The morphology of *Chrysochromulina rotalis* sp. nov. (Prymnesiophyceae, Haptophyta), isolated from the Skagerrak

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**SARSIA**


*Chrysochromulina rotalis* sp. nov. was isolated from surface water in the Skagerrak (58°11’N, 09°06’E) using the serial dilution culture method. The cells are saddle shaped with the appendages inserted subapically on the ventral side of the cell. The coiling haptonema measures 2-4 times the length of the flagella when extended. The cells contain two golden brown chloroplasts. The periplast is covered by two types of scale with dimorphic scale faces. The scales constituting the outer layer bear a short spine supported by four struts, the inner layer scales lack protrusions.

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**INTRODUCTION**

The haptophyte genus *Chrysochromulina* Lackey has a worldwide distribution and more than 50 species have been described, 38 of them have been reported from Scandinavian waters (e.g. Throndsen 1969; Leadbeater 1972a, 1972b; Espeland & Throndsen 1986; Kuylenstierna & Karlson 1997; Eikrem & al. 1998). In addition a number of undescribed forms or species has been observed in the area (e.g. Eikrem & al. 1998; Jensen 1998). Representatives of the genus are widespread and the number of species worldwide may exceed 100 (Thomsen & al. 1994). Representatives of the genus are widespread and the number of species worldwide may exceed 100 (Thomsen & al. 1994). The first species of *Chrysochromulina, C. parva* Lackey was described from freshwater (Lackey 1939) and further species were not added until the 1950s when Parke and Manton began to describe marine species of *Chrysochromulina* (e.g. Parke & al. 1955, 1956, 1958, 1959). They introduced the use of electron microscopy in haptophyte research and ultrastructural details of the scales covering the cell body became the main taxonomic criterion for species identification in *Chrysochromulina*. However, they stressed the importance of light microscopic observations and in fact some *Chrysochromulina* species may be identified in the light microscope (e.g. *C. parkeae* Green & Leadbeater and *C. spinifera* (Fournier) Pienaar & Norris). Since then a number of additional authors have contributed to the now long list of formally described *Chrysochromulina* species (e.g. Green & Leadbeater 1972; Hällfors & Niemi 1974; Estep & al. 1984; Hällfors & Thomsen 1985; Moestrup & Thomsen 1986; Kawachi & Inouye 1993).

The cells of the *Chrysochromulina* species have two golden brown chloroplasts, two flagella, a haptonema and a scale covered periplast. The cells may vary in size from c. 4 µm (e.g. *C. apheles* Moestrup & Thomsen and *C. minor* Parke & Manton) to c. 25-30 µm (e.g. *C. birgerii* Hällfors & Niemi and *C. parkeae*). Their shape is quite variable and they may be spherical, oblong or saddle shaped. In some species the haptonema may be very long (e.g. *C. campanulifera* Manton & Leadbeater and *C. cymbium* Leadbeater & Manton) and in most species it has the ability to coil. In a few species (e.g. *C. spinifera* and *C. parkeae*) it is short (c. 2-5 µm) and rigid. The appendages may be inserted apically or subapically and in the saddle shaped species they are inserted ventrally.

The saddle shaped species of the genus *Chrysochromulina* all possess long coiling haptonemata. The group has been reviewed recently (Eikrem & Moestrup 1998) and at present 13 described species are known (Eikrem & al. 1998), *C. rotalis* sp. nov. included. The present species is one of a number of previously undescribed species encountered in Skagerrak in 1990, as part of a project under the Norwegian Research Programme on Harmful Algae.
MATERIAL AND METHODS

Chrysochromulina rotalis was isolated from a serial dilution culture (Throonsen 1978) inoculated with surface water collected in the Skagerrak (58°11’N, 09°06’E; off Hirtshals, Denmark) on 30 June 1990.

The culture was grown in IMR 1/2 medium (Eppley et al. 1967) enriched with 10 nM selenite at about 15 °C under white fluorescent light with a quantum flux of about 100 µmol photons m⁻² s⁻¹ and 16:8 h L:D cycle.

The cells were studied live under a Nikon Microphot FX fitted with phase contrast and differential interference contrast optics and electronic flash. Stained whole-mounts were prepared according to Moestrup (1984) and some of the preparations were shadowed with gold-palladium in an Edwards Speedivac 12 E6 coating unit.

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The cells were left overnight in 2 % aqueous uranyl acetate. Thereafter the cells were rinsed in distilled water and dehydrated in an ethanol series starting at 30 % and gradually rising to 96 %. The dehydration was concluded with 4 × 10 min in 100 % ethanol and 2 × 10 min in propylene oxide. The pellets were left over night in a 1:1 mixture of propylene oxide and Epon (Burke & Geiselman 1971) embedding resin. Finally the cells were given 3x1h in undiluted Epon before they were polymerized at 50 °C for 12 h. The sectioning was carried out on a Sorvall Ultra Microtome MT 5000.

Sections and whole-mounts were viewed in a Jeol 1200ex and 100cx at the EM laboratories for Biosciences, Department of Biology, University of Oslo.

RESULTS

Diagnosis: Cells saddle shaped measuring 4-6 µm with two chloroplasts. Two flagella, usually equal 8-18 µm. Haptonema 22-80 µm long, coiling. Periplast covered with dimorphic scales, in two layers; outer layer scales slightly elongate, 450-570 × 350-470 nm, with spine supported by four decurrent struts, spine approximately equaling the scale radius; inner layer scales slightly elongate, 400-550 × 320-460 nm, without protrusions. Proximal scale faces with concentric fibrils and distal faces with radiating ribs.


Holotype: Fig. 8; (EM graph from a section of Embedding no. 15, block P16, at Dept. Biology, University of Oslo, Norway).

Type locality: Skagerrak; 58°11’N, 09°06’E; off Hirtshals, Denmark

Etymology: rotalis = having wheels (referring to the scales)

Observations

The cells of C. rotalis are typically saddle shaped with two golden brown parietal chloroplasts. Each chloroplast has a pyrenoid (not illustrated) which can barely be seen in the light microscope. The outline of the cell is apple shaped in ventral or dorsal views (Figs 1, 3 & 4), the length and width being almost equal, measuring 4-6 µm. In lateral view (Fig. 2) the outline is bean shaped, the dorsiventral axis being approximately half of the cell length. As can be seen in Fig. 2, the cell axis is kept at an angle of approximately 30° to the plane of the haptonema and flagella. The two flagella are usually equal to subequal in length, measuring 8-18 µm. The haptonema is long, normally 22-80 µm when extended (Figs 1-3), and easily visible also when coiled (Fig. 4 arrowhead). During normal swimming the flagella beat homodynamically along the sides of the cell, pushing the cell with the haptonema pointing forward (Figs 2 & 3), or they are reversed while the haptonema is dragged behind the cell (Fig. 1). This is also the case when the haptonema attaches temporarily to the cover slip surface.

The periplast is covered with two types of scales; plate scales and spine scales (Figs 5-8). The distal faces of both types have concentric fibrils which may be revealed in negative stained (Fig. 7) or shadow cast preparations (Fig. 6). The proximal faces have radiating ribs most easily seen in shadow cast preparations (Figs 5 & 6), but also evident in sections; in Fig. 9 showing a glancing section of an embedded scale both the concentric pattern of the proximal side and the radial pattern of the distal side can be seen. The scales of the outer layer bear short spines approximately equal to or shorter than the radius of the scales (Figs 6 & 8). The spines are supported by four decurrent struts.
DISCUSSION

Disregarding the spine, the scales of *Chrysochromulina rotalis* resemble those of the saddle shaped *C. simplex* (Estep & al.) Birkhead & Pienaar, as we know the latter from Scandinavian waters, i.e., with scales of one size only (figs 11-13 in Birkhead & Pienaar 1995; fig. 35 in Eikrem & al. 1998). The type described by Estep (Estep & al. 1984) as well as the emended diagnosis by Pienaar and Birkhead include larger scales with two central perforations in addition to the small scales. Cells with this kind of scale (Estep & al. 1984 fig. 15) have only been observed once in Norwegian waters (fig. 49a in Eikrem & al. 1998). This organism may represent another
morphotype, perhaps another species. The question of species delimitation is further complicated in the Prymnesiophyceae by the different morphology of various stages in the life cycle of a species. This has been demonstrated for coccolithophorids e.g., species of *Hymenomonas* (Rayns 1962), *Coccolithus* (Parke & Adams 1960) and *Emiliania* (Braarud 1963), and has also been shown for *Chrysochromulina polylepis* Manton & Parke (Edvardsen & Paasche 1992).

The saddle shaped species share a number of ultrastructural features (Eikrem & Moestrup 1998) and also according to genetical information they cluster in one group (e.g. Simon & al. 1997; Medlin & al. 1997, including *C. rotalis* as P 16). The ornamentation of the scales covering the periplast is the character used in species identification, but the variation in scale ornamentation between the saddle shaped species is considerable (e.g., Eikrem & Moestrup 1998) and may not be used as a unifying character of the group. Similarity in scale ornamentation may indicate a close relationship between species, and in the case of *C. cymbium* and *C. strobilus* Parke & Manton which only differs in minor details of scale structure (Leadbeater & Manton 1969 a,b), this has been confirmed by genetic analysis (Medlin pers. commn). On the other hand, large differences in scale morphology do not necessarily indicate a remote relationship between species as exemplified by the marked differences in scale morphology between the two alternate stages of *C. polylepis*. This emphasizes the importance of using several criteria when characterizing a species, and future investigations should preferably be based on unialgal/clonal cultures which will provide material for fine structural and genetic analysis, as well as for experimental physiology and aut-ecology.

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