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The Life History and Cell Cycle of *Kryptoperidinium foliaceum*, A Dinoflagellate with Two Eukaryotic Nuclei

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*Kryptoperidinium foliaceum* is a binucleate dinoflagellate that contains an endosymbiont nucleus of diatom origin. However, it is unknown whether the binucleate condition is permanent or not and how the diatom nucleus behaves during the life history processes. In this sense, it is also unknown if there is a sexual cycle or a resting stage during the life history of this species, two key aspects necessary to understand the life history strategy of this dinoflagellate. To answer these questions, life history and cell cycle studies were performed with the following results: (i) *Kryptoperidinium foliaceum* has a sexual cycle and in the dinoflagellate strains studied, the binucleate condition is permanent. Sexuality in the host was confirmed by the presence of fusing gamete pairs and planozygotes in clonal cultures (revealing homothallism), but signs of meiosis in the endosymbiont were not observed. The endosymbiont nucleus likely fuses first, because fusing gamete pairs were found to have two dinoflagellate nuclei but only one endosymbiont nucleus. After complete gamete fusion, the planozygotes had apparently normal endosymbiont and dinoflagellate nuclei. (ii) Asexual division studies using flow cytometry showed that the S phase in the endosymbiont (diatom) nucleus starts 6—8 h later than in the host nucleus, but there was no evidence of mitosis in the former. (iii) Sexual and asexual cysts were formed in culture. Neither cysts from natural samples nor those formed in culture exhibited a dormancy period before germination.

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**Key words:** *Kryptoperidinium foliaceum*; binucleate; cell cycle; cysts; dinoflagellate; endosymbiont; life history; sexual reproduction.

Introduction

Eukaryotic mitochondria and plastids are unique in that they evolved from free-living prokaryotes through endosymbiosis (Bhattacharya et al. 2007). The endosymbiosis hypothesis (Margulis 1970; Martin and Kowallik 1999), originally formulated to explain the origin of mitochondria and plastids, states that plastids most likely originated from a single primary endosymbiotic event between a eukaryotic host and a cyanobacterium, thereby giving rise to plants, green and red algae, and glaucophytes. On several occasions, these
primary algae were themselves taken up by another eukaryote and permanently integrated in what is referred to as secondary endosymbioses. One algal group that contains a secondary plastid thought to have been derived from the red algal plastid lineage is the dinoflagellates. Their secondary plastid, called the peridinin plastid, is distinguished by the pigment peridinin and by three bounding membranes rather than the four common to most secondary plastids (see Falkowsky et al. 2004; Yoon et al. 2005 for a review). However, other hypotheses suggest that the peridinin plastid evolved from a heterokont alga through a tertiary endosymbiosis (Bodyl and Moszczyński 2006). Dinoflagellates are unique among eukaryotic algae in that they have taken the process of endosymbiosis one step further, since in several independent lineages the peridinin plastid has been replaced either with a successive secondary plastid or with a plastid from another secondary alga, resulting in tertiary plastids. Examples are the dinoflagellate groups of haptophyte (Karenia, Karlodinium, Takayama) (Yoon et al. 2002), green algal (Lepidodinium) (Watanabe et al. 1990), and diatom origin (Kryptoperidinium, Durinskaia, Galeidinium, and Dinothrix as well as some species presently included in Peridinium [P. quinquecorne] and Gymnodinium [G. quadriloba-tum]). The diatom endosymbiont is related to either centric (P. quinquecorne) (Horiguchi and Takano 2006) or pennate (Kryptoperidinium) (Chesnick et al. 1996, 1997) groups. The tertiary endosymbiosis is more or less stable, depending on the dinoflagellate group, although knowledge of the symbiont—host system is limited, mainly restricted to genera such as Kryptoperidinium. An endosymbiosis is considered to be stable if there is genetic integration of the endosymbiont and host, meaning transfer of essential endosymbiont genes to the host nucleus and protein re-targeting (Bhattacharya et al. 2007). For example, Dinophysis has long been considered to have a cryptophyte endosymbiont chloroplast, but the latter was very recently found to have been acquired through kleptoplastidy (Koike et al. 2005; Minnhagen and Janson 2006; Takishita et al. 2002).

The degree to which symbiosis has evolved differs depending on the particular genus and on the organelle. In some genera, such as Karenia, Alexandrium, and Karlodinium, chloroplast genes have been permanently transferred to the host nucleus (Hackett et al. 2004; Ishida and Green 2002; Patron et al. 2006). In Durinskaia and Kryptoperidinium, mitochondrial genes of both host and endosymbiont are expressed (Imanian and Keeling 2007), but there is also a relict endosymbiont nucleus that may be in the process of being eliminated (Kempton et al. 2002). This suggests that Kryptoperidinium foliaceum is in an intermediate stage of endosymbiont reduction (Imanian et al. 2007).

Although chloroplast symbiosis is clearly stable in K. foliaceum-like dinoflagellates, there are conflicting data about the nature of the nuclear symbiotic relationship. While some studies point to an obligate association of the endosymbiont nucleus with its host, others suggest that the association is transient (Kempton et al. 2002; Wolny et al. 2004). However, most of the available data on endosymbiosis are derived from genetic studies, whereas the behavior of the endosymbiont—host system during the dinoflagellate life cycle remains poorly understood. These aspects were the focus of the present study.

Little is known about the life cycles of dinoflagellates with respect to tertiary endosymbiosis. Chesnick and Cox (1987, 1989) described gamete fusion and zygote formation in D. baltica, the only species of the group for which a sexual life cycle has been, at least partially, elucidated: during zygote formation, the dinokaryotic nuclei fuse first followed by plasmogamy of the symbionts; the endosymbiont nuclei fuse later, after migrating to a central position. It was suggested that these nuclei are products of meiosis, but since the isolated zygotes either died or ceased development under culture conditions, the sexual route could not be completely confirmed, nor has a cyst stage been described. The existence of fusing gamete pairs was also reported for K. foliaceum (named as Glenodinium foliaceum in Blanchard-Babilot 1972), although neither progress of the fusion process nor a ploeyzogote stage was observed. Therefore, two important questions related to the life cycle of the two nuclei-bearing species remain unanswered:

(i) Is there mitosis in the endosymbiont nucleus? Dinoflagellates are haploid eukaryotes with permanently condensed chromosomes (see Rizzo 1991 for review) whereas diatoms are diploid and chromatin condensation should precede mitosis. However, this latter process has never been seen in the group of two-nuclei-bearing dinoflagellates and only a small condensation phase was depicted for D. baltica (Chesnick and Cox 1987, 1989); therefore, amitosis (division of the nucleus by means of constriction without the chromosome...
condensation typical of mitosis) has been proposed for both *D. baltica* (Tippit and Pickett-Heaps 1976) and *K. foliaceum* (Blanchard-Babillot 1972). The DNA content of the endosymbiont nucleus of *K. foliaceum* (35 pg) is unusually large compared to the nuclei of its possible free-living chromophyte relatives. A previous study showed that DNA content is relatively constant within a given strain but not within the species (Kite et al. 1988). To study the stability and a possible cell cycle of the nuclear endosymbiont (DNA synthesis), the host and endosymbiont were analyzed using flow cytometry and observed by light and electron microscopy.

(ii) Is there efficient sexuality (i.e., a viable planozygote) during the life cycle of *K. foliaceum*? If so, is there sexuality in both the endosymbiont and the dinoflagellate? Since diatoms are diplonts, chromatin condensation and chromosome reduction should accompany meiosis in the endosymbiont nucleus. To verify this process, the nuclei of culture-formed planozygotes were stained and then observed by fluorescence microscopy.

## Results

### Morphology

#### Vegetative Motile Stage

The study of tabulation in *K. foliaceum* is difficult such that most of the descriptions found in the literature are not very accurate and perhaps represent more than one species. To avoid possible misinterpretations in identifying the organism, the morphology of the strains used in this study is as follows: The plate formula of our strains is $3', 2a, 7'', 4c, ?s, 5''', 2'''$ (Fig. 1a, b). The cells were extremely flattened dorsoventrally so that the plates were observed on their ventral or dorsal sides. The sutures of some plates were on the edge but other plates were folded, with a ventral side and a dorsal side. Plates having two sides were: apicals $2'$ and $3'$, precingulars $2''$ with only a small dorsal side and $6''$, postcingulars $2'''$ and $5'''$, the last one with a small dorsal side, and the two antapicals. The shape of three plates was very peculiar. Sa was quite large with an indentation in the epitheca and thus easily confused with a precingular plate. Plate $7''$ was L-shaped, very narrow, much thicker than the other plates of the theca, and formed the right edge of the epitheca (Fig. 1a). The first apical plate $1'$ was very large and kidney-shaped, with the concavity formed by Sa. Our strains are characterized by a conspicuous red stigma centered in the area of their flagellar pores.

#### Non-Motile Stage (Cysts)

The non-motile benthic stage (cyst) (18—35 μm long, 14—35 μm wide) is round, flattened, and has one or more reddish spot(s) (Figs 7a, 8d and 9a).

### Growth Rate in Culture

*K. foliaceum* had the highest division rates at the two highest temperatures tested (Fig. 2): at 19°C, 0.24±0.02 divisions day$^{-1}$, and at 23°C, 0.23±0.01 divisions day$^{-1}$. The division rate was significantly lower (0.18±0.01 divisions day$^{-1}$, ANOVA $p<0.05$) at 15°C.
**K. foliaceum** DNA Content and Cell Cycle by Flow Cytometry

The haploid DNA content of synchronized *K. foliaceum* cultures was comparable to the DNA content of other dinoflagellates, particularly *Alexandrium minutum* (Fig. 3a), and was estimated to be ~30 pg per cell, based on a relative comparison with the DNA content of that species. However, depending on the sampling time, two haploid DNA contents were found, one of which corresponded to the *K. foliaceum* endosymbiont (Fig. 3b—d). Also, cell-cycle analyses showed two S phases and two peaks of G2M cells (Fig. 3e), suggesting that the endosymbiont nucleus enters S phase 6—8 h later than the host nucleus.

**K. foliaceum** Life Cycle

The dominant life-cycle stage of *K. foliaceum* is a motile form with one longitudinal flagellum (Fig. 4a, b) and a typical dinoflagellate nucleus (dinokaryon) that is almost spherical and with permanently condensed chromosomes. There is also a thin and elongated endosymbiotic nucleus, in which the chromosomes are not distinguishable and sometimes this nucleus appear to be double branched (Fig. 4c, d). At some stages of development, there seems to be a small area connecting the two nuclei, as shown in Figure 4c.

Large differences in cell size, mainly in sexually induced cultures, were recorded (Fig. 4e). Small cells, some only 13 μm in length, were stained to confirm that they contained dinokaryon and endosymbiont nuclei (Fig. 4f). The relative percentages of cells in each size fraction and the changes of the population size over time were determined by observing the cells in a Coulter counter. Table 1 shows the average increase in cell size over time. At the beginning of the exponential phase of growth, two size groups were identified but the size distribution became unimodal in later samplings, although with a larger standard deviation and a shift to larger cells (Table 1). The sizes of the cells during the different life-cycle stages were followed using an inverted light microscope (Table 2).

**Asexual Cycle**

Cell division takes place in both the motile and non-motile states (Fig. 5). In the motile state, division results in the formation of two motile cells. As shown in Figure 5a and b, division occurred simultaneously and the dinokaryon and the endosymbiont nuclei acquired a similar bi-lobed shape (Fig. 5b). Non-motile cysts were abundant in clonal cultures and also underwent division (Fig. 5c—e). The multiple rounds of division in some cases yielded up to 6 cells, which were consecutively released in <24—48 h (Fig. 5f).

**Sexual Cycle**

Sexuality (fusing gamete pairs and planozygotes) was observed in clonal strains (reflecting homothallism). Fusing gamete pairs (Fig. 6a, b) were commonly detected in exponentially growing cultures. The gametes attached through the sulcal region at different angles of their respective cingula, ranging from a perpendicular (Fig. 6a) to an almost parallel orientation (Fig. 6b), and were of very different sizes (Table 2). In the fusing gamete pairs studied, the endosymbiont nuclei were in the process of fusion (Fig. 6c) or had already fused (Fig. 6d), whereas the dinoflagellate nuclei remained separated (Fig. 6c, d). Planozygotes were recognizable by the presence of two longitudinal flagella (Fig. 6e, f) and by their large size (Table 2). The endosymbiont nucleus of the planozygotes was spherical in shape, whereas the dinoflagellate nucleus was more elongated and larger than in the vegetative stages (Fig. 6g, h). The cysts were irregularly shaped, with a well-defined double wall and a large reddish spot (Fig. 7a). Some cysts had one dinokaryon and one endosymbiont nucleus (Fig. 7b), and liberated one cell, with one or two longitudinal flagella (Fig. 7c). Therefore, this cyst morphology matched both sexual and asexual cysts. Other cysts contained multiple cells (Fig. 7d) with up to four dinokaryon and four endosymbiont nuclei (Fig. 7e, f).

The sexual stages were also identified by scanning electron microscopy, which showed fusing gametes attached at different angles through the sulcal region, although their individual morphology was indistinguishable from that of the vegetative stage (hologamy) (Fig. 8a). Planozygotes were identified by the presence of two longitudinal flagella (Fig. 8b, c). The morphology of the cysts was highly variable. Some exhibited the morphological characteristics of vegetative stages, i.e., sulcal and cingular regions, whereas others were flattened and lacked any vegetative attributes (Fig. 8d).

**Field Study**

Sediment samples contained 3.3 ± 0.6 cysts of *K. foliaceum* per ml of wet sediment (Fig. 9a, b).
Figure 3. (a) Flow cytometry chart comparing the haploid DNA contents of *K. foliaceum* (thin line) and *Alexandrium minutum* (thick line). (b—d) Flow histograms of samples at different phases of the cell cycle. Dark and white areas are shown when the adjustment of the MODFIT program is better considering the presence of two haplo-diploid cycles (dark for the dinoflagellate nucleus and white for the endosymbiont nucleus). (e) Percentages of cells in S (squares) and G2/M (triangles) phases during a diel cell cycle. Data points are averages and vertical bars are standard errors/SD of the mean. The black bar represents the dark period of the diel cycle. The running mean indicates the expected values when the average value of the variable in a specific number of previous periods is considered.
Whereas all of the cysts seemed to be unicellular with respect to their external appearance (Fig. 9a, b), nuclear staining consistently revealed two dinokaryon and two endosymbiont nuclei (Fig. 9c). All germlings studied presented one longitudinal flagellum (Fig. 9d) and one dinokaryon and one endosymbiont nucleus (Fig. 9e), although in some germlings the shape of the endosymbiont nucleus was similar to that observed in planozygotes from cultures (Fig. 9f). Cysts needed 1—2 days to germinate, with 100% germination frequency and 100% surviving germlings.

**Figure 4.** Vegetative stages of *K. foliaceum* (strain Baiona B9). External appearance (a, b) and nuclear morphology (c, d) of vegetative stages, D is showing the dinokaryon and E the endosymbiont nucleus; (e) comparison of sizes among culture cells; (f) nuclear morphology of the smaller cell shown in (e). Scale bars: 10 μm.
Discussion

Our results can be summarized in three main points; (i) in the studied strains the binucleate condition is permanent, (ii) evidence of neither mitosis nor meiosis (i.e., chromosomes condensation) was found in the endosymbiont nucleus, (iii) flow cytometry analyses suggest the existence of an S-phase in the endosymbiont nucleus.

Flow cytometric analyses indicated that the dinoflagellate nucleus contains around 30 pg DNA/cell, whereas relatively minor amounts were measured in the endosymbiont nucleus. These estimates were based on a comparison with the DNA content of the dinoflagellate species *Alexandrium minutum*, whose DNA content was previously shown to be almost identical to that of our *K. foliaceum* strain (Figueroa et al. 2007). Our results were obtained by a different methodology than Kite et al. (1988), which may explain why these authors reported higher DNA content values for *K. foliaceum* than determined in this study.

Table 1. Volumes of cells, as determined using a Coulter counter, at different times of the growth curve.

<table>
<thead>
<tr>
<th>Growth (cell ml⁻¹)</th>
<th>Volume (µm³ ml⁻¹ µm⁻¹)</th>
<th>Number of cells (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Group 1 average SD</td>
<td>Group 2 average SD</td>
</tr>
<tr>
<td>1429</td>
<td>208</td>
<td>14 2.0</td>
</tr>
<tr>
<td>2146</td>
<td>197</td>
<td>14 2.0</td>
</tr>
<tr>
<td>5044</td>
<td>1527</td>
<td>14 2.1</td>
</tr>
<tr>
<td>8700</td>
<td>890</td>
<td>16 4.5</td>
</tr>
<tr>
<td>12,945</td>
<td>1727</td>
<td>18 5.5</td>
</tr>
<tr>
<td>16,583</td>
<td>527</td>
<td>19 5.9</td>
</tr>
</tbody>
</table>

Two modes of size (groups 1 and 2) were observed at the beginning of the exponential phase.

Table 2. Sizes of *Kryptoperidinium foliaceum* life-cycle stages (n ≥ 10) as determined by optical microscopy.

<table>
<thead>
<tr>
<th>Life-cycle stage</th>
<th>Length range (µm)</th>
<th>Width range (µm)</th>
<th>Length (µm) (average ± SD)</th>
<th>Width (µm) (average ± SD)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Vegetative cells</td>
<td>13—38</td>
<td>10—30</td>
<td>26 ± 8</td>
<td>22 ± 7</td>
</tr>
<tr>
<td>Gametes</td>
<td>14—23</td>
<td>12—23</td>
<td>17 ± 4</td>
<td>16 ± 4</td>
</tr>
<tr>
<td>Planozygotes</td>
<td>34—45</td>
<td>31—38</td>
<td>40 ± 4</td>
<td>33 ± 4</td>
</tr>
<tr>
<td>Cysts</td>
<td>18—35</td>
<td>13—33</td>
<td>29 ± 5</td>
<td>26 ± 4</td>
</tr>
<tr>
<td>Germlings</td>
<td>31—33</td>
<td>26—33</td>
<td>32 ± 2</td>
<td>28 ± 4</td>
</tr>
</tbody>
</table>

The size of the endosymbiont nucleus was observed to be variable depending on culture growth conditions, since very small cells with noticeable smaller endosymbiont nucleus (Fig. 4) mainly occurred in our study after sexual induction in nutrient-deficient medium or in early phases of the growth curve, during which two size groups were identified using a Coulter counter. This method allowed us to corroborate the large differences in sizes observed by light microscopy.

A variable endosymbiont DNA content among different strains was reported by Kempton et al. (2002). These authors found that *K. foliaceum* may actually comprise two different but similar species, since two morphologically and genetically distinct groups were identified. Although we were not able to confirm the original description of Paulsen, the peculiar 7" was not included in previous descriptions of the species (Biecheler 1952; Lebour 1925; Lindemann 1924) and it was not until recently that plates of different strains were photographically described (Kempton et al. 2002). Our strains are similar to strains CCAP 1116/3 and CSIRO 291, based on figure 1C and D in Kempton et al. (2002), but different from UTEX LB 1688, CCMP 1326 and their South Carolina isolates, as shown in figure 1A and B, and E, in that report. Although the strains of this group were all binucleated, strains with or without an endosymbiont nucleus were described in the other group, formed by UTEX LB 1688 and CCMP 1326 and the South Carolina and Florida isolates (Kempton et al. 2002). The presence of an eyespot seems to be a variable feature of...
Kryptoperidinium, as it was not observed in two (UTEX LB 1688 and CCMP 1326) of the four strains studied by Kempton et al. (2002). This variability is supported not only by the fact that not all the strains have an eyespot, but also because strain UTEX LB 1688, used by Dodge and Crawford (1969) to describe the eyespot, was later claimed by Kempton et al. (2002) to lack one. Although we cannot rule out the possibility of a cryptic species within the genus, these findings, when considered together with our results on the variability in the size of the endosymbiont nucleus,

**Figure 5.** Dividing stages of *K. foliaceum* (strain Baiona A3). External morphology (a) and nuclear development (b) of a mobile stage in division, showing a completed mitosis. (c—f) Non-mobile dividing stages. Outer morphology (c) and nuclei (d) of a two-cell dividing cyst. Six-cell dividing cyst (e) and progressive germination (f). Scale bars: 10 μm.
may suggest that gene transfer from the diatom to the dinoflagellate has already been accomplished and that the endosymbiont nucleus is no longer functional and in the process of being evolutionarily discarded (Bhattacharya et al. 2007).

Other observations support the conclusion that mitosis and meiosis are lacking in the endosymbiont nucleus. As in D. baltica (Tippit and Pickett-Heaps 1976) and K. foliaceum (Blanchard-Babillot 1972), no chromatin condensation or mitotic spindle was observed in the endosymbiont nucleus, suggesting that it does not undergo mitosis, although there may be a phase of DNA synthesis (S phase) albeit delayed by 6—8 h relative to the dinoflagellate nucleus. A sexual process in the endosymbiont would imply endosymbiont meiosis simultaneous with dinoflagellate mitosis yielding haploid gametes. This possibility cannot be excluded because very small cells contained very small endosymbiont nuclei; nonetheless, there was no evidence of meiosis, since neither chromatin condensation nor chromosome segregation occurred. The complete sequence of endosymbiont fusion was described by Chesnick and Cox in their study of D. baltica (Chesnick and Cox 1989). The lack of meiosis in the endosymbiont may be compatible with the typical sexual behavior of diatoms, since sexuality in these organisms is much less frequent than in dinoflagellates and usually only occurs about once a year, when there has been a critical reduction in size such that it is necessary to restore the normal cellular dimensions (see, for example, the revision of Lewis (1984)).

Among the organisms with an endosymbiont-like origin, those most related to K. foliaceum are D. baltica and G. rugatum (Horiguchi and Takano 2006). However, the morphology and life cycle (a dominant non-motile phase and a brief motile phase) of G. rugatum are completely different from those of other such species (Tamura et al. 2005). It is still unknown whether there is sexuality in the life cycle of these two-nuclei-bearing organisms. As mentioned above, Chesnick and Cox (1987) reported signs of sexuality in D. baltica, such as gamete fusion and zygote formation; however, as the viability of the zygotes was not determined, the existence of a sexual cycle could not be established. We observed planozygote encystment in K. foliaceum, thus confirming the existence of an effective sexual route in this strain.

Sexual reproduction is widespread among eukaryotes (Bell 1982), although it entails several

Figure 6. Sexual cycle of K. foliaceum (strain Baiona B9). Isogamous (a) and anisogamous (b) fusing gamete pairs. (c, d) Nuclear morphologies of fusing gamete pairs. (e, f) Planozygote external morphologies, showing the two longitudinal flagella (arrows). (g, h) Nuclear morphologies of planozygotes. Scale bar: 10 μm.
costs that are avoided in asexual reproduction, for example, the need to find a mate. Conversely, in environments where perturbations are common, sexual recombination confers several benefits as it provides new genotypes that allow the population to adapt to the changing conditions (Hamilton 1980; Lively 1989) and to restore damaged DNA through recombinational repair (Bernstein 1983). This line of reasoning is consistent with the role of a sexual route in the life cycle of *K. foliaceum*, an organism that resides in shallow estuaries, that are highly and rapidly variable systems due to the strong effects of tides and river flow on their hydrological structure (Trigueros and Orive 2000). These habitats are common for other phylogenetically related species, such as *G. quadrilobatum* (Horiguchi and Pienaar 1994), *D. capensis* (Pienaar et al. 2007), and *P. quinquecorne* (unpublished data), and may explain homothallism as the mating-type system of *K. foliaceum*. In homothallism, sexual zygotes can be produced by self-fertilization, i.e., by the fusion of genetically identical gametes. This ability makes sex energetically cheaper and facilitates more frequent sexual events within a population, because individuals do not lose energy in finding a partner with a compatible mating type. However, since homothallism implies the recombination of identical genotypes, the ability of sexual recombination to produce new genotypes is lost. Nonetheless, evidence for the advantages of sexual recombination comes from recent work showing that the fitness effects of sex in homothallic systems are not the result of recombination but rather are due to the fact that sex promotes the accumulation of fewer deleterious mutations (Bruggeman et al. 2003) and breaks up genetic association among loci (linkage disequilibria) (Otto 2003). In addition, if sexuality enables a resting stage, it provides other benefits, such as survival during unfavorable conditions, species dispersal, reproduction, and regulated seasonal succession (Anderson and Rengefors 2006; Anderson and Wall 1978; Anderson et al. 1984; Rengefors and Anderson 1998; 

**Figure 7.** Encystment and germination of a culture cyst (strain Baiona A3). External morphology (a) and nuclei (b) of a culture cyst; (c) germling showing the two longitudinal flagella. Outer morphology (d) of a cyst containing four endosymbiont and four dinoflagellate nuclei (e, f). Scale bars: 10 μm.
Although not previously reported for *K. foliaceum*, we were able to document the formation of cysts in culture and in natural samples. Two factors are important regarding sexual cyst formation in a given species; the sexual or asexual nature of the cysts and the dormancy period. *K. foliaceum* produced sexual and asexual cysts with identical external morphologies. It is thought that maintenance of the haplo-diplontic life cycle is advantageous when the environment offers two different ecological niches, each of which can be occupied by one of the two stages (Stebbins and Hill 1980; Steidinger and García 2006). This becomes irrelevant when there is no relation between morphology and ploidy and a given morphology (the cyst) can occur in different ploidy levels (Lubchenco and Cubit 1980). However, this system can be maintained under conditions that favor short-lived individuals and thus promotes the selection of two short phases rather than a single long phase within a cycle (Richerd et al. 1993). Similarly, the requisite period of dormancy is 1–2 days, which is much shorter than that recorded for several other dinoflagellates. For example, in *Alexandrium* species, the dormancy period ranges from 1 to 12 months (Anderson 1980; Dale et al. 1978; Figueroa et al. 2006; Montresor and Marino 1996; Pérez et al. 1998; Walker and Steidinger 1979). By contrast, the cysts of *Gymnodinium catenatum* Graham germinate within 2 weeks or less (Bravo and Anderson 1994), and *Peridinium gatunense* Nygaard cysts excyst within 12 h after formation (Pfiester 1977). Similar to what we observed, these short time intervals do not provide a true dormancy condition. The length of the dormancy period is a factor that strongly determines the bloom pattern of a species. Long dormancy periods are related to long-term survival and seasonal successions because the organism is thus able to remain dormant in bottom sediments during temperature extremes, with seasonal germination inoculating vegetative cells into the water column only when the temperature and light conditions are suitable for growth (Anderson 1997). A short dormancy period indicates rapid cycling between benthic and planktonic stages (Hallegraeff et al. 1998), which may be crucial for survival when short-term variability has a much greater impact than the seasonal transition. This is consistent with the usual conditions of the *K. foliaceum* natural habitat. Thus, the life cycle described here serves as a clear model of adaptation to a challenging environment, such as an estuary, where the effect of diurnal or semi-diurnal tides plays a major role in controlling the main hydrological parameters and hence, the intensity, distribution, and species composition of blooms.
Methods

Culture isolation: *Kryptoperidinium foliaceum* was isolated from a recurrent bloom in the estuary of Miño River (NW Spain). The bloom occurs during the summer months along the estuarine front, which is located, depending on the tides, from the mouth to a few km up-river. A strong saline vertical gradient of 4–20 psu (practical salinity units) was measured during high tides at the internal-most stations (Fig. 10a). Vegetative cells of *K. foliaceum* were present in high concentrations, coinciding with the front (Fig. 10b), but concentrations as high as 15 × 10^4 cells l^-1 were detected during falling tides, with salinities of 19–24 psu during ebb tides.

Culture maintenance and growth: Three strains, Baiona A3, Baiona B1, and Baiona B9, were employed in this study. All three derived from single cells and therefore were clonal. These cultures were deposited in the culture collection of the Spanish Institute of Oceanography in Vigo (Contact address: Instituto Español de Oceanografía, Centro Oceanográfico de Vigo, Subida á Radio Faro, 50, E36390 Vigo, Spain, http://www.vi.ieo.es). The cultures were grown at 19 °C with a photoperiod of 12:12 h light:dark (L:D) cycle. Illumination was achieved with fluorescence tubes, which provided a photon irradiance of 90 μmol photons m^-2 s^-1. Culture stocks were maintained in Erlenmeyer flasks filled with 50 ml of L1 medium (Guillard and Hargraves 1993), without the addition of silica, prepared with Atlantic seawater and adjusted to a salinity of 10 psu by the addition of sterile Milli-Q water. Duplicate Erlenmeyer flasks were filled with 50 ml of L1 medium, inoculated with 200 cells of each parental strain ml^-1, and sampled every 2 days. Strain Baiona B9 was cultured at three different temperatures (15, 19, and 24 °C) and adapted at each temperature over a period of 1 month. Subsamples were fixed with Lugol’s solution for cell enumeration in Sedwick–Rafter chambers; at least 200 cells were counted.

Morphology of the vegetative stage: Morphology was assessed in live or formalin-fixed cultured cells. The plate pattern was identified by staining the cells with Fluorescent Brightener 28 (Sigma, St. Louis, MO, USA) following a modification of the technique reported by Fritz and Triemer (1985).

Size of vegetative stages: The cells were counted and their sizes measured with a Multisizer 3 Coulter counter (Beckman) and by using a 63 × lens mounted on an inverted microscope (Axiovert Zeiss 135, Oberkochen, Germany).

Planozygote behavior: Strain Baiona B9 was self-crossed at 19 °C in 10 ml of medium without added nitrates (L–N). Duplicate sterile polystyrene Petri dishes (35-mm diameter, Iwaki, Tokio, Japan) were inoculated with exponentially...
growing cells (3000—5000 cells ml\(^{-1}\)) to a final concentration of 700 cells ml\(^{-1}\). Thirty planozygotes were individually isolated and separately transferred to 96-well tissue culture plates (Iwaki, Tokio, Japan) filled with 0.4 ml of L1 medium at 10 psu. These cultures were placed under the same conditions as those previously described for culture maintenance. Planozygote evolution was monitored at least daily and photographically documented with a digital camera (CANON EOS-D60, Tokio, Japan). The size of these cells was determined using a 63× lens mounted on an inverted microscope (Axiovert Zeiss 135, Oberkochen, Germany) and a black and white analogical video camera (Sony AV/C-D5CE; Sony Co., Tokyo, Japan) mounted on the microscope together with a camera adapter and an Image IPplus analyzer (Media cybernetics, Berkshire, UK).

**Staining of sexual-stage nuclei:** Sexual stages in different phases of development (\(n \geq 5\)) were stained in order to visualize their nuclei. Fusing gamete pairs, planozygotes, cysts, and germlings from the self-cross of strain Baiona B9 in L—N medium were individually isolated and fixed for 10—15 min in 2% paraformaldehyde in 0.01 M PBS (phosphate-buffered saline, Sigma, St. Louis, USA), pH 7.4. The fixed cells were washed in several drops of PBS buffer and then stained with 1:100 Sybr green (Molecular Probes, Eugene Oregon, USA) in 0.01 M PBS, pH 7.4, for 30 min, washed again, and observed by fluorescence microscopy (LEICA DMLA, Wetzlar, Germany) with excitation filters of BP 450—490 nm and a long-pass emission filter (LP 515).

**Dormancy period and excystment of wild and culture cysts:** Thirty wild and 30 newly formed culture cysts were individually isolated into 96-well polystyrene plates (Iwaki, 6.4 mm diameter) and checked for excystment every 2 days. Excystment was defined as the complete emergence of the protoplast from the cyst even if the germling remained non-motile (Anderson and Wall 1978). From the cysts isolated on a given date, the percentage that had germinated was determined by dividing the number of germinated cysts by the number of non-germinated ones. The development of the germinated cells (planomeiocytes) was assessed at least daily and they were photographed as explained above.

**Diel division cycle (synchronization experiment):** To study the cell cycle in *K. foliaceum*, the clonal strain Baiona B9 was incubated in duplicate at a density of 5000—5500 cells ml\(^{-1}\) in replete L1 medium in 0.5-l Erlenmeyer bottles. At this cell density (exponential growth phase) and after the completion of a 24-h 12L:12D cycle, the incubators lights were turned off for a period of 48 h during which the cultures were synchronized by maintaining them in darkness. After the light conditions had been restored, duplicate samples for cell counting and flow cytometric analysis were taken every 2 h during 24 h.

**Study of the dinoflagellate and endosymbiont cell cycle by flow cytometry:** DNA analyses were done following the method described by Figueroa et al. (2006, 2007). Culture aliquots of 45 ml were fixed in 1% paraformaldehyde for 10 min at room temperature and washed with PBS.
(Sigma-Aldrich, USA, #P4417), pH 7. After centrifugation (4500 rpm, 20 min), chlorophyll was extracted by resuspending the pellet in 3 ml cold methanol and storing the suspension for 12 h at 4 °C. The cells were washed twice in PBS and resuspended in staining solution (PBS, 3 μg propidium iodine ml⁻¹ and 1.1 μg RNase ml⁻¹) for at least 2 h before analysis with a Becton Dickinson FACScalibur bench machine with a laser emitting at 488 nm. Samples were run at low flux (approx. 18 μl min⁻¹) and data were acquired in linear and log modes until around 10000 events had been recorded. The internal standard consisted of a solution containing 10⁶ particles ml⁻¹ of yellow-green 0.92-μm Polysciences latex beads, with 10 μl added per sample. Fluorescence emission of propidium iodide was detected at 617 nm. ModFit LT (Verity software House, Topsham, ME, USA) was used to determine the distribution of the DNA fluorescence of the population, computing the peaks, their ratios, and the coefficients of variation (CVs). To compare the trends of the different treatments with respect to cell-cycle pattern, the percentages of cells in the G1, S, and G2/M phases were adjusted to yield polynomial curves and the delimited areas were integrated over the diel cycle.

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**References**


**Bernstein H** (1983) Recombinational repair may be an important function of sexual reproduction. BioScience 33: 326—331


**Biecheler B** (1952) Reserches sur les Péridiniens. Bull Biol Fr Belg Supplément 36: 1—149


**Bruggeman J, Debets GM, Hoekstra RF** (2003) Sex slows down the accumulation of deleterious mutations in the homothallic fungus *Aspergillus nidulans*. Genetics 164: 479—485


Guillard RRL, Hargraves PE (1993) Stichocystis immobillis is a diatom, not a chrysophyte. Phycologia 32: 234—236


Lebour M (1925) The Dinoflagellates of Northern Seas. Marine Biological Association of the United Kingdom, Plymouth


Lindemann E (1924) Der Bau der Hu¨ lle bei Heterocapsa und Kryptoperidinium (Stein) n. nov. [zugleich eine vorl. Mitteilung]. Bot Arch 5: 114—120

Lively CM (1989) Adaptation by a parasitic trematode to local populations of its snail host. Evolution 43: 1663—1671


Wall D (1971) Biological problems concerning fossilizable dinoflagellates. Geosci Man 3: 1—15


