THE CYST OF *UROSTYLA GRANDIS*  
(HYPOTRICHIDA  UROSTYLIDAE) : ULTRASTRUCTURE  
AND EVOLUTIONARY IMPLICATIONS

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ABSTRACT

Ultrasructural study of *Urostyla grandis* cysts shows that the cyst wall is constituted of three layers: a fibrous ectocyst, a structurally homogeneous endocyst and a granular layer that exhibits a dispersed granular or fibrillar structure. The maintenance of only a few kinetosomes and scattered bundles of subpellicular microtubules, and other particularities distinguish the *Urostyla grandis* cyst, both KR and NKR cysts. It is proposed to amplify the WALKER and MAUGEL classification by an additional group. Based on comparative features, some of the phylogenetic relations are discussed.

Key words: cyst, ultrastructure, hypotrich, *Urostyla*. classification.

RÉSUMÉ

L'examen, en microscopie électronique, du kyste d'*Urostyla grandis* a révélé que sa paroi se compose de trois couches : un exokyste fibrillaire, un endokyste à structure homogène et une couche granulaire avec une structure granulaire ou fibrillaire. Le maintien de quelques cinétosomes, microtubules subpelliculaires et d'autres particularités permettent de distinguer ces kystes des kystes KR et NKR. On propose d'ajouter un nouveau groupe à la classification de WALKER et MAUGEL. On commente quelques relations phylogéniques reposant sur des caractéristiques comparatives.


INTRODUCTION

According to the ultrastructural characteristics displayed by the cysts of hypotrich ciliates, WALKER and MAUGEL in 1980, proposed a general classification for these of two groups:


NKR cysts (no kinetosome-resorbing cysts): *Dyophris scutum* (WALKER and MAUGEL, 1980).

Only a few species of the Oxytrichidae have been examined since 1980 (CALVO et al., 1983; GUTJERREZ et al., 1983; MATSUSAKA and HONGO, 1984; VERNI et al., 1984), but all of them exhibit KR cysts; so apparently, KR cysts may be a feature of the Oxytrichidae. In the same way, WALKER and MAUGEL (1980) suggest that the NKR cysts seem to be a characteristic of the Euplotidae.

In this paper, we describe the ultrastructural topography and fine structure of the cyst of the hypotrich ciliate *Urostyla grandis*. Special emphasis has been placed on the fact that the *Urostyla grandis* cyst displays characteristics between KR and NKR cysts and we suggest the
possibility of the establishment of an additional group, including species showing similar resting cysts. These ultrastructural variations may be a new criterion offered to taxonomists to resolve both taxonomic and phylogenetic problems in the Order Hypotrichida.

**MATERIALS AND METHODS**

*Urostyla grandis*, a hypotrich ciliate, was cultivated at 20 ± 1 °C in mineral water and fed *Chlororogonium* sp.

In cultures of *Urostyla grandis*, encystment occurs when food becomes a limiting factor. In order to induce this process, to obtain resting cysts, vegetative cells were transferred from the maintenance culture to mineral water without adding *Chlororogonium* sp.

**Staining for light microscopy**

The nuclear apparatus was stained by using the Feulgen procedure (TORRES, 1977).

**Electron microscopy**

For scanning electron microscopy (SEM), the cysts were fixed for 30 minutes at 4 °C in cacodylate buffer 0.05 M pH 7.2, containing a 1:2 mixture of 1.5 o/o glutaraldehyde and 2 o/o osmium tetroxide. The fixed material was dehydrated by increasing concentrations of acetone, was air-dried, then gold-coated and finally examined in a Jeol scanning electron microscope.

For transmission electron microscopy (TEM), the cysts were fixed for 40 minutes at 4 °C in a mixture 1:2 of 1.5 o/o glutaraldehyde and 2 % osmium tetroxide in a 0.05 M cacodylate buffer pH = 7.2. The fixed material was embedded in 2 o/o agar blocks and dehydrated by increasing concentrations of acetone and embedded in Spurr resin (SPURR, 1968). Ultrathin sections were cut with a Reichert-Jung ultramicrotome, double stained with uranyl-acetate and lead citrate and then examined with a Siemens Elmiscop 102 electron microscope.

Alternatively, the cysts were stained with ruthenium red (Lu FT, 1971) and then processed like those mentioned above.

**RESULTS**

**Light microscopy**

The *Urostyla grandis* cysts are spherical and average 22-24 µm in diameter. The volume of the cyst is smaller than that of the vegetative cell. The cytoplasm exhibits an homogeneous aspect and in older cysts it is possible to observe a dark granule, in the center, that stands out clearly from the rest of the cytoplasm (Fig. 1). Cysts possess the normal vegetative nuclear complement consisting of a lot of macronuclei and a few micronuclei (Fig. 2).

**Electron microscopy**

In electron micrographs obtained by SEM, the cyst surface appears irregular with wrinkles that give a rough

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FIG. 1. — Photomicrograph of a resting cyst of *Urostyla grandis* "in vivo". ec, cytoplasm; g, central granule; w, cyst wall. (1400 x).

FIG. 2. — Resting cyst of *Urostyla grandis* showing a lot of macronuclear fragments. Stained by Feulgen. (1400 x).

FIG. 3. — Scanning electron micrograph of a resting cyst of *Urostyla grandis*. (1400 x).

FIG. 4. — Transmission electron micrograph of a cyst showing cyst wall, remaining subpellicular microtubules and mucocysts. ec, ectocyst; en, endocyst; gl, granular layer; mt, subpellicular microtubules; mu, mucocyst. (27 000 x).

FIG. 5. — Transmission electron micrograph of the cyst wall stained with ruthenium red. ec, ectocyst; en, endocyst; gl, granular layer: (1. - outer ectocyst — 2. - inner ectocyst). (33 000 x).

FIG. 6. — Transmission electron micrograph. Cyst wall and cyst pellicle. ec, ectocyst; en, endocyst; gl, granular layer; p, pellicle. (45 000 x).

FIG. 7. — Transmission electron micrograph showing a pair of kinetosomes beneath the cyst wall. k, kinetosomes. (30 000 x).

FIG. 8. — Mitochondrial cluster. (16 000 x).

FIG. 9. — Cyst cytoplasm showing single mitochondria closely associated to RER and Golgi apparatus. Note transition vesicles. mi, mitochondria; G, Golgi apparatus; tv, transition vesicles; R, rough endoplasmic reticulum. (50 000 x).
Our observations obtained by TEM of *Urostyla grandis* cyst show that the wall consists of three layers, like the *Dyophris scutum* cyst (WALKER and MAUGEL, 1980). On the other hand, the morphology of these layers is basically similar to that of the three inner layers of *Oxytrichid*s cysts (mesocyst, endocyst and granular layer). According to this result, we could say that the *Urostyla grandis* cyst seems to lack the ectocyst present in all *Oxytrichid*s cysts described.

No data have been reported indicating structural differences between the cyst pellicle and the vegetative cell pellicle. However, in *Urostyla grandis* cyst, the outer most membrane unit, which always appears in the vegetative state, has never been observed. It may be due to the degradation of this structure during the encystment process; although another possibility also exists. This membrane unit can remain intact but is masked by the endocyst. Studies in progress, in our laboratory, seem to indicate that the second option is the most probable one and that it is between this membrane-unit and the rest of the pellicle where the material that will form the granular layer will be left. This layer shows a variety of appearances, but, in every case, the cytoplasm matches it. The significance of this fact is as yet unknown.

In most of the ultrathin sections a few remnants of ciliature, consisting of one, two or several kinetosomes can be observed. This characteristic distinguishes the *Urostyla grandis* cyst from the KR cyst, where the ciliature disappears almost completely, and the NKR cyst, in which the ciliature stays almost intact (including AZM). Another singular characteristic of these cysts is the fact that the subpellicular microtubules are reduced to scattered bundles beneath the pellicle.

Since the numerous macronuclear fragments observed in the vegetative cell can be found in the cyst, it is possible to conclude that macronuclear fusion does not take place during the encystment process. No appreciable changes are noticeable in the macronuclear chromatin arrangement; although, occasionally, a small increase of the condensation degree could be seen.

In agreement with the classification proposed by WALKER and MAUGEL (1980) (KR and NKR cysts), all the species belonging to the *Oxytrichidae* family, whose cysts have been studied, exhibit KR cysts. In addition, WALKER and MAUGEL (1980) suggest that the NKR cysts may be a feature of the *Euflagellidae*. However, this classification loses its validity in the *Urostyla grandis* cysts because they show mixed characteristics of both types. Some observations of other species of *Urostylids* reveal that their cysts exhibit the same characteristics described in this paper (unpublished data). For these reasons, we think it is necessary to amplify the classification with a third group, which includes all the species whose cysts: (1) exhibit a three-layered wall, (2) resorb alia cilia but maintain a few kinetosomes and microtubules, (3) display mitochondrial clusters, (4) do not fuse macronuclei, (5) rate cyst volume/vegetative cell volume = 0.25-0.40 and (6) rate cyst wall thickness/cyst radius = 0.08 ± 0.02.

The encystment can be considered as a process of cell differentiation to defend itself against possible environmental challenges. Two general types of hypotrichs cysts (KR and NKR) exist as an adjustment for survival in different environmental conditions. As suggested by different authors, the main adaptive advantages of the formation of NKR cysts seem to be their rapid ability of encystment and their lower energy required for this
process, as a consequence of the conservation of all their cortical structures. With regard to the KR cysts, their higher resistance to extremal values of the enzymatic parameters (due to thicker wall) is an advantage which withstands the loss of valuable cortical markers such as cilia and kinetosomes. In respect to this, the Urostyla grandis cyst appears to be situated half-way between the two, since its resistance properties are lower than the KR cysts, but the ciliary structures that remain in the resting form (kinetosomes and subpellicular microtubules) may be useful indicators of morphogenetically determinative positions for the excystment process.

Up to the present time, the formation of the cyst wall in a few ciliates has been assumed to be the result of mucocysts discharge (CHEISSEN and MOSEVICH, 1962; HOLT and CHAPMAN, 1971; ME ARDLE et al., 1980; REPAK and PFISTER, 1967), at least partially. But, in hypotrichs, the appearance of mucocysts in the cyst has never been reported. In Urostyla grandis (and other species of Urostylidae family, unpublished data) it is possible to observe a lot of these organelles, both in the vegetative cell and the resting form, although their functional role in the encystment process still remains obscure. At this time, in our laboratory, we are studying the structure and function of mucocysts in encysting Urostyla grandis.

If we assume that the cyst described in this report is the general type of the Urostylidae cysts, in our opinion, sufficient data exist to establish a correlation between the morphogenetic division patterns and the cyst types. So, we suppose that the four-layered cyst present in the Oxytricha which exhibits a type of morphogenetical pattern characterizing evolved forms of hypotrichs is derived from the three-layered cyst present in the Urostylids whose morphogen etical pattern characterizes the more primitives hypotrichs. Therefore, the knowledge of the ultrastructure of the cysts support the idea of a lineal phylogeny of Stichotrichina with Sporadotrichina, as proposed by several authors (CORLISS, 1979; MARTIN et al., 1983). However, we also think that the available data on the Euplotidae cysts are not sufficient to draw conclusions about their uncertain origin nor to establish phylogenetic relations although intuitive reasons exist to consider this family as divergent evolutionary line inside the Order Hypotrichida.

REFERENCES