1 glucose, and 10 HEPES), into which the GABA receptor agonists and cust_2.3.1 stgtL_1 t glucose; and 10 HEPES); into which the GABA receptor agonists and antagonists were ejected using pressure. Current- and voltage-clump recordings were performed with the discontinuous single-deturbance method using a SEC-101 amplifier (NPI Electronic; Tamm, Germany). The recording electrodes were filled with 3 M KC1 and the electrode resistance was 40-80 M/2m solution. The figure only shows one of the 14 VS3 neurons. In this preparation the dendrites are detached from the slits and the axons are cut about 200 µm from the somata.

GABA-RECEPTOR AGONIST AND ANTAGONIST EFFECTS ON THE VS-3 NEURONS



This experiment demonstrates that the vertebrate GABA_A receptor blocker, bicuculline did not inhibit the VS-3 neuron's response to GABA. Both recordings were done when the neuron was voltage-clamped to -100 mV.



Imidazole-4-acctic acid (I4AA) is a partial agonist of vertebrate GABA_A receptors and an antagonist of GABA_C receptors. When added to the perfusing solution of the VS-3 neurons it inhibited the GABA response (adove), but a rapid application of I4AA alone induced an inward current (right). Responses to I4AA were not consistent in all neurons, and we therefore assume that there may be more than one type of ionotropic GABA receptors on the VS-3 neurons. All recordings were performed when the neurons were voltage-clamped to -100 mV.





Muscimol, an agonist of ionotropic GABA receptors produced a similar inward current to GABA. The GABA or muscimol induced current was significantly reduced or completely blocked when the Cl² channel inhibitor picrotoxin was applied. The recordings shown here were performed when the neurons were voltage-clamped to 1-00 mV.



Application of low concentrations of the vertebrate GABAg receptor agonist baclofen (above) did not affect a spider VS-3 neuron. Middle trace shows the response of the same neuron to GABA application. However, when a high concentration of baclofen was applied to the bath solution it sometimes produced similar inward current as GABA (lower trace). This current may have been produced via an endogenous route. All recordings were performed when the neurons were voltage-clamped to -100 mV.

SUMMARY AND CONCLUSIONS

The peripherally located somata of the spider VS-3 mechanosensory neurons receive an extensive and complex synaptic innervation. Many of the efferent neurons are immunoreactive to GABA (Fabian-Fine et al. 1999) and here we show that they also respond to GABA applied via the bath solution. The major findings of this work were:

1. Bath application of GABA and the ionotropic GABA-receptor agonist muscimol induce an increase in membrane conductance and a depolarization similar to the primary afferent depolarization (PAD) that occurs in many mechanosensory afferent terminals. This depolarization inhibits the action potential discharge.

2. The inward current underlying the depolarization reverses at a potential of about -20 mV.

3. The GABA induced current is insensitive to the vertebrate GABAA receptor blocker bicuculline, but is inhibited by the chloride channel blocker picrotoxin.

4.14AA, which is an antagonist of vertebrate GABA_C receptors and a partial agonist of GABA_A receptors had both of these actions on the VS-3 neurons: It acted as an agonist when applied rapidly and as an antagonist when added to the perfusion.

5. Baclofen, a specific agonist of GABAB receptors, was only effective when applied at very high concentrations, and only in some of the neurons suggesting that this may

Taken together, these findings indicate that the VS-3 neuron somata have ionotropic GABA receptors that have more similarities to the vertebrate GABA_C than GABA_A receptors. However, based on the findings that the individual neurons responded to some agonists and antagonists in different ways, it is likely that there are several types of GABA receptors in the VS-3 neurons and their distribution may vary between neurons. When these receptors are activated by GABA it will result in a complete or partial inhibition of neuronal activity with an mechanism very similar to that described in the axon terminals of other mechanosensory neurons. Based on previous voltage-channels the cells from firing action potentials. In contrast to the presynaptic inhibition of zone terminals, which only inhibits the action of individual axonal branches, the GABA induced inhibition of the neuronals show of the prevents in the number and type of efferent neurons that are active and each neuron in a slit may be modulated separately, creating a very complex system for processing the mechanical information detected by the VS-3 organ.