The Nervous Systems of Basally Branching Nemertea (Palaeonemertea)

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Abstract

In recent years, a lot of studies have been published dealing with the anatomy of the nervous system in different spiralian species. The only nemertean species investigated in this context probably shows derived characters and thus the conditions found there are not useful in inferring the relationship between nemerteans and other spiralian taxa. Ingroup relationships within Nemertea are still unclear, but there is some agreement that the palaeonemertean species form a basal, paraphyletic grade. Thus, palaeonemertean species are likely the most informative when comparing with other invertebrate groups. We therefore analyzed the nervous system of several palaeonemertean species by combining histology and immunostaining. 3D reconstructions based on the aligned slices were performed to get an overall impression of the central nervous system, and immunohistochemistry was chosen to reveal fine structures and to be able to compare the data with recently published results. The insights presented here permit a first attempt to reconstruct the primary organization of the nemertean nervous system. This comparative analysis allows substantiating homology hypotheses for nerves of the peripheral nervous system. This study also provides evidence that the nemertean brain primarily consists of two lobes connected by a strong ventral commissure and one to several dorsal commissures. During nemertean evolution, the brain underwent continuous compartmentalization into a pair of dorsal and ventral lobes interconnected by commissures and lateral tracts. Given that this conclusion can be corroborated by cladistic analyses, nemerteans should share a common ancestor with spiralian lineages; probably with a simple brain consisting of paired medullary, frontally commissurized and reinforced cords. Such an organization resembles the situation found in presumably basally branching annelids or mollusks.

Introduction

Nemerteans are a spiralian taxon, of which most described species are marine benthic hunters. The majority of the animals are nocturnal predators able to follow their prey [1], [2], [3] and catch it with a unique structure, the eversible proboscis. Due to their lifestyle, nemerteans possess well-developed nervous systems and sensory structures, enabling them to interact with their environments. Since nearly all nemerteans are predators, their central nervous system has been considered quite uniform, composed of a four-lobed brain and two lateral medullary cords which originate from the ventral lobes of the brain and extend the whole length of the animal until they converge at the posterior tip of the animal [4], [5]. The paired dorsal and ventral lobes of the brain are interconnected by a commissural tract above or below the rhynchocoel. The central nervous system is composed of a central neuropil that is homogeneously surrounded by neuronal cell somata. The neuronal cell somata may be separated by an extracellular matrix, the inner neurilemma, from the neuropil and/or the entire brain may be enclosed by an outer neurilemma [4], [6], [7]. A set of minor nerves, such as the cephalic, esophageal and proboscidial nerves originate from different parts of the brain, while additional ventral and dorsal nerves are only found in certain groups. These minor nerves are not covered by neuronal cell somata (perikarya) and belong to the peripheral nervous system.

Nemerteans possess a number of sensory organs, which are mostly located on the tip of the head. These organs are called the frontal and cerebral organs and are supposed to be involved in chemoreception [8], [9], [10], [11], [12]. Photoreception is perceived through pigmented eyes, of which up to hundred may be present in one animal [5]. Adult palaeonemerteans generally lack eyes although larval eyes have been reported for two species (e. g. Procephalothrix simulus [12] and Carinoma tremaphoros [13]).

Formerly, nemerteans were classified in three higher taxa: Palaeo-, Hetero- and Hoplonemertea [4], [15]. Palaeonemerteans were regarded as the basal most branching taxon of nemerteans [15], [16] [17]. Traditionally, four different taxa belonged to this group, the Tubulanidae, Cephalothricidae, Carinomidae and Hubrechtidae [4], [5], [14]. New evidence however, suggests that Hubrechtidae are the sister group of Heteronemertea, because they possess a pilidium larvae and a cerebral organ which is typical for heteronemerteans [16].

Recent molecular studies provide some evidence that the remaining palaeonemerteans could be a monophyletic group [17]. Morphological characters supporting this result, however, are missing. Cephalothricidae have been suggested as the basal most
branching nemertean group due to the absence of cerebral organs and the position of the nervous system inside the musculature [18]. Thus, palaeonemerteans form a basal grade rather than a basally branching clade of nemerteans, a conclusion that has also been drawn by earlier morphologists [4] and molecular studies [15], [16], [17].

The palaeonemertean nervous system shows the highest variation among nemertean taxa [4], [5] and potentially provides a rich source of characters. In recent years, neuroanatomy, by means of immunohistochemistry, has yielded promising data that contributes to unraveling the phylogeny of certain metazoan groups, e.g [19], [20], [21], [22], [23]. The only representative of nemerteans investigated with this method so far, Lineus viridis [24], probably shows derived characters and data concerning basal branching nemerteans is still lacking. Studies on the phylogeny of Nemertea and the relationship they share with other lophotrochozoans need to focus particularly on basally branching nemertean taxa. Therefore, we started a comparative analysis of the nemertean nervous system [25]. In this study, the first of a series of papers, we are presenting detail information on the neuroanatomy of different species of the palaeonemerteans.

Figure 1. Schematic drawings of the nemertean central nervous system. a: Dorsal view of the central nervous system, the neuronal cell somata are shown in bright blue. b: Transverse section of the brain. The brain is composed of a central neuropil (gray), which is surrounded by neuronal cell somata (bright blue). The neuronal cell somata may be separated from the neuropil by an inner neurilemma. The whole brain may be enclosed by an outer neurilemma. The two halves of the brain are interconnected by a dorsal (dct) and a ventral commissural tract (vct) which form a ring around the rhynchocoel (ry), dl dorsal lobe of brain, pr proboscis, vl ventral lobe of brain.

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Additional slices that were used to generate 3D reconstructions of the whole central nervous system are made freely available in MorphDBase, a morphological database [26] (www.morphdbase.de), in terms of data transparency. The respective hyperlink is given at the beginning of each description. Immunochemistry was chosen to reveal the fine structure of the nervous system. Antibody subsets representing standard markers that are used by many researchers and allow comparability of the morphological elements are also chosen here. Given that palaeonemerteans are likely not monophyletic and comprise an assemblage of basally branching nemerteans, some hypotheses on the primary organization of the nemertean nervous system can be derived and evaluated with respect to certain outgroups.

Results
General remarks
The neuroanatomical nomenclature used in this paper largely follows the terminology provided in the review by Richter et al. [21]. In order to define the specific terms of nemertean neuroanatomy, some general remarks will be given prior to
describing the nervous system in each of the different species studied (figs. 1 & 2). The main components of the nemertean nervous system are the central nervous system consisting of a brain and nerve cords, the peripheral nervous system that includes nerves and plexus, and finally the sensory structures. While the central nervous system is characterized by a clear separation between neurites (neuropil) and their perikarya (somata), nerves are bundles of neurites with a few nuclei interspersed and are surrounded by extracellular matrix (ecm). Plexus are meshworks of neurites, and sensory organs are specialized, larger aggregations of several sensory cells.

Central nervous system

Brain. An outer neurilemma of ecm may surround the entire central nervous system. The somata are located peripherally and externally to the central neuropil; somata and neuropil may be separated by an inner neurilemma consisting of ecm (fig. 1b). The brain is always bilaterally symmetrical and may contain subunits, or compartments like lobes which connect contralaterally by commissural tracts and ipsilaterally by lateral tracts.

Nerve cords. Nerve cords are locally-restricted, longitudinal strands of neuropil surrounded by the somata of their neurons. A pair of medullary cords is ventro-laterally located. They extend throughout the body and originate from the posterior ventrolateral margin of the brain. The distribution of the somata relative to the medullary cord neuropil varies among the taxa (fig. 2b). Cephalic nerve cords originate from the anterior margin of the brain.

Neurons. The somata are grouped peripherally to the neuropil and differ in their morphology and staining affinities. According to the structure of their perikarya (somata), four different size classes have been established [4], [27]. Somata of class 1 neurons (S1) are small and often have a variable shape, while somata of class 2 neurons (S2) are spherical to pear-shaped and are twice as large as S1, and somata of class 3 neurons (S3) are spherical to drop-shaped with large nuclei, a prominent nucleolus and are twice as large as S2. Class 4 cells are known as neurochord cells, but it is unknown whether these are neurons at all.

Peripheral nervous system and sensory structures

The nemertean peripheral nervous system consists of several minor nerves, such as the cephalic nerves, the proboscis nerves, dorsal and ventral nerves, esophageal nerves and nerve plexus (figs. 1 & 2).

Nerves. The cephalic nerves originate at the anterior margin of the brain and run towards the anterior tip of the animal. The ventral nerve is a single, median nerve that originates from the posterior margin of the ventral commissural tract and initially runs ventrally to the rhynchocoel and then more posteriorly, ventral to the gut. The dorsal nerve is a single, median nerve located dorsal to the rhynchocoel. It originates from the posterior margin of the dorsal commissural tract and runs posteriorly. A second dorsal nerve may be located ventral to the dorsal nerve. It originates from the dorsal nerve and runs posteriorly, directly above the rhynchocoel. Bundles of neurites regularly connect both nerves. A pair of esophageal nerves originates from the caudal margin of the ventral commissural tract or from the ventral lobes. Both nerves are commissurized anterior to the mouth opening, encircle the foregut, turn ventro-laterally and run posteriorly on either side.
of the gut (figs. 1a, 2b). In basal nemerteans the proboscis nerves originate from the dorsal margin of the ventral commissural tract. A dorsal nerve (dn) originates from the dorsal commissural tract (dct), a ventral nerve (vn) from the ventral commissural tract. The branching esophageal nerves (en) originate from the ventral nerve and surround the mouth opening (mo). The lateral medullary cords (mc) originate in the ventral lobes of the brain (br). Letters on the right (a–f) refer to the histological sections in figure 5. The nerve plexus originating in the dorsal nerve (see fig. 5e) is omitted.

Sensory structures. Nemerteans possess several anterior sensory organs, such as the frontal organ, eyes, and cerebral organs (fig. 1a). The eyes are visible by their black pigment, whereas cerebral and frontal organs are defined by their morphology and position. Frontal organs consist of one to three densely ciliated pits located on the anterior tip of the head. Cerebral organs are tubular epidermal invaginations with distinct morphological differences among the lining cells and a close nervous connection to the brain. If present, they are always posterior to the brain in palaeonemertean species. Beside the sensory organs, isolated sensory cells are distributed across the entire body surface.

Procephalothrix filiformis and Cephalothrix linearis (Cephalothricidae)

Cephalothricid species are characterized by a long and bluntly pointed head [5] that is termed rostrum. Their brain is located in the rostrum of the animals, far anterior to the mouth opening (figs. 3a, b; 4), and embedded in the head musculature. The nervous system in the two species studied is almost identically organized (see https://www.morphdbase.de?P_Beckers_20130201-M-15.1 for Procephalothrix filiformis stack and https://www.morphdbase.de?P_Beckers_20130201-M-14.1 Figure 4. Procephalothrix filiformis, schematic drawings of the central nervous system based on 3D-reconstruction of 501 aligned 0.5 µm sections, dorsal (a) and lateral (b) view. a: The nervous system is composed of neuropil (np, gray) which may be surrounded by cell somata (cs, blue). Cephalic cords (cc) are circular arranged around the head of the animals. The paired proboscisial nerves (pn, yellow) originate from the ventral commissural tract (vct). A dorsal nerve (dn) originates from the dorsal commissural tract (dct), a ventral nerve (vn) from the ventral commissural tract. The branching esophageal nerves (en) originate from the ventral nerve and surround the mouth opening (mo). The lateral medullary cords (mc) originate in the ventral lobes of the brain (br). Letters on the right (a–f) refer to the histological sections in figure 5. The nerve plexus originating in the dorsal nerve (see fig. 5e) is omitted. b: Central nervous system lateral view. The dorsal lobe (dl) of the brain and the ventral lobe (vl) of the brain are connected by several lateral tracts (lt).

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for *Cephalothrix linearis* stack. For this reason immunhistological methods were only applied to *P. filiformis*.

Central nervous system

**Brain.** The central part of the brain neuropil is separated from the neuronal somata by an inner neurilemma consisting of extracellular matrix (ecm), which was indicated by blue coloration in the Azan staining [figs. 5d, 6a; [7b], [8a]]. An outer neurilemma is not visible. The brain consists of four distinct, lobe-like compartments that anteriorly are juxtaposed to the head lacuna [figs. 5c,d; 7b,d]. The dorsal lobes are located several micrometers anterior to the ventral lobes. A small dorsal commissural tract connects both dorsal lobes [fig. 5c]. It originates at the anterior-dorsal margin of each lobe, next to the origin of the dorsal cephalic nerve cord. The dorsal lobes are more voluminous than the ventral ones. Each dorsal lobe is bean-shaped, and tapers posteriorly to form a small process that ends in a layer of neuronal cell somata. On each side of the brain, a series of at least five lateral tracts connects the dorsal lobe to the ventral lobe. When numbered along the anterior-posterior axis, the third lateral tract is the broadest of these tracts. The ventral lobes are connected by a prominent ventral commissural tract, which is located posterior to the dorsal commissural tract and originates at the anterior-ventral margin of each ventral lobe, next to the insertion of the first lateral tract [fig. 9b]. The neurites from each commissural tract are horizontally arranged. Each ventral lobe also tapers posteriorly and is confluent with the medullary cords [figs. 5e; 9a, 7d].

**Medullary cords.** The medullary cords are ventro-laterally located and embedded into the longitudinal muscle layer. They travel along the animal to merge at its posterior end [figs. 9d]. The

Figure 5. *Procephalothrix filiformis*, light micrographs of Azan stained transverse sections of the brain. **a:** Frontal part showing the huge layer of neuronal cell somata (cs). **b:** Four cephalic cords (cc) extend towards the tip of the head. **c:** A dorsal commissural tract (dct) connects the two halves of the brain (br). The neuronal cell somata are separated from the neuropil of the brain by an inner neurilemma (in). The brain is embedded in a layer of longitudinal muscles (lm), which lie underneath a layer of circular muscles (cm). **d:** The brain (br) is divided into dorsal (dl) and ventral (vl) lobes in its posterior part and composed of a central neuropil (np) which is surrounded by cell somata (cs). The somata are separated from the neuropil by an inner neurilemma (in), and two proboscisial nerves (pn) are present. **e:** A dorsal nerve (dn) arises from the dorsal commissural tract inside the musculature and migrates dorsally, a ventral nerve (vn) from the ventral commissural tract. Neurites (ne) of the dorsal nerve extends as a roof-like structure around the rhynchocoel. mc: medullary cord. **f:** The lateral medullary cords (mc) are embedded into the longitudinal muscle layer, and the dorsal nerve (dn) is located subepidermally. bv: blood vessel, fg: foregut, vn: ventral nerve.

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neuropil of the medullary cords is capped by two crescent-shaped layers of neuronal somata, which form a dorsal and a ventral cap in transverse sections (fig. 5f; 7d). The neuronal cell somata are separated from the neuropil by an inner neurilemma. Some neurites, however, leave the neuropil and arborize between the cell somata (fig. 5e). The medullary cords are interconnected by serially arranged small circular neurites that show immunoreactivity against an antiserum directed against serotonin in *P. filiformis* (fig. 9e).

**Cephalic nerve cords.** In *P. filiformis* and *C. linearis*, a massive layer of neuronal somata is located frontal to the brain, underneath the outer circular and the inner longitudinal muscle layers (figs. 5a). The somata surround four strands of neuropil, and are separated by an inner neurilemma (fig. 5b, 7a). In the literature they have been termed cephalic nerves [5]. According to the definitions above, however, these four strands are part of the central nervous system and must be termed cephalic nerve cords. These four nerve cords are arranged in pairs, a dorsal and ventral one, which are so tightly apposed to the head lacuna that their lumen seems to be compressed (fig. 5a). The dorsal cords are anteriorly bifurcated, while the ventral cords simply taper anteriorly. The dorsal cephalic nerve cords originate from the antero-dorsal margin of the dorsal lobes, directly underneath the origin of the dorsal commissural tract. The ventral cephalic nerve cords originate from the antero-ventral margin of the dorsal lobes, anterior to the first lateral tract that interconnects the dorsal and ventral lobes. The somata of the cephalic nerve cords in *P. filiformis* barely show any immunoreactivity against an antiserum directed against FMRFamid or serotonin (figs. 9a, b).

**Neurons.** In both cephalothricid species, the neurons can be classified according to their size, as proposed by Bürger [4], with class 2 neurons having somata and nuclei that are twice as large as class 1 neurons and somata and nuclei are half as large as class 3 neurons. All differ in structure, shape and Azan staining ability (figs. 6a–c). In *P. filiformis* cell bodies of class 1 neurons are slender and beaked with orange staining cell bodies and small, red nuclei that appear condensed (diameter: 0.92±0.15 μm, n = 12). These neurons are rare, but evenly distributed all over the brain (fig. 6a). Somata and nuclei of class 2 neurons are globular, having cell bodies almost twice as large as that of class 1 neurons (fig. 6b). Azan stains the cell body and nuclei of class 2 neurons bright to grayish orange. Class 2 neurons are the most prevalent cell type throughout the brain and the entire central nervous system (diameter: 1.85±0.21 μm, n = 11). Class 3 somata are globular, having a grayish Azan coloration, while the nuclei stain dark orange (diameter: 4.05±0.67 μm, n = 13). Class 3 neurons are evenly distributed throughout the brain, although they are less frequent than class 2 neurons (fig. 6c). All size classes differ significantly in diameter (p<0.001).

In *C. linearis*, class 1 neurons are small and have nuclei stained red (figs. 8a–c). The greatest accumulation of class 1 neurons is found in the cephalic nerve cords, although they are also found rarely but regularly distributed all over the brain (fig. 8a) (diameter: 2.86±0.41 μm, n = 10). Cell bodies of class 2 neurons are drop-shaped, stain bright purple with darker nuclei (figs. 8b, c),

**Figure 6. Procephalothrix filiformis, light micrographs of Azan stained neurons. a:** Transverse section, overview of the brain, position of the three different types of neurons (S1–S3). **b:** Higher magnification of somata of type 1 and 2 neurons (S1, S2). Nuclei of S1 dye orange and the cell bodies are beak shaped. Cell bodies of S2 are circular and enlarged. Nuclei dye orange. **c:** The nuclei of cell somata type 3 (S3) also stain orange. The cell bodies are circular but more prominent than in S1–S2. in inner neurilemma, np neuropil.

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and are found primarily in the posterior ventral part of the brain, arranged in clusters (diameter: 5.38 ± 0.86 μm, n = 11). The nucleus of class 3 neurons shows the same bright purple coloration as its soma (fig. 8c), but contains a large, red nucleolus (diameter: 10.68 ± 1.49 μm, n = 13). Class 3 neurons are found in the middle of the brain, just behind the ventral commissural tract. In *P. filiformis* the majority of neurons belong to type 2 neurons, whereas in *C. linearis*, most neurons belong to type 1 and type 2. In the posterior part of the medullary cords, the layer of neuronal somata decreases in thickness.

Although Azan staining reveals an enormous layer of cell somata in the brain, staining *P. filiformis* with an antiserum against FMRFamid marks only very few cells (figs. 9a, b). The neuropil is not only restricted to the central part of the brain (clearly separated by *ecm* from the somata), but is also branching into the neuronal cell somata layer (fig. 5d).

**Minor nerves and peripheral nervous system**

In the *P. filiformis* head, four minor nerves show immunoreactivity against an antiserum directed against FMRFamid and serotonin. These nerves extend from the tip of the head to the posterior end of the animal (figs. 9c; 10a). A circular commissural nerve connects these nerves (fig. 9c). In both cephalothricid species, a single **dorsal nerve** originates from the posterior margin of the dorsal commissural tract and runs posteriorly within the outer longitudinal muscle layer. More posteriorly, it passes the outer circular muscle layer to run underneath the basal lamina of the epidermis. The dorsal nerve then merges into the connection between the lateral medullary cords at the posterior end of the animal (figs. 5d–e; 7d). The **ventral nerve** has two roots, which branch off where the ventral lobes are connected by the ventral commissural tract and run medially for a few micrometers before fusing and running posteriorly ventral to the rhynchocoel. A few micrometers in front of the mouth, this nerve splits into two branches that surround the forae gut. Both branches fuse behind the mouth and run posteriorly ventral to the gut as a single ventral nerve (figs. 4a; 5e; 9a). **Esophageal nerves** are lacking in both cephalothricid species. In *P. filiformis* staining against an antiserum directed against FMRFamid reveals two additional minor nerves, which originate behind the mouth (fig. 9a) and travel to the posterior end of the animal. In both species, two **proboscial nerves** arise from the posterior margin of the ventral commissural tract (fig. 4). The nerves run posteriorly on either side of the proboscis and are occasionally joined by a proboscial plexus. In *P. filiformis*, the longitudinal proboscis nerves show immunoreactivity with an antiserum directed against FMRFamid, but not against serotonin (figs. 5d; 9c, e).

In *P. filiformis* a bright gray coloration in Azan stained sections clearly indicates a **nervous plexus** that arises from the dorsal nerve and forms a roof-like structure around the rhynchocoel (fig. 5c). Immunoreactivity with an antiserum directed against FMRFamid provides strong evidence that the neurites of the intraepidermal plexus are arranged in a regular, grid-like way (fig. 10b). A series of circular neurites connect the medullary cords (fig. 9c). The arrangement of the neurites of other plexus could not be reconstructed due to a lack of signal.
Sensory structures

Although adult *P. filiformis* react strongly to changes in the illumination (photonegative) (unpubl. observations), there are no pigmented photosensory structures present. In the very tip of the head, a cluster of cell shows immunoreactivity with an antiserum directed against FMRFamid (fig. 10c). There are many cells distributed inside the epidermis and all over the body, which show immunoreactivity with an antiserum directed against FMRFamid and serotonin. The cells are bottle-shaped and may represent sensory cells (fig. 9d). A combination of immunostaining and electron microscopy should be employed to test this. Neither a frontal organ nor cerebral organs are present in either cephalothricid species.

**Carinoma mutabilis** (*Carinomidae, “Palaeonemertea”*)

The nervous system of this species is located inside the longitudinal muscle layer; the brain lies anterior to the mouth; (see: https://www.morphdbase.de/?P_Beckers_20130201-M-10.1 for *C. mutabilis* stack).

Central nervous system

**Brain.** The entire brain is surrounded by an outer neurilemma; an inner neurilemma is not present (figs. 11, 12b, c; 13a). The central neuromere of the brain is surrounded by a layer of neuronal cell somata (figs. 12b, c; 13a). The *Carinoma mutabilis* brain is situated anteriorly in front of the mouth opening. Like in the cephalothricid species, it is surrounded by the head musculature. The brain is not externally visible. It consists of paired dorsal and ventral lobes (fig. 12c). Three commissural tracts connect the dorsal lobes, the posterior two being small, closely associated and widely separated from the third prominent anterior dorsal commissural tract. The lateral portion of the anterior dorsal commissural tract and the anterior margin of the dorsal lobe give rise to cephalic nerves (figs. 11; 12a). The dorsal- and ventralmost of these nerves branch off to give rise to further cephalic nerves until a complete ring of more than thirty cephalic nerves is formed. They are located within the head musculature. The dorsal commissural tract also gives rise to a pair of posteriorly directed dorso-lateral nerves. On either side, they turn latero-ventrally and merge with the dorsal lobe where the second dorsal commissural tract originates (fig. 11). Directly posterior the third commissural tract, a horn-like extension projects from the inner margin of the dorsal lobe. Posterior to this structure, the dorsal lobe tapers to form a small process that ends within a cup of neuronal cell soma. The dorsal lobes of the brain are more voluminous than the ventral ones and are located anterior to them. The ventral lobe
are located slightly ventro-median to them and slightly oblique to the main body axis. On either side the ventral lobe is broadly connected to the dorsal one, but several small bundles of ECM and occasionally muscle fibers, pierce this connection and cause the impression of several closely packed lateral tracts connecting both lobes (fig. 12b). Both ventral lobes are connected by a broad ventral commissural tract located between the origin of the first and the second dorsal commissural tract (fig. 12b). The neurites of the ventral commissural tract are horizontally arranged.

**Medullary cords.** The ventral lobes taper caudally and give rise to the lateral medullary cords. The medullary cords run posteriorly between the outer longitudinal muscle layer and the inner circular muscle layer. The neuronal cell somata of the medullary cords are located dorsally and ventrally to the neuropil (fig. 12d). The location of the medullary cords was observable until they entered the gonad region.

**Neurons.** The somata of the neurons fall into two size classes, with somata and nuclei of class 2 neurons being twice as large as those of class 1 neurons (figs. 13a, b). As the nuclei of class 1 neurons stain red, and a cytoplasmic surrounding is hard to detect, the cell somata of class 1 neurons must be very small (fig. 13b) (diameter: 1.75 ± 0.32 μm, n = 12). These neurons are distributed all over the body and the medullary cords (mc) are connected by serially arranged circular nerves (scn). Note that the proboscis nerves show no immunoreactivity against serotonin. pr proboscis.

Figure 9. *Procephalothrix filiformis*, confocal laserscanning (cLSM) micrographs of differently immunostained whole mounts. a: Anti-FMRF. The cephalic cords (cc) originate in the lateral aspects of the brain (br). The lateral medullary cords (mc) extend the full length of the animal. There are two protocerebral nerves (pn) that run opposed to each other, along both sides of the proboscis. The esophageal nerves (en) originate at the ventral nerve (vn) and branch shortly before the mouth opening (mo). b: Anti-FMRF. Only few neuronal cell somata (cs) of the brain (br) are immunoreactive against FMRF. c: Anti-FMRF. Four minor nerves (arrow) unite in the tip of the animals’ head (arrowhead). The nerves are interconnected by a circular nerve (asterisk) d: Anti-FMRF. The dorsal nerve (dn) is connected to the medullary cords (mc) in the very posterior part of the animal; bottle shaped sensory cells (sc) are distributed all over the body. e: Anti serotonin. The medullary cords (mc) are connected by serially arranged circular nerves (scn). Note that the proboscis nerves show no immunoreactivity against serotonin. pr proboscis.

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A third type of cell is restricted to the dorso-median region of the dorsal lobe between the first and the second dorsal commissural tracts. There is also a ventro-median patch of these cells at the posterior margin of the ventral commissural tract and the origin of the esophageal nerves. The bodies of these cells are at least ten times more voluminous as those of the neurons, and contain nuclei at least four times larger. These cells possess a very prominent orange staining nucleolus. Dorsally these cells are in close contact with blood vessels, so they may represent glandular cells rather than neurons (figs. 13a, c).

Minor nerves and peripheral nervous system

The cephalic nerves form a complete ring of longitudinal neurites with occasionally interspersed nuclei. Close to their origin, these nerves are interconnected, but are increasingly isolated from each other by an ecm as they run anteriorly. More than 35 of these nerves can be seen in transverse sections of the head region (fig. 12a). A dorsal nerve originates from the first dorsal commissural tract. Anteriorly the dorsal nerve is located in the musculature, but turns dorsally to extend beneath the epidermal basal lamina to the posterior end of the animal (figs. 12b–d). Very fine rings of neurites occasionally connect the dorsal nerve and the medullary cords. A ventral nerve is not present. The esophageal nerves arise from the posterior margin of the ventral commissural tract, turn medially and merge a few micrometers in front of the mouth. Posterior to the point of merging they bifurcate and surround the mouth on either side (fig. 12c). Posterior to the mouth, both nerves have two branches, all six of theses running parallel to the esophagus. The paired proboscical nerves originate from the antero-dorsal margin of the ventral commissural tract and run posteriorly on either side of the proboscis (fig. 12d). Both are occasionally connected by a circular bundle of neurites.

Sensory structures

No sensory structures such as eyes or cerebral organs are found in C. mutabilis. At the tip of the head, a medio-dorsal series of pits and slits with prominent sensory cilia and underlying neurons may represent the frontal organ. The regionally restricted large cells found in the dorso-median portion of the brain (figs. 13a, c) may represent glandular cells and might have a neurosecretory function.

Tubulanus superbus and Tubulanus polymorphus (Tubulanidae, “Palaeonemertea”)

Tubulanus species are characterized by a sharply demarcated head [5], (figs. 3c, d). The anterior nervous systems of Tubulanus superbus and Tubulanus polymorphus is almost identically organized, although both species differ in size (see: https://www.morphdbase.de?P_Beckers_20130201-M-9.1 for Tubulanus superbus stack, https://www.morphdbase.de?P_Beckers_20130201-M-11.1 for Tubulanus polymorphus stack).
For this reason immunohistological methods are only shown for *T. polymorphus*.

**Central nervous system**

**Brain.** The brain of both *Tubulanus* species is composed of a central neuropil, which is surrounded by neuronal cell somata; a prominent outer neurilemma is present, while an inner neurilemma is missing (figs. 14b, 15a). In both species the brain is located ventro-laterally and anterior to the mouth, but is not externally visible (figs. 3d, e). The brain consists of four lobes that are located directly underneath the epidermal basal lamina (figs. 14a, 15b). The dorsal lobes are larger than the ventral lobes. They are connected by a series of dorsal commissural tracts, of which the anteriormost and the posteriormost are larger than the remaining ones (fig. 16). On either side of the brain, the dorsal and ventral lobes are connected by a continuous layer of neurites, which has a slightly smaller diameter than either of the lobes, making the lobes hard to distinguish (figs. 17b, 18a). Each connection represents a single broad lateral tract. It is, however, restricted to the region anterior to the posteriormost dorsal commissural tract. Each dorsal lobe has a posterior extension that does not connect to the ventral part of the brain and terminates in a layer of neuronal somata. Both ventral lobes are connected by a broad and prominent ventral commissural tract that lies posterior to the anteriormost dorsal commissures (figs. 14b, 16, 17b). The neurites of the ventral commissural tract are horizontally arranged (fig. 18a).

**Medullary cords.** Each ventral lobe is confluent with a lateral medullary cord, which runs posteriorly between the subepidermal basal lamina and the outer circular muscle layer (figs. 14d, 17d). In *T. superbus*, the neuronal somata of the medullary cords form an inverted “C” to cover the lateral, dorsal and ventral aspect of the neuropil (fig. 14d). Bundles of neurites branch into the layer of somata. In *T. polymorphus*, the neuronal somata of the medullary cords form two caps, which dorsally and ventrally cover the neuropil (fig. 17d). In this species, bundles of neurites branch into the layer of somata like in *T. superbus*.

**Neurons.** In both species, the neurons of the central nervous system fall into three size classes (figs. 15, 18). Like the cephalothricid species, somata and nuclei of class 2 neurons are twice as large as those of class 1 neurons and half as large as those of class 3 neurons. In *T. superbus*, the nuclei of class 1 neurons stain dark purple, are arranged in clusters and have small, elongated cell bodies (figs. 15a, b) (diameter: 2.90 ± 0.44 μm, n = 14). The nuclei of class 2 neurons stain purple and their cell bodies are pyriform (figs. 15a, c). These neurons are less frequent than class 1 neurons, and occur evenly all over the brain (diameter: 5.56 ± 0.72 μm, n = 13). The nuclei of class 3 neurons also stain purple and contain an orange staining nucleolus (figs. 15a, d) (diameter: 12.40 ± 2.04 μm, n = 11). These neurons are only found in small numbers. The prevailing neuron type in the medullary cords is class 1 neurons.
In *T. polymorphus*, the nuclei of class 1 neurons stain dark purple, their cell bodies are small, elongated and appear in clusters (figs. 18a, b) (diameter: 2.29 ± 0.56 \( \mu \)m, \( n = 13 \)). The nuclei of class 2 neurons stain orange and their cell bodies have a circular appearance (figs. 18a, b) (diameter: 4.36 ± 0.45 \( \mu \)m, \( n = 10 \)). These cells are found in small numbers only, but are evenly distributed all over the brain. Cell bodies of class 3 neurons are 50% larger than those of class 2 neurons, having nuclei that are much larger than nuclei of the former two neuron types and stain bluish purple with a prominent orange nucleolus (figs. 18a, c). The prevailing neurons in the medullary cords are class 1 neurons. All size classes differ significantly in diameter (\( p < 0.001 \)).

Figure 12. *Carinoma mutabilis*, light micrographs of Azan stained transverse sections of the brain. a: The cephalic nerves (cn) are circularly arranged around the inner margins of the head. b: The brain (br) is composed of a central neuropil (np) surrounded by cell somata (cs). The whole brain is encircled by an outer neurilemma (on). The two proboscidial nerves (pn) originate from the ventral commissural tract (vct). c: A dorsal nerve (dn) arises from the dorsal commissural tract, the esophageal nerves (en) originate from the ventral commissural tract, the brain is in its posterior part divided into a dorsal (dl) and ventral (vl) lobe. Dorso-lateral nerves (dln) arise from the dorsal commissural tract and merge with the dorsal lobes. d: The ventral lobes are confluent with the lateral medullary cords (mc), two proboscidial nerves are present (pn). Note the different positions of the dorsal nerve (dn1–dn3). e: Higher magnification of the anterior head region, showing the location of the cephalic nerves (cn). dn dorsal nerve, mo mouth opening.

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Minor nerves and peripheral nervous system

In both species cephalic nerves originate from the anterior margin of the ventral and the anteriormost dorsal commissural tract as well as from the ventral and dorsal lobes. While running to the anterior they form a complete ring, between the basal lamina and a layer of longitudinal muscles (figs. 14a, 16, 17a). An ecm (neurilemma) separates each nerve from its neighbor. The cephalic nerves are not covered by neuronal cell somata. The dorsal nerve of *T. superbus* originates in one of the small dorsal commissural tracts and runs posteriorly underneath the basal lamina of the epidermis (fig. 14d). In *T. polymorphus*, this nerve takes the same course, but arises from the posterior margin of the
posteriormost dorsal commissural tract. A ventral nerve is not present. In *T. superbus*, two esophageal nerves (fig. 14d) originate from the posterior margin of the ventral commissural tract, although in *T. polymorphus* these nerves branch off the ventral lobes next to the origin of the ventral commissural tract. In both species, they run posteriorly, pass the mouth and follow the course of the foregut running on either of its sides. Anterior to the mouth, both nerves are connected by a commissure (figs. 16; 17c). In both species, two proboscis nerves arise from the ventral commissural tract more central to the origin of the esophageal nerves. The proboscis nerves run opposite to each other along both sides of the proboscis and are occasionally connected by a proboscidial plexus. The proboscis nerves arise from the ventral commissural tract. (figs. 14b; 16; 17c).

Immunostaining reveals several nerve plexus in *T. polymorphus* (figs. 19a–d). The neurites of the intraepidermal plexus form a regular, ladder-like meshwork (fig. 19b). The neurites of the subepidermal plexus show immunoreactivity with an antiserum directed against FMRFamid (fig. 19a). The commissural plexus is well developed (fig. 19d) and connects the two medullary cords. The neurites of the latter two plexus are horizontally arranged. The neurites of the intrastomatogastric nerve plexus are arranged in an irregular net-like manner (fig. 19c).

Sensory structures

Cerebral organ. Both *Tubulanus* species possess a pair of cerebral sense organs located in the mid-posterior part of the brain (fig. 16). Each cerebral organ consists of a simple ciliated tube that invaginates from a lateral epidermal pore (fig. 20a). The tube-like duct ends next to the posterior margin of the dorsal lobe. Some of the somata next to the duct belong to type 3 neurons in *T. superbus* and type 1 neurons in *T. polymorphus*, indicating that some of the ciliated cells lining the tube are sensory (fig. 20b). The inner section of the cerebral organ is connected to the dorsal lobe of the brain by small branches of neurites (fig. 20b). In both species there is no contact to the blood vessels.

**Callinera grandis** (Tubulanidae, “Palaeonemertea”)

*Callinera grandis* has a subepidermal nervous system; the brain is located anterior to the mouth opening and is not visible from the exterior (fig. 3c). (see: https://www.morphdbase.de:P_Beckers_20130201-M-12.1 for *Callinera grandis* stack).

Central nervous system

Brain. As in the other species studied, the brain is composed of neuropil, which is surrounded by a layer of neuronal cell somata (figs. 21, 22b, c; 23a–c). The entire organ is encased in an outer neurilemma which cannot be differentiated from the epidermal
basal lamina, so that the brain is subepidermal (fig. 22c). An inner
neurilemma separates the somata of the neurons from the neuropil
(fig. 22d). The brain of Callinera grandis is located ventro-laterally
and close to the mouth opening. Anteriorly the brain is juxtaposed
merely separated from the blood space by a small muscle layer (fig. 22c).

The brain consists of four lobes that are connected by a broad and continuous lateral tract, so that, like in both Tubulanus species, dorsal and ventral lobes can hardly be
discriminated as distinct morphological structures (fig. 22c). The terms dorsal and ventral lobes, however, will be kept for the remaining description.

In front of the brain, a pair of lateral bundles of neurites gives
rise to anteriorly directed cephalic nerves as well as to the proboscidial nerves (fig. 23c). Anterior to the brain, the dorso-medial and the ventral-medial sections of the animal initially contain no obvious neuronal structures. As they travel anteriorly, the dorsal- and ventral-most cephalic nerves branch into additional nerves so that a massive ring of parallel cephalic nerves is established about 40 μm in front the brain (figs. 21, 22a).

The lateral bundle of neurites which gives rise to the cephalic nerve is posteriorly connected to the dorsal lobes and to the ventral lobes. The latter connection, however, is less prominent. Both dorsal lobes are located anterior to the ventral ones and are connected by two small dorsal commissural tracts. The posterior section of the dorsal lobe shows a lobe-like dorsal extension that consists of one posteriorly and one anteriorly directed horn. The latter gives rise to the posterior dorsal commissural tract.

Both ventral lobes are connected by a broad and prominent ventral commissural tract (fig. 22c). On either side, this tract originates where the small bundle of neurites connects the crescent shaped origin of the cephalic nerves to the ventral lobes. About 50 μm caudal to the posterior margin of the ventral commissural tract, three successive, small commissural tracts connect the ventral lobes. The neurites of the ventral commissural tracts are horizontally arranged.

A cluster of neurons occurs at the heights were the two esophageal nerves branch off the minor ventral commissural tract (fig. 22d). These neurons are associated with the esophageal nerves as they follow the course of the latter. The neurons are located dorsally to the nerves and ventrally to the proboscidial muscles and follow the esophageal nerves until they divide into a pair of nerves on each side along the foregut. At this point the neurons have contact to the neurons of the medullary cords.

Medullary cords. The ventral lobes taper caudally and give rise to lateral medullary cords. Located directly underneath the epidermal basal lamina, the medullary cords run to the posterior (fig. 22f). Several small rings of neurites connect both medullary cords. The somata of the medullary cord neurons are apposed to the exterior face of the cords to form a lateral, crescent cap (fig. 22f).

Neurons. Like in most of the previously described species, the neurons fall into three size classes. Somata and nuclei of class 2 neurons are twice as large as those of class 1 neurons and half as large as those of class 3 neurons (figs. 23a–c). Cell bodies of class 1 neurons are small and circular, with bright red cytoplasm and a purple nuclei (figs. 23a, b) (diameter: 2.82 ± 0.29 μm, n = 17). These cells are primarily found on the dorsal lobe of the brain. Class 2 neurons are pear shaped, have purplish cytoplasm and...
their nuclei stain slightly darker (figs. 23a, c) (diameter: 4.49±0.62 µm, n = 11). These cells are found on the lateral and ventral parts of the brain. Cell bodies of class 3 neurons stain bright purple, as does the nucleus; their prominent nucleolus is orange (figs. 23a–c) (diameter: 11.26±1.29 µm, n = 12). All size classes differ significantly in diameter (p<0.001).

Minor nerves and peripheral nervous system

The cephalic nerves from a complete ring of longitudinal, partly-interconnected neurites with occasionally interspersed nuclei that are completely isolated from each other by an ecm (figs. 21, 22a). Since these nerves are subepidermal, their ecm and the epidermal basal lamina form a massive layer of extracellular material frontal to the brain. More than 60 of these nerves can be seen in transversal sections of the head region (fig. 22 a). A dorsal nerve arises from the first dorsal commissural tract of the brain. It runs to the posterior, embedded between the basal lamina of the epidermis and a layer of circular muscles (fig. 22d). This nerve is interconnected with the circular bundles of neurites that connect both medullary cords, thus forming a dorsal connection between the rings of neurites. A ventral nerve is not present. Each esophageal nerve has two roots that arise from the lateral portion of the first two small ventral commissural tracts that posteriorly follow the large and prominent anterior ventral tract (figs. 21, 22e). The roots merge on either side of the animal to form a pair of esophageal nerves, which run dorso-medially to merge anterior to the mouth. Ten to 15 µm posterior to this junction, both nerves separate again and surround the mouth on either side. During their course around the mouth, each esophageal nerve bifurcates. Both branches run in parallel laterally and ventro-
Carinina ochracea (Tubulanidae, “Palaeonemertea”)  

The nervous system of C. ochracea is basiepidermal; the brain is located anterior to the mouth and is externally visible (figs. 3f, 24), (see: https://www.morphdbase.de/P_Beckers_20130201-M-13.1 for Carinina ochracea stack)

Central nervous system  

Brain. The brain of Carinina ochracea is basiepidermal, resting on the epidermal basal lamina and an outer neurolamina is missing (figs. 24, 25b). The central part of the brain neuropil is partly separated by an inner neurolamina from the neuronal cell somata, which form the outer layer of the brain (fig. 25b). Fibers of extracellular matrix branch into the brain neuropil (fig. 25d). In contrast to the previous species, the brain does not consist of four discernable lobes. Instead, either side of the brain consists of a pair of exteriorly bulged and crescent-shaped lobes (fig. 25b). Both lobes are connected by a small dorsal commissural tract and a broad ventral commissural tract (figs. 24, 25b). The neurites of the commissural tracts are horizontally arranged, with both commissures located on the same level, so that in transverse sections the brain appears to be U- or ring-shaped, depending on whether the dorsal commissure is sectioned or not (figs. 24, 25b, 27a, b). The dorsal commissural tract gives rise to several anteriorly directed and initially interconnected cephalic nerves. The dorso-frontal margin of the brain is distinctly bifurcated with one branch being oriented anterior-ventrally and the other anterior-dorsally. Both branches also give rise to cephalic nerves, but those originating from the anterior-ventral branch are rather short (fig. 25a). A small, globular extension originates from the dorsal-caudal part of each lobe, tapering quickly and ending broadly in a layer of neuronal cell somata. The ventral commissural tract is slightly crescent-shaped, so that its anteriormost point is located further frontally than the dorsal commissural tract. A few short cephalic nerves originate from the anterior margin of the ventral commissural tract.

Medullary cords. The medullary cords arise from the ventral margin of the each brain lobe. They run posteriorly exterior to the epidermal basal lamina and thus are basiepidermal. The neuronal cell somata cover the neuropil of the medullary cords dorsally and ventrally like a cap (fig. 25d).

Neurons. Like in Carinoma mutabilis, the somata of the neurons fall into two size classes, with somata and nuclei of class 2 neurons being twice as large as those of class 1 neurons (fig. 26). Cell bodies of class 1 neurons are small and circular with nuclei that stain purple (fig. 26a) (diameter: 2.58±0.46 μm, n = 14). The nuclei of class 2 neurons also stain purple (figs. 26a, b) (diameter: 6.11±0.74 μm, n = 10). The different types of neuronal cell somata cannot be discriminated with the applied immunostaining techniques (figs. 27a–d). Class 1 neurons prevail throughout the central nervous system and are characteristic for the medullary cords. A third type of neurons is found around the terminal part of the canal of the cerebral sense organ and will be described below (fig. 26c). The two size classes differ significantly in diameter (p<0.001).

Minor nerves and peripheral nervous system  

The ring of longitudinal cephalic nerves initially forms a basiepidermal basket in the animal’s head (figs. 24, 25a). The dorsal cephalic nerves originate from the dorsal commissural tract,
the lateral ones from both anterior branches of the dorsal lobe and the ventral ones from the ventral commissural tract. All cephalic nerves are initially interconnected. As they travel, they become separated from each other by an ecm (figs. 24; 25a). The lateral cephalic nerves are much shorter than the dorsal ones, so that merely dorsal and ventral cephalic nerves are found in the anterior part of the head. A dorsal nerve arises from the dorsal commissural tract and runs posteriorly, directly opposed to the epidermal side of the basal lamina (basiepidermal) (fig. 25d). A second dorsal nerve runs posteriorly underneath the ecm of the rhynchocoel. Both dorsal nerves are repeatedly interconnected by small neurites. A ventral nerve is not present. Posterior to the ventral commissural tract, two esophageal nerves branches off of each ventral lobe. Both esophageal nerves turn inward, surround the mouth and run posteriorly on either side of the foregut. Two proboscis nerves arise from the posterior margin of ventral commissural tract (figs. 24, 25b). These nerves run posteriorly on either side of the proboscis. A proboscidal plexus regularly interconnects both nerves.

Several different nerve plexus are present in *C. ochracea* (figs. 28a–c). The neurites of the intraepidermal plexus show immunoreactivity with an antiserum directed against FMRFamid and form a ladder-like meshwork (fig. 28b). A subepidermal nerve plexus becomes visible when stained with an antiserum directed against FMRFamid and against α-tubulin (figs. 28a, c, d). Distinct spherical neurite concentrations within this plexus give rise to neurites that connect it to the lateral medullary cords (figs. 28c, d). Whether these are circular or not could not be determined due to the size of the animals and the methods used. The neurites of the intrastomatogastric nerve plexus show a diffuse net-like pattern (fig. 28e). A commissural or a rhynchocoel plexus was not discernable.

**Sensory structures**

**Cerebral organ.** The cerebral sense organs are paired, tube-like structures or ducts that extend inward and terminate dorsally above the dorsal globular extension of the brain. They originate on either side of the animal from a shallow epidermal pit that is located on the level of the terminal section of the dorsal lobe (figs. 24; 29a–c). The cerebral organ initially runs medially, enters the dorsal lobe’s neuronal somata, turns dorsally and runs posteriorly for a few micrometers finally passing the somata layer to blindly end dorso-laterally (figs. 25d, 29a). The entire cerebral organ is lined by ciliated and sensory cells; the density of the latter, however, increasing towards the inner part of the organ. Immunostaining with anti-α-tubulin shows a very brightly stained structure, which presumably represents densely arranged cilia (figs. 29b, c). Azan staining reveals cell somata that differ from those of the brain and to those of the medullary cords (fig. 26c), since they are elongated, twice as large as class 2 neurons and their nuclei stain purple. Anti-α-tubulin immunostaining also shows that the sensory cells are connected to the posterior part of the dorsal lobe of the brain by small branches of neurites (figs. 29a–c). There is no contact between the cerebral organ and the blood vessels. Pigmented eyes and frontal organs are absent in this species.

**Sensory cells.** Many sensory cells are distributed all over the animal. Most of these intraepidermal cells originate from the...
subepidermal nerve plexus (fig. 28a). The cells presumably have a tactile function.

Discussion

Most textbooks simplify the nemertean central nervous system as being quite uniform, consisting of a four lobed brain and two lateral medullary cords [5], [27], [28], [29], while others describe the nemertean nervous system as being primarily ring-shaped [30]. The present study, however, shows a remarkable diversity in the organization of the central nervous system among palaeonemertean species and corroborates a previous comparative analysis [31]. While the CNS is strictly bilaterally symmetrical, four lobes can clearly be identified in only in a few species. In addition, the peripheral nervous system also differs amongst different groups and often even between closely related species. Palaeonemerteans most likely represent a grade of basally branching nemertean groups [4], [5], [14], [15], [16], but this is not unanimous [17]. Comparing their nervous systems, thus, could reveal some important insights into the primary organization of the nemertean nervous system. The present study covers representatives of the three larger palaeonemertean groups Cephalothricidae, Carinomidae and Tubulanidae. Since the monophyly of the latter is questionable [32], we analyzed representatives of each genus except for Carinesta. In the subsections below we will comparatively evaluate the central nervous system in nemerteans in terms of topology, structure, types of neurons and extracellular components as well as their peripheral nervous system and their sensory organs.

Brain and central nervous system

Topological position. The topological position of the central nervous system (CNS) differs between the palaeonemertean species. Like in other cephalothricids [4], [31], [33], [34], [35], [36] the CNS is located inside the musculature of both cephalothricid species studied. There is only very limited information on the nervous system of carinomids, however, Friedrich [37], mentioned that the anatomy of carinomids is rather uniform throughout the group, so we assume the intramuscular position of the CNS in Carinoma mutabilis is characteristic for Carinomidae. The course of the medullary cords, however, differs between carinomids and cephalothricids. While they are completely intramuscular and run within the longitudinal muscle layer in cephalothricid species, this is only the case in the posterior region of carinomid species; anteriorly they are situated underneath the outer circular muscle layer. This observation contradicts [38] who claims that the medullary cords are initially located outside the circular muscles in Carinoma species. In Tubulanus, Callinera, and Carinesta species, the brain and medullary cords are subepidermal, located between the subepidermal ECM and the underlying outer circular muscles [31], [34], [35], [36], [38], [39]. In Carinina ochracea, like in other Carinina species they are found basiepidermally [31], [36], [38].

Burger [4] and Iwata [18] assumed the position of the CNS as having important implications for considerations of nemertean...
phylogeny. Bürger [4] supposed that a basiepidermal nervous system, like that of C. ochracea was characteristic for the nemertean ancestor and assumed that the CNS migrated inward during nemertean evolution, so that a subepidermal nervous system represents a derived condition in Nemertea. Iwata [13], on the other hand, supposed the Cephalothricidae to be the basal most branching taxa, due to the location of the CNS inside the musculature and the lack of a cerebral organ. This interpretation suggests that any other topological position of the CNS must therefore be derived.

Presently, we are unable to provide support for either traditional interpretation of CNS evolution since the position of nemerteans within the Spiralia is unresolved and any position of the CNS found within the nemerteans can also be found in at least one of the possible outgroups. In platyhelminthes [40], sipunculids [41] and kamptozoans [42], the nervous system is located subepidermally. In basal molluscs [43] the nervous system is located in the longitudinal musculature and in brachiopods [44] and phoronids [45] it is located basiepidermally. In annelids the nervous system may be located basiepidermally (Oxenia fusiformis, [46]), or in the musculature (Nereis species, [47]). Despite an ongoing debate about the phylogeny of annelids [48,49], the heterogeneous position of the nervous system within the Spiralia needs a thorough cladistic analysis. The presently known topological differences, however, may indicate that this character does not contain a signal for unraveling the higher-level phylogeny.

**Structure.** A brain lobe has been defined above as voluminous compartment of the brain that is connected to other such compartments by tracts. The heteronemertean L. viridis has paired dorsal and ventral lobes, each of which is more or less completely surrounded by a matrix and thus a clearly identifiable morphological unit [24]. Identification of these lobes is difficult, if such an extracellular surrounding is missing. Despite the lack of an ECM that demarcates the lobes, however, their larger volume compared to the commissural and lateral tracts allows identification of all four brain lobes in non-tubulanid palaeonemerteans. In cephalothricid species, several lateral tracts connect dorsal and ventral lobes on each side of the brain. In Callinara grandis, and in both of the Tubulanus species examined, dorsal and ventral lobes from each body side are connected by an anterior-posterior broad, but medio-laterally narrow lateral tract, so that slight differences in the volume allow the identification of all four brain lobes. This is especially evident, when taking a look at 3D-reconstructions of the brain. If these are not available, some ambiguity will remain when describing the structure of the brain and lead to differing descriptions of the brain in different species of the same genus (see table 1).

![Figure 19. Tubulanus polymorphus, confocal laserscanning (cLSM) micrographs of differently immunostained vibratome sections. a: Transverse section, anti-FMRF (red), anti-α-tubulin (green) showing the subepidermal (snp) and the stomatogastric (sgp) nerve plexus. b: Horizontal section, anti-FMRF showing the regular arranged neurites (ne) of the subepidermal nerve plexus. c: Horizontal section, anti-FMRF. The neurites (ne) of the intrastomatogastric nerve plexus (isgp) are arranged in a regular ladder-like fashion. d: Transverse section, anti-serotonin. The lateral medullary cords (mc) are interconnected by a commissural plexus (cnp). The commissural plexus surrounds the mouth opening (mo).](http://www.plosone.org/figure/19/)

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In Carinina ochracea however, no criteria exist to identify dorsal and ventral lobes. Instead the entire brain consists of two laterally bulged lobes that are connected by a dorsal and at least one ventral commissural tract. The same has been stated for Carinesta species [39], [36]. Given that tubulanids are a basal grade within the nemerteans, as indicated by some molecular studies [16], [17], the bi-lobed, ring-shaped brain found in Carinina and Carinesta species [39], [50], [31], [36] could represent the primary condition of the nemertean brain. Moreover, its basiepidermal position in Carinina would then be in accordance with Bürger’s [4] assumption of such a position representing a primary condition in nemerteans. This assumption implies that the brain becomes increasingly regionalized with its dorsal part becoming prominent and separated during the nemertean evolution – an evolutionary scenario that has already been argued for eloquently by [31]. This scenario requires interpretation of the brain structure in Collinera and Tubulanus as intermediates. The bi-lobed, ring-shaped brain without clearly identifiable dorsal and ventral lobes in Carinina species could, however, also be explained by condensation of the brain and loss of brain compartments, making the brain structures in Collinera and Tubulanus again intermediates. Although this alternative explanation presently seems unlikely to us (see below), a final decision will require a cladistic analysis of the nemerteans that includes additional anatomical characters. Complex internal compartmentalization, like the one found in a number of annelid representatives that contain mushroom bodies, has not been observed in nemerteans. [51,52]

Medullary cords have been described in Platyhelminthes [40], basal molluscs [53], [54], [55], [56], [57] and polychaetes [47], [58]. The arrangement of the neuronal cell somata surrounding the neuropil of the medullary cords is not described in these studies, although they are present according to classical contributions [40], [43], [47]. As longitudinally arranged nerve cords, these parts of the nervous system can certainly be homologized, but it remains to be resolved which cords of the tetraneural mollusk and the orthogonal platelminth nervous system are homologous to the nemertean medullary cords. This, as well as an evaluation of the polychaete medullary cords needs thorough cladistic analyses based on morphological matrices that contain detailed information on the nervous system. A serial arrangement of circular nerves connecting the medullary cords is found in Procephalothrix filiformis. In Carinina ochracea, the neuropil of the medullary cords is connected to the neurites of the subepidermal nerve plexus with regular arranged branches of neurites showing immunoreactivity against α-tubulin. In addition, at the spots where the neurites of the subepidermal nerve plexus branch off, a spherical concentration of neurites occurs. Since a serial arrangement of particular nerves is also present in platyhelminthes [40], annelids [47] and basal molluscs [43], we suggest this as an element of the last common ancestor of the Spiralia or a subgroup within this clade.
Neurons. The neuronal cell somata surrounding the brain neuropil appear to be different in all investigated palaeonemerteans. Despite their different staining, the neurons of palaeonemertean species fall in three size classes, with the second and the third being almost twice as large as their previous ones. The third, largest class, which is missing in Carinina ochracea, always possesses a prominent nucleolus and a large nucleus. Its morphology implies high metabolic rates. The first class of cells (type 1 neurons) predominates in all nervous systems. Neurochord cells (type 4 according to Burger [4]) are missing in the palaeonemertean groups except for Cephalothricidae. Although Bianchi [59] discriminated six types of neuronal cell somata in the heteronemertean Cerebratulus marginatus, the three types of neuronal cell somata in Lin vulvula (Heteronemertea) described by [25] most parsimoniously represent the primary condition in nemerteans. Since the neurons in basal mollusks have also been grouped in three classes by size differences [60], this could hint at an ancient pattern in nerve cell organization in Spiralia. Presently, the data are too scattered to draw any conclusion, but it may be fruitful to include neuron size into a cladistic analysis.

Neurilemma. The neuronal cell somata of the brain may be separated from the neuropil by an extracellular matrix, also termed inner neurilemma [4]. Within the Cephalothricidae, Carinina ochracea has an inner neurilemma, while it is lacking in Callinera grandis and Tubulanus. Little is known about the function of the extracellular matrix in the invertebrate nervous system. This ECM may be involved in neurite formation, maintaining structure and function as well as in repair processes [61], [62]. An extracellular matrix surrounding or encapsulating the brain has been described for basal molluscs [53], for certain platyhelmintes [40] and polychaetes [47]. Although this seems to indicate that an ECM surrounding the brain is characteristic either for a common stem species of the Spiralia or for a group within the Spiralia including annelids, mollusks and nemerteans, the outer neurilemma cannot be differentiated from the epidermal basal lamina in basipidermal and subepidermal nervous systems. The outer neurilemma, thus, might be nothing more than a reinforced subepidermal matrix and therefore overestimated in evolutionary considerations. Nevertheless, an inner neurilemma, which is present in some palaeonemertean species, is unique and seems to be of phylogenetic importance.

Minor nerves and peripheral nervous system

Cephalic nerves differ in their arrangement and position between taxa. In Cephalothricidae there are four cephalic cords; cephalic nerves were not found. In the remainder of the investigated palaeonemerteans (Tubulanus, Callinera grandis, Carinoma mutabilis, Carinina ochracea), they are circularly arranged around the head lacuna. Since the heteronemertean Lin vulvula also possess an anterior ring of several cephalic nerves, such an organization must represent a primary condition in nemerteans, making the nerve cords in cephalothricid species derived, probably correlated with the expansion of the preoral body region that characteristic for this group.

A dorsal nerve is present in all palaeonemertean species. The dorsal nerve of Cephalothricidae and Carinoma mutabilis originates in the dorsal commissural tract inside the musculature. The nerve then runs dorsally and finally lies just underneath the basal lamina of the epidermis. A ventral nerve is only present in Cephalothricidae. It has two roots that branch off the ventral lobe at the origin of the ventral commissure, like the esophageal nerves of the remaining palaeonemertean species. The ventral nerves encircle the mouth like the esophageal nerves, however they differ by running below the gut as single nerve. Since a ventral nerve is lacking in the remaining palaeonemerteans and esophageal nerves are lacking in Cephalothricidae, the origin, position and initial course clearly indicate a homology of the ventral nerve and the esophageal nerves. This hypothesis is supported by Gibson’s [35] remark that the ventral nerve (foregut nerve) supplies the foregut. Since a ventral nerve is lacking in Cephalothricidae, the origin, position and initial course clearly indicate a homology of the ventral nerve and the esophageal nerves. This hypothesis is supported by Gibson’s [35] remark that the ventral nerve (foregut nerve) supplies the foregut. All investigated palaeonemerteans possess paired proboscidial nerves, which also originate in the ventral commissural tract of the brain.

Palaeonemerteans possess nerve plexus in different parts of their body. These plexus may be well developed and can clearly be shown by immunohistochemistry. The arrangement of the...
neurites found there differs between nerve plexus but not between species.

Sensory structures

Nemerteans perceive chemical substances with different chemosensitive sensory structures, which are situated on the head of the animals. These are called the frontal and cerebral organs. As shown in studies of a hoplonemertean [11] and cephalothricid palaeonemertean [12], these sensory structures are designed for a hunting life style. Frontal organs are not present in the species investigated here, although they have been described for Procephalothrix simulus (Palaeonemertea) [63].

The cerebral organs of *Tubulanus* species consist of simple epidermal tubes that open laterally to the environment and terminate exterior to the basal lamina in the epidermis. The inner part of the tube’s epithelium is lined with sensory cells, which are

![Figure 22. Callinera grandis, light micrographs of Azan stained transverse sections of the brain. a: The cephalic nerves (cn) are composed of neuropil (np) and neuronal cell somata (cs), a cephalic gland (cg) is located dorsally. b: The anterior region of the brain (br) is covered by an enormous layer of cell somata (cs). c: The brain is composed of a central neuropil (np) and a surrounding layer of cell somata (cs). The two halves of the brain are connected by ventral (vct) and dorsal (dct) commissural tracts. d: Posteriorly the brain is divided into a ventral (vl) and dorsal (dl) section. A dorsal nerve (dn) arises from the dorsal commissural tract, and the paired esophageal nerves (en) arise from the ventral commissural tract. Shortly anterior to the foregut a concentration of neurons (gn) occurs which are associated with the esophageal nerves. The somata are separated from the neuropil by an inner neurilemma (in), and the whole brain is enclosed by an outer neurilemma (on). e: The ventral lobes of the brain are confluent with the lateral medullary cords (mc). mo: mouth opening, pr: proboscis. f: The lateral medullary cords (mc) run to the posterior of the animal. The cell somata (cs) cover the neuropil in a C-shaped manner. fg foregut. doi:10.1371/journal.pone.0066137.g022](image-url)
connected to the brain. In *Carinina ochracea*, the canal of the cerebral organ is elongated and runs dorsally but it also terminates exterior to the basal lamina of the epidermis. In contrast to the conditions found in *Tubulanus*, the duct of the cerebral organ in *C. ochracea* passes through the neuronal cell somata of the brain since the brain in *C. ochracea* is basiepidermal. Because of the similarities in the morphology and staining affinities, we assume a common origin of the cerebral organ and not convergent evolution.

In *Carinina ochracea*, several sensory cells originate in the subepidermal plexus and proceed into the outer margins of the epidermis. These cells may have a tactile function. In *Procephalothrix filiformis*, a cluster of cells showing immunoreactivity against an antiserum directed against FMRFamid is present in the tip of the animal’s head. These cells might represent sensory cells.

Ciliated sensory structures that may have a chemo-sensitive function are found in Platyhelminthes, Mollusca and in Annelida. Platyhelminthes possess ciliated sense organs on the tip of the head, which are supposed to have a chemoreceptive function [30]. In most anelids species, nuchal organs are present. These are small chambers situated on the prostomium of the animals. The chamber is lined with sensory cells that are connected to the brain by neurites [64], [65], [66]. Aplacophoran molluscs possess a sensory organ on the tip of their head that is also supposed to have a chemoreceptive function [67]. The function, morphology and

![Figure 23. Callinera grandis, light micrographs of Azan stained neurons. a: Transverse section, overview of the brain position of the three different types of neurons (S1–S3); all nuclei stain purple. b: Higher magnification of somata of type 1 and 3 neurons (S1, S3). Cell bodies are circular; those of S1 are arranged in clusters, those of S3 are most prominent. c: Higher magnification of somata of type 2 and 3 neurons. S2 cell bodies are pear shaped and slightly enlarged. The neurites (ne) of the brain branch into the cell somata layer. np neuropil. doi:10.1371/journal.pone.0066137.g023](image-url)
The nervous system is composed of neuropil (np, gray) which may be surrounded by cell somata (cs, blue). Cephalic nerves (cn) form basiepidermal basket, the paired proboscisial nerves (pn, yellow) originate from the ventral commissural tract (vct). A dorsal nerve strand (dn) originates from the dorsal commissural tract (dct). A second dorsal nerve (2dn) runs underneath the dorsal nerve. The cerebral organs (co, green) gain contact to the environment via small canals (ca, red), which open laterally. The cerebral organ is connected to the dorsal section of the brain. The branching esophageal nerves (en) originate from the ventral commissural tract (vct). The lateral medullary cords (mc) originate in the ventral part of the brain. The letters on the right (a–d) refer to the histological sections in figure 25. doi:10.1371/journal.pone.0066137.g024

position of these sensory structures are comparable to the frontal organs of nemertians. In older studies, a connection between the frontal organs and brain has been described [5], but the present study failed to analyze this connection in detail, because neither Azan nor in the immunostaining revealed sufficient information.

The homology of the cerebral organs within the nemertians is still a matter of debate [16], due to strong positional differences among the taxa. If they are homologous however, the plesiomorphic position would be posterior to the brain, since an anterior position is only described for hoplonemertean species and according to recent phylogenetic analysis [16], [17], Hoplonemertea are derived within the Nemertea. Specific correspondences between the cerebral organs and cephalic sensory organs in other spiralian species are thus far unknown, since innervations pattern and ultrastructural details are largely unknown for the cerebral organs.

**Conclusions**

Our comparative survey of the palaeonemertean nervous system reveals that each brain consists of two halves in which the neuronal perikarya (cell bodies, somata; Ganglienzellkörper [36], Ganglienzellen sensu [4]) are opposed and form a layer of neuronal somata. Lateral fiber masses (neuropil), can be discriminated into dorsal and ventral compartments, each of which is called a lobe. Our study also gives rise to the idea of an increasing compartmentalization of the nemertean brain during the course of evolution leading to prominent brain lobes in Hoplo- and especially Heteronemertea. This had two evolutionary consequences: (1) the medial portion of fiber masses became smaller and partitioned, so that it is identified as lateral tracts rather than being part of a large lateral fiber mass, and (2) the lobes dominate the histological section in such a way that each lobe has been termed ganglion by some researchers [4], [34]. Especially in heteronemertians, this impression is strengthened by the fact that the lobes are separated from each other by a neurilemma. This idea, however, depends on the phylogenetic position and the mono- or paraphyly of the palaeonemertians.

If the palaeonemertians were monophyletic as implied by recent molecular studies [17], the idea of an increasing compartmentalization of the brain and the translocation of the nervous system into deeper tissue layers during nemertean evolution would either have to be modified or cannot be held: either (1) the intraepidermal nervous system with a rather simple brain represents the primary condition in nemertians so that compartmentalization and translocation of the nervous system into deeper tissue layers during the course of evolution leading to prominent brain lobes in Hoplo- and especially Heteronemertea. In the latter case our idea of an increasing compartmentalization cannot be held upright. The former case requires assuming that brain compartments and translocation of the nervous system into deeper tissue layers evolved repeatedly within nemertians and, thus, is presently not parsimonious.

Given that palaeonemertians are a basally branching grade, as implied by older molecular studies [15,16] and presently missing morphological support for a taxon Palaeonemertea, the nemertean brain primarily consisted of two bilaterally symmetrical lobes, interconnected by a ventral and one or several small dorsal commissures. A pair of medullary cords originates from the ventral portion of the brain. According to this assumption the primary organization of the nemertean nervous system is retained in Carinina ochracea. Provided that the brain expanded dorsally and became dorsally commissurized along with the evolution of a proboscis, nemertians and their spiralian sister taxa should have had at least two longitudinal medullary cords connected by frontal commissures that were laterally reinforced and, thus, resemble the
Figure 25. *Carinina ochracea*, light micrographs of Azan stained transverse sections of the brain. 

a: Frontal region showing the circularly arranged cephalic nerves (cn). br: blood vessel. 
b: The brain (br) is composed of a central neuropil (np) which is surrounded by cell somata (cs). A ventral commissural tract (vct) connects the two halves of the brain, and an outer neurilemma (on) encloses the whole brain. (pn): proboscidial nerves. 
c: The canals (co) of the cerebral organs open laterally, where the brain divides into a dorsal (dl) and ventral (vl) sections. Two proboscidial nerves (pn) are present. 
d: The canals of the cerebral organs (co) end dorsally. A dorsal nerve strand (dn) arises from the dorsal commissural tract, the ventral part of the brain is confluent with the lateral medullary cords (mc). Note the ecm (arrow) branching into the neuropil of the brain. mo mouth opening.
doi:10.1371/journal.pone.0066137.g025

Figure 26. *Carinina ochracea*, light micrographs of Azan stained neurons. 
a: Cell bodies of type 1 neurons (S1) are not enlarged while those of type 2 neurons (S2) are; nuclei stain purple. 
b: Neuronal cell somata of the medullary cord (mc) are of the second type described for the brain. 
c: The canal of the cerebral organ ends in a layer of special neuronal cell somata (sco), which are not present in the brain.
doi:10.1371/journal.pone.0066137.g026
nervous system of basally branching mollusks and annelids. Actually, basiepidermal nervous systems are known from polychaete annelids, phoronids and brachiopods, the molluscan nervous system is always subepidermal (intramuscular). The phylogenetic position of the nemerteans is still unresolved among spiralians and lophotrochozoans [68]. In order to complete the information and to solve this potential conflict, we are presently analyzing the nervous systems of the two remaining subgroups, the Pilidiophora (Heteronemertea + Hubrechtidae) and the Hoplonemerteans and potential outgroup taxa, using the same methods and providing a coding scheme. A final cladistic analysis will help to elucidate the evolution of the nemertean nervous system and test the different hypotheses on its primary design.

Materials and Methods

Animals

Carinoma mutabilis Griffin, 1898 was found in March 2007 at Friday Harbor, Cattle Point, San Juan Island, WA, USA. Procephalothrix filiformis Johnston, 1828 was found on the isle of Sylt in February 2009. Callineura grandis Bergendal 1903 and Carinina ochracea Sundberg, Chernyshev, Kajihara, Kanneby, Strand 2009 were found in Pouldohan/Brittany France in September 2009. Cephalothrix linearis Rathke, 1799; Tubulanus polymorphus Renier 1804; Tubulanus superbus Kolliker 1845 were found in Roscoff/Brittany France in March 2010.

We neither used endangered species nor were the investigated animals collected in a protected area. All animals were collected with the permission of the local marine biological stations.

Histology

One or two specimen of each species were anesthetized in a refrigerator using a solution of 7.5% MgCl₂ in seawater, and fixed overnight in Bouin’s fixative modified after Dubosque- Basí. The animals were completely dehydrated in an ethanol series, followed by incubation in methylbenzoat and butanol. Afterwards the animals were preincubated in Histoplast (Thermo Scientific, Dreieich, Germany) at 60°C for three days with several medium changes and finally embedded in Paraplast (McCormick Scientific, Richmond, USA). Different incubation times were used depending on the size of the animal.

Sections 5 µm thick were made using a Reichert-Jung Autocut 2050microtome (Leica, Wetzlar) and transferred to glass slides coated with albumen-glycerin.

Sections were stained with Carmalaun, subsequently differentiated with sodium phosphotungstate (5%), washed in distilled water and stained with aniline blue orange G. Afterwards the
sections were embedded with Malinol (Waldeck, Münster, Germany).

Azan stains the neuropil of the nervous system gray. The nuclei of the neuronal cell somata stain red, sometimes with a tinge of orange or purple. The extracellular matrix (ECM, called neurilemma in the central nervous system) stains blue, while musculature stains orange.

**Immunohistochemistry**

Specimen of one species of each genus (table 2) except for *C. grandis* and *C. mutabilis* due to the lack of material were anesthetized in a refrigerator using a solution of 7.5% MgCl\(_2\) in seawater. The worms were fixed overnight in a 4% paraformaldehyde (Electron Microscope Sciences, Hatfield, PA) in seawater (0.1 M) solution. After fixation, the head or middle regions of the animals were washed several times in cold phosphate buffered saline (PBS 0.01

**Figure 28. Carinina ochracea, confocal laserscanning (cLSM) micrographs of differently immunostained vibratome sections.**

- **a:** Anti-FMRF, horizontal section. A sensory cell (sc) originates in the subepidermal plexus (snp) and proceeds into the epidermis (ep).
- **b:** Anti-FMRF, horizontal section. The neurites (ne) of the intraepidermal nerve plexus are arranged in a ladder-like manner.
- **c:** Anti-α-tubulin, sagital section. The lateral medullary cords (mc) are interconnected to the subepidermal plexus (snp) by branches of neurites (ne).
- **d:** Higher magnification of **c.** Note the concentration of neurites in the subepidermal nerve plexus (arrows).
- **e:** Anti-FMRF, horizontal section. The neurites (ne) of the intrastomatogastric nerve plexus are irregular arranged.
- **f:** Anti-FMRF, horizontal section. Multipolar neurons (cs) are distributed in the intraepidermal nerve plexus.

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M). They were then embedded in a gelatine/albumin medium and the blocks were hardened overnight in a 14% formalin solution in the refrigerator. Each sample was cut into sections 60 mm thick with a vibratome (Micron HM 650 V, Thermo Scientific, Dreieich, Germany). The sections were then washed 5 times for 20 minutes in PBS with 0.1% Triton X-100 (Sigma Aldrich, St. Louis, MO, USA) and pre-incubated overnight in a blocking solution of PBS (0.01 M) containing 0.5% TX and 5% normal swine serum (Jackson ImmunoResearch, West Grove, PA). Primary antibodies were added into the blocking solution and incubated overnight at room temperature. The primary antibodies anti- FMRFamide (ImmunoStar, Hudson, WI) and anti-serotonin (Sigma-Aldrich, Saint Louis, MO, USA) were both used at a dilution of 1:10000. Antibodies against acetylated α-tubulin (Sigma Aldrich, St. Louis, MO, USA) were used at a dilution of 1:500. After incubation with primary antibodies, the sections were again washed 5 times for 20 minutes in PBS with 0.1% TX and incubated overnight at room temperature. The primary antibodies anti-FMRFamide (ImmunoStar, Hudson, WI) and anti-serotonin (Sigma-Aldrich, Saint Louis, MO, USA) were both used at a dilution of 1:10000. Antibodies against acetylated α-tubulin (Sigma Aldrich, St. Louis, MO, USA) were used at a dilution of 1:500. After incubation with primary antibodies, the sections were again washed 5 times for 20 minutes in PBS with 0.1% TX and incubated overnight with the secondary antibody conjugated to fluorophore (Cy3- conjugated goat anti-rabbit; Jackson ImmunoResearch, West Grove, PA) at a dilution of 1:2000. Secondary antibodies against acetylated α-tubulin (Alexa 633, Life Technologies, Darmstadt) were added at a dilution of 1:250. Subsequently the sections were rinsed again in several changes of PBS containing 0.1% TX and then mounted on chrome alum/gelatine-coated glass slides under glass cover slips using Elvanol (mounting medium after [69]).

For whole animal immunostaining, animals were relaxed in a 7.5% MgCl2 solution and fixed overnight in 4% paraformaldehyde (Electron Microscope Sciences, Hatfield, PA). Animals were postfixed for 20 minutes in methanol and then washed in several steps of PBS (0.01 M). Afterwards they were washed with PBS containing Triton X at a dilution of 0.1% and preincubated in a blocking solution with NSS (Jackson ImmunoResearch, West Grove, PA). The primary antibodies for anti-FMRFamide and serotonin were added at a dilution of 1:5000. After the incubation, the primary antibodies were washed out with PBS containing Triton X at a dilution of 0.1%. The secondary antibodies (Cy3 for anti-FMRFamide and anti Serotonin, and Alexa 633 for α-tubulin) were added at a dilution of 1:2000 and incubated for 24 hours. Afterwards the animals were washed in PBS containing 0.1% Triton X and pure PBS. Then the animals were mounted in Murray clear (2 parts benzyl benzoate, 1 part benzyl alcohol) after having been treated with methanol and isopropanol.

**Analysis and 3D reconstruction**

Azan stained slices were analyzed with an Olympus microscope (BX-51) and photographed with an Olympus camera (Olympus cc12), which was equipped with the dot slide system (2.2 Olympus, Hamburg). Afterwards the slices were aligned with imod [70] and imod align [http://www.q-terra.de/biowelt/3drekon/guides/imod_first_aid.pdf]. 3D reconstructions were performed with Fiji (1.43b) [71]/Trakem [72] and Amira (4.0). The schematic drawings were performed according to the 3D reconstructions with Illustrator and Photoshop (CS4).

Immunohistochemically treated samples were analyzed with a Leica confocal laser scanning microscope (TCSSPE, excitation wavelength 488 nm, detection range 500–630 nm was used to detect...
Table 1. Elements of the nervous system of basally branching nemerteans.

| taxon                        | lateral lobe | dorsal lobe | ventral lateral lobe | dorsoventral commissure | cephalic cords | medullary cords | cephalic nerves | dorsal nerve | ventral nerve | esophag. nerve | probscid nerve | cerebral organ | Author          |
|-----------------------------|--------------|-------------|----------------------|-------------------------|----------------|----------------|---------------|-------------|---------------|----------------|----------------|----------------|----------------|
| Cephalothrix linearis       | -            | 1 pair      | 1 pair               | 1                       | 4              | 1 pair         | -             | 1           | -             | -              | 2              | -              | this study      |
| Procephalothrix filiformis  | -            | 1 pair      | 1 pair               | 1                       | 4              | 1 pair         | -             | 1           | 1             | -              | 1              | 2              | -              |
| Carinina ochracea           | 1 pair       | -           | -                    | 1                       | 2              | -              | 1             | 1           | -             | 1              | 2              | 1              | this study      |
| Tubulanus superbus         | 1 pair       | -           | 1 pair               | -                       | 1 pair         | >5             | 1             | 1           | -             | -              | 2              | 2              | this study      |
| Tubulanus polymorphus      | -            | 1 pair      | 1 pair               | 1                       | 4              | 1 pair         | >5            | 1           | 1             | -              | 2              | 2              | 1              | this study      |
| Callinera gisandis         | -            | 1 pair      | 1 pair               | 1                       | 4              | 1 pair         | >5            | 1           | -             | -              | 2              | 2              | -              | this study      |
| Carinoma mutabilis         | -            | 1 pair      | 1 pair               | 3                       | 1              | -              | 1 pair        | >35         | 1             | -              | 2              | 2              | -              | this study      |
| Carinina grata             | -            | 1 pair      | 1 pair               | 1                       | 1              | -              | 1 pair        | ?           | 1             | -              | -              | 2              | 2              | -              |
| Carinina johnstoni         | 1 pair       | -           | -                    | 1                       | 1              | -              | 1 pair        | ?           | 1             | -              | -              | 2              | 2              | 1 pair         |
| Callinesta dellichiajai     | 1 pair       | -           | -                    | 1                       | 1              | -              | 1 pair        | ?           | 1             | -              | 2              | 2              | 2              | -              |
| Callinera blanchardi       | 1 pair       | -           | -                    | 1                       | 1              | -              | 1 pair        | several     | 1             | -              | 2              | 2              | -              | this study      |
| Callinera quaterfasciata    | 1 pair       | -           | -                    | 2                       | 1              | -              | 1 pair        | several     | 1             | -              | 2              | 2              | -              | -              |
| Cephalothrichella gaimardi | -            | 1 pair      | 1 pair               | 2                       | 1              | -              | 1 pair        | 4           | 1             | -              | 2              | 2              | 2              | -              |
| Callinesta orientalis      | 1 pair       | -           | -                    | 2                       | 2              | -              | 1 pair        | >5          | 1             | -              | -              | 2              | 2              | -              |
| Tubulanus lucidus          | 1 pair       | -           | -                    | 2                       | 2              | -              | 1 pair        | >5          | 1             | -              | -              | 2              | 2              | -              |
| Carinesta uchidai          | 1 pair       | -           | -                    | +                       | 1              | -              | 1 pair        | ?           | 1             | -              | 2              | 2              | 2              | -              |
| Procephalothrix fasciculus | -            | 1 pair      | 1 pair               | 1                       | 4              | 1 pair         | -             | 1           | -             | -              | -              | -              | -              | -              |
| Procephalothrix arenarius  | -            | 1 pair      | 1 pair               | 1                       | 4              | 1 pair         | -             | 1           | 1             | 1              | -              | -              | -              | -              |
| Carinina tubulanoides      | 1 pair       | -           | -                    | 1                       | 1              | -              | 1 pair        | +           | 1             | 1              | -              | 2              | 2              | -              |
| Tubulanus norvegicus       | 1 pair       | -           | -                    | 1                       | ?              | -              | 1 pair        | ?           | 1             | 1              | -              | 2              | 2              | -              |

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Table 2. Staining protocol of the palaeonemertean species investigated.

| Species                  | Histology | Immunostaining |
|--------------------------|-----------|----------------|
| Procephalothrix filiformis | x         | x x x          |
| Cephalothrix linearis    | x         |                |
| Carinoma mutabilis       | x         |                |
| Tubulanus superbus       | x         |                |
| Tubulanus polymorphus    | x x x x   |                |
| Callinera grandis        | x         |                |
| Carinina ochracea        | x x x     |                |

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Alexa 488 fluorescence. Excitation wavelength 543 nm, detection range 555–700 nm was used to detect Cy3 fluorescence. Excitation wavelength 633 nm, detection range 650–800 nm was used to detect Alexa 633 fluorescence using the LAS AF 1.6.1 software. Stacks were loaded into Fiji [ImageJ 1.43k plugin] [71] and further processed, using the “maximum projection” tool of the ImageJ software (Macophonics 1.44p). The images were finally processed with Fiji [71] and Adobe Photoshop (CS4). Figures were arranged using Adobe Illustrator (CS4).

Statistical analysis

Neuron diameters were measured with Cell^P^D (3.2 Olympus). For analyses, the SPSS 15.0 statistical package was used. Data of each species and each class of neurons were tested for normal distribution with Shapiro-Wilk test. Since the data were normally distributed, parametric statistics (ANOVA) were used to test whether the size classes of each species differ significantly.

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Author Contributions

Conceived and designed the experiments: PB. Performed the experiments: PB. Analyzed the data: PB TB. Contributed reagents/materials/analysis tools: PB RL. Wrote the paper: PB TB RL.

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