Identification of tick periviscerokinin, the first neurohormone of Ixodidae: Single cell analysis by means of MALDI-TOF/TOF mass spectrometry

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Abstract

The first peptidergic neurohormone from the ticks Ixodes ricinus and Boophilus microplus has been identified by using a combination of immunocytochemistry and mass spectrometric analysis of single cells. The novel peptide (Ixori-PVK, PALIPFPRV-NH₂) shows a high sequence homology with members of the insect periviscerokinin/CAP2 b peptides that in insects are involved in the regulation of water balance. The function of this peptide in ticks is still unknown, but these pests consume large amounts of blood in a single blood meal which is a challenge for the regulation of diuretic processes. Thus, the novel peptide may be involved in one of the key physiological processes in ticks. High energy collision-induced dissociation was successfully used to distinguish between Leu/Ile ambiguities in single cell preparations. This is the first successful de novo sequencing of a peptide from single cell preparations of arthropods.

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Keywords: MALDI-TOF/TOF mass spectrometry; Neuropeptide; Periviscerokinin; Boophilus; Ixodes; Tick; Single cell analysis

Members of the family Ixodidae, commonly known as ticks, cause great economic losses of livestock; an estimated damage of several million dollars per year is reported for the United States [1]. As obligatory blood feeders, ticks consume large amounts of blood in a single blood meal which is a challenge for the regulation of diuretic processes. The necessity to remove excessive water after a blood meal has also been observed in a number of insects such as mosquitoes [2–4], blood-feeding bugs (e.g., Rhodnius prolixus) [5,6], fleas [7], and lice [8]. In the insects, diuresis is known to be regulated by several neuropeptides which influence secretion and/or reabsorption processes in the Malpighian tubules and the rectum [9]. Among these neuropeptides, are the insect kinins and periviscerokinin/CAP2 b-related peptides which may occur with multiple forms in a single species [10–14]. However, nothing is known about the hormonal control of excretion in ticks and no neuropeptides have been unambiguously identified from any tick so far. Recently, a receptor with sequence similarity to an insect kinin receptor (Drosophila melanogaster, CG 10626) [15] has been identified from the Southern Cattle Fever tick Boophilus microplus (CG10626) [16] and shown to functionally respond to various kinin peptides [17].

In this study, we searched for peptidergic neurohormones that may influence the water balance of ticks by screening the nervous system for peptide homologs to the insect kinins and periviscerokinin/CAP2 b-related peptides. Antisera against these neuropeptides revealed the expression of immunoreactive substances in a few neurons. We
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used the advantages of MALDI-TOF/TOF mass spectrometry to perform an analysis of those neurons that were found to contain immunoreactive material. This strategy resulted in the identification of a tick neuropeptide, which is a member of the periviscerokins, a peptide family that had been found only in insects so far. High energy collision-induced dissociation was used to distinguish between Leu/Ile in single cell preparations. As shown in our study, the novel peptide is expressed in another tick as well.

Materials and methods

Animals. Southern Cattle Fever ticks (B. microplus) were maintained at the Cattle Fever Tick Research Laboratory (USDA-ARS, Mission, TX), where all preparations on this pest were performed as well. The *Ixodes ricinus* was caught in Jena (Germany) by sweeping a net in grass fields. Only unfed adult ticks were used throughout the experiments.

Dissection and sample preparation for mass spectrometry. Isolated central nervous tissue of adult ticks was fixed with microneedles in a chamber containing insect saline. A number of putative neurosecretory neurons could be identified due to their slight bluish color (Tyndall effect). After selection of the neurons of interest, the ganglionic sheath was partially removed and the respective neurons were individualized. Without any enzyme treatment, cells were removed step by step using an uncoated glass capillary and transferred to a stainless steel sample plate for MALDI-TOF mass spectrometry. Subsequent sample preparation was performed as described in Neupert and Predel [18].

MALDI-TOF/TOF tandem mass spectrometry. MALDI analysis was performed on the ABI 4700 proteomics analyzer (Applied Biosystems, Framingham, MA). Due to the nature of the samples all acquisitions were taken in manual mode. Initially the instrument was operated in reflectron mode, in order to determine the parent masses. A laser energy of 3800 was typically employed for ionizing the neuropeptides. For the tandem MS experiments, the CID acceleration was 1 kV in all cases. The number of laser shots used to obtain a spectrum varied from 500 to 5000, depending on signal quality. In order to change the net amount of activation energy imparted to the primary ions, the collision gas (atmospheric air) pressure was increased. Three gas pressures were employed: none, 10 mTorr, and 50 mTorr. The fragmentation patterns from these three different settings were used to determine the sequence of the peptide. The fragmentation data obtained in these experiments was handled using the Data ExplorerT software package.

Immunocytochemistry. Dissected nervous system was fixed for 24 h at 4 °C with 4% formaldehyde in phosphate-buffered saline (PBS), pH 7.2. Subsequently, preparations were washed in PBS–4% Triton X-100 and PBS–1% Triton X-100 for 24 h, respectively. Then, the preparations were incubated for 3 days at 4 °C in anti-PeA-PVK-2 serum [19] (1:4000) diluted with PBS–1% Triton X-100 containing 0.25% bovine serum albumin and normal goat serum. Following washing in 0.1 mol/L Tris–HCl, 3% NaCl, and 1% Triton X-100 (pH 7.6) for 24 h, the Fluorochrome-labelled Cy3 (Cy3; Amersham, Braunschweig, Germany) secondary antibodies were directly used as a mixture in PBS–bovine serum albumin (2.5 mg/ml) at a concentration of 1:3000 for 3 days. The preparations were washed again for 24 h in 0.1 mol/L Tris–HCl, 3% NaCl, 1% Triton X-100 (pH 7.6), and then incubated at 4 °C in anti-leucokin-1 (DPAFNSWG) serum (1:4000) diluted with PBS–1% Triton X-100 containing 0.25% bovine serum albumin and normal goat serum. After 3 days, the samples were washed (24 h) in 0.1 mol/L Tris–HCl, 3% NaCl, and 1% Triton X-100 (pH 7.6), the Fluorochrome-labelled secondary antibodies Cy2 were directly used as a mixture in PBS–bovine serum albumin (2.5 mg/ml) at a concentration of 1:3000 for 3 days. Finally, the preparations were washed again 24 h in 0.1 mol/L Tris–HCl, 3% NaCl, and 1% Triton X-100 (pH 7.6) and transferred to glycerol. For visualization, tissues were dehydrated in ethanol, cleared in methyl salicylate, and mounted in entellan (Euromex microscopes Holland).

Results and discussion

The search for peptidergic hormones in arthropods as tiny as ticks is generally hampered by the small size of the central nervous system that contains low amounts of bioactive substances. In addition, working on ticks provides the researcher with further problems. First of all, and despite their great economic importance, not a single peptide hormone is known from ticks and bioassays to perform conventional neuropeptides purification strategies are not available. This paucity of information was confirmed by a current search for tick neuropeptides in GenBank that only retrieved sequences for two putative salivary secreted neuropeptide-like proteins from the tick *Ixodes pacificus*. Second, the synganglionic tick nervous system does not exhibit well-separated neurohemal release sites as observed in insects, where neurohormones are accumulated and easily accessible for structural elucidation [12]. Having in mind the positive experiences with the analysis of single identified neurons in an insect [18], we decided to perform the following strategy for the identification of putative diuretic hormones. Antisera raised against insect diuretic peptides, namely leucokin-1 [21] and Pea-periviscerokinin-2 [22], were used to locate neurons in the CNS of *I. ricinus* which may express peptides that are sequence related to the insect kinins and periviscerokinins. Indeed, this procedure yielded several immunoreactive cells, a majority of these neurons were immunostained by both antisera (Fig. 1). Due to their slightly bluish color (Tyndall effect), we succeeded in detecting a number of these neurosecretory neurons in...
the untreated central nervous system of *I. ricinus*. The most distinctive somata (T1) were found in the cheliceral ganglion where four neurons are immunostained on each side. Two of these neurons (20–25 μm) showed exclusively leucokinin-immunoreactivity and developed extensive and segmentally arrayed arborizations (Fig. 2), resembling the histamine-immunoreactive projections in spiders [23]. In the next step, these large somata were individually dissected and directly prepared for mass spectrometric analysis. Resulting mass spectra showed different ion signals in the mass range of 600–3000 Da with the 1395.69 Da signal cokin-immunoreactivity and developed extensive and segmentally arrayed arborizations (Fig. 2), resembling the histamine-immunoreactive projections in spiders [23].

![Fig. 2](image1.png)

**Fig. 2.** Microphotograph of leucokinin-like immunoreactivity in the synganglion of *I. ricinus* with extensive and segmentally arrayed arborizations that resemble the histamine-immunoreactive projection in spiders (black arrows). Large anterior neurons (T1) were first isolated and analysed by mass spectrometry (white arrows). Scale bar: 50 μm.

![Fig. 1](image2.png)

**Fig. 1.** Double-immunofluorescence staining in whole-mounts of the nervous system from the European hard tick *I. ricinus* by means of an anti-PVK-2 serum (red fluorescence) and an anti-leucokinin-1 serum (green fluorescence). Five cell groups are immunoreactive (T1–T6), four of these groups (T2–T6) and nearly the total network of arborizations throughout the synganglion are colocalized. Two neurons of the cheliceral ganglion (T1: 20–25 μm) contain only leucokinin-immunoreactivity. CG, cheliceral ganglion; OP, opisthosomal ganglion. Scale bar: 50 μm. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this paper.)

![Fig. 3](image3.png)

**Fig. 3.** MALDI-TOF mass spectrum of a T1 neuron from the cheliceral ganglion (CG) of the synganglion from *I. ricinus*. The most abundant ion signal at 1395 Da, as well as further less prominent ions were subjected to fragmentation but did not yield any fragments. Thus, the substances in these T1 neurons are either not fragmenting well or not peptides at all.

![Fig. 4](image4.png)

**Fig. 4.** Representative MALDI-TOF mass spectra of identified homolog cells (T4/T5) (*Boophilus*: cell group) from (A) central European hard tick *I. ricinus* L. and (B) southern cattle tick *B. microplus* C. In both spectra, a mass peak at [M+H]+: 1008.63 Da (arrow) was observed which was subsequently fragmented. Remaining, ion signals differed somewhat between the species.
being the most abundant one (Fig. 3). However, no fragment could be obtained from any of these substances, even under conditions of collision-induced dissociation (CID). Control experiments with identified insect neurons verified that the overall signal intensity of the parent ions was sufficient to lead to a number of fragment ions (not shown).

Thus, the observed substances in these leucokinin-immuno-reactive neurons either do not fragment well or are not peptides at all. Subsequently, neurons of the opisthosomal ganglion (Fig. 1: T5/T6; 10–15 μm) were analyzed using the same technique as described above. Some of these preparations yielded mass spectra with a distinctive ion signal at 1008.63 Da (Fig. 4A). This substance was chosen for fragmentation and produced clear fragments under conditions of low energy fragmentation as well as high energy CID (Fig. 5). The following sequence could be reconstructed from the calculation of mass differences between successive fragments of the different ion series: PA(L/I)(L/I)PFPRV-....

### Table 1

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Using the information obtainable from the CID spectra, even the assignment of the L/I ambiguities was possible [14,24] (Fig. 5, inset). The novel peptide is clearly related to the insect periviscerokinins (see Table 1) and is designated as I. ricinus periviscerokinin (Ixori-PVK). This is the first successful de novo sequencing of a peptide from single cell preparations of arthropods, and particularly the correct assignment of Leu/Ile from such a tiny sample is an excellent verification of the opportunities associated with modern mass spectrometry. Fragmentations of synthetic Ixori-PVK yielded spectra that are nearly identical with those of native PVKs from single cell preparations.

Homologue cells of another tick species, B. microplus, were subjected to mass spectrometric analysis (Figs. 4B and C). B. microplus, which transmits several diseases known collectively as ‘tick fever,’ is a very small tick (roughly 1–2 mm body length). Nevertheless, mass spectra taken from the respective neurosecretory neurons in the opisthosomal ganglion verified the expression of Ixori-PVK in this species as well (not shown).

A screening of the anterior leucokinin-immunoreactive cells of the chelicer ganglion of B. microplus also revealed the abundant ion signal at 1395.69 Da, which altogether indicates the close phylogenetic relationship between the studied tick species.

In summary, the single cell approach, as described in this study, allowed the elucidation of the first peptidergic neurohormone from arachnids. The origin of another peptide recently isolated from 10,000 synganglia of Amblyomma testudinarium [25] awaits confirmation since it is very closely related to mammalian hemorphins and may result from contamination due to the impossibility of the complete removal of the esophagus from the synganglion. Ixori-PVK belongs to the periviscerokinin/CAP₂ peptide family, which was previously known only in insects. The effects of periviscerokinins in water balance are well established [26] and it thus seems to be possible that the tick PVK may also regulate diuresis in these blood-feeding arthropods. This function, however, has yet to be established. In the future, the sequence information may aid in the development of mimetic analogs capable of disrupting such critical physiological processes in ticks.
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References


