Ultrastructural Observation of *Pseudoamphisiella lacazei* Reveals More Morphological Features and Provides Taxonomical Evidence

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Abstract

Ultrastructure of *Pseudoamphisiella lacazei* was investigated by transmission electron microscopy (TEM) and scanning electron microscopy (SEM) with an improved sample preparation method. The major results included: 1) A vacuole layer which had been considered absent in this species was found, which indicated that the cortex layers should thus contained three layers: a smooth membrane, a vacuole layer, and a fiber layer. 2) Extrusomes were found to distribute among the cortex vacuoles, and they were supposed to be a type of mucocyst which were similar to the mucocyst of Urostylida based on their shape and structure. 3) Some bacteria were observed to be attached to the fiber layer as if they were endosymbionts of *P. lacazei*. In order to analyze the features and taxonomical position of *Pseudoamphisiella*, morphological features of all four species of *Pseudoamphisiella* were compiled and compared. Our results suggested that ultrastructure observation could supplement and correct the gross and micro-level observation and provide additional evidence to the taxonomical argument.

Keywords: Ciliate; Electron microscopy; Morphology

Received: April 10, 2017; Accepted: April 25, 2017; Published: May 02, 2017

Introduction

Hemberger placed *Pseudoamphisiella* in Family Amphiellidae, but Song transferred it to a new family *Pseudoamphisiellidae* based on the origin of the right marginal cirri and the absence of front-o-terminal cirri, making the group sufficiently different from other hypotrichs (Order Hypotrichida) [1-3].

The main features of genus *Pseudoamphisiella* according to Song [2] include a long elliptical body with one left and one right marginal cirri row; frontal, buccal and caudal cirri are present; extremely well-developed transverse cirri; two parallel and widely-spaced ventral rows that derive from numerous fronto-midventral transverse cirri anlagen; and conspicuous extrusomes which are commonly located in an outer alveolar layer. This genus contains four known species, *P. lacazei* [2] (formerly *Holosticha lacazei*), *P. alveolata* [4] Song and Warren (formerly *Holosticha alveolata*) [5], *P. quadrinucleata* Chen et al. and *P. elongate* Li et al. [6,7]. This new genus has been widely investigated using various methods, including living observations, protargol impregnation, and small subunit ribosomal rRNA (SSU rRNA) gene sequencing [3-9]. *Pseudoamphisiella lacazei* was selected to be the type species of this genus. This free-living species has been recorded in marine habitats in the northern China seas. It swims relatively rapid but with long stationary periods, and feeds on bacteria, debris, small ciliates or flagellates [2,10]. However, no ultrastructure has not yet been described.

In this paper, some new ultrastructure characteristics of *P. lacazei* cortex and cytoplasm observed by transmission electron microscopy (TEM) and scanning electron microscopy (SEM) were added to the diagnosis of this species. We also compared the available morphological features of living and protargol staining observations of all four species of *Pseudoamphisiella*. 
Materials and Methods

Microorganisms

Pseudoamphisiella lacazei, provided by Prof. Weibo Song, Ocean University of China, was collected from coastal waters of the Yellow Sea. Samples were cultured in Petri dishes with boiled seawater with the salinity about 31%, water temperature 18 and pH 8.1 [2]. Some rice grains were added to promote the growth of bacteria.

Method of SEM observation

Samples were prepared according to the method described by Gao et al. [11]. The ciliates were fixed in 1% KMnO₄ and 100% glacial acetic acid at room temperature for 15 min. After washing with double distilled water for several times, washing off the cytoplasm, the samples were then conventionally dehydrated, freeze dried and gold sprayed. The samples were then observed by a HITACHI S-3400 SEM.

Method of TEM observation

Samples were prepared according to the method described by Gu et al. [12]. They were fixed by 2% OsO₄ and 5% glutaraldehyde at 4°C for 30 min. After washing with the cacodylate buffer, followed by conventional dehydration, the samples were embedded with Epon 812 and successively polymerized at 40°C for 20 h and at 60°C for 48 h. The embedded samples were then ultrathin-sectioned and stained with uranyl acetate and citrate lead before they were observed under a FEI TECNAI G² TEM.

Results

SEM ultrastructures of Pseudoamphisiella lacazei (Figure 1)

The Conventional SEM method can only illustrate the external morphology of samples (Figure 1a). But a new method allows us to peel off a piece of cell membrane to observe the inner surface of the cortex [11] (Figure 1b). By using this new method, the

![Figure 1](http://www.imedpub.com/insights-in-aquaculture-and-biotechnology)
following new characteristics of \textit{P. lacazei} are revealed by the current study: (1) The cortex is conspicuous and contains two parts: an inner fiber layer (\textbf{Figures 1c, 1d and 1f}) and an external vacuole membrane (\textbf{Figure 1e}). (2) Numerous oval extrusomes distributed among the vacuoles (\textbf{Figure 1e}) and retained on the outer surface of the cell after extending from the cortex (\textbf{Figure 1f}). (3) High numbers of bacteria are attached closely to the fiber layer and are intertwined with these fibers (\textbf{Figures 1c and 1d}).

**TEM ultrastructures of \textit{Pseudoamphisiella lacazei} (\textbf{Figure 2})**

TEM observation showed that the vacuole membrane consists of two parts: a smooth membrane outside and an intensive vacuole layer beneath it (\textbf{Figures 2a and 2c}). The fiber layer, which appears to be only one cell-layer thick, covers the inner surface of these vacuoles (\textbf{Figure 2a}).

The extrusomes are distributed among those vacuoles (\textbf{Figures 2b-2e}). Their shape under TEM is long-oval, with one end slim, the other end round (\textbf{Figure 2d}). The average size is about 1.17 $\mu$m \times 0.41 $\mu$m. These extrusomes can be observed to have two parts, which we tentatively name “head” and “body”. The length ratio of head and body is about 1:5. The head is wrapped in a dark cap-shaped structure (\textbf{Figure 2c}). The body part looks like a sac filled with filamentous matters (\textbf{Figures 2c and 2d}). A cup-shaped extrusome was observed as it had ejected the filamentous matters and the head were lost or indented into the body (\textbf{Figure 2e}).

The cytoplasm is not shown very dense. There were a few macronuclei and many mitochondria distributed in it (\textbf{Figure 2h}), but most of the space of cytoplasm was filled with various vacuoles, including different food vacuoles (\textbf{Figures 2f and 2g}) and some lysosomes (\textbf{Figure 2i}).

**Discussion**

\textbf{Supplementary description of \textit{Pseudoamphisiella lacazei} and its modified diagnosis}

As the type species of \textit{Pseudoamphisiella}, the morphological features of \textit{P. lacazei} have been described on gross and...
microscopic levels [13] as follows: Body is usually 150-300 × 45-100 μm in vivo, elongate-shaped with both sides parallel and posterior end of cell broadly rounded; it’s left margin of anterior region usually ear-shaped and extrusomes bar-shaped. There are three frontal, two buccal and 8-11 caudal cirri. 20 rows transverse cirri that extends to anterior 1/2 of body, two ventral rows with 14-16 and 16-21 cirri respectively, 21-34 left and 20-31 right marginal cirri and 8-11 dorsal kineties. AZM composes of 39-56 membranelles. These extrusomes in *P. lacazei* are very similar in shape, structure and class to those in *Urostyla grandis*, the type species of Urostylida [10]. This result supports that the Family Pseudoamphisiellidae should be assigned to the order Urostylida [8,10].

The following new features are reported at the ultrastructural level in this paper, which should be added to the diagnosis of this species: Three structure layers constitute the cortex including an instant membrane, multiple vacuole structure layer and a fiber layer; only one style of extrusome is present, which should be mucocyst according to Hausmman’s description [14].

**Morphological features and taxonomy of Pseudoamphisiella species**

There are four species belonging to Pseudoamphisiella, *P. alveolata*, *P. elongate*, *P. quadrinucleata* and *P. lacazei*. We compiled their morphological features reported by different researchers since 1996 in **Tables 1 and 2**. All these features were obtained based on in vivo observation and protargol staining experiments, using a light microscope. Some morphological features observed under light microscope were used to identify species, such as: (1) The shape and size of extrusome. In *P. lacazei*, it is bar-like, 2 μm long compared to bar-like, 2–3 μm long in *P. alveolata* and *P. quadrinucleata*, rod-shaped in *P. elongate*. (2) The vacuole layer is absent in *P. lacazei*, but is present in the other three species. Obviously, these features are illustrated better by SEM and TEM which has been confirmed in the current study (**Figures 1 and 2**). However, our SEM and TEM observation confirmed that the vacuole layer does exist but is covered by a smooth membrane from outside and a fiber layer from inside (**Figures 1e and 1f**).

According to Berger [10] and Shao et al. [8], the Family Pseudoamphisiellidae should be treated as a peripheral group in the order Urostylida, based on its morphological and morphogenesis features. Recently some researchers found that Pseudoamphisiellidae may represent an ancestral form for both Discocephaline and Urostylida species [15] according to the small subunit ribosomal RNA sequence. We recommend the Pseudoamphisiellida should be placed in the order Urostylida.

In summary, ultrastructure observation can supplement and correct the micro-level observation and provide some evidence to the taxonomical argument.

**Table 1** Cirri features of Pseudoamphisiella species.

<table>
<thead>
<tr>
<th>Species</th>
<th>No. of AM</th>
<th>No. of BC</th>
<th>No. of FC</th>
<th>No. of cirri in ventral row 1</th>
<th>No. of cirri in ventral row 2</th>
<th>No. of LMC</th>
<th>No. of RMC</th>
<th>Pretransvers</th>
<th>No. of TC</th>
<th>No. of CC</th>
<th>No. of DK</th>
<th>No. of Mac.</th>
<th>No. of Mic.</th>
<th>References</th>
</tr>
</thead>
</table>

Note: Data based on protargol-stained specimens; –: Not mentioned; Mac: Macronuclei; Mic: Micronuclei; DK: Dorsal Kineties; TC: Transverse Cirri; CC: Caudal Cirri; RMC: Right Marginal Cirri; LMC: Left Marginal Cirri; FC: Frontal Cirri; BC: Buccal Cirri; AM: Adoral Membranelles
Table 2 Other morphological features of Pseudoamphisiella species.

<table>
<thead>
<tr>
<th>Species</th>
<th>Extrusome (size, shape, μm)</th>
<th>Alveolar layer (size, shape, μm)</th>
<th>Food vacuoles</th>
<th>Body size (μm × μm)</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>P. quadrinucleata</em></td>
<td>2–3</td>
<td>Present 5</td>
<td>Difficult to recognize</td>
<td>160–220 × 60–100</td>
<td>Shen et al. [6]</td>
</tr>
<tr>
<td><em>P. alveolata</em></td>
<td>Bar-like</td>
<td>Present 3</td>
<td>Not observed</td>
<td>120–200 × 50–70</td>
<td>Hu and Suzuki [4]</td>
</tr>
<tr>
<td><em>P. alveolata</em></td>
<td>Bar-like</td>
<td>Irregular polygonal structure</td>
<td>Difficult to recognize</td>
<td>120–240 × 50–80</td>
<td>Song and Warren [5]</td>
</tr>
<tr>
<td><em>P. lacazei</em></td>
<td>2</td>
<td>Absent</td>
<td>Contain small ciliates and flagellates</td>
<td>120–300 × 40–80</td>
<td>Song et al. [3]</td>
</tr>
<tr>
<td><em>P. lacazei</em></td>
<td>Thin (inconspicuous)</td>
<td></td>
<td>Large and few</td>
<td>200–300 × 70–100</td>
<td>Song [2]</td>
</tr>
</tbody>
</table>

Note: Data based on *in vivo* observation; –: Not mentioned

Bacteria

A large number of bacteria were observed within *P. lacazei* cell. They are embedded within the fibers but not in food vacuoles. There can be two possible explanations for this phenomenon: (1) The bacteria were taken in as food but were not digested. This might have been due either to the accidental bursting of the phagosome membrane or to the inactivated or lost lysosome receptors [16]. (2) A daring speculation is that the bacteria might be the endosymbiont of *P. lacazei*. The phenomenon appears to support the hypothesis that the endosymbiont of the bacterium and the ciliate probably originates from phagocytosis.

Conclusion

1. According to the mucosyts of *P. lacazei*, the Family Pseudoamphishiellidae should be assigned to the order Urostylida.

2. According to the ultrastructure results of the cortex of *P. lacazei*, which consists of two layers of structures, the vacuole layer outer and a fiber layer from inner.

3. Large numbers of bacteria were attached to the fiber layer of *P. lacazei*, two possible explanations for this phenomenon; a daring speculation is that the bacterium might be the endosymbiont of *P. lacazei*.

Acknowledgement

This work was supported by the National Natural Science Foundation of China (No. 31471950, 31101613, and No. 30970311), Heilongjiang Natural Science Foundation (No.C2015021), and Heilongjiang Postdoctoral Funding Project “Study on toxicology of intermediate-type filament system and other ultrastructures of ciliates”. Our thanks also due to Wang Xuedong and Cui Lin, Electron Microscopy Laboratory, Northeast Agriculture University, for technical assistance.
References


