Newly-discovered muscle in the larva of *Patella coerulea* (Mollusca, Gastropoda) suggests the presence of a larval extensor

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**Abstract**

The development of the muscular system of the gastropod mollusc *Patella* has been thoroughly studied. As a result, two larval retractors, the main and accessory larval retractor, had been described in the larva of *Patella*. These muscles were supposed to be responsible for the retraction of the larval body into the shell. Previously no larval extensors, which would be responsible for the extension of the larval body out of the shell, have been described. Using cell-lineage tracer injection and phalloidin staining of muscles, a newly-discovered muscle is herein identified in the larva of *Patella coerulea*. A functional model is presented in which two muscles are responsible for the extension of the larva, the newly-discovered muscle together with the previously recognized but inaccurately named accessory larval retractor. In our model, the latter muscle does not seem to function as a retractor but rather as an extensor.

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**Introduction**

For many decades, the gastropod mollusc *Patella* has been used as a model system to study various aspects of early development. For instance, the cell-lineage has been described by Patten (1886), Wilson (1904), Smith (1935), van den Biggelaar (1977), and Dictus and Damen (1997). The formation of the 3D-macromere, which acts as an organizer and is responsible for the correct dorsoventral patterning of the embryo, has been described by e.g. van den Biggelaar and Guerrier (1979), Arnold et al. (1983), Verdonk and van den Biggelaar (1983), Martindale et al. (1985), and Damen and Dictus (1996). The analysis of gene function during early development has been described by van der Kooij et al. (1996, 1998), Damen et al. (1994), Damen and van Loon (1996), Klerkx et al. (2001), Lartillot et al. (2002), Lespinet et al. (2002), Nederbragt et al. (2002a, b, c), and Wanninger and Haszprunar (2001). The formation of the musculature has been described by Smith (1935), Crofts (1955), and Wanninger et al. (1999).

In the first half of the previous century, Smith (1935) and Crofts (1955) studied muscle morphogenesis in *Patella* with the use of a traditional technique, i.e. microscopical observation of serial sections. Much later, Wanninger et al. (1999) studied the formation of the musculature in the larva of *Patella* using a much more recently introduced technique. They stained larvae of *Patella* with fluorescently-labeled phalloidin, which binds to F-actin, in order to visualize the musculature using confocal laser scanning microscopy (CLSM). With this technique, muscle development can be analyzed at a much higher resolution and even enables a three-dimensional analysis of muscle localization. Using CLSM, Wanninger et al. (1999) showed that the formation of the principal larval and adult muscles in *Patella* takes place between about 28 and 52 h...
after first cleavage. The muscles they described are the main larval retractor, the accessory larval retractor, the velum muscle ring, the pedal plexus and the anlagen of the left and right adult shell muscles (Fig. 1). Detailed comparison of both approaches demonstrates that the traditional approach of Smith (1935) and Crofts (1955) is quite prone to misinterpretations and even may result in missing certain details of muscle formation.

The 48 to 52 h old post-torsional larva of *Patella* is a well-developed larva with a head, a foot, a shell and a prototroch (Fig. 1). It is capable of retraction into its larval shell. The main larval retractor and the accessory larval retractor are supposed to be responsible for this process, although this has not been stated explicitly for *Patella* (Wanninger et al., 1999). Besides retraction, the animal must be able to extend out of its larval shell. Despite that the larva is obviously able to do so, neither muscles nor other mechanisms responsible for this action have been described in the literature.

As part of a larger study (Damen and Dictus, submitted) various early cleavage-stage blastomeres, viz., 2b, 3a, 3b, 3c, 3d, 3A, 3B, 3C, 4d and 4D, of *Patella coerulea* were injected with a fluorescent marker to study their cell-lineage. Injection of one of the blastomeres, the 2b-micromere, resulted in labeling of an unknown bundle of fibers in 48 to 52 h old post-torsional larvae. After injection of the 3a- and 3b-micromeres small parts of this bundle of fibers were labeled as well. Since this unknown bundle of fibers strongly resembled a muscle, 48 to 52 h old post-torsional larvae were stained with fluorescently-labeled phalloidin. In a number of these larvae, besides the muscles described above, the unknown bundle of fibers was labeled as well, demonstrating that the unknown fibers are indeed muscle fibers.

Based on its location and orientation, together with the fact that no explanation for the extension of the larva out of its shell is given in the literature, it is hypothesized that the newly-discovered bundle of muscle fibers is involved in the extension of the larval body out of the shell. A hypothetical model is presented in which the newly-discovered muscle, together with the accessory larval “retractor”, are responsible for the extension of the larval body out of the shell. As a result, these two muscles...
form the antagonists of the main larval retractor. Reanalysis of literature data suggests that a structure similar to the previously undescribed muscle, and thus a potential larval extensor, is present in at least one other gastropod and that the previously undescribed muscle is homologous to the circular muscle fibers in the Müllers larva of the platyhelminth Hoploplana.

**Materials and methods**

**Obtaining embryos**

Adult specimens of the limpet *Patella coerulescens* (Mollusca, Gastropoda) were obtained from the Mediterranean coast near Trieste (Italy) and near Banyuls-sur-Mer (France). Embryos were obtained as described before (van den Biggelaar, 1977; Damen and Dictus, 1996; Dictus and Damen, 1997). Before injection of cell-line tracer, embryos were treated with acidified MPFSW (pH 3.9) for 2-3 min to remove the surrounding jelly layer. Embryos that were not injected but only cultured to stain the musculature with phalloidin were also dejetilised. To reduce microbial infection, embryos were usually transferred to antibiotic-containing MPFSW (60 mg penicillin G and 50 mg streptomycin per liter MPFSW) about 6 to 7 h after first cleavage. Some embryos were cultured in the absence of antibiotics. During culturing and all experimental procedures embryos and larvae were kept at 17 to 18°C.

**Cell-lineage, quadrant identification and cell-lineage tracer injection**

The early development and cell-lineage of *Patella* have been described before (see e.g., Wilson, 1904; Smith, 1935; van den Biggelaar 1977; Dictus and Damen, 1997). The nomenclature employed is that of Wilson (1892) and Conklin (1897). For clarity, figure 2 shows some stages of the development of *Patella*.

The cell-lineage tracer tetramethylrhodamine dextran (MW 10,000 Da, D-1868, Molecular Probes Europe, Leiden, Netherlands; TMR-dextran) was injected using high pressure-injection as described previously (Damen and Dictus, 1994; Dictus and Damen, 1997). TMR-dextran was dissolved at a concentration of 10% (w/v) in aqua dest or in injection buffer (10 mM Hepes, 150 mM KCl, 1 mM CaCl₂, 10 mM EGTA, pH 7.0).

Blastomeres, viz., 2b, 3a, 3b, 3c, 3d, 3A, 3B, 3C, 4d and 4D, were injected at specific stages of development. For instance, the 2b-micromere was injected at the 16-cell stage. Embryos of *Patella* are radially symmetrical up to about 60 min after the fifth cleavage (32-cell stage). Therefore, quadrant identity (A-, B-, C-, or D-quadrant) cannot be determined until after this moment (van den Biggelaar 1977; van den Biggelaar and Guerrier, 1979; Damen and Dictus, 1996). By analyzing injected larvae retrospectively, the identity of injected blastomeres was determined and embryos in which the 2b-micromere was injected were identified. The 3a- and 3b-micromeres were injected at the 32-cell stage. Since at about 60 min after fifth cleavage embryos are still at the 32-cell stage, and the quadrants can be denominated at this stage, 3a- and 3b-micromeres could be identified before injection.

**Selection and fixation**

Abnormal larvae at 48 to 52 h after first cleavage, i.e., larvae that do not possess eyes, a prototroch, a shell, a foot and an operculum or that are not able to retract into their shell, were discarded. The muscles of normal larvae of 48 to 52 h after first cleavage were relaxed by adding drops of 0.75 M MgCl₂ to the MPFSW. Subsequently, the larvae were fixed in 4% formaldehyde in 0.1 M phosphate buffer (pH 7.4) for 1 h and rinsed in buffer (3 times for 10 min).

**Phalloidin staining and mounting**

In order to visualize the musculature, most of the fixed larvae were stained with fluorescently-labeled phalloidin (Alexa Fluor 488 phalloidin, 300 U, A-12379, Molecular Probes Europe, Leiden, Netherlands), that binds to F-actin (modified after Wanginger et al., 1999). Larvae were permeabilized for 1 h in 0.1 M phosphate buffer (pH 7.4) to which 0.2% (v/v) Triton X-100 was added to allow penetration of phalloidin. A stock solution of phalloidin was prepared by dissolving one vial of Alexa Fluor 488 phalloidin (300 U) in 1.5 ml methanol. This stock solution was kept at -20°C. Some 5 or 10 μl of the stock solution were put in a small glass petri dish and the methanol was allowed to evaporate. Larvae were taken in 200 μl of the 0.1 M phosphate buffer (pH 7.4) to which 0.2% Triton X-100 was added and put on the dry phalloidin. The petri dish was gently shaken to dissolve the dried phalloidin after which the larvae were incubated for 1 h. Subsequently, the larvae were washed in 0.1 M phosphate buffer (3 times for 10 min) and dehydrated in a graded series of ethanol (70%, 80%, 90%, 96%, 100%, 100%, 100%; each step 5 min). After transferring the larvae to a 1:1 (v/v) mixture of ethanol and Murray's (1:2 v/v benzylalcohol and benzylbenzoate), they were mounted in convex polished slides in Murray's to which 0.1% of the antifading agent n-propyl gallate (Sigma, St. Louis, Mo, USA) was added.

**Confocal laser scanning microscopy (CLSM)**

All larvae were observed in a Leica upright confocal laser scanning microscope [Leica DM-RBE (microscope), Leica TCSNT (confocal laser), Leica, Heidelberg, Germany]. One, or in most cases two or even three z-series were recorded at different angles. All z-series consisted of 72 images and were transformed into stereo images that were analyzed with a pair of red-green stereo glasses.
Results

Fluorescent cell-lineage tracer injection

Early cleavage-stage blastomeres, viz., 2b, 3a, 3b, 3c, 3d, 3A, 3B, 3C, 4d, and 4D, were injected as part of a large study to trace the cell-lineage of the musculature in *Patella coerulea* (Damen and Dictus, submitted). Figure 2 shows drawings of early cleavage-stage embryos in which the localization of some injected blastomeres is indicated. Injection into one of these blastomeres, the 2b-micromere (yellow in Fig. 2A), resulted in a complex pattern of labeling in 48 to 52 h old post-torsional larvae (Fig. 3). Analysis with CLSM and the subsequent analysis of z-series revealed two large, ciliated prototroch cells and a ring of fluorescent cells that bordered the prototroch at the post-trochal side (Fig. 3A). Medially, a strand of cells was labeled. This strand ran from a ventral position, between the two labeled main prototroch cells, in a dorsal direction towards the dorsal side of the prototroch and curved in a posterior direction towards a position near the visceral mass (Fig. 3). Adjacent to this strand, other cells were labeled at the level of the prototroch. In addition, some labeling was seen on the inside of the right-lateral side of the shell. Most prominently labeled was a post-trochally located transverse band of fibers that was connected to the lateral shell walls. This band of fibers was U-shaped with the base of the U at the ventral side of the larva. This transverse-oriented band of fibers strongly resembled muscle fibers. After injection of cell-lineage tracer into the 3a- and 3b-micromeres (yellow in Fig. 2B), small areas in the right, respectively the left part of the U-shaped transverse band of fibers were labeled as well (results not shown).

Phalloidin staining

In order to verify whether the transverse-oriented structure formed from the progeny of the 2b-micromere is indeed a muscle, 48 to 52 h old larvae were stained with phalloidin and analyzed. The fluorescence of the main larval retractor, accessory larval retractor, pedal plexus, velum muscle ring, and the anlagen of the left and right adult shell muscles was clearly visible. In addition, an unknown muscle was detected (Fig. 4). The morphology of this muscle was exactly like that of the other muscles, although the phalloidin staining of this unknown muscle was weaker. This newly-discovered muscle is located between the two lateral shell walls and is U-shaped. The base of the “U” is located near

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*Fig. 2. Schematic drawings of several stages in the development of Patella, modified after van den Biggelaar (1977). (A) Lateral view of a 16-cell stage embryo. The first quartet micromeres (1a-1d) have divided and the second quartet micromeres are formed (2a-2d). (B) Lateral view of a 32-cell stage embryo. In this view the first (1a-1d progeny), second (2a-2d progeny) and third (3a-3d) quartet micromeres as well as the third generation macromeres (3A-3D) are visible.*
the hinge of the operculum. The fluorescence of this muscle co-localized with the fluorescence of the transverse-oriented band of fibers observed after tracing the 2b-progeny with TMR-dextran.

**Discussion**

**Newly-discovered muscle**

The results above describe a hitherto unknown structure in the 48 to 52 h old post-torsional larva of *Patella coerulea* that strongly resembles a muscle.
Based on the morphology, the positive staining with phalloidin, and the resemblance to muscle fibers, it is concluded that the structure is indeed a muscle. Based on its location, this muscle has been named transverse muscle.

Analysis of cell-lineage data demonstrates that the transverse muscle is derived from the 2b-micromere. In addition, the 3a- and 3b-micromeres form small parts of the transverse muscle. In contrast to the other muscles in phalloidin-stained larvae, the transverse muscle exhibited weaker staining. This indicates that these muscle fibers are not so heavily filled with F-actin containing myofibrils, and this may explain why the transverse muscle has not been observed before by Smith (1935), Crofts (1955), and Wanninger et al. (1999).
Hypothetical model for larval extension

The discovery of an undescribed muscle, the transverse muscle, allows us to present a hypothetical model to explain the extension of the larval body out of the larval shell (Fig. 5). Fig. 5A shows a fully retracted larva with a closed operculum. The larva retracts into its shell and closes the operculum when it is disturbed, e.g. by “teasing” the larva. Fig. 5B shows a larva that is partly retracted. This is an intermediate state between the fully extended and the fully retracted state that is also obtained when larvae are treated with MgCl₂, to relax the muscles. When larvae are left undisturbed for some time, they can become fully extended (Fig. 5C). Since treatment with MgCl₂ immobilizes the musculature, the situation of Fig. 5B represents the resting situation in which no muscular activity is present.

In the 48 to 52 h old post-torsional larva of *Patella coerulea*, contraction of the transverse muscle, which inserts on the two lateral shell walls, will decrease the volume that the larval body can occupy within the shell. Since the body of the larva functions as a ‘hydrostatic skeleton’, this will result in extrusion of the larval body out of the shell (see Fig. 5). The function of the transverse muscle may be compared to squeezing meat out of a sausage that is opened on one side. In addition to the transverse muscle, we think that the muscle previously known as the accessory larval retractor (alr) may also be involved in the extension process. We observed that the so-called accessory larval retractor consists of four fibers (see Fig. 1A and 4C). All four fibers reached far into the mantle and did not reach the pedal region or the velum. Since the fibers of the accessory larval retractor all end in the mantle, the accessory larval retractor cannot be involved in the retraction process of the larva. Because of its location and insertion area, we propose the opposite: the accessory larval retractor is involved in the extension process of the larva. Observation of numerous partly and fully extended larvae demonstrated that in these larvae the greater part of the shell is empty, i.e., not filled with the larva body. This can also be seen in various papers, e.g., Patten.

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**Fig. 5.** Hypothetical model explaining the extension of the larval body out of the larval shell. In this model, the main larval retractor (mlr) pulls the larval body back into the shell. The accessory larval retractor (alr), together with the transverse muscle (new), are the antagonists of the main larval retractor and are responsible for the extension of the larval body out of the shell. Note that in the schematic drawings only the muscles involved in retraction and extension are depicted. The gray area represents the larval body that functions as a ‘hydrostatic skeleton’. (A) Fully retracted larva. The main larval retractor (mlr) is contracted. The accessory larval retractor (alr) and the transverse muscle (new) are relaxed (extended). (B) Partly retracted larva. The main larval retractor (mlr), accessory larval retractor (alr), and the transverse muscle (new) are all partly contracted. In this intermediate state, the main larval retractor (mlr) on the one hand, and the accessory larval retractor (alr) and the transverse muscle (new) on the other hand, are in equilibrium. (C) Fully extended larva. The main larval retractor (mlr) is relaxed (extended). The accessory larval retractor (alr) and the transverse muscle (new) are contracted. For legend see figure 1. Scale bar: 50 μm.
(1886), Crofts (1955), and Wanninger et al. (1999). In Fig. 5 [compare to e.g., Fig. 3A in the paper of Wanninger et al. (1999)], the insertion area of the accessory larval retractor at the larval shell is visible. Since the fibers of the accessory larval retractor insert ventrally on the shell and project into the mantle, and the mantle is connected to the rim of the shell, contraction of the accessory larval retractor will result in extrusion of the larval body out of the shell. Since we expect that the accessory larval retractor is responsible for the major displacement of the larval body out of the shell, this muscle fulfills the function of a main larval extensor. The transverse muscle, which is expected to play a moderate role in the displacement of the larval body, fulfills the function of an accessory larval extensor. As a result, these two muscles form the antagonists of the main larval retractor. According to this model, the accessory larval retractor should be renamed the main larval extensor, whereas the transverse muscle may also be called the accessory larval extensor. Since no accessory larval retractor is present in this model, the main larval retractor may be renamed larval retractor.

Until now, no satisfactorily explanation for the extension of the larval body out of the shell has been given in the literature. The model presented above, which incorporates a completely different function of the accessory larval retractor, together with a function of the transverse muscle, solves this problem.

Phylogenetic considerations

Although the transverse muscle has not explicitly been described before in any gastropod, reanalysis of literature data strongly suggests that a similar muscle may also be present in at least one other gastropod, i.e., Haliotis. Degnan et al. (1997) described the musculature of the Haliotis larva using BODIPY-phallacidin. In their figure 1D, fibers are visible at the exact location where the transverse muscle is expected to be located. These fibers more or less radiate from a point between the foot and the visceral mass, close to the hinge of the operculum. These fibers are not named in the paper of Degnan and associates (1997) but may very well be the fibers of the transverse muscle. Future investigations will have to provide evidence that the transverse muscle is indeed a larval extensor and whether a transverse muscle is a universal muscle in gastropods or is only present in some species.

In Patella coerulea, the transverse muscle is almost entirely formed from the progeny of the 2b-micromere. The circular muscle fibers in the Müllers larva of the platyhelminth Hoploplana are also derived from the 2b-micromere (Henry et al., 2000). Possibly these circular muscle fibers are homologous to the transverse muscle of Patella coerulea. Further ontogenetic research, especially in species located in a phylogenetic position between the platyhelmints and the gastropods, can shed more light on the possible homology of the circular muscle fibers in Hoploplana and the transverse muscle in Patella coerulea.

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