Independent specialisation of myosin II paralogues in muscle vs. non-muscle functions during early animal evolution: a ctenophore perspective

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Abstract

Background: Myosin II (or Myosin Heavy Chain II, MHCII) is a family of molecular motors involved in the contractile activity of animal muscle cells but also in various other cellular processes in non-muscle cells. Previous phylogenetic analyses of bilaterian MHCII genes identified two main clades associated respectively with smooth/non-muscle cells (MHCIIa) and striated muscle cells (MHCIIb). Muscle cells are generally thought to have originated only once in ancient animal history, and decisive insights about their early evolution are expected to come from expression studies of Myosin II genes in the two non-bilaterian phyla that possess muscles, the Cnidaria and Ctenophora.

Results: We have uncovered three MHCII paralogues in the ctenophore species Pleurobrachia pileus. Phylogenetic analyses indicate that the MHCIIa / MHCIIb duplication is more ancient than the divergence between extant metazoan lineages. The ctenophore MHCIIa gene (PpiMHCIIa) has an expression pattern akin to that of "stem cell markers" (Piwi, Vasa, . . ) and is expressed in proliferating cells. We identified two MHCIIb genes that originated from a ctenophore-specific duplication. PpiMHCIIb1 represents the exclusively muscular form of myosin II in ctenophore, while PpiMHCIIb2 is expressed in non-muscle cells of various types. In parallel, our phalloidin staining and TEM observations highlight the structural complexity of ctenophore musculature and emphasize the experimental interest of the ctenophore tentacle root, in which myogenesis is spatially ordered and strikingly similar to striated muscle formation in vertebrates.

Conclusion: MHCIIa expression in putative stem cells/proliferating cells probably represents an ancestral trait, while specific involvement of some MHCIIa genes in smooth muscle fibres is a uniquely derived feature of the vertebrates. That one ctenophore MHCIIb paralogue (PpiMHCIIb2) has retained MHCIIa-like expression features furthermore suggests that muscular expression of the other paralogue, PpiMHCIIb1, was the result of neofunctionalisation within the ctenophore lineage, making independent origin of ctenophore muscle cells a likely option.

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Background

Myosins are a tremendously diverse family of actin-binding ATP-dependent molecular motors that appeared and diversified early during eukaryotic evolution [1-5]. All myosins share a homologous myosin head domain containing the ATPase and actin-binding activities, while particular combinations of additional domains define the various myosin “classes” or “types”. In their recent comprehensive analyses of myosin diversity, Odrontiz and Kollmar [4] identified 35 myosin classes at the eukaryote scale, of which 3 were present in the last common eukaryotic ancestor. The most intensively studied myosins, class-II myosins (also called “conventional myosins”, myosin II, or MHCII) are characterised by the insertion of a glycin (at position 507) in the head domain, the presence of a SH3 domain N-terminal to the head domain, and a long C-terminal tail mostly comprised of a coiled-coil domain. Class II myosins originated in unikonts, i.e. eukaryotes ancestrally bearing a single flagellum or no flagellum, including the amoebozoa, fungi and holozoans (e.g. choanoflagellates and multicellular animals or Metazoa) [5].

Myosin II is a phylogenetically well-defined and diversified class [6], to which notably belong the well-known myosins that provide the physical force for muscle contraction in animals. Beyond these crucial functions in animal muscle cells, a wide range of non-muscle myosin II functions is documented. Myosin II is for example involved in amoeoboid motility in the unicellular life stage of the amoebozoan Dictyostelium discoideum [7]. In metazoans, various myosin II proteins play pivotal roles in cytokinesis [8,9], cell migration [10-12], cell-cell adhesion [12,13], or cell polarity [11,14,15]. Therefore, myosins II regulate fundamental aspects of cellular shape morphology [16-18], cytokinesis [19], cell differentiation [20,21] and more generally, many aspects of cellular behaviour [22]. Whatever the context (muscle or non-muscle), MHCII are integrated within macromolecular complexes notably through direct interaction with smaller proteins called “myosin light chains” (MLC) (which lack a head domain).

Previous analyses of myosin II proteins in bilaterian animals have recognised two phylogenetic groups, the first containing genes expressed in smooth muscle cells and in non-muscle cells, while genes of the second group are specifically expressed in striated (skeletal or cardiac) muscle cells [2,23]. This major dichotomy has been thought to reflect independent evolutionary origin from non-muscle cells for each major type of muscle cells, i.e. smooth muscle cells and striated muscle cells [1,24], through independent co-option of different myosin II paralogues. However, this suggestion was based on data from bilaterian animals only and remained highly speculative.

Outside from the bilaterian clade, muscle cells are present in two animal phyla, Cnidaria and Ctenophora, and this cell type is classically viewed as a synapomorphy (shared derived character) of the Eumetazoa (Cnidaria + Ctenophora + Bilateria), together with nerve cells. Most cnidian muscle cells are in fact multifunctional myoepithelial cells [25], integrated within the ectoderm and endoderm [26], although there are some reported instances of mesogleal muscle cells in cnidarians [27-30]. The myoepithelial cell of cnidarians typically comprises a contractile portion (the muscle fibre or myoid) attached to a globular cellular body involved in other functions (e.g. body protection, glandular secretion, fluid circulation through ciliary beating, etc.) [30]. In contrast, ctenophores are commonly considered to have true muscle cells, i.e. their muscle fibres lack a cellular body with filament-free cytoplasm ([31] but see [32] for an alternative point of view). In addition, part of the ctenophore musculature is located in the mesoglea and develops from a mesodermal-like germ layer [33,34]. Therefore, comparative studies using ctenophores have a great potential for improving our understanding of the early evolution of muscle cells and muscular protein families.

The phylogenetic position of ctenophores is still debated. Some recent phylogenomic analyses have placed them as the sister group to all other metazoans [35,36], but this result was probably due to an artefact of long-branch attraction [37-39]. According to the phylogenomic analyses of Philippe et al. [37], ctenophores and cnidarians form a coelenterate clade, sister-group to the bilaterians, within monophyletic Eumetazoa.

Ctenophores are marine animals with a highly original biradially-symmetrical body plan and featuring unique anatomical traits (Figure 1) [26,31]. Their main distinctive feature is a locomotory system consisting of eight distinctive meridional rows of comb plates (swimming paddles), each made of the many fused giant cilia of “polster cells”. At their aboral pole, ctenophores possess an apical sensory organ involved in equilibration and flanked by two elongated ciliated areas called polar fields. There are two distinct nerve nets extending throughout the body, the epithelial (or polygonal) nerve net and the mesogleal nerve net [40]. The gastrovascular system, of mainly endodermal origin, opens at one extremity by the mouth and at the other by two anal pores. The ramified gastro-vascular canal system allows water circulation and distribution of nutrients throughout the body. Ctenophores are hermaphrodite, with paired male and female gonads housed in the walls of eight endodermal meridional canals, placed under each of the comb rows. Most ctenophores catch prey by using a pair of long and contractile tentacles which bear lateral branches or tentillae on their oral side. The epidermis of
tentacles and tentillae is densely covered with adhesive cells called colloblasts, to which the prey adheres. Tentacles can extend from and retract into a tentacular sheath whose epithelial lining is continuous with the outer epidermis.

In the ctenophore order Cydippida, to which belong our model species *Pleurobrachia pileus*, two fundamental categories of muscle cells can be distinguished: parietal muscle fibres are located in the basal part of the ectodermal epithelium, above the basal lamina (Figure 2A, B, E), while mesogleal muscle fibres are positioned below the basal lamina and run through the mesoglea (Figure 2C, F) [31]. In addition, a special mesogleal musculature is housed in the core of tentacles and tentillae (Figure 2D, G). The very particular “mesogleal giant smooth fibres” of ctenophores belonging to the order Beroida (which lack the parietal musculature [31]) have been the subject of specific studies [41-44], while other muscle cell types, and muscles of other ctenophores, are much less known. Ctenophore muscles are considered to be of the smooth type due to the absence of a striation pattern on electronographies (except in the tentillae of the cydippid ctenophore *Euplokamis* [31,45]), but their mesogleal muscle cells are multinucleated [31], which represents a fundamental difference with the mononucleated smooth muscle cells of bilaterians.

In tentaculate ctenophores, myogenesis is particularly intense throughout the life span in the thickened tentacle base (tentacle root), where histogenesis continuously compensates for the loss of tentillae and tentacle pieces that are damaged upon feeding. Experimental data indicate that 36 hours are sufficient for regeneration of an entire tentacle [46]. Putative stem cells of the colloblasts and muscle cells have been recently characterised in the tentacle root of the ctenophore *Pleurobrachia pileus* by expression analyses of *Piwi* and *Vasa* genes and DNA-labelling experiments (Figure 3 in [47]). The muscle putative stem cells and progenitors are localised along a median ridge in the symmetry plane of the tentacle root on its internal face [47-51]. Thanks to these characteristics, the tentacle root is a particularly suitable model to investigate myogenesis in ctenophores.

We studied the expression of three paralogous class II myosin (*MHCII*) genes in the adult of the cydippid ctenophore *Pleurobrachia pileus*. In parallel, the complex organisation of muscle fibres throughout the *Pleurobrachia pileus* body was investigated using phalloidin staining to help understand the gene expression patterns. One of the *MHCII* paralogues was specifically expressed in muscle cells (parietal and mesogleal) while the other two had non-muscle expression. Several evolutionary scenarios are proposed to interpret the ctenophore *MHCII* expression data in light of our reconstruction of the *MHCII* gene phylogeny and of the known functions of bilaterian *MHCII* genes.

**Methods**

**Specimen collection**

Adult specimens of *Pleurobrachia pileus* were collected in Villefranche-sur-Mer and in Roscoff (France) between March and June using specific plankton nets. They were
Figure 2 (See legend on next page.)
kept at 16°C in filtered natural seawater, under continuous water circulation and fed daily with *Artemia* nauplii.

**Blast searches and phylogenetic analyses**

MHCII sequences were retrieved by TBLASTN searches using *Mus musculus* MHCII sequences on two *Pleurobrachia pileus* transcriptome assemblies: (i) Phrap assembly of a collection of about 36,000 ESTs, sequenced from animals collected in Villefranche-sur-Mer (France) by the Sanger method at the Genoscope (Evry, France) (see [47] for details), and publicly available in dbEST (http://www.ncbi.nlm.nih.gov/nucest?term=pleurobrachia) and (ii) Newbler assembly of one run of 454 sequencing of total RNA extracted from mixed embryonic, larval, and adult stages with starting material obtained from Roscoff (France).

**Figure 2** *Pleurobrachia pileus* (ctenophore) muscle cells. (A-D) Transmission electron microscopy (TEM) views of *Pleurobrachia pileus* muscle cells. (A, B) Sections through the external epithelium showing a parietal muscle cell (Mu) in transverse section (A) and in longitudinal section (B). (C) Section showing a portion of mesoglea with a mesogeleal muscle fibre (Me f) cut obliquely. (D) Tentacle muscle cells (Mu) in transverse section. (E-G) Pictures showing the aspect of the main muscle cell types on phalloidin-stained preparations. (E) Parietal muscle fibres. (F) Close-up of a mesogeleal muscle fibre showing its ramified extremity and its multinucleate organisation. (G) Multiple parallel muscle fibres in a tentacle (Tc) and two tentillae (Tt). BL: Basal Lamina; Me: Mesogelea; Me f: Mesogeleal muscle fibre; Mu: Muscle fibre; Nu: Nucleus; Tc: Tentacle; Tt: Tentilla. Scale bars: A, B: 1 μm; C: 2 μm; D: 5 μm; E, F, G: 20 μm.

**Figure 3** Phylogenetic analysis of *Pleurobrachia pileus* MHCII sequences. Amino-acid sequences were analysed using the Maximum likelihood (ML) method. Numbers associated with the branches are ML bootstrap values (1000 replicates). Sequences from *Pleurobrachia pileus* are indicated in bold font and green colour. The MHCIIa clade is indicated in blue and the MHCIIb clade in purple. The letter between parentheses after the name of mouse genes of the MHCa clade indicate whether the gene is expressed in non-muscle cells (n) or in smooth muscle cells (s).

Abbreviations for species names: Aae: *Aedes aegypti*; Aqu: *Amphimedon queenslandica*; Asu: *Ascaris suum*; Cow: *Capsaspora owczarzaki*; Ddi: *Dictyostelium discoideum*; Dme: *Drosophila melanogaster*; Hma: *Hydra magnipapillata*; Lgi: *Lottia gigantea*; Lpe: *Loligo pealei*; Mbr: *Monosiga brevicollis*; Mmu: *Mus musculus*; Nve: *Nematostella vectensis*; Ppi: *Pleurobrachia pileus*; Sar: *Sphaeroforma arctica*; Sro: *Salpingoeca rosetta*; Tad: *Trichoplax adhaerens*. 
The alignment was constructed using published sequences and sequences retrieved by TblastN searches on public databases for a representative taxon set. Sequences were aligned using MUSCLE [52]. Ambiguous regions of the alignment were removed using Gblocks [53]. The initial alignment (before removal of ambiguously-aligned regions) is provided in Additional file 1. This alignment contains the head domain of several myosins belonging to classes other than class II (outgroups). After pruning of ambiguously-aligned sites, two final sub-alignments were extracted to take into account the absence of sequence data for the head domain on public databases for a representative taxon set. Sequences and sequences retrieved by TblastN searches on public databases were used to account for the absence of sequence data for the head domain. Two final sub-alignments were extracted to take into account the absence of sequence data for the head domain in one of the ctenophore sequences (PpiMHCIIb2): (i) a first alignment containing the outgroup sequences (non-class II myosins). The PpiMHCIIb2 gene was excluded from this alignment because its partial sequence contains only the tail and therefore it has no alignable residues with non-class II myosins (alignment in Additional file 2; resulting tree shown in Additional file 3); (ii) a second alignment, in which non-class II myosin sequences have been eliminated, thus allowing inclusion of PpiMHCIIb2 (Alignment in Additional file 4; resulting tree, rooted on ameobozoan MHCII, shown in Figure 3).

Maximum-likelihood analyses were performed using the PhyML program [54], with the WAG model of amino-acid substitution and a BioNJ tree as the input tree. A gamma distribution with eight categories was used. The gamma shape parameter and the proportion of invariant sites were optimised during the searches. The statistical significance of the nodes was assessed by bootstrapping (1000 replicates).

In situ hybridisation

Adults were fixed at 4°C in 4% paraformaldehyde in 50% seawater and 50% PBST (10 mM Na₂HPO₄, 150 mM NaCl, pH 7.5, 0.1% Tween 20), for 1 hour then washed three times in PBST and dehydrated through a graded series of ethanol and stored in methanol at −20°C. The in situ hybridisation (ISH) protocol was as described in [55] but colour was developed with NBT/BCIP (Roche Diagnostics, Meylan, France). After ISH, samples were stained with DAPI (1 μg/ml) for 15 min to visualise DNA, and then washed 3 times for 15 min in PBST. Before mounting in Citifluor solution (Oxford instruments SAS, Saclay, France), animals were dissected. Longitudinal and transverse sections were performed to clarify the precise distribution of the gene expression patterns. For longitudinal views of the median expansion of the tentacle root, the lateral expansions were removed with forceps. Negative controls (with a sense probe and without any RNA probe) performed in parallel showed no staining after extensive revelation, except occasionally in a few isolated cells in the general epithelium (between adjacent comb rows, around the apical organ). For this reason, staining of isolated epithelial cells occasionally obtained with MHCII probes was not taken into account.

Immunofluorescence

Animals were fixed in 4% paraformaldehyde in 50% seawater and 50% PBST for 30 min, at room temperature, then samples were washed several times in PBST, dehydrated through a graded series of ethanol and stored in methanol at −20°C. After re-hydration to PBS, samples were permeabilised with Triton-X100 (0.2% in PBS, then 0.01% in PBS, 10 min at room temperature). After blocking with 1% bovine serum albumin, samples were incubated with the rat monoclonal anti-tyrosylated α-tubulin or YL1/2 antibody (1:1000 dilution in PBS-Triton-X100 0.01%, BSA 1%), (Morphosys AbD Gmbh, Düsseldorf, Germany) for 4 hours at room temperature. After washing with PBS Triton-X100 0.01%, samples were incubated overnight at 4°C with the Alexa Fluor 568 goat anti-rat IgG secondary antibody. Dilutions of primary and secondary antibodies were made using 1X PBS containing 0.01% Triton-X100.

Phalloidin staining

For phalloidin staining, specimens were not dehydrated after fixation. They were incubated for 45 min in a dilute Phalloidin-TRITC (Sigma-Aldrich, St-Quentin-Fallavier, France) solution (10 μg/ml in PBST) and rinsed three times in PBST. All specimens were finally stained with DAPI (1 μg/ml) for 15 min for DNA visualisation, and then washed three times for 15 min in PBST. They were micro-dissected and mounted in Citifluor solution (Oxford instruments SAS, Saclay, France).

EdU labelling of DNA-replicating cells

EdU (ethynyl deoxyuridine) is a thymidine analogue similar to the classical BrdU but quicker and easier to use [56]. EdU incorporation assays were done using the Click-it EdU Alexa Fluor 488 Imaging Kit from Invitrogen (Cergy-Pontoise, France). The protocol was as described in [47]. We performed 12 hours of pulse and no chase to visualise proliferating cells in the aboral region.

Sections for light and transmission electron microscopy (TEM)

Two types of sections (cryosection and ultra-thin section) were performed. For cryosectioning, tentacle roots extracted after whole mount ISH were incubated for two days in PBST 1X, 15% sucrose at 4°C, then for 2 hours in PBST 1X, 15% sucrose, 7.5% pig gelatine. Then, blocks were frozen at −65°C in 2-methyl-butane. Cryosectioning was done on a Leica CM3050 S cryostat or on a Jung Frigocut 2800E cryostat, at a thickness of 14 μm.
Slices were mounted in Citifluor (Oxford instruments SAS, Saclay, France).

*Pleurobrachia pileus* living specimens for ultra thin sections were fixed for 10 minutes at room temperature in 3% glutaraldehyde, 0.1 M sodium cacodylate pH 7.3, 0.3 M sodium chloride, 0.05% OsO₄ (modified after [57]). They were then transferred to the same solution without OsO₄ for 2 hours. Specimens were then post-fixed for 1 hour in 1% OsO₄, 1.5k-ferricyanide, 2.5% NaHCO₃ pH 7.2 and 0.25 M sodium chloride (modified after [58]). Finally, material was dehydrated through an ethanol series, and embedded in Spurr. Sections were done using a Leica Ultracut R ultra-microtome, at a thickness of 60 nm for ultra thin sections.

**Imaging**

Pictures of *Pleurobrachia pileus* entire specimen and tentacle root observed in toto were acquired on a stereo-microscope Olympus SZX12 using a Q-imaging QICAM with Image pro software (MediaCybernetics, Bethesda, MD). All fluorescence and DIC images were acquired on an Olympus BX61 microscope using a Q-imaging Camera with Image Pro plus software (MediaCybernetics, Bethesda, MD). To enhance some details, Higauss filter was used on some pictures (from Image Pro). TEM images were acquired on a Jeol JEM-1400 equipped with a Morada (SIS) camera at the “Centre Commun de Microscopie Appliquée” (CCMA) (Université de Nice Sophia-Antipolis, Faculté des Sciences, Nice).

**Results**

**Ancestral duplication of class II Myosin Heavy Chain genes in metazoans**

By Blast searches on *Pleurobrachia pileus* ESTs and phylogenetic analyses we could identify three ctenophore Myosin Heavy Chain II genes (MHCII). Although these searches were conducted on relatively extensive transcriptomic data, and myosin genes are expected to be expressed at a high level, we cannot exclude the existence of additional paralogues in the *Pleurobrachia pileus* genome. Of these three identified *Pleurobrachia pileus* paralogues, one (*PpiMHCIIa*) falls in a clade containing all bilaterian MHCII genes expressed in smooth muscle cells and/or non-muscle cells (MHCIIa clade: blue colour in Figure 3), while the two remaining genes (*PpiMHCIIb1* and *PpiMHCIIb2*) branch with bilaterian MHCII genes expressed in striated muscle cells (MHCIIb clade, highlighted in purple colour in Figure 3).

We conducted two phylogenetic analyses with different outgroup gene samplings, providing essentially similar results: an analysis rooted using the MHCII sequence of the amoebozoan *Dictyostelium discoideum* (Figure 3) and another analysis, rooted with sequences of myosin classes V, VII and X (Additional file 3) (see Methods for explanations).

The presence of genes from ctenophore, cnidarians (*Hydra magnipapillata* and *Nematostella vectensis*) but also sponge (*Amphimedon queenslandica*) and placozoan (*Trichoplax adhaerens*) in both MHCII clades indicates that the duplication happened before the last common ancestor of metazoans. Other holozoans (*Monosiga brevicollis, Salpingoea rosetta, Capsaspora owczarzaki* and *Sphaeroforma arctica*) each have one orthologue of MHCIIa but no MHCIIb gene, suggesting that the MHCIIa/MHCIIb duplication occurred in the last common ancestor of holozoans and was followed by independent MHCIIb losses in the various unicellular holozoan lineages. An alternative explanation could be that the duplication took place in a metazoan ancestor, but that sequences of MHCIIb are misplaced in the tree due to a higher subsequent sequence divergence. In vertebrates, the subdivision of MHCII genes into MHCIIa and MHCIIb orthology groups is clearly correlated with functional specialisation, MHCIIb genes being associated with striated muscle cells and MHCIIa genes being expressed in either non-muscle or smooth muscle cells. In addition, independent diversification within both groups occurred in vertebrates (Figure 3, Additional file 3). Finally, the two ctenophore MHCIIb paralogues *PpiMHCIIb1* and *PpiMHCIIb2* are quite divergent but they clearly originated from a ctenophore-specific duplication within the MHCIIb group (Figure 3).

The three MHCII paralogous genes are differentially expressed in the *Pleurobrachia pileus* tentacle root

In the tentacle root, *PpiMHCIIa* is expressed in the putative stem cells/undifferentiated progenitors of colloblasts and muscle cells. The morphology of the tentacle root (in internal, external and longitudinal views) is summarised in Figure 4A-A”.

The expression of *PpiMHCIIa* closely looks like that of the “stem cell genes” investigated in Alié et al. [47]. On the internal side of the tentacle root, the *PpiMHCIIa* antisense RNA probe stains three longitudinal lines running from the oral to the aboral pole of the tentacle root (Figure 4B): one median line and two lateral lines. In longitudinal section (Figure 4C), the median staining appears confined to the distal extremity or ridge of the median expansion, although it is more diffuse towards both poles. In transverse sections of the tentacle root after whole-mount *PpiMHCIIa* ISH, transcripts appear clearly confined to the three ridges of the tentacle root expansions (Figure 4D, E), where putative stem cells are localised according to a previous study [47]: muscle stem cells in the median ridge and colloblast stem cells in the lateral ridges. Transcript distribution appears more diffuse when the sectioning plane is closer to the oral pole.
Figure 4 (See legend on next page.)
Figure 4 Expression of PpiMHCIIa, PpiMHCIIb1 and PpiMHCIIb2 in the tentacular apparatus. (A) Schematic drawing of the tentacle root in internal view. The three coloured horizontal planes materialise the three different planes of transverse cryosections corresponding to panels D, I, N (in green), E, J, O (in orange) and F, K, P (in purple). (A’) The tentacle root in external view (corresponding to the side of tentacle insertion). The dotted lines delineate the longitudinal dissections to remove tentacle root lateral expansions in order to obtain the preparations shown in (C, H, M). (A”) Schematic drawing of the tentacle root in longitudinal section, after removal of the lateral expansions following the dotted lines in (A’). Horizontal lines indicating the three planes of cryosections as in (A): (B, G, L) Internal views of isolated tentacle roots showing PpiMHCIIa (B), PpiMHCIIb1 (G), PpiMHCIIb2 (L) expressions. (C, H, M) Longitudinal sections of the tentacle root stained with PpiMHCIIa (C), PpiMHCIIb1 (H), PpiMHCIIb2 (M) anti-sense probes. The aboral pole is at the top in panels (B, C, G, H, L, M; D, F, I, K, N, P) Transverse cryosections of whole-mount ISH for the three genes, with sectioning plane indicated by the colour of the surrounding line according to the colour code outlined in panel (A). The black arrowhead in (G–J) points to a stained line above the median ridge which appears to be more or less in continuity with the two layered bands labelled M Tcl. (Q) YL1/2 (anti-tyrosylated-a-tubulin, in red) and DAPI (in blue) counterstaining of the region boxed in (N, R). Higher magnification of the region indicated by the box in (Q). (S) YL1/2 (red) and DAPI (blue) counterstaining of the region boxed in (P). (T–W) Expression of PpiMHCIIb1 gene in tentillae. (T) Whole mount ISH of tentacle and tentillae. (U) Higher magnification view of the region boxed in (T). (V) Transverse cryosection of whole-mount ISH of a tentilla. Two symmetrical muscle fibres are stained. (W) YL1/2 (in red) and DAPI (in blue) counterstaining of (V). The YL1/2 staining in colloblasts is certainly due to non-specific fixation of the antibody on the sticky colloblast granules. (X) Schematic drawing of a tentilla in transverse section. Coll: Colloblasts; F: Forming tentillae; LR: Lateral Ridge; MR: Median Ridge; M tcl: Tentacle; Tt: Tentilla. Scale bars: B, C, G, H, L, M: 100 μm; D–F, I–K, N–P: 200 μm; Q: 100 μm; R, S: 10 μm; T: 50 μm; U, V, W: 25 μm.

(See figure on previous page.)

The PpiMHCIIb1 gene is expressed in the differentiating muscle cells of the tentacle root. In internal view (Figure 4G), transcripts are detected in two neighbouring but disconnected regions, corresponding respectively to the muscle differentiation zones of the tentacle and of the tentillae: (i) two wide bands (M Tcl in Figure 4G) positioned on both sides of the median expansion; (ii) an additional domain towards the oral pole (M tt in Figure 4G), where the tentillae are produced. In the longitudinal view section, these two expression domains are also clearly distinct (Figure 4H), with the first here appearing as a wide fan-shaped stained area converging to the base of the tentacle (M tcl in Figure 4A”, H), in addition to a thin stripe located just above the median ridge (black arrowheads in Figure 4G–I). These two PpiMHCIIb1 expression domains are also shown in transverse section (sectioning planes indicated in Figure 4A): expression in differentiating muscle cells of the tentacle axis is visible on the two aboral-most sections (M tcl in Figure 4I and 4J) and expression in differentiating muscle cells of tentillae is visible on the two oral-most sections (M tt in Figure 4J and 4K).

This interpretation of PpiMHCIIb1 expression patterns in the tentacle root is supported by our TEM observations, which identify four distinct stages of tentacle muscle differentiation from stem cells to fully differentiated muscle cells (Figure 5B–E; summary drawing in transverse section in Figure 6). The first step corresponds to cell multiplication in the median ridge (Figure 5B and see EdU incorporation assays described in [47]). The 2nd step is represented by spindle-shaped mononucleate progenitors found laterally and aligned in an oblique orientation (Figure 5C–C’; light orange in Figure 6). These cells are connected to each other by cytoplasmic bridges (Figure 5C; white arrowheads). A more advanced cell stage (step 3) is represented by multinucleated cells (up to 8 nuclei per cell counted), located closer to the symmetry plane of the tentacle root (Figure 5D; dark orange in Figure 6), indicating that progenitors have undergone cell fusion. These cells contain well-developed endoplasmic reticulum and many mitochondria, but are still devoid of myofilaments. Finally, mature muscle cells are observed in two symmetric and median bundles (red in Figure 6), externally surrounded by the two preceding less differentiated cell populations. These differentiated muscle cells are multinucleated and contain dense longitudinal myofilaments as well as many mitochondria (Figure 5E). The first expression domain of PpiMHCIIb1 described above closely matches the differentiation zone of tentacle muscle (orange colours in Figure 6), an interpretation also supported by the strong expression observed for the muscle differentiation marker gene Tropomyosin in exactly the same zone (Additional file 5).

The formation of tentillae is an independent process occurring at the oral pole of the tentacle root. The...
second domain of PpiMHCIIb1 expression (M tt in Figure 4G, H, J, K) corresponds to the production zone of the tentillae, and therefore the gene is also expressed in differentiating muscle cells of the tentillae. More aborally, PpiMHCIIb1 shows a striped pattern in the youngest formed tentillae (F tt in Figure 4H).

In the mature tentacular apparatus, PpiMHCIIb1 is expressed in the muscle masses housed within the tentillae, but not in those of the tentacle itself (Figure 4T: note the absence of staining in the tentacle, Tcl). In longitudinal and transverse sections of mature tentillae, PpiMHCIIb1 expression is seen in two parallel bundles corresponding to the position of muscle fibres (Figure 4U-X). These muscle bundles are separated by an unstained central nerve cord, revealed by counter-staining using the structural marker YL1/2 (anti-tyrosylated-α-tubulin antibody) (Figure 4W, 4X), known to stain nerve cells with high intensity in Pleurobrachia pileus [40]. The expression pattern of PpiTropomyosin in the tentilla is identical to that of PpiMHCIIb1 (compare Additional file 5D with Figure 4V), supporting the muscular nature of PpiMHCIIb1-expressing cells. The fact that PpiMHCIIb1 and PpiTropomyosin genes (see Tcl in Additional file 5C) are expressed in the mature muscles of tentillae but not of the tentacle (although these genes are expressed in the differentiating muscle of both) indicates that these two kinds of contractile structures have muscle fibres of different types. It is therefore clear that in spite of their topological relationships, tentillae are not just ramifications of the tentacle but instead are original organs, both ontogenetically and structurally.

Figure 5 Stages of tentacle muscle differentiation observed using TEM. (A) DAPI staining of a transverse section of the tentacle root. The light blue line delineates the tentacle root. (B-E) TEM images of myogenesis stages in the median expansion of the tentacle root. (B) TEM view of the region boxed in yellow in (A), showing the muscle stem cells / progenitors. (C) TEM view of the region boxed in orange in (A) showing spindle-shaped muscle cell progenitors. (D) Higher magnification of the region boxed in (C). White arrowheads point to cytoplasmic connections and plasma membrane interdigitations. (E) TEM view of the region boxed in red in (A), showing a muscle cell in differentiation (delineated by the white dotted line), and resulting from the fusion of several muscle cell progenitors, as indicated by the multinucleate organisation of the cell. These cells exhibit a developed endoplasmic reticulum, many mitochondria and are devoid of myofilaments. (F) TEM view of the region boxed in purple in (A), showing a mature muscle cell, with aligned nuclei (Nu) and distinct myofilaments (Myo). White holes in (B-D) are fixation artefacts. LR: Lateral Ridge; Mi: Mitochondria; MR: Median Ridge; Myo: Myofilaments; Nu: Nucleus. Scale bars: A: 100 μm; B: 10 μm; C, C’: 2 μm; D: 5 μm; E: 2 μm.

Figure 6 Dynamic model of myogenesis in the Pleurobrachia pileus tentacle root. Drawing of a tentacle root transverse section at the aboral level illustrating the proximal-distal distribution of myogenesis stages. The four recognised stages along the muscle cell lineage are indicated in bold.
The expression pattern of the other MHCIIb paralogue \textit{PpiMHCIIb2} is sharply different from that of \textit{PpiMHCIIb1} in the tentacle root, as it seems to associate with neural cells instead of muscle cells. In internal and longitudinal views, \textit{PpiMHCIIb2} expression is visible as a thin stripe above the median ridge, which becomes broader at the oral pole and does not reach the oral extremity (Figure 4L and M). In transverse sections, transcripts were detected in a median group of cells located between the two bundles of differentiated muscle cells (N tcl in Figure 4N and O). Counter-staining of these transverse sections using the anti-tyrosylated-α-tubulin antibody (YL1/2) reveals that the \textit{PpiMHCIIb2}-expressing zone corresponds to a nerve plexus (Figure 4Q). A closer examination of this area reveals the presence of two symmetrical layers of nerve cells sandwiched between tentacle muscles (Figure 4R, Figure 6). This nerve plexus also runs through tentacle and tentillae (data not shown); it is depicted (white area and black line in the median expansion) in Figure 6.

Towards the oral pole, \textit{PpiMHCIIb2} is also expressed in the production zone of the tentillae (N tt and F tt in Figure 4M), in the form of a thin external stripe of positive cells at the interface between the lateral and median expansions (N tt in Figure 4O and P). YL1/2 and DAPI counterstaining reveal the presence of densely packed neural cells with small nuclei on a thin layer corresponding to the zone of \textit{PpiMHCIIb2} expression (Figure 4S, counterstaining of the region boxed in Figure 4P). At the level of the youngest formed tentillae, \textit{PpiMHCIIb2} is expressed in a striped pattern, probably corresponding to the nerve cords of each tentilla (Ftt on Figure 4M and N). However, mature tentillae do not show any \textit{PpiMHCIIb2} expression signal, suggesting that \textit{PpiMHCIIb2} is expressed in a sub-population of neural cells, or at a particular stage of neuron differentiation.

Other sites of \textit{PpiMHCIIa} expression are associated with putative stem cells and proliferating progenitors

The expression of \textit{PpiMHCIIa} is consistently associated with undifferentiated dividing cells in the adult \textit{Pleurobrachia pileus}. Within the aboral sensory complex, \textit{PpiMHCIIa} is expressed in eight patches of cells at the proximal extremities of the polar field marginal and central zones (Figure 7A and A'). These cells present a high nucleocytoplasmic ratio and are densely packed (data not shown). The four internal-most patches (black arrowheads in Figure 7A) correspond to the four stem cell pools identified by Alié et al. [47]. Therefore, \textit{PpiMHCIIa} expression encompasses these stem cell populations but is not restricted to them. In fact, the distribution of \textit{PpiMHCIIa} expression in the aboral sensory complex closely matches the distribution of proliferating cells as revealed by Edu incorporation experiments with a 12-hour pulse and no chase (see Figure 7B' in [47], and Figure 7A" in this paper).

The \textit{PpiMHCIIa} gene is strongly expressed in both female and male gonads. In \textit{Pleurobrachia pileus}, eight meridional canals located under the eight comb rows contain the gonads in their walls. The male and female germlines are spatially segregated at both sides of each canal. Expression is observed in the entire female gonad (Figure 7B and B') where young and mature oocytes as well as nurse cells are stained (Figure 7B'). In the male gonad, transcripts are restricted to spermatogenesis stages, in the most peripheral part of the gonad, while the internal-most part of the canal wall (corresponding to mature stages \textit{i.e.} spermatozoids) is not stained (Figure 7B', B'). To conclude, \textit{PpiMHCIIa} is expressed in the germ cells of \textit{Pleurobrachia pileus}, at all stages of oogenesis and during spermatogenesis but not in mature spermatozoids.

Within the comb rows, \textit{PpiMHCIIa} is expressed in the forming combs and at both extremities of the mature combs (Figure 7C-G'). Recognisable by their small size and characteristic position (see [47]), oral and aboral baby combs are entirely stained (Figure 7C, C', D, D', E). In mature combs, transcripts are restricted to two crescent shape areas at both extremities of each comb (Figure 7F, G). The observation of DAPI counterstaining indicates that positive cells possess a round nucleus with prominent nucleolus, whereas the unstained poster cells have elongated and twisted nuclei (Figure 7G'). These expression sites correspond to progenitors of the polster cells (ciliated cells of the combs) as identified in a previous work [47].

\textit{PpiMHCIIb1} is specifically expressed in various kinds of muscle cells

\textit{PpiMHCIIb1} is expressed in both kinds of muscle cells described in \textit{Pleurobrachia pileus}: parietal and mesogleal muscle fibres. Using phalloidin-stained preparations, we could reconstruct the complex arrangement of these muscle fibres throughout the \textit{Pleurobrachia pileus} body (Figure 8 and Additional file 6). An important observation (Figure 8) is that the parietal musculature consists mainly of fibres placed in a circular orientation (= latitudinal fibres) and present throughout the body wall, whereas longitudinal parietal fibres are restricted to a few particular regions of the body: (i) aborally, the four areas delineated by the four pairs of adjacent ciliated grooves; (ii) in the tentacular plane, two symmetrical muscle bands extending from each tentacle sheath opening to the mouth region; (iii) in the oral region, the area comprised between the oral extremity of the comb rows and the mouth. At least for the two latter sites, the presence of longitudinal fibres is clearly linked to a functional requirement for strong and relatively rapid contractions allowing changes in mouth orientation and shape during feeding [59]. The arrangement of mesogleal fibres is more complex and is only outlined in Figure 8 for those located in the equatorial
These mesogleal fibres can notably connect two adjacent meridional canals (looped fibres on Figure 8 and Additional file 6 B), a meridional canal and the external epithelium (Figure 8) or the apical organ and the tentacle sheath (Additional file 6 H).

After a short time of ISH-signal development, PpiMHCIIb1 expression in the parietal musculature is detected only in longitudinal fibres (Figure 9 A, C-D). Notably, PpiMHCIIb1 is strongly expressed in the two symmetrical muscle bands that run from the tentacle sheath openings to the mouth (Figure 9 C-D). A neural condensation called the juxta-tentacular nerve cord [40] is located in the centre of these muscle bands (seeYL1/2 immunostaining and phalloidin staining in Figure 9 E). There is also strong PpiMHCIIb1 expression in the numerous short longitudinal fibres that surround the mouth opening (Figure 9 B). When the ISH signal is allowed to develop for a longer time, the whole parietal musculature (including the latitudinal fibres throughout the body wall) becomes stained (not shown). The observed difference in terms of staining intensity between latitudinal and longitudinal parietal fibres might reflect either different levels of PpiMHCIIb1 gene expression, or an optical effect due to the smaller diameter and looser organisation of latitudinal fibres compared to longitudinal ones (well visible in Figure 9 E).

Figure 7 PpiMHCIIa expression outside the tentacle root. (A) PpiMHCIIa expression in polar fields. Eight patches of PpiMHCIIa are identified. The black arrowheads point to the four internal-most patches (thought to correspond to stem cells) and the white arrowheads indicate the 4 external-most patches (thought to correspond to proliferating cells but not stem cells). (A') Summary of the PpiMHCIIa expression pattern (blue patches) in polar fields. (A') EdU labelling in the aboral sensory complex after a 12 h pulse and no chase. (B) Expression of PpiMHCIIa in the gonads (male on the left and female on the right). Spermatoids (Spz) are not stained. (B') DAPI counter-staining of panel (B). (B'') Oocyte (Oo) surrounded by its nurse cells, both expressing PpiMHCIIa. (C) PpiMHCIIa expression at the aboral extremity of a comb row (the black arrow points to an aboral baby comb). (C') DAPI counter-staining of panel (C). The white arrow points to the aboral baby comb. (D) PpiMHCIIa expression in the youngest baby comb at the oral extremity of a comb row. (D') DAPI counter-staining of panel (D). (E) PpiMHCIIa expression in the six youngest combs located at the oral extremity of a comb row of another specimen. (F) PpiMHCIIa expression in a mature comb. (G) Detailed view of the lateral extremity of a mature comb showing the localised expression of PpiMHCIIa. (G') DAPI counter-staining of panel (G). ao: apical organ; Cpf: Central zone of polar field; Mpf: Marginal zones of polar field; Ps: Polster cells; Spg: Spermatogenesis stages; Spz: Spermatoids. Scale bars: A, A', C, C': 50 µm; B, B': 100 µm; B': 10 µm; D, D': 5 µm; E, F: 25 µm; G, G': 20 µm.
observation of phalloidin and DAPI stained preparations revealed that parietal muscle cells are multinucleated (like the mesogleal muscle cells) (Figure 9F).

PpiMHCIIb1 is also expressed in mesogleal muscle fibres (Figure 9G-J). Strong PpiMHCIIb1 staining is notably observed in mesogleal muscle fibres attached to the meridional canals (Figure 9G, H). Some of the PpiMHCIIb1 expressing mesogleal fibres are shorter and connect the two paragastric canals to the mouth (Figure 9I, J). Phalloidin and DAPI staining confirmed that the mesogleal fibres are multinucleated (with elongated nuclei) (Figure 2F). These observations furthermore indicate that mesogleal muscle fibres are on average much thicker than parietal muscle fibres and are ramified at both extremities where they attach to the epidermis or to the meridional canals (Figure 2F).

Finally, our data on PpiMHCIIb1 expression provide interesting insights into the nature of an enigmatic type of fibres (here called “inter-comb fibres”) tightly associated with the combs (Figure 9K). These fibres are conspicuously stained with phalloidin and they extend medially from one comb to the following one, with shorter lateral fibres occurring only on the oral side of each comb (Figure 9K). These structures have been observed previously by Chun [49], erroneously interpreted as extensions of the polster cells in Hormiphora by Samassa [60], and more recently observed by Hernandez-Nicaise [61]. Although they closely resemble muscle fibres in phalloidin-stained preparations, inter-comb fibres do not show any PpiMHCIIb1 expression and recent observations suggest that they do not contract following stimulation (L. Moroz, personal communication). Therefore, they are probably not muscular but could represent supporting fibres, possibly maintaining the mechanical cohesion of the comb row against the tension generated by comb beating.

**Outside the tentacle root, PpiMHCIIb2 is expressed in various types of non-muscle cells**

PpiMHCIIb2 is expressed in different cell types in Pleurobrachia pileus but apparently not in muscle cells. Close examination of the floor of the aboral organ reveals the presence of two distinct sites of expression: two peripheral spots close to the epithelial papillae (Figure 10A, black arrowheads; Figure 10A'), and two paired internal stripes, located between the balancers (Figure 10A, white arrowheads; Figure 10A'). Cells surrounding the anal pores are also stained by PpiMHCIIb2 (Figure 10B).

Moreover, PpiMHCIIb2 is also expressed in the female gonad (within the body walls of the meridional canals), but here the staining is restricted to growing oocytes and their surrounding nurse cells (Figure 10C-C') unlike PpiMHCIIa which is expressed throughout the ovary (Figure 7B). Oocyte and nurse cells are known to belong to the same cell lineage [33,62,63]. Another site of PpiMHCIIb2 expression are the paragastric canals, where staining is restricted to particular clusters of cells (of unknown nature) on the edges of the canals (Figure 10D).

In addition, PpiMHCIIb2 has a complex expression pattern within mature combs, in (i) both lateral extremities of the comb (Figure 10E, E'), which might reflect PpiMHCIIb2 expression in polster cell stem cells/progenitors, (ii) a subterminal spot (star in Figure 10E') and (iii) two small symmetrical groups of cells in the centre of each comb (Figure 10E; black arrowhead). These latter two cell sub-populations have not been previously distinguished morphologically. A last expression site of PpiMHCIIb2 is observed in the ectodermal epithelium along the tentacular plane (white arrowheads in Figure 10E). This stripe of PpiMHCIIb2 expression is located close to the juxtatentacular nerve cord and of the longitudinal muscle expressing PpiMHCIIb1 and PpiTropomyosin (Figure 9C-E). However, the PpiMHCIIb2-expressing cells in this region are not muscle cells but round-shape epithelial cells of unclear nature (data not shown).

**Discussion and conclusions**

**Conserved expression of MHCIIa genes in the germ line, in stem cells and in proliferating cells**
The bilaterian orthologues of PpiMHCIIa are expressed in smooth muscle cells or in non-muscle cells [64-67]. Some of the vertebrate members of this group of myosin II are classically known for their role in the contractile
activity of smooth muscle cells, but other members are essential for mitosis, in all cell lineages, being involved in the formation of the contractile ring during cytokinesis. Recent findings demonstrate that smooth/non-muscle myosin II also play an active role in various other cellular activities such as generation of cell polarity, cell migration and cell-cell adhesion (for a review see [11]).

In Pleurobrachia pileus, the expression pattern of PpiMHCIIa is strikingly similar to that of the "stem cell markers": Vasa, Piwi, Bruno and PL10 [47] and the SOX genes PpiSOX2 and PpiSOX12 [55]. Its expression territories include the male and female gonads and several spatially restricted pools of stem cells / progenitors involved in somatic cell renewal, within the comb plates, aboral sensory complex and tentacle root. Expression of PpiMHCIIa throughout the ctenophore germ line (except in the most mature stages of spermatogenesis) suggests a conservative role for non-muscle myosin II (MHCIIa clade) in gametogenesis, as proposed in different studies of vertebrate meiotic chromosomes [69,70]. In somatic territories, PpiMHCIIa is consistently expressed in the various populations of stem cells that were recently identified in the adult ctenophore using a combination of gene expression data and DNA-label incorporation and long-term retention assays [47]. This is particularly obvious in the tentacle root, where PpiMHCIIa expression is restricted to the three parallel ridges known to house stem cells. It is furthermore significant that PpiMHCIIa expression in the tentacle root is not limited to muscle stem cells (located in the median ridge) but also includes the colloblast stem cells (in the lateral ridges), suggesting a link with stemness/undifferentiated state/cell proliferation rather than with muscle cell identity. This conclusion finds additional support from PpiMHCIIa expression in stem cell pools located at both extremities of the combs (and involved in renewal of the ciliated "polster cells") as well as in the four internal stem cell pools previously identified around the apical organ.

When examined in detail, some aspects of the PpiMHCIIa expression pattern further suggest that beyond
stem cells, this myosin is associated with cell proliferation, including in progenitors that are already committed to undergo differentiation. In the aboral sensory complex, after a 12 h EdU pulse and no chase, eight discrete patches of nuclei are labelled around the apical organ (in the proximal region of the polar fields) (Figure 7B” in [47] and Figure 7A” in this paper). When a 5-day chase is added after the pulse, only four of these patches remain labelled. A likely explanation for this observation is that cells of the four other patches have become differentiated and therefore were proliferating progenitors but not stem cells. Whereas genes like Piwi and Vasa tend to have their transcripts highly concentrated in the four stem cell patches [47], PpiMHCIIa is equally expressed in eight patches whose position matches the eight labelled cell groups obtained after EdU pulse and no chase. PpiMHCIIa expression might also extend to proliferating progenitors in other tissues (tentacle root, combs) but the continuous spatial gradient of cell differentiation from stem cells in these tissues makes it more difficult to distinguish expression in stem cells vs. in progenitors. The suggestion that PpiMHCIIa expression

Figure 10 PpiMHCIIb2 expression outside the tentacle root. (A) PpiMHCIIb2 expression in the apical organ. White arrowheads point to paired structures localised near the balancers (b) and black arrowheads point to paired structures located near the epithelial papillae (ep). (A’) Summary of the PpiMHCIIb2 expression pattern in the apical organ. (B) PpiMHCIIb2 expression in the anal pores (Ap). (C) PpiMHCIIb2 expression in the female gonad. (C’) Higher magnification of the region boxed in (C), showing an oocyte (Oo) surrounded by its nurse cells (nc), both expressing PpiMHCIIb2. (D) Expression of PpiMHCIIb2 in a paragastric canal. (E) Expression of PpiMHCIIb2 along a comb row and in the neighbouring epithelium. White arrowheads indicate the tentacular plane, where a line of PpiMHCIIb2 expressing cells is observed. (E’) Higher magnification of the region boxed in (E). The black arrowhead points to the symmetrical expression spots in the middle of the comb. The star points to the “subterminal spot” of staining near the extremity of the comb. ao: apical organ; Ap: Anal pore; b: balancer cells; Cpf: Central zone of polar field; ep: epithelial papillae Mpf: Marginal zones of polar field; nc: nurse cell; Oo: Oocyte. Scale bars: A: 20 μm; B, D, E: 50 μm; C, E: 100 μm; C’, D’: 10 μm.
is associated with all kinds of proliferating cells in *Pleurobrachia pileus* is consistent with its involvement in the movements of the contractile ring upon cytokinesis in bilaterian models.

Finally, whereas vertebrate smooth muscle cells express *MHCIIa* genes [64-67], in ctenophore *PpiMHCIIa* is apparently not expressed in muscle cells, even if all of them are said to be of the smooth type [31]. The only case of *PpiMHCIIa* expression in a muscle cell lineage concerns the muscle progenitors along the median ridge of the tentacle root, but as explained above this has probably nothing to do with muscular identity but rather with stemness and/or cell proliferation.

**Independent functional specialisation of MHCII myosins and the possible convergent origin of ctenophore muscles**

At least some of the non-muscular functions played by MHCIIa proteins are certainly very ancient since myosins are cellular motors involved in various cellular processes (e.g. cytokinesis) throughout eukaryotes including members of unicellular lineages [71,72]. The MHCIIa/MHCIIb duplication predated the divergence of sponges and placozoans and therefore occurred well before the origin of muscle cells, implying that both MHCII sub-families were initially working in a non-muscular context. The ctenophore data furthermore suggest that *MHCIIa* genes were not involved in muscle contraction but had retained the pleiomorphic non-muscular expression in the last common ancestor of ctenophores and bilaterians, although muscle cells are generally thought to have been present in this ancestor ([37,73]. Expression of some MHCIIa myosins in the smooth muscle cells of vertebrates is certainly the result of a late co-option event, in consistence with the idea that vertebrate smooth muscle cells were independently evolved [1,24]. Accordingly, in scallop (a mollusc) smooth and striated muscles of the foot express two different splicing variants of the same MHCII gene, closely related to vertebrate striated muscle myosin II (MHCIIb) [74], indicating that functional specialisation of myosins according to muscle cell types (smooth vs. striated) occurred independently through different paths in different bilaterian lineages.

The expression of one of the ctenophore MHCIIb paralogues, *PpiMHCIIb2*, shows several striking similarities to that of *PpiMHCIIa* and other metazoan myosins of the MHCIIa subfamily. First, in contrast to the other parologue *PpiMHCIIb1* (expressed in muscle cells of any kind), *PpiMHCIIb2* has strictly non-muscular expression. Like *PpiMHCIIa*, this gene is expressed in putative stem cells and/or progenitors cells at both extremities of the comb rows, and also in the germ line. Here *PpiMHCIIb2* is expressed in a more restricted territory than *PpiMHCIIa* (i.e. only within large oocytes and their surrounding nurse cells in the ovary). This might reflect functional specialisation between *PpiMHCIIa* and *PpiMHCIIb2* during gametogenesis, perhaps in cell proliferation for the former vs. formation of the cytoplasmic bridges that link the oocyte and the nurse cells (both deriving from common precursors [33,63]) and/or cytoplasmic transport from the nurse cells to the oocyte, as documented for non-muscle myosin II in *Drosophila* oogenesis [75,76], for the latter. Additionally, distinct sub-populations of nerve cells express *PpiMHCIIb2* in at least two regions of the body: a nerve plexus in the symmetry plane of the tentacle root, and several territories in the epithelial floor of the apical organ, exactly matching the previously-reported distribution of antivasopressin immunoreactive neuro-sensory cells (see Figure 6D-F in [40]). This neural expression of *PpiMHCIIb2* reminds the involvement of some *MHCIIa* genes in the nervous system of various vertebrates and *Drosophila*, where they promote synaptic vesicle assembly and movements through interactions with F-actin partners, neuronal migration, neuronal morphogenesis and growth cone motility [77-80].

The apparently odd MHCIIa-like expression pattern of *PpiMHCIIb2* can be easily explained if ctenophore muscle cells evolved independently from those of cnidarians and bilaterians. Under this scenario, when the ctenophore lineage originated, the two single-copy genes *MHCIIa* and *MHCIIb* would have still retained the pleiomorphic non-muscular functions.

Then MHCIIb duplicated in ctenophores, and one of the resulting paralogues (*PpiMHCIIb1*) underwent neo-functionalisation in the context of a newly-evolved type of contractile fibre cell. Yet it is possible to imagine alternative scenarios, not assuming independent muscle cell origins. If muscle cells emerged only once in a eumetazoan ancestor with MHCIIb correlatively becoming exclusively muscular, then the observed non-muscular expression of ctenophore *PpiMHCIIb2* must be interpreted as derived, and we would face a unique and rather weird instance of a formerly muscular myosin having shifted to non-muscular function. Furthermore, in that case, shared expression features between *PpiMHCIIb2* and *PpiMHCIIa* find no explanation and must be considered fortuitous. A last plausible scenario assumes that in the last common ancestor of Eumetazoa, MHCIIb had pleiotropic expression in both non-muscle cells (pleiomorphic role) and newly acquired muscle cells (apomorph role). Then bilaterians lost the non-muscular MHCIIb expression, whereas in the ctenophore lineage the MHCIIb duplication was followed by functional specialisation of the two copies, one taking over the muscular function and the other the non-muscular roles, a scenario conforming to the...
duplication-degeneration-complementation model [81]. It is finally quite clear that, albeit non-parsimonious with respect to the taxon phylogeny, the hypothesis of convergent muscle origins offers the most straightforward explanation to the observed ctenophore myosin expression patterns.

It is nevertheless important to underline that data from cnidarians will be essential to evaluate these scenarios but are insufficient at the moment. In the hydro-medusa Podocoryne carnea, a MHCIib group gene seems to be specifically expressed in the striated muscle layer of the medusa, located on the inner side of the bell [82], but expression data for cnidarian MCHIIa genes are critically lacking.

The originality and interest of ctenophores for studying muscle differentiation
A first striking characteristic of the ctenophore musculature is the diversity of muscle cell types and the complexity of their spatial arrangement (see Figure 8), as exemplified by our TEM observations combined with expression analyses of the MHCIib1 gene and phalloidin staining. The existence in ctenophores of two fundamentally distinct musculatures, epithelial (i.e. parietal muscles) and mesogleal, parallels the similar distinction recently confirmed using immunofluorescence [40] for the nervous system, and represents a fundamental difference with cnidarians. While the later are fundamentally epithelial animals, ctenophores have a fully developed mesogleal compartment with proper nervous and muscle systems, a condition more alike that of the triploblastic bilaterians. Furthermore in ctenophores, within each of the two main muscle categories, there is a significant diversity of muscle fibre subtypes differing by their size, position within the body, orientation, organisation and mode of formation (documented only for the tentacle root).

Among these ctenophore muscles, those of the tentacular apparatus (comprised of the tentacle and its attached tentillae) represent an autonomous muscle system not only from a functional point of view, but most importantly in terms of ontogeny and cell lineages. The ctenophore tentacle root is a highly promising model for studying cell differentiation and particularly myogenesis. Indeed, it is the only site of muscle cell production within the body where progenitors are easily identifiable and thereby are amenable to experimental approaches, whereas for parietal muscle cells and for other mesogleal muscle cell, we do not know where are the reservoirs of stem cells and progenitors. TEM observations and histological cryosections, combined with expression analyses of the three MHClI and Tropomyosin genes lead us to propose an original myogenic model implying spatially ordered regionalisation of muscle cell differentiation stages (Figure 6). This model is furthermore consistent with recent analyses of the progenies of somatic stem cells based on stem cell gene expression and EdU DNA labelling and long-term retention [47]. Whereas the main axis (median expansion) of the tentacle root is essentially a centre of muscle production, the two lateral expansions are mostly dedicated to the differentiation and maturation of colloblasts (adhesive cells), a ctenophore-specific cell type [31,47,48]. At the oral pole of the tentacle root, we identified an additional and independent centre, responsible for the production of tentilla muscles. Thus, the tentacle root contains at least three autonomous cell lineages, each with its particular niche of stem cells (note that the “aboral external cell masses” described in [47], probably represent a fourth cell lineage, of unknown identity).

The high renewal rate of tentacle cells (with complete tentacle regeneration in 36 hours [46]) makes this system highly suitable not only for studying regeneration processes but also for analysing the role of key regulators, such as myogenic genes, yet unknown in ctenophores. The spatial segregation of muscle cell stages in the tentacle root in transverse section should facilitate the identification of genes involved in the various steps of myogenesis. Therefore, the tentacle root is a promising experimental system to study stem cell regulation, cell specification and cell differentiation in ctenophores.

The limits of the “smooth” vs. “striated” classification of muscle cell types
The mode of muscle cell differentiation described here in the ctenophore tentacle root is strikingly similar to that of striated skeletal muscles in vertebrates. In both cases, muscle development passes through four main stages: multiplication of myoblasts, alignment, fusion into myotubes, and finally maturation of the multinucleated muscle cells. In the ctenophore tentacle root, these myogenesis stages are distributed in four distinct areas along the proximal-distal axis on both sides of the tentacle root symmetry plane (Figure 6). A different mechanism operates in vertebrates for smooth muscle cells, which are mononucleated and whose formation does not involve a phenomenon of myoblast fusion. It is therefore surprising that although ctenophore muscle cells are of the smooth type (they lack recognisable sarcomeres), their multinucleate condition and their mode of development is more alike vertebrate striated muscle cells than vertebrate smooth muscle cells. Expression of MHCIib is another shared characteristic of ctenophore muscle cells and vertebrate striated muscle cells. The evolutionary significance of these observations remains unclear: ctenophore muscles could have acquired these characteristics by convergence, or the similarities might reflect ancestral features of eumetazoan muscle cells, consistent with a scenario suggested previously in which
smooth vertebrate muscle cells would be a derived (vertebrate-specific) cell type [1,83].

In any case, these observations imply that a classification of muscle cell types based solely on presence or absence of a striation pattern (i.e. of Z discs allowing transverse or oblique alignment of the myofilaments) captures only part of the variation existing among animal muscles. A more comprehensive classification of muscle cell types at the eumetazoan taxonomic scale therefore remains to be elaborated, by integrating not only cytological characteristics but also molecular and developmental features. In this perspective, it will be crucial to obtain data from a wide sampling of early-branching metazoans concerning the expression and function of myogenic and muscle differentiation genes, including the various forms of myosins.

Additional files

**Additional file 1: Alignment of myosin amino-acid sequences.** This file contains the initial alignment (before site pruning by Gblocks) in graphic view. Note that for myosins of classes other than class II (outgroups), only the sequences of the motor domain have been included (the C-terminal part of these proteins is not alignable with the C-terminal part of myosins II). For *Saccharomyces cerevisiae* MHCl, the portion of the protein sequence downstream from position 1300 has been likewise omitted for the same reason.

**Additional file 2: Final sequence alignment (after Gblocks) including non-class II myosins.** The corresponding tree is shown in Additional file 3, see Methods for details. The alignment is provided in Phylip 4 format.

**Additional file 3: Phylogenetic analysis rooted on non-class II myosins.** Amino-acid sequences were analysed using the Maximum likelihood (ML) method. Numbers associated with the branches are ML bootstrap values (1000 replicates). Sequences from *Pleurobrachia pileus* are indicated in bold and green. Branches within the MHClA clade are in blue and within the MHClB clade in purple. The tree was rooted with sequences myosins classes V, VII and X (outgroups). The letter between parentheses after the name of mouse genes of the MHClA clade indicate whether the gene is expressed in non-muscle cells (n) or in smooth muscle cells (s). The PpiMHCIIb2 gene was excluded because its partial sequence contains only the tail and therefore it has no alignable residues with non-class II myosins (only the head being homologous between class II and non-class II myosins). Abbreviations for species names: *Aae*: *Aedes aegypti*; *Aq*: *Amphimedon queenslandica*; *Asu*: *A. suum*; *Cow*: *Capsaspora owczarzaki*; *Ddi*: *Dicyostelium discoideum*; *Dme*: *Drosophila melanogaster*; *Hma*: *Hydra magnipapillata*; *Lgi*: *Lottia gigantea*; *Lpe*: *Loligo pealei*; *Mbr*: *Monosiga brevicollis*; *Mmu*: *Mus musculus*; *Nve*: *Nematostella vectensis*; *Ppi*: *Pleurobrachia pileus*; *Sar*: *Sphaeroma arctica*; *Sce*: *Saccharomyces cerevisiae*; *Sro*: *Salpingaeca rosetta*; *Td*: *Trichoplax adhaerens*.

**Additional file 4: Final sequence alignment (after Gblocks) including only class II myosins.** The corresponding tree is shown in Figure 3 (see Methods for details). The alignment is provided in Phylip 4 format.

**Additional file 5: Expression pattern of *PpiTropomyosin* gene in *Pleurobrachia pileus* muscles.** (A) Internal view of a dissected tentacle root stained with the *PpiTropomyosin* antisense probe (CU420922.1). (B) Longitudinal view of the tentacle root (after removal of lateral expansions) showing *PpiTropomyosin* expression in tentacle and tentilla muscle progenitors. Forming tentilla are also stained. (C) Transverse cryosection (according to the dotted line on (B)) of a tentacle root after *PpiTropomyosin* whole-mount ISH. The tentacle base is also sectioned transversally (Tcl). Note that the musculature of the tentacle sheathing is also stained (TS). (D) Transverse cryosection of a tentilla after *PpiTropomyosin* whole-mount ISH (compare with Figure 4V). (E) Expression of *PpiTropomyosin* in longitudinal muscle fibres (LM) in the tentacular plane. (F) Expression of *PpiTropomyosin* in mesogleal muscle fibres connected to a meridional canal. (G) Higher magnification of *PpiTropomyosin* expression in a mesogleal muscle fibre. C: Comb; Coll: Collabolasts; F: Tentilla; LM: Longitudinal Muscle fibres; MC: Meridional Canal; Me: Mesogleal muscle fibre; MR: Median Ridge; M:tc: Tentacle Muscle progenitors; M:tt: Tentilla Muscles progenitors; Mu: Muscle fibres; Ots: Orifice of tentacle sheath; Tc: Tentacle; TS: Tentacular Sheath; Tt: Tentilla. Scale bars: A, B, C, E: 200 μm; F: 100 μm; D: 25 μm; G: 10 μm.

**Additional file 6: Pleurobrachia pileus musculature (phalloidin staining).** (A) Drawing of the arrangement of *Pleurobrachia pileus* musculature, with areas corresponding to pictures (B-I) indicated by black boxes. (B-I) Phalloidin staining of muscle fibres in selected regions of the body (boxes in (A)). (B) A large mesogleal muscle fibre connecting two meridional canals. Inter-comb fibre cells are also visible. (C) Dense parietal muscle fibres along a ciliated groove. (D) Parietal muscle fibres in the aboral region, in the pharyngeal plane. (E) Parietal muscle fibres in the aboral region, in the tentacular plane. (F) Parietal muscle fibres in the epidermis between two comb rows. All fibres have circular orientation. (G) Parietal muscles fibres in the epidermis between two comb rows in the tentacular plane: a dense band of longitudinal fibres is superimposed on the circular fibres. (H) Epithelial muscle fibres around the opening of the tentacle sheath. Note that some mesogleal fibres are visible connecting the apical organ area to the tentacle sheath. (I) Epithelial muscle fibres fibres around the oral extremity of the comb row. (J) Higher magnification of epithelial muscles fibres at the oral extremity of the comb row. The light coloured structure visible at the top of the picture is the oral extremity of the meridional canal underlying the comb row. C: Comb; Cg: Ciliated groove; CR: Comb row; F: Inter-comb fibrous cell; M: Meridional canal; Me: Mesogleal muscle fibre; Ots: Opening of tentacle sheath. Scale bars: B-I: 100 μm.

**Authors’ contributions**

The work presented here was carried out in collaboration between all authors. CD performed the experiments, analysed the data and drafted the paper. EQ and MM designed analytical strategies and revised the paper. AA, PC and CD carried out the TEM investigations. MJ assisted CD in the in-situ hybridisation and immunofluorescence experiments and revised manuscript. HLG provided all facilities. All authors have read and approved the final manuscript.

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