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Studies on woloszynskioid dinoflagellates VI: description of *Tovellia aveirensis* sp. nov. (Dinophyceae), a new species of Tovelliaceae with spiny cysts

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A new species of *Tovellia*, *T. aveirensis*, is described on the basis of light (LM) and scanning electron microscopy (SEM) of motile cells and resting cysts, complemented with transmission electron microscopy (TEM) of flagellate cells and phylogenetic analysis of partial sequences of the large subunit ribosomal RNA gene. Both vegetative cells and several stages of a life cycle involving sexual reproduction and the production of resting cysts were examined in cultures established from a tank in the University of Aveiro campus. Vegetative cells were round and little compressed dorsoventrally; planozygotes were longer and had a proportionally larger epicone. Chloroplast lobes were shown by TEM to radiate from a central, branched pyrenoid, although this was difficult to ascertain in LM. The amphiesma of flagellate cells had mainly 5 or 6-sided vesicles with thin plates, arranged in 5–7 latitudinal series on the epicone, 3–5 on the hypocone. The cingulum had 2 rows of plates, the posterior row extending into the hypocone and crossed by a series of small projecting knobs along the lower edge of the cingulum. A line of narrow amphiesmal plates extended over the cell apex, from near the cingulum on the ventral side to the middle of the dorsal side of the epicone. Eight or 9 narrow amphiesmal plates lined each side of this apical line of plates (ALP). Resting cysts differed from any described before in having numerous long, tapering spines with branched tips distributed over most of the surface. Most mature cysts showed an equatorial constriction. Neither cysts nor motile cells were seen to accumulate red cytoplasmic bodies in any stage of the cultures. The phylogenetic analysis placed, with high statistical support, the new species within the genus *Tovellia*; it formed a clade, with moderate support, with *T. sanguinea*, a species notable for its reddening cells.

**Key words:** cyst, dinoflagellates, LSU rDNA, phylogeny, taxonomy, Tovelliaceae, *Tovellia aveirensis*, ultrastructure

Introduction

Recent studies on the so-called woloszynskioid dinoflagellates, traditionally characterized by having a cell cover of numerous thin amphiesmal plates, revealed a heterogeneous, polyphyletic assemblage (Lindberg et al., 2005; Moestrup et al., 2008). The consequent reclassification of woloszynskioid species led to the establishment of several genera distributed over different families: *Tovellia* Moestrup, K. Lindberg & Daugbjerg, *Jadwigia* Moestrup, K. Lindberg & Daugbjerg, *Esoprotodinium* Javornicky (a probable synonym of *Bernardinium* Chodat) and *Opisth Haoalax* Calado are currently included in the family Tovelliaceae (Lindberg et al., 2005; Calado et al., 2006; Calado, 2011; Fawcett & Parrow, 2012; Borghiella Moestrup, Gert Hansen & Daugbjerg and *Baldinia* Gert Hansen & Daugbjerg are placed in the Borghiellaceae (Hansen et al., 2007; Moestrup et al., 2008, 2009a); *Biecheliar* Moestrup, K. Lindberg & Daugbjerg and *Biecheliopsis* Moestrup, K. Lindberg & Daugbjerg are placed in the Suessiaceae (Moestrup et al., 2009a, 2009b). These extensive taxonomic changes are supported by molecular data and by morphological differences in eyespot structure, organization of the cell apex and type of resting cyst.

Members of the Tovelliaceae typically possess an extraplastidial eyespot composed of pigment globules not bounded by membranes (eyespot type C in Moestrup & Daugbjerg, 2007). In addition, in *Tovellia* and *Jadwigia*, a straight or slightly curved line of narrow plates provided with a row of knobs (apical line of plates, ALP, sensu Lindberg et al., 2005) is present on the epicone and lined on each side by a row of amphiesmal plates narrower than the average hexagonal or pentagonal vesicles of the amphiesma. *Tovellia* species produce resting cysts...
Tovellia aveirensis, sp. nov.

with an equatorial constriction (sometimes interpreted as a paracingulum), axial horns and pre- and postcingular protuberances or scattered short spines; cysts with equatorial constriction and axial horns have also been demonstrated in *Opisthoxoaulax vorticella* (F. Stein) Calado (Lindberg et al., 2005; Moestrup et al., 2006; Calado 2011). In contrast Jadwigia and *Esotropondium/Bernardinium* produce smooth, round cysts (Lindberg et al., 2005; Calado et al., 2006).

The present work describes a new species of *Tovellia* with a distinctive type of cyst, slightly constricted in the middle and ornamented by numerous processes that are variously branched near the tip. The morphology of swimming cells and cysts is described on the basis of light (LM) and scanning electron microscopy (SEM), and the general internal fine structure of swarmers is shown in transmission electron microscopy (TEM). A phylogenetic analysis based on partial nuclear-encoded large subunit ribosomal RNA gene sequences (LSU rDNA) corroborates the taxonomic assignment of the new species. This is the first species of *Tovellia* to be reported from Portugal (Pandeirada et al., 2013).

Materials and methods

Biological material

The organism described in this work was found in a net sample (mesh size 25–30 μm) collected from a clean water tank at the University of Aveiro Campus, Aveiro, Portugal, on 12 October 2009. Several swimming cells were transferred to one culture well with L16 medium (Lindström, 1991) supplemented with vitamins according to Popovský & Pflieger (1990) and maintained at 18°C with 12:12 light:dark photo-period. One month later, five cysts were observed in the well. These were re-isolated separately to five wells with the same medium and placed under the same temperature and light conditions. Three cysts germinated and gave rise to three culture lines.

Light microscopy (LM)

Light micrographs of motile cells and cysts were taken with a Zeiss Axioplan 2 imaging light microscope (Carl Zeiss, Oberkochen, Germany) equipped with a DP70 and a ColorView IIIu Olympus camera (Olympus Corp., Tokyo, Japan). Asexual reproduction was recorded with a JVC TK-C1481BEG colour video camera (Norbain SD Ltd, Reading, UK) mounted on a Leitz Biomed light microscope (Leica, Oberkochen, Germany). A clear visualization of amphiesmal vesicles (by removing the outer membrane) of the swimming cells was obtained with the following procedure: 1.5 ml of culture was fixed for 25 min by adding 1 ml of a fixative mixture made of saturated HgCl₂ and 2% osmium tetroxide in a proportion of 1 : 5. After fixation, cells were retained on Isopore polycarbonate filters with 8-μm pore size (Millipore Corp., Billerica, Massachusetts, USA) and washed with distilled water for 10 min. Dehydration of filters with the cells was performed with a graded ethanol series with an overnight stop in 70% ethanol at 4°C. Dehydration was completed the following day and the material was critical-point-dried in a Baltec CPD-030 (Balzers, Liechtenstein). The dried filters were glued onto stubs, sputter-coated with gold-palladium and examined with a Hitachi S-4100 (Hitachi High-Technologies Corp., Tokyo, Japan) scanning electron microscope.

Cysts were prepared in a similar way with a different fixative proportion: 1 ml of culture was added to 0.5 ml of fixative mixture made of saturated HgCl₂ and 2% osmium tetroxide in a proportion of 1 : 3, for 15 min.

Transmission electron microscopy (TEM)

Swimming cells from culture were picked up, transferred to 2% glutaraldehyde in phosphate buffer 0.1 M, pH 7.2, and fixed for 1 h 15 min. After being washed in the same buffer, cells were embedded in 1.5% agar blocks and post-fixed overnight in 0.5% osmium tetroxide in the same buffer. The agar blocks with the cells were rinsed in phosphate buffer and distilled water. Following dehydration through a graded ethanol series and propylene oxide they were embedded in Spurr’s resin. The resin blocks were cured for about 14 h at 70°C. Cells were sectioned with a diamond knife on an EM UC6 ultramicrotome (Leica Microsystems, Wetzlar, Germany). Ribbons of sections 70 nm thick were picked up with slot grids and placed on Formvar film. Sections were stained with uranyl acetate and lead citrate and examined using a Zeiss EM 10A (Carl Zeiss, Oberkochen, Germany) transmission electron microscope operated at 60 kV.

DNA extraction and PCR amplification of LSU rDNA

Ten ml of exponentially growing culture line MSP14c was harvested by centrifugation at 1201 g for 10 min. Most of the supernatant was discarded and the cell pellet resuspended in about 100 μl growth media. The material was transferred to a 1.5 ml Eppendorf tube and allowed to freeze for 2 days at −18°C. After this time 30 μl of the cell pellet was extracted using the CTAB method of Doyle & Doyle (1987) with a few modifications as previously outlined by Daugbjerg et al. (1994). PCR amplification of partial LSU rDNA (nearly 1400 base pairs) was performed as stated in Hansen & Daugbjerg (2011). The PCR amplified fragments of LSU rDNA were purified as described in Craveiro et al. (2013). 30 ng of DNA was used for sequence determinations in both directions using external primers (i.e. D1R and ND28–1483R) and internal primers (i.e. D3A, D3B and D2CR). See Daugbjerg et al. (2013) and Scholin et al. (1994) for primer sequences. For sequencing we used the service provided by Macrogen, Korea.

Single cell PCR partial amplification of LSU rDNA

For comparison between culture lines, single cells or pairs of cells of a different strain, MSP14b, were isolated, using a micropipette under a Leitz Labovert FS inverted microscope, into 0.2-ml PCR tubes and frozen at −8°C for 3 days before
PCR reactions. Cell DNA constituted the template to amplify about 1500 base pairs (bp) of the LSU rRNA gene using the terminal primers D1R (Scholin et al., 1994) and 28-1483R (Daugbjerg et al., 2000). These were added to the PCR tubes with the isolated cells, followed by illustra™ puReTaq Ready-To-Go PCR Beads (GE Healthcare UK Ltd) containing all other chemicals necessary for PCR amplification, and amplified in a Biometra–TProfessional Trio thermocycler. Thermal cycling for PCR amplification was as outlined in Moestrup et al. (2008), but with a longer final extension step of 10 min (rather than 6 min). DNA fragments were loaded on a 1% agarose gel, run for 20 min at 90 V and viewed under a UV light table (Molecular imager chemiDoc XRS System from Bio-Rad Laboratories, Inc.). PCR products were purified using the QIAquick PCR Purification Kit (Qiagen), following the manufacturer’s recommendations, and sent to Macrogen Europe (Amsterdam, the Netherlands) for sequence determination in both directions. The sequencing primers used were D1R, D2CR, D3A, D3B and ND28-1483R. The sequences obtained were identical to that of MSP14c.

Alignment and phylogeny

Sequence alignment was optimized based on the secondary structure of the LSU rRNA molecule as proposed by De Rijk et al. (2000) and further edited by eye using Jalview, ver. 2.8 (Waterhouse et al., 2009). Our data matrix comprised 1173 base pairs including introduced gaps. This corresponded to a DNA fragment that covered variable domain D1 to 8 base pairs of the variable domain D6 (sensu Lenaerts et al., 1989). However, domain D2 was omitted, as this is known to be highly divergent and therefore a challenging task to align unambiguously. To infer the phylogeny of Tovellia aveirensis we used both Bayesian (BA) and maximum likelihood (ML) analyses. For BA we used MrBayes ver. 3.2.2 (Ronquist & Huelsenbeck, 2003) and for ML we used PhyML ver. 3.0 (Guindon & Gascuel, 2003). In BA we used 20×10⁶ generations and a tree was sampled every 1000th generation. The BA analysis was run on a local computer. In order to evaluate the burn-in value we plotted the LnL values as a function of generations and a tree was sampled every 1000th generation. The burn-in value we plotted the LnL values as a function of generations and a tree was sampled every 1000th generation in BA analysis was run on a local computer. In order to evaluate tree topologies was evaluated using bootstrapping with 1000 replications.

Comparative studies using ultrastructural characters and phylogenetic reconstructions based on molecular data (e.g. Van de Peer et al., 1996) have indicated that Ciliata and Apicomplexa comprise the sister group to the Dinophyceae. Thus, we used four species of ciliates (Euplotes aediculatus, Spatidium amphoriforme, Tetrahymena thermophila, T. pyriformis), seven species of apicomplexans (Cryptosporidium parvum, Eimeria tenella, Hammondia hammondi, Neospora canium, Sarcocystis neurona, Theileria parva, Toxoplasma gondii) and the Perkinsid Perkinsus andrewsi to polarize the ingroup of dinoflagellates. The ingroup consisted of a diverse assemblage of dinoflagellates representing 39 genera and 64 species.

We used PAUP* (ver. 4.0b10) to calculate the sequence divergence estimate between all pairwise comparisons involving three species of Tovellia, Esotropondium gemma and Jadwigia applanata.

Results

Tovellia aveirensis Pandeirada, Craveiro, Daugbjerg, Moestrup & Calado, sp. nov. (Figs 1–21)

Description: Vegetative cells ovoid or nearly spherical, slightly compressed dorsoventrally or not at all. Cingulum descending, displaced about one cingulum width. Epicone broadly round, slightly longer than wide and often somewhat longer than the obliquely flattened hypocone. Cells 25–34 µm long, 17–24 µm wide and 14–21 µm thick. Chloroplast lobes yellowish-green, radiating from the cell centre towards the periphery. Nucleus in the hypocone. Eyespot nearly rectangular in ventral view, located in the sulcal area. Cell cover mainly formed by pentagonal or hexagonal amphiesmal vesicles roughly arranged in latitudinal series, 5–7 series on the epicone and 3–5 on the hypocone. Cingulum with 2 series of vesicles, the anterior vesicles abutting the sharply defined anterior cingulum edge whereas the roughly hexagonal vesicles of the posterior row extend into the hypocone over the rounded posterior cingulum edge. A row of knobs marked the posterior edge of the cingulum. The initial part of the cingulum showed additional two or three nearly hexagonal vesicles intercalated between the two regular vesicle rows. A line of narrow vesicles started near the proximal end of the cingulum, on the ventral side, and extended over the apex of the cell. Cysts yellowish-brown, ornamented by branched processes that were usually absent in the equatorial area, which was often slightly constricted. Nuclear-encoded partial LSU rRNA gene sequence = GenBank accession KF819359.

Holotype: SEM stub with critical point dried material from culture line MSP14c, fixed to display the amphiesmal vesicles, deposited at the University of Aveiro Herbarium registered as AVE-A-T-4. Figures 8–13 illustrate cells from this stub.

Isotype: SEM stub with critical-point-dried material from culture line MSP14c, containing swarvers, mostly retaining outer membranes, and mature cysts, deposited at the University of Aveiro Herbarium registered as AVE-A-T-5. Figures 22–25 illustrate cysts from this stub.

Type locality: Freshwater (conductivity about 300 µS) tank at the University of Aveiro Campus, Aveiro, Portugal (40°38’4.52”N, 8°39’30.21”W), collected on 12 October 2009.
Etymology: Latin aveirensis, ‘from Aveiro’, in reference to both the city and the University of Aveiro Campus, where the species was found.

**General morphology of motile cells**

Motile cells are illustrated in LM and SEM in Figs 1–13. Vegetative cells were ovoid or nearly spherical, not compressed dorsoventrally at cingulum level or only slightly so (Figs 1–3, 8–11). The cingulum was located slightly below the middle of the cell and had the distal (right-hand side) end about one cingulum width below the proximal end (Figs 1, 8). The epicone was broadly round and slightly longer than wide (Figs 1, 2, 8). In side view the hypocone appeared obliquely flattened (Figs 3, 10). Cells were 25–34 μm long (n = 30), 17–24 μm wide (n = 30) and 14–21 μm thick (n = 19). A bright-red, trough-shaped eyespot (rectangular in ventral view) underlay the full width of the sulcus (3–3.5 μm in the proximal half) along up to 4 μm of its length, although it was usually smaller in non-vegetative cells (Figs 1, 4, 6, 7). The sulcus widened somewhat in its posterior half (Figs 8, 11). Vegetative cells displayed numerous yellowish-green chloroplast lobes near the surface (Figs 1, 3). A radial arrangement of chloroplast lobes was barely discernible in optical sections of the epicone (Fig. 2). The roundish to transversely ellipsoid nucleus occupied a large portion of the hypocone (Figs 3, 5). Vegetative cells swam with a regular, continuous motion, usually rotating around the longitudinal axis. Smaller, roughly spherical cells, 16.5–20 μm long (n = 5), 12–15.5 μm wide (n = 5), about 12 μm thick (n = 2), commonly appeared in the cultures; they had fewer chloroplast lobes, to the point of sometimes appearing nearly colourless (Figs 4, 5), and occasional accumulation bodies (Fig. 5, arrowhead). These rapidly swimming small cells were seen fusing with one another to yield the formation of planozygotes and are therefore referred to as gametes. Planozygotes, identified by the presence of two longitudinal flagella (Fig. 7), were usually more elongated than vegetative cells (Figs 6, 7). They were 33–49 μm long and 24–37 μm wide (n = 9). Planozygotes increased in size over time and eventually became brownish. Larger, darker planozygotes swam slowly, sometimes rotating on the
Figs 8–11. *Tovellia aveirensis*, motile cells, SEM. **Fig. 8.** Ventral view. The ALP is indicated by arrowheads. The vertical arrows point to the row of knobs at the posterior edge of the cingulum. Two additional vesicles intercalated between the two regular series of cingular plates are marked at the proximal end of the cingulum (asterisks). Note the ventral ridge (vr) and the longitudinal flagellum. **Fig. 9.** Dorsal view. The ALP (arrowheads) is separated from the cingulum by about five series of plates. The sharply delineated anterior edge of the cingulum is marked by arrows. **Fig. 10.** View from the right-ventral side showing the slanting ventral face of the hypocone. The arrowhead indicates the proximal end of the ALP. The inset shows a higher magnification of the row of knobs across cingular plates. Inset scale bar = 2 µm. **Fig. 11.** Approximate antapical view showing sulcal plates (sp) arranged in a single line. The asterisk marks the antapex, where a slightly larger plate appears surrounded by latitudinal series of hypocone plates. The ventral ridge (vr) is seen rising above the cingulum-sulcus area. All scale bars except inset = 5 µm.
Figs 12–13. *Tovellia aveirensis*, motile cell apex, SEM. **Fig. 12.** The ALP (arrowheads) extends over the epicone and is separated from the cingulum by three series of plates. **Fig. 13.** Detail of the ALP. Sutures between individual plates of the ALP are indicated by arrowheads. A row of small knobs (arrow) ornaments each plate of the ALP. The narrow thecal plates bordering the ALP are marked with asterisks. Scale bar in Fig. 12 = 3 μm, in Fig. 13 = 1 μm.

Figs 14–17. *Tovellia aveirensis*, TEM of motile cell. **Fig. 14.** Longitudinal section through the middle of the cell, showing chloroplast lobes (c) radiating from central pyrenoid complex (Py). The somewhat curved nucleus (n) is visible in the hypocone (nu, nucleolus). A few trichocysts (t) and oil droplets (o) are visible in the peripheral cytoplasm. Starch grains (s) accumulate between chloroplasts lobes and near the antapex. Arrowheads indicate a pusular tube in the central cytoplasm. **Fig. 15.** Detail of central pyrenoid complex showing scattered thylakoid lamellae. **Fig. 16.** Pusular tube with diverticula. **Fig. 17.** Sulcal area in approximately grazing section, with the eyespot (e) and the longitudinal microtubular root (LMR/r1). Scale bars: Fig. 14 = 10 μm; Figs 15, 16 = 0.5 μm; Fig. 17 = 2 μm.

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**Figs 14–17. Tovellia aveirensis**, TEM of motile cell. **Fig. 14.** Longitudinal section through the middle of the cell, showing chloroplast lobes (c) radiating from central pyrenoid complex (Py). The somewhat curved nucleus (n) is visible in the hypocone (nu, nucleolus). A few trichocysts (t) and oil droplets (o) are visible in the peripheral cytoplasm. Starch grains (s) accumulate between chloroplasts lobes and near the antapex. Arrowheads indicate a pusular tube in the central cytoplasm. **Fig. 15.** Detail of central pyrenoid complex showing scattered thylakoid lamellae. **Fig. 16.** Pusular tube with diverticula. **Fig. 17.** Sulcal area in approximately grazing section, with the eyespot (e) and the longitudinal microtubular root (LMR/r1). Scale bars: Fig. 14 = 10 μm; Figs 15, 16 = 0.5 μm; Fig. 17 = 2 μm.
same spot, and finally developed into walled cysts (see below).

Structure of the amphiesma

The cell cover was formed by many amphiesmal vesicles containing thin thecal plates, mainly pentagonal or hexagonal, roughly arranged in latitudinal series, 5–7 on the epicone and 3–5 on the hypocone (Figs 8–12). The strict latitudinal arrangement was disrupted by occasional plates intercalated between series. In the precingular series, plates were mainly pentagonal with the posterior edges aligned to form the sharply defined anterior border of the cingulum (arrows in Fig. 9). The cingulum had two series of plates: an anterior row of pentagonal plates adjacent to the distinct upper cingulum edge, and a posterior row of mainly hexagonal plates that extended into the hypocone for about half of their length (Figs 8–10). A transverse row of knobs around the middle of the posterior row of plates surrounded the cell, marking the posterior edge of the cingulum (Figs 8–10). Most cells had five–eight knobs per plate (n = 10). Variations were observed, such as the presence of two rows of knobs in some plates, or knobs fused, scattered or disposed irregularly (not shown). On the left, proximal part of the cingulum, there were usually one–two additional, nearly hexagonal vesicles (Fig. 8, asterisks); similar vesicles more seldom appeared intercalated in more distal parts of the cingulum (not shown). A straight or slightly curved apical line of narrow plates (ALP) extended past the apex of the epicone, from the ventral to the dorsal side of the cell (Figs 8–10, 12, 13). The ALP was separated from the cingulum on the ventral side by one plate (Figs 8, 10) and by three–five plates on the dorsal side (Figs 9, 12). Plates of the ALP were 0.15–0.35 μm wide (n = 37) and three–seven times longer, and displayed an axial row of small knobs (Fig. 13). The ALP was lined on each side by a row of narrow amphiesmal plates 0.45–1.3 μm wide (n = 30) (Figs 12, 13). Eight to nine plates were present in each of the rows of plates bordering the ALP, as estimated from SEM views in which most of the structure was visible (n = 10). The antapex of the cells did not show a thickened or otherwise prominent, clearly antapical plate. A heptagonal plate is indicated in Fig. 11 (asterisk), which seems to be surrounded by regularly arranged hexagonal plates, but we did not identify a similar plate in other cells. The sulcal area was lined by three plates in a single row (Fig. 11). Directly anterior to the sulcus a linear elevated area above the exit pore of the cingular flagellum seemed to mark the beginning of the cingulum; on the basis of its position and appearance this was interpreted as a ventral ridge (Figs 8, 11, vr).

General internal fine structure

The general internal structure of a cell is shown in longitudinal section in Fig. 14; size and shape of this cell suggest it was a planozygote. Chloroplast lobes radiated from a central pyrenoid complex toward the periphery (Fig. 14). The deeply lobed pyrenoid area was crossed by scattered thylakoid lamellae (Fig. 15). The nucleus and a nucleolus are visible in the hypocone in Fig. 14. The pusular system was located centrally, at cingulum level (Fig. 14, arrowheads); it was formed by a tube some 250 nm wide with numerous diverticula about 160 nm long and constricted at the base (Fig. 16). An eyespot was located underneath the sulcus, beneath the longitudinal microtubular root (LMR, r1 sensu Moestrup, 2000), with the appearance of fused oil droplets not surrounded by a membrane (eyespot type C sensu Moestrup & Daugbjerg, 2007).
(Fig. 17). Oil droplets (Fig. 14, o) were more prominent in the anterior cytoplasm and starch grains were present in groups between chloroplast lobes (Fig. 14, s). Scattered trichocysts were visible in the peripheral cytoplasm (Fig. 14, t).

Cysts

Different stages of cyst development are shown in Figs 18–21. Mature cysts were elongate-ellipsoid and usually displayed a transverse constriction a little off the middle, in a position correspondent to the cingulum of planozygotes, here interpreted as a paracingulum (Figs 20–22, 24). Cyst length varied between 33 and 44 μm and width between 21 and 28 μm (n = 32). Cyst contents were usually yellowish-brown with green and colourless areas. Traces of the eyespot were sometimes visible in cysts (Fig. 20, arrow), and brown accumulation bodies were often present (Fig. 21, arrowheads). The cyst wall had a rough surface and no traces of amphiesmal plates (Figs 22–24). Mature cysts were ornamented by tapering wall processes, which were usually absent from the paracingulum area (20, 21, 22–24). Most processes were branched near the tip in a way that was reminiscent of the antlers of a deer (Fig. 25). Full
grown processes were 7.5–10 μm long (n = 41). Cysts in earlier stages of development were ovoid and often did not show a paracingulum (Figs 18, 19). Initial stages showed only small protuberances (Fig. 18) whereas intermediate stages were ornamented by short, but already branched processes (Fig. 19).

Asexual reproduction

Asexual reproduction occurred in the immobile stage (within a division cyst), usually giving origin to four cells. Cells about to divide stopped on the bottom of culture wells, lost their flagella, increased in size and became more round. Cleavage furrows eventually became visible in the peripheral cytoplasm. An advanced division stage is shown in Fig. 26, with four cells already formed inside the division cyst. The daughter cells in this stage appeared already formed, with visible furrows and outward-facing eyespots, but no traces of flagella. Segregation and subsequent release of daughter cells usually occurred over a few minutes, during which cells slowly slid apart and emerged through an opening in the somewhat inflated cover of the division cyst (Figs 27, 28). Shortly after, flagella became visible and cells started to swim, one after another (Figs 28, 29). Division cysts with only two cells were also seen, but daughter cell release from those cysts was not observed.

Molecular phylogeny

The phylogenetic relationships of Tovellia aveirensis are shown in Fig. 30. Tovellia species clustered in a single clade highly supported by Bayesian (posterior probability, pp = 1.0) and maximum likelihood analyses (bootstrap support, BS = 100%), thus confirming the monophyly of the genus. Tovellia aveirensis formed a sister taxon to T. sanguinea. However, this relationship received high Bayesian support (pp = 0.97) but only moderate bootstrap support (67.6%). Tovellia coronata formed a sister taxon to the T. sanguinea/T. aveirensis clade. The family Tovelliaceae received high support from Bayesian analysis (pp = 1.0) but only 70.3% in ML bootstrap support. Within Tovelliaceae Esoptrodinium gemma formed a sister taxon to Tovellia, and Jadwigia a sister taxon to Tovellia spp. and E. gemma (Fig. 30). Except for the early branching of Moestrupia oblonga (pp = 0.85 and BS < 50%) the topology of the deepest lineages was unresolved in the phylogenetic analyses conducted here (Fig. 30). However, the class Dinophyceae received high support (pp = 1.0, BS = 99.8%).

Sequence divergence estimates were based on 1027 base pairs, including introduced gaps (Table 1). Despite the supposedly close relationship between Tovellia spp., Esoptrodinium gemma and Jadwigia planulata (all members of the family Tovelliaceae) the highly variable domain D2 could not be aligned unambiguously. Hence, it was excluded prior to sequence divergence estimations. As expected the highest sequence divergences were seen in all pairwise comparisons between Jadwigia/Esoptrodinium and any of three Tovellia species (ranging from 12.4–16.5% based on P values and 13.7–18.8% based on Kimura-2-parameter values). The lowest divergence estimate was seen when comparing Tovellia coronata and T. aveirensis (~4% in both calculations). The divergence between T. sanguinea and T. aveirensis was 6.3% or 6.7% and between T. sanguinea and T. coronata it was 5.1% or 5.3%, depending on the calculation method (P values versus K-2-p model) (Table 1).

Discussion

Morphology and taxonomic affinities

The tovelliacean affinity of live swimming cells of the species we report on is mainly suggested by the prominent, trough-shaped eyespot combined with the firm, thin cell cover that gives them a woloszynskioid appearance. The confirmation of the type C eyespot (Moestrup & Daugbjerg, 2007) and the demonstration of a tubular pusule with diverticula, both by TEM observations, firmly place the new species in the

Figs 26–29. Tovellia aveirensis, asexual reproduction, LM. The images were prepared from still frames of a video recording and are marked with the time elapsed between the moments they were recorded. Fig. 26. Division cyst with four daughter cells already formed inside. Fig. 27. Beginning of cell separation. The four cells started to slide apart and emerging from the division cyst (the cyst cover is marked with an arrowhead in Fig. 29). Cells did not have developed flagella at this stage. Fig. 28. The first cell swam away from the group (arrow). The flagella of the remaining cells also begin to be visible in the video recording at this point (although they are indistinct in the still image). Fig. 29. Only one cell still remains within in the division cyst cover (arrowhead); it swam away seconds later. All figures at the same scale. Scale bar = 10 μm.

Table 1
Fig. 30. Molecular phylogeny of *Tovellia aveirensis* sp. nov. (in bold font) and 63 other dinoflagellate species inferred from Bayesian analysis of nuclear-encoded LSU rDNA sequences. Four ciliates, seven apicomplexans and *Perkinsus* formed the outgroup. The first numbers at internal nodes are posterior probabilities ($\geq 0.5$) from Bayesian analysis (BA) and the last numbers are bootstrap values ($\geq 50\%$) from maximum likelihood (ML) with 1000 replications. Filled circles illustrate the highest possible support in BA and ML (1.0 and 100\%, respectively). GenBank accession numbers are written in parentheses. The family Tovelliaceae is marked in grey.
Tovellia aveirensis and T. sanguinea have two rows of narrow plates bordering the ALP whereas in Jadwigia the ALP is in contact with plates that are not fundamentally different from other amphiblesmal plates on the epicone (Lindberg et al., 2005). The two rows of narrow plates lining the ALP of the species described herein clearly point to Tovellia.

Another link to Tovellia is the morphology of the resting cysts produced in cultures of T. aveirensis. The different cyst types produced by woloszyńskioids were among the first pieces of evidence suggesting the group was polyphyletic (Stosch, 1973), and cyst morphologies have been used as generic level characters in the group (Lindberg et al., 2005; Moestrup & Daugbjerg, 2007). Woloszyńskioid resting cysts that are clearly bipolar and show an equatorial constriction or paracingulum have only been found in Tovellia, in Opisthaoaulax and in the marine suessiacean species Polarella glacialis Montresor, Procaccini & Stoecker (Montresor et al., 1999; Moestrup et al., 2009a; Calado, 2011). Vegetative cell characters and habitat separate Opisthaoaulax and Polarella from T. aveirensis.

### Comparison with other Tovellia species

Seven species are currently classified in the genus Tovellia (see comparative overview in Table 2). The strongly flattened motile cells of T. leopoliensis (Woloszyńska) Moestrup, K. Lindberg & Daugbjerg, and the sharply pointed or projecting apices of both this species and T. apiculata (Stosch) Moestrup, K. Lindberg & Daugbjerg contrast with the overall round appearance of T. aveirensis cells (Woloszyńska, 1917; Stosch, 1973). Swimming cells and cysts of both T. coronata (Woloszyńska) Moestrup, K. Lindberg & Daugbjerg and T. sanguinea Moestrup, Gert Hansen, Daugbjerg, Flaim & d’Andrea typically display cytoplasmic accumulations of red pigment, sometimes to the point of large populations of these species conferring a strong red colour to the water (Flaim et al., 2004; Lindberg et al., 2005; Moestrup et al., 2006).

Neither swimming cells nor resting cysts of T. aveirensis were ever seen to acquire a red colour regardless of the age or state of growth of the batch cultures examined. The little known T. glabra (Woloszyńska) Moestrup, K. Lindberg & Daugbjerg was originally described as a variety of T. coronata from which it differed only in not having a prominent, punctate antapical plate (Woloszyńska, 1917); Thompson (1951) described T. coronata-like cells without such a plate, therefore identifiable as T. glabra, without noting red cytoplasmic bodies. Thompson’s (1951) illustrations of T. glabra show cells shorter and more broadly rounded epicones than in T. aveirensis, and Woloszyńska’s (1917) concept of the taxon included larger amphiplesmal plates, distributed over a smaller number of latitudinal rows, than shown here for T. aveirensis.

### Table 2

<table>
<thead>
<tr>
<th>Tovellia aveirensis</th>
<th>Tovellia sanguinea</th>
<th>Tovellia coronata</th>
<th>Esoptrordinium gemma</th>
<th>Jadwigia applanata</th>
</tr>
</thead>
<tbody>
<tr>
<td>T. aveirensis</td>
<td>–</td>
<td>6.3</td>
<td>4.0</td>
<td>16.5</td>
</tr>
<tr>
<td>T. sanguinea</td>
<td>6.7</td>
<td>–</td>
<td>5.1</td>
<td>16.5</td>
</tr>
<tr>
<td>T. coronata</td>
<td>4.1</td>
<td>5.3</td>
<td>–</td>
<td>15.2</td>
</tr>
<tr>
<td>E. gemma</td>
<td>18.8</td>
<td>18.8</td>
<td>17.2</td>
<td>–</td>
</tr>
<tr>
<td>J. applanata</td>
<td>16.4</td>
<td>16.2</td>
<td>14.8</td>
<td>13.7</td>
</tr>
</tbody>
</table>

Reliable identification of Tovellia species from observations on motile cells remains a demanding task that usually requires detailed observation of the cell cover. This is nearly impossible to accomplish in live cells, and even with SEM observations of...
## Table 2. Comparative overview of characters described for *Tovellia* species and *Jadwigia applanata*.

<table>
<thead>
<tr>
<th>Jadwigia and Tovellia species</th>
<th>length × width (µm)</th>
<th>cell shape</th>
<th>red pigment bodies</th>
<th>nucleus</th>
<th>chloroplast arrangement</th>
<th>plate series in epicone, cingulum, hypocone</th>
<th>apical line of plates (ALP)</th>
<th>cyst</th>
<th>Sources</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Jadwigia applanata</em></td>
<td>24–34 × 24–32</td>
<td>ovoid to round, compressed dorsoventrally</td>
<td>absent</td>
<td>longitudinally elongate, along right side of epicone</td>
<td>parietal</td>
<td>6–7, 1, 5–6</td>
<td>lateral series absent, line long, oblique</td>
<td>spherical, smooth</td>
<td>1</td>
</tr>
<tr>
<td><em>T. apiculata</em></td>
<td>25–46 × 18–36</td>
<td>round to obpyriform, slightly compressed dorsoventrally, apices pointed</td>
<td>absent</td>
<td>laterally elongate (sausage-shaped), dorsal in hypocone</td>
<td>radiating from cell centre (pyrenoid?)</td>
<td>6–7, 1, 4–5</td>
<td>long</td>
<td>bipolar with axial, pointed horns and scattered protuberances numerous branched processes</td>
<td>2</td>
</tr>
<tr>
<td><em>T. aveirensis</em></td>
<td>25–34 × 17–24</td>
<td>ovoid to nearly spherical</td>
<td>absent</td>
<td>roundish to transversely ellipsoid in hypocone</td>
<td>radiating from central pyrenoid complex</td>
<td>5–7, 2, 3–5</td>
<td>long</td>
<td>1, 3</td>
<td></td>
</tr>
<tr>
<td><em>T. coronata</em></td>
<td>25–30 × 24–32</td>
<td>nearly spherical</td>
<td>often present</td>
<td>roundish to transversely ellipsoid in hypocone</td>
<td>parietal</td>
<td>4, 1, 4 (antapical plate distinct)</td>
<td>medium length bipolar with axial, pointed horns and scattered protuberances</td>
<td>1, 3</td>
<td></td>
</tr>
<tr>
<td><em>T. glabra</em></td>
<td>19–35 × 14–32</td>
<td>nearly spherical to slightly compressed dorsoventrally</td>
<td>presumably as in <em>T. coronata</em></td>
<td>presumably as in <em>T. coronata</em></td>
<td>parietal</td>
<td>series presumably as in <em>T. coronata</em></td>
<td>presumably as in <em>T. coronata</em></td>
<td>unknown</td>
<td>3, 4</td>
</tr>
<tr>
<td><em>T. leopoliensis</em></td>
<td>c. 40 × c. 35</td>
<td>strongly compressed dorsoventrally, apices pointed</td>
<td>absent</td>
<td>laterally elongate (horseshoe-shaped), near cell middle elongated, low in hypocone</td>
<td>parietal?</td>
<td>8, 1, 6</td>
<td>incompletely described bipolar with axial, pointed horns and scattered shorter spines</td>
<td>3</td>
<td></td>
</tr>
<tr>
<td><em>T. nygaardii</em></td>
<td>20–28 × 20–24</td>
<td>ovoid to spherical, slightly compressed dorsoventrally</td>
<td>absent</td>
<td>radiating from cell centre (pyrenoid?)</td>
<td>parietal?</td>
<td>3, 1, 2</td>
<td>unreported bipolar with axial, pointed horns</td>
<td>5</td>
<td></td>
</tr>
<tr>
<td><em>T. sanguinea</em></td>
<td>c. 24 × c. 19</td>
<td>ellipsoid-elongate</td>
<td>mainly present</td>
<td>roundish to transversely ellipsoid in hypocone</td>
<td>radiating from central pyrenoid complex</td>
<td>5, 1, 2 (antapical plate distinct)</td>
<td>short, only near apex bipolar with axial, pointed horns</td>
<td>6</td>
<td></td>
</tr>
<tr>
<td><em>T. stoschii</em></td>
<td>30–37 × 28–35</td>
<td>ovoid-conical, compressed dorsoventrally</td>
<td>absent</td>
<td>triangular, at base of epicone</td>
<td>parietal</td>
<td>7–8, 2, 6–7 (antapical plate distinct)</td>
<td>medium length–long</td>
<td>unknown</td>
<td>7</td>
</tr>
</tbody>
</table>

1, Lindberg *et al.* (2005); 2, Stosch (1973); 3, Wókoszyńska (1917); 4, Thompson (1951); 5, Christen (1958); 6, Moestrup *et al.* (2006); 7, Shyam & Sarma (1976).
adequately prepared material uncertainty may arise from individual variation and from the lack of comparative SEM studies on several of the species. A constant feature of cells of *T. aveirensis* observed in SEM was a line of knobs across the posterior row of cingular plates that seemed to mark the posterior edge of the cingulum. This has not been demonstrated in other species of *Tovellia* and is therefore suggested as an additional marker of the identity of *T. aveirensis*.

**Asexual reproduction**

Asexual reproduction has not been described for all species presently included in the genus *Tovellia*. However, all available reports of this process indicate that it occurs through the division of non-motile cells, usually into two–eight zoospores (Woloszyńska, 1917; Christen, 1958; Stosch, 1973; Shyam & Sarma, 1976). Asexual reproduction of *T. aveirensis* is generally similar to that described for other *Tovellia* species, in particular to *T. apiculata* (Stosch, 1973) and *T. stoschii* (Shyam & Sarma, 1976). In all cases where successful release of daughter cells from division cysts of *T. aveirensis* was observed, the number of cells was four. Observations of division cysts containing only two cells was never followed by the release of those cells, and neither did we observe division cysts with a number of cells higher than four. This suggests that, under the culture conditions adopted, division cysts of *T. aveirensis* produce mainly four offspring cells and that a second round of cell division was likely to occur in the cysts with only two cells.

**Phylogeny**

In spite of a smaller sequence divergence between *T. aveirensis* and *T. coronata* than between *T. aveirensis* and *T. sanguinea* the phylogenetic analysis grouped the latter two species in a poorly supported clade. Both in the arrangement of amphiasmal plates, including the organization of the ALP, and in the tendency for accumulating red cytoplasmic bodies, the species *T. coronata* and *T. sanguinea* (Lindberg et al., 2005; Moestrup et al., 2006) seem more closely related than either to *T. aveirensis*. One structural aspect in which the new species approaches *T. sanguinea* is the radial arrangement of chloroplast lobes in the epicone, converging to a central pyrenoid complex, which is reportedly absent in *T. coronata* (Lindberg et al., 2005; Moestrup et al., 2006). However, the limited number of species for which LSU rDNA sequences are available renders any conclusions about the close affinities between species of *Tovellia* premature.

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**References**


Tovellia aveirensis, sp. nov.


