New approaches to studying morphological details of intramolluscan stages of *Angiostrongylus vasorum*

Novas abordagens para estudo morfológico dos estágios intramoluscos de *Angiostrongylus vasorum*

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**Abstract**

*Angiostrongylus vasorum* is a pulmonary artery parasite of domestic and wild canid. On molluscs, intermediate host, first stage larvae (L1) are found after the first day of infection, in the 8th L2 and in the 30th L3. It was evaluated L1, L2 and L3 recovered by Baermann technique from *Achatina fulica* infected with 1000 L1. Fifty larvae/stage were incubated with antibodies anti-β-tubulin, anti-α-tubulin, anti-α-actin, anti-β-actin and anti-collagen, and then with Alexa 633. Fifty larvae/stage were observed with picrosirius red and Oil Red O. It was also observed in the anterior region of L1 the beginning of the chitinous stems development, in the initial portion of the intestine and genital primordium. In L2 anterior region, the papillae, chitinous canes juxtaposed to the mouth and intestines bigger than L1. The L3 musculature is well defined, next to the chitinous stems, there are two round distally arranged from each other. It was observed the whole extension of the intestine genital primordium and intense cellularity in the L3 distal portion. With the picrosirius red the L1, L2 and L3 musculature could be observed, as the nerve ganglia on L3. Oil Red O revealed that L1, L2 and L3 store energy in lipid droplets.

**Keywords:** *Angiostrongylus vasorum*, *Achatina fulica*, morphology, antibodies, picrosirius red, Oil Red O.

**Resumo**

*Angiostrongylus vasorum* é um parasito de artérias pulmonares dos canídeos domésticos e silvestres. Nos moluscos, hospedeiros intermediários, encontram-se no primeiro dia após a infecção, larvas de primeiro estágio (L1), ao 8º L2 e ao 30º L3. Avaliou-se L1, L2 e L3 recuperadas pela técnica de Baermann de *Achatina fulica* infectada com 1000 L1. Cinquenta larvas/estágio foram incubadas com anticorpos anti-β-tubulina, anti-α-tubulina, anti-α-actina, anti-β-actina e anti-collagênio e, em seguida, com anticorpo Alexa 633. Observaram-se também 50 larvas/estágio com picrosirius red e Oil Red O, na região anterior da L1, o início do desenvolvimento de hastes quitinosas, a porção inicial do intestino e o primórdio genital. Na região anterior de L2, papilas, bastões quitinosos juxtapostos à boca e ao intestino maior que em L1. A musculatura de L3 é bem desenvolvida, próximo às hastes quitinosas, há duas estruturas redondas dispostas distalmente uma da outra. Observaram-se também toda a extensão do intestino, o primórdio genital e a intensa celularidade na porção distal da L3. Com o picrosirius red observou-se a musculatura de L1, L2 e L3, assim como, gânglios nervosos na L3. Oil Red O revelou que L1, L2 e L3 armazenam energia em gotículas lipídicas.

**Palavras-chave:** *Angiostrongylus vasorum*, *Achatina fulica*, morfologia, anticorpos, picrosirius red, Oil Red O.
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**Introduction**

*Angiostrongylus vasorum* affects the right ventricle of the heart and the pulmonary arteries in dogs, red foxes and other canids (Mozzer & Lima, 2015). In humans, *A. cantonensis* causes eosinophilic meningitis and *A. costaricensis* triggers abdominal eosinophilic ileocolitis (Thiengo et al., 2013). The complete life cycle of these nematodes takes place in their definitive hosts, which are usually rodents. The eggs settle in the intestinal submucosa, while the adults lodge in the mesenteric arteries.

*Achatina fulica* the African giant snail was introduced in Brazil in 1980, currently its present in 23 Brazilian states including Minas Gerais (Barçante et al., 2005; Thiengo et al., 2013). It was found naturally infected with *A. cantonensis*, in Pernambuco, Rio de Janeiro, Santa Catarina, Espírito Santo and Amazonas (Thiengo et al., 2013). The eosinophilic meningitis cases in Espírito Santo are related to the ingestion of these molluscs. *Achatina fulica* also can be infected with *A. costaricensis* and *A. vasorum* (Neuhauss et al., 2007; Coaglio et al., 2016). In Brazil *A. fulica* are found naturally infected with other nematodes of importance in the veterinary field, *Aelurostrongylus abstrusus*, a cat lung parasite, *Rhabditis* sp. and *Strongyluris* sp., parasites found in the intestine of amphibians and reptiles (Oliveira et al., 2010). Knowing this, the participation of *A. fulica* in the context of public health must be proposed, as there are reports of natural infection related to this molluscs in several regions of Brazil, as such, it is important to know the development of the different stages of mollusc nematode parasite, as the knowledge gathered could help in creating control strategies.

Treatment of angiostrongyliasis with nematicides generally does not produce good results, and new therapies are extremely necessary. Although recent “omic” data on parasitic nematodes accelerate the identification of potential drug targets (Mccoy et al., 2017), the validation of these targets in functional experiments remains a challenge because few techniques allow functional characterization. Therefore, the use of different morphological markers for parasitic nematodes may help morphological observations of these nematodes and may contribute to “omic” studies.

In the present study, we analyzed the morphology and morphometry of L1, L2, and L3 larvae of *A. vasorum* recovered from *Achatina fulica* by assessing the cytoskeleton, connective tissue and lipid distribution of these larvae using cytoskeleton and collagen antibodies and staining with picrosirius red and Oil Red O, respectively.

**Materials and Methods**

**Obtaining intramolluscan stages of *Angiostrongylus vasorum***

The strain of *A. vasorum* that was used is maintained in dogs at the experimental kennel of the Institute of Biological Sciences of the Federal University of Minas Gerais (ICB/UFMG) (Lima et al., 1985). To obtain L1 larvae, feces collected from the dogs that maintain the strain were subjected to the Baermann technique. This study and maintenance of the strain in animal hosts was approved by the ethics committee for animal use of UFMG (CEUA/UFMG) under the number 147/2011.

Then, 1 mL of water containing 1000 L1 was placed in each of specimen of infection containers of width 5 cm and height 5 cm. Specimens of the mollusc *Achatina fulica* of mean size of 4 cm were individually introduced into each infection container (one per container) so that they would be exposed to L1. They were left there for 24 h, at the room temperature of 23 °C (RT). After this period, these molluscs were transferred to a rectangular terrarium containing autoclaved soil supplemented with calcium carbonate. Water and lettuce were provided *ad libitum*. The terrariums were cleaned three times a week.

The remaining larvae in the infection containers were quantified as described by Pereira et al. (2006). Fifty molluscs were used to recover L1 (1 day post-infection, dpi), 50 to recover L2 (8 dpi) and 50 to recover L3 (30 dpi). The L1, L2 and L3 larvae were recovered under a Baermann’s technique, as proposed by Coaglio et al. (2016).

**Morphological evaluation of *A. vasorum* larval stages**

350 larvae per stage (L1, L2 and L3) were recovered with the help of a stereoscopic microscope (25x) and were stored in a 2 mL polypropylene tube containing 300 μl of PBS. Then, 900 μl of 4% paraformaldehyde was added and the samples were incubated for 24 h at 4 °C. To remove excess fixative, the samples were washed 3x with water by means of centrifugation at 180 g for 10 min. At the end of this, the larvae were resuspended in 900 μl
Details of Angiostrongylus vasorum intramolluscan water and used as described below. 50 larvae per stage (L1, L2 and L3) were used as a control that was detected by means of light microscopy.

To allow small pores to be formed in the cuticle of the nematodes, the tubes with the larval stages were placed in a 10 ml beaker with 95% ethyl alcohol for 4 min at -80 °C. Subsequently, the tubes were placed in water for 10 min at RT, followed by slight agitation on a microplate shaker for 30 min at RT. After this period, the samples were centrifuged at 450 g for 1 min. After this procedure, 0.5% saponin was added and the samples were incubated for 10 min at RT under gentle shaking. They were then washed 3x with water by means of centrifugation at 180 g for 10 min.

Next, each sample received 200 μl of BTB (borate triton β-mercaptoethanol solution (Borate buffer/25 mM; Triton X-100/0.5%; β-mercaptoethanol/3% in the final concentration) and was incubated for 1 h at RT, under gentle shaking (Biomixer Ts-2000A shaker). Subsequently, the samples were washed with BTB by means of centrifugation at 220 g for 2 min and, following the removal of BTB, each sample was resuspended in borate triton (BT) solution (Borate buffer/25 mM; Triton X-100/0.5% in the final concentration).

Following the removal of excess BT as described above, the cytoskeleton content and distribution of the larvae (L1, L2 and L3) was investigated. Fifty larvae of each stage were incubated individually with 50 μl of each of the following primary antibodies (Santa Cruz Biotechnology) diluted 1:500 in PBS containing bovine serum albumin (BSA): anti-β-actin (sc 69879), anti-α-tubulin (sc 5286), anti-β-tubulin (mouse-sc 55529) and anti-collagen (Cat. No. sc-9855, Santa Cruz Biotechnology, CA). The samples were incubated overnight at 4 °C under gentle agitation. They were then centrifuged at 220 g for 2 min. The supernatant was discarded and 250 μl of the blocking solution (4% BSA/PBS) was added for a further 1 h 30 min at 4 °C, under gentle shaking. Excess dye was removed by centrifuging the samples at 220 g for 2 min. The procedure was repeated four times. Then, 50 μl of the secondary antibody conjugated to Alexa 633 (goat anti-mouse, Invitrogen/Cat. No. A-21052) diluted at 1:500 in PBS was added to the samples and incubated for 2 h in the dark at RT, under gentle shaking. To remove the excess antibodies, the samples were washed 4x with PBS at 220 g for 2 min. The samples were mounted between slides and coverslips and images were captured using an epifluorescence microscope model Axiovert 200M APOTOME (Carl Zeiss, Germany).

To investigate the lipid content and distribution, 50 larvae of each stage were incubated with the Oil Red O dye at a ratio of 3:1 for 24 h at RT. The larvae were mounted between a slide and a coverslip and observed under an Olympus® BX41 light microscope coupled to an Olympus® DP12 digital camera.

Following this, picrosirius red (Junqueira et al., 1979) (3:1 in PBS) was added to 50 larvae of each stage for 24 h at 4 °C. The larvae were then centrifuged at 220 g in PBS 2x to remove the excess dye. The samples were mounted between slides and coverslips, and images were obtained via epifluorescence microscopy with Apotome, as described above, because picrosirius red can be detected at the red wavelength.

Results

Morphological characterization of intramolluscan larval stages of Angiostrongylus vasorum

First-stage larvae (L1)

The L1 specimens that were recovered were thin and transparent, measuring 332 ± 12 μm in length and 11 ± 1 μm in width. The anterior end was rounded and the tail was curved ventrally with an unguiform appendage, as seen using light microscopy (Figure 1A).

Anti-α-tubulin antibodies showed the nerve ring, the intestinal region and the anal opening (Figure 1B). Although some structures were not identified through this antibody, there were two oval structures next to the chitinous rods that were not detected by means of light microscope. These structures measured 5 μm², with an average of 2 μm in length and 0.5 μm in width (Figure 1B). The nerve ring was located 70 μm from the anterior end of the larva and was 6 μm in length and 7 μm in width (Figure 1B). The width of the initial portion of the intestine was 6 μm and the total length of the intestine was 158 μm (Figure 1B).

Anti-β-tubulin antibodies labelled the chitinous rods and intestine well, while the nerve ring was poorly shown by them. The chitinous rods measured 7 μm in length and the anal opening was located 25 μm from the final
The distance from the tip of the tail to the slit that is present in this region was 4 μm (Figure 1C).

Using anti-collagen antibodies, collagen fibers were detected in the structures of L1, including the intestine, chitinous rods and nerve ring (Figure 1D). The distance from the sheath of L1 to the wall of the body was 1 μm (Figure 1D). When labeled with anti-α-tubulin, anti-collagen and anti-β-tubulin, L1 showed prominence of the posterior portion, especially the intestine (Figure 1B-D).

Initially, with epifluorescence microscopy without Apotome, staining with picrosirius red only revealed a few structures. However, after using the Apotome, it was possible to detect the chitinous rods, the nerve ring, the esophagus and the musculature around it, the intestine and the genital primordium of L1. Picosirius red staining showed an oval-shaped genital primordium measuring 6 μm in length and located at 69 μm from the anal opening and 99 μm from the tip of the tail. The L1 musculature was 2 μm thick and had small oval-shaped cellular structures of 3 μm in length (Figure 2A). Staining with Oil Red O showed intense presence of lipids throughout the body of L1 (Figure 2B).

Second-stage larvae (L2)

The L2 larvae were located in the tissues of the intermediate host and were characterized by being slightly mobile, arched (‘C’ shaped) and brownish in color due to the presence of granules inside the intestinal cells. They measured about 420 ± 23 μm in length and 12 ± 2 μm in width and had two sheaths that occupied the entire interior space, as shown by light microscopy (Figure 3A).
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Figure 2. First-stage larva of Angiostrongylus vasorum. (A) L1 stained with picrosirius red: yellow arrowhead indicates the chitinous rods; red arrowhead indicates the genital primordium; blue arrowhead indicates the general appearance of muscle cells; (B) L1 stained with Oil Red O, showing lipid droplets (dark spots).

Anti-α-tubulin antibodies revealed an esophagus of 147 μm in length. When dilated, the esophagus measured 12 μm in width and 27 μm in length. The nerve ring in L2 had a rectangular shape, measuring 15 μm in width, and was located 80 μm away from the tip of the anterior portion. The anti-α-tubulin antibodies also showed that the intestine as a compact mass of 27 μm in width, occupying almost the whole width of the larva, differently from L1, whose intestine was observed as a filament (Figure 3B).

Anti-β-tubulin antibodies marked the intestine more intensely than the esophagus. Every segment of the intestine of L2 was formed by small rounded cells (Figure 3C). Only the gut became labelled with anti-β-actin antibodies and, when the Apotome was used, we observed two small papillae in the anterior portion, about 2 μm wide and 5 μm apart from each other. The oral opening measured 15 μm (Figure 4C).

The esophageal wall stained with picrosirius red (Figure 4A). This morphological detail was not observed when L2 were incubated with the antibodies. The intestine was not shown with this dye when epifluorescence was used alone; however, after using the Apotome attachment, we were able to view the structures of the intestine, along with the esophageal composition, the excretory pore (30 μm in length), the genital primordium and the anal opening (Figure 4A). Picrosirius red staining also revealed that the genital primordium was not as well-defined in L2 as it is in L1. Rather, it was a cluster of cells, 17 μm in length and 6 μm in width, located 180 μm away from the end of the tail (Figure 4A). Lipids were recognized by means of the Oil Red O dye, throughout the body of L2 (Figure 4B).

Third-stage larvae (L3)

L3 appeared free from sheaths and was clearer and more transparent than the other stages, with two chitinous rods arranged longitudinally at the anterior end. The tail ended with a digitiform appendix, measuring about 470 ± 25 μm in length and 19 ± 1 μm in width, as shown by light microscopy (Figure 5A).

Anti-collagen antibodies also revealed the nerve ring (Figure 5B).

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Anti-α-tubulin antibodies revealed the chitinous rods, nerve ring, esophagus and intestine. The genital primordium and anal opening became labelled less intensely than was observed when the other stages were analyzed. The lateral cell layer was composed of rounded cells (Figure 5C and 5D). The 13 μm-long chitinous rods were 2 μm apart and presented a bulb at their base measuring 1.5 μm in length and 1 μm in width. The nerve ring was 64 μm away from the anterior end of the larvae, measuring 18 μm in length by 8 μm in width (Figure 5D). The length of the esophagus was 192 μm and it had a dilation in the final portion, which was 46 μm long and 20 μm wide (Figure 5D). The initial width of the intestine (19 μm in length) could be noted in the portion that attached to the esophagus: 260 μm in length and 20 μm in width.

Regarding anti-β-tubulin antibodies, α-tubulin marked the chitinous rods, nerve ring intestine and the genital primordium (Figure 5C and 5D). The Oil Red O dye weakly stained the gut (Figure 5E).
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Staining with picrosirius red dye revealed nerve ganglia, which were oval-shaped structures spaced symmetrically from the esophagus (Figure 6A and 6B). The dye also showed that nerve cells existed in the anterior region and were similar to the amphidial pouch found in some adult nematodes (Figure 6A). Furthermore, the genital primordium was 156 μm away from the end of the tail; it was 21 μm in length by 6 μm in width and was composed of 15 rounded cells (Figure 6B). The excretory pore (32 μm in length) extended into the esophagus and was located 66 μm away from the oral opening (Figure 6B). The labeling also revealed the full extent of the intestine (Figure 6B). In the final median portion of the intestine, oval-shaped cells were observed, symmetrically spaced 17 μm from one another (Figure 6B).

Table 1 shows the morphological information on the intramolluscan larval stages of A. vasorum. Although the chitinous rods were present from the L1 stage onwards, they were twice as large in L3 as those found in the other two stages. The nerve ring was three times as long in L3 (18 μm) as in L1 (6 μm), but only 3 μm longer than in L2. The genital primordium developed greatly between the stages: it was 2.5 times smaller in L1 than in L2 and 3.5 times bigger in L3 than in L1. L2 showed the greatest distance between the genital primordium and the tail. The esophagus was more developed in L3.

Figure 4. Second-stage larva of Angiostrongylus vasorum. (A) L2 stained with picrosirius red: the genital primordium is indicated as between the yellow arrowheads; the distance from the genital primordium to the final portion of the tail is indicated as between the red arrowheads; the excretory pore is indicated by the blue arrowhead; (B) L2 stained with Oil Red O; dark spots show lipid droplets; (C) Intestine viewed using the Apotome attachment: the anti-collagen antibody is indicated as between the red arrowheads; the anti-β-tubulin antibodies are indicated as between the green arrowheads and the anti-β-actin antibodies are indicated as between the blue arrowheads. Two spherical structures at the anterior end of the larva that were revealed with anti-β-actin antibodies are indicated as between the yellow arrowheads.
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**Figure 5.** Third-stage larva of *Angiostrongylus vasorum*. (A) L3 under the light field microscope; (B) L3 with anti-collagen antibodies: red arrowhead indicates the nerve ring; (C) Posterior portion of L3 with anti-α-tubulin and anti-β-tubulin antibodies: blue and yellow arrowheads indicate genital primordial cells; (D) Anterior portion of L3 with anti-α-tubulin, indicated by blue arrowheads: anti-β-tubulin antibodies are indicated by yellow arrowheads; the arrowheads indicate the distance between the anterior end and the nerve ring and the distance between the nerve ring and the beginning of the gut; (E) Dark spots indicate lipid droplets stained with oil red O.

**Table 1.** Measurements (in μm) of the main structures of the larval stages of *Angiostrongylus vasorum* in the snail *Achatina fulica*.

| *Angiostrongylus vasorum* stage | Chitinous rods | Nerve ring | Distance from ring to the anterior region | Esophagus | Genital primordium | Distance from the genital primordium to the tail |
|--------------------------------|----------------|------------|----------------------------------------|-----------|-------------------|-----------------------------------------------|
| L1                             | 7 ± 0.2        | 6 ± 0.4    | 70 ± 1                                 | 109 ± 0.8 | 6 ± 0.3           | 99 ± 2                                       |
| L2                             | 6 ± 0.3        | 15 ± 0.8   | 80 ± 3                                 | 147 ± 1   | 15 ± 0.4          | 170 ± 4                                      |
| L3                             | 13 ± 0.5       | 18 ± 0.9   | 64 ± 4                                 | 192 ± 2   | 21 ± 0.7          | 156 ± 5                                      |
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Discussion

Angiostrongylus vasorum undergoes two molts inside its molluscs hosts, developing from L1 to L2 and then from L2 to L3. The larval stages of A. vasorum recovered from A. fulica were similar to those found in L1 of Arion sp., Omalonyx matheroni and Subulina octona, since L1 showed a mean length of 335 μm, L2 had a mean length of 420 μm and L3 had a mean length of 485 μm, respectively (Guilhon & Cens, 1973; Bessa et al., 2000; Mozzer et al., 2011). The morphological details presented during the development of the digestive and reproductive tracts, which begins at the larval stages of A. vasorum, have not been well studied so far. Here, we described these details for the L1, L2 and L3 larval stages, which are not well observed through optical field microscopy.

Figure 6. Third-stage larva of Angiostrongylus vasorum stained with picrosirius red. (A) Yellow arrowheads indicate the oval-shaped cellular layer spaced symmetrically along the esophagus and nerve ganglia; (B) The chitinous rods are indicated as between yellow arrowheads; the genital primordium is circled; blue arrowheads indicate the anal opening; red arrowheads indicate the tip of the tail.
In L1, the antibodies and dyes used here revealed the oral opening, the chitinous rods and a pair of rounded structures close to these, along with the nerve ring, esophagus, intestine and anal opening. Although anti-β-actin antibodies presented the least intensity, they detected the intestine and nerve ring well. We also observed prominent areas throughout the interior of L1. Using picrosirius red, we were able to detect the initial formation of the chitinous rods in the anterior region of L1. The morphology revealed here suggests that the chitinous rods are part of the oral cavity, thus indicating that the oral cavity is in the process of formation. This structure seems to be part of the digestive system of L1, which would thus be a narrow, almost closed stoma. However, in L3, a dilation between the two structures was observed. These structures were defined as chitinous rods and the dilation between them gives a stomal appearance to this apparent oral opening. The esophagus detected in L1 has a rhabditiform shape that is characteristic of the first-stage larvae, as shown by Rosen et al. (1970). The L1 intestine was well characterized at the 1st dpi, in the final third of the larvae, ending in the anal opening. This observation is different from what was presented by Lv et al. (2009), who found that the intestine of the L1 larvae of A. cantonensis had a dilated appearance only five days after infecting Pomacea canaliculata. The tail of L1 has an unguiform appendage shape, which is the main characteristic differentiating the L1 stage of Metastrongylidae from other parasitic nematodes (Rosen et al., 1970).

By using these markers in L2, we were able to show a pair of rounded structures, as well as the nerve ring, esophagus, gut (which presented as a compact mass occupying virtually all of the posterior portion of the larva), genital primordium and anal opening. In the oral opening, two protuberances that were not present in the other stages were found. The morphology of L2 is the least studied so far, probably due to the difficulty in recovering these larvae from the molluscs' tissues. Nevertheless, it is known that L2 has two sheaths: one that originated from L1 and is being eliminated; and the other that is under development to form the sheath in L3 (Guilhon & Afghahi, 1969; Bessa et al., 2000; Coaglio et al., 2016).

The oral opening, the chitinous rods (anti-α-tubulin, anti-β-tubulin and picrosirius red), the nervous system (anti-α-tubulin, anti-β-tubulin, anti-collagen and picrosirius red) the excretory pore (picrosirius red), the esophagus (picrosirius red), the intestine (anti-α-tubulin and anti-β-tubulin), the genital primordium (picrosirius red), the anal opening (picrosirius red) and lipid reserves in the intestine (Oil rRed O) were all detected in L3. In the final portion of the tail, after the anal opening, we observed a set of cells with symmetrical sizes that may form the structures that will arise in the future stages of the parasite in the definitive host. The esophagus formed a bulb close to the intestine.

Morphological information regarding structures such as the chitinous rods, nerve ring, excretory pore, genital primordium and esophagus has been reported for L1, L2 and L3 larvae of some metastrongyloids during intramolluscan development (López et al., 2005; Rebello et al., 2013; Coaglio et al., 2016; Colella et al., 2017). However, that information did not have the degree of detail presented here. Indeed, the present study reveals novel observations regarding chitinous rods in L1 and L2, nerve ganglia in L3, spherical structures located near the mouth region of L2 and lipid reserves, intense cellularity and body cavities in all three stages.

The infecting stage is L3. Angiostrongylus vasorum is characterized by the presence of a pair of chitinous rods near the mouth. Our data suggest that these structures are present from the L1 stage onwards but only reach full development in L3 when they show a space between each other. They may constitute the stoma in the adult parasite. The parasites A. cantonensis, A. costaricensis and A. abstrusus also present structures similar to the chitinous rods found in A. vasorum that are characteristic for L3. To date, there have not been any reports from any morphological studies characterizing the internal morphology of intramolluscan nematodes (Ash, 1970; Hata & Kojima, 1990; Lv et al., 2009; Ohlweller et al., 2010). The muscle structure that forms the intestine consists of a thick cellular assembly that ends in the anal opening. Also, the body musculature of L3 is denser than that of L1. Picrosirius red strongly labelled the final portion of the intestine of L3 and therefore may be considered to be a useful tool for studying intramolluscan nematode development. The tail ends with a digitiform appendix, which is characteristic of L3 that is common to A. vasorum, A. costaricensis and A. cantonensis (Ash, 1970; Guilhon & Cens, 1973; Hata & Kojima, 1990; Bessa et al., 2000).

This study was the first to use antibodies to better characterize the structures of A. vasorum larvae. Other studies have used a similar strategy with antibody markers to characterize the structures of Schistosoma mansoni (Collins et al., 2011), Echinostoma paraensei (Souza et al., 2011), Ascaris suum (Fellowes et al., 1999; Williamson et al., 2009) and Caenorhabditis elegans (Albeg et al., 2011). Our study pioneered the use of picrosirius red to morphologically characterize parasites, since this dye had so far only been used to observe the production of collagen in histological tissue sections. For example, picrosirius red had been used on mouse liver to analyze its response when infected...
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with *Schistosoma mansoni* (Lenzi et al., 1999) and on rat liver infected with *Plasmodium berghei* (Haque et al., 2011). This low-cost dye successfully stained L1, L2 and L3, thereby revealing almost all the structures that were also observed using antibodies. Staining with picrosirius red and labelling with cytoskeleton and collagen antibodies provided a more realistic view than did optical field microscopy, because these tools showed details about how the larvae are actually structured. The dyes and antibodies used here generated more detailed information about the larval structures. Therefore, future studies using these tools may help expand the current knowledge about the biological role of the structures of the larval stages of *A. vasorum*.

Staining the larvae of *A. vasorum* with Oil Red O showed the main lipid reserve sites at each larval stage. Saturated lipids are the most important energy reserves in nematodes and account for 11-67% of the dry weight of the parasite’s juvenile phase, whereas neutral fats such as triglycerides are the main form of energy storage in adults (Andaló et al., 2011). The lipid reserves observed in L1 were uniformly distributed throughout the body. Lipid droplets are the main storage sites for neutral lipids (Xu et al., 2012). The lipid reserve of L2, which remains immobile in the host’s tissue, was found in the whole body. Mendonça et al. (2008) found similar results when staining L2 of *A. costaricensis* with Oil Red O. L3, which is the most active larval stage, spends more of its energy reserves than do the other two stages. Consequently, L3 has fewer lipid droplets concentrated in the intestine than are observed in the other larval stages.

**Conclusions**

Using anti-β-tubulin, anti-α-tubulin, anti-collagen and anti-β-actin antibodies, along with picrosirius red and Oil Red O staining, we revealed here unprecedented details regarding the internal morphology and morphometry of the intramolluscan larval stages of *A. vasorum* that should help expand the current understanding about the biology of this parasite. The picrosirius red dye revealed the musculature of *A. vasorum* and nerve ganglia in L3. The picrosirius red dye was better for morphological and morphometric analysis of *A. vasorum* L1, L2 and L3 in *A. fulica*, and it was also a cost-effective dye. Staining L1, L2 and L3 of *A. vasorum* with Oil Red O revealed that they store energy in the form of lipid droplets that change their main location as the larvae progress through their development.

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