Encystment of the red tide flagellate *Chattonella antiqua* (Raphidophyceae): cyst yield in batch cultures and cyst flux in the field

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ABSTRACT: In order to clarify the encystment conditions for the red tide flagellate *Chattonella antiqua*, cyst yield in batch cultures under a variety of environmental treatments and cyst flux in natural populations were monitored. In laboratory culture experiments, attempts were made to form 'small cells' (gametes) and cysts under nutrient-replete conditions, but they could not be formed without N- or P-depletion. Once small cells were formed by nutrient depletion, encystment was affected by environmental conditions. Cyst production was highest under continuous darkness and decreased with increasing light intensity. The optimum temperature range for encystment was 21.6 to ≥ 26.6 °C, broader than that for maximum growth rate. Cyst production increased linearly with increase in motile cell concentration, indicating that the efficiency of encystment was independent of motile cell concentration. Re-addition of nutrients to N- or P-depleted cultures did not affect cyst production. In the field, cyst flux of *Chattonella* spp. together with environmental variables were monitored throughout the blooming period of *C. antiqua* in the Seto Inland Sea, Japan. Cysts were formed mainly below a depth of 15 m when nutrients were exhausted in the *C. antiqua* habitat (0 to 10 m) and the population was decreasing. Based on laboratory culture experiments and field observations, a simple model for encystment in the field was proposed. Following the development of the bloom, nutrients in the habitat of *C. antiqua* were exhausted and small cells were formed due to N- or P-depletion. Since small cells have a tendency to sink, they descended to the lower layer (> 15 m) where environmental conditions were more favorable for encystment than in the upper layer (i.e. lower irradiance, optimal temperature, and replete nutrients do not affect encystment), and cysts were formed through the fusion of small cells below 15 m. However, the possibility that small cells are formed without nutrient depletion cannot be completely ruled out, so the above model is not conclusive.

INTRODUCTION

In the last 2 decades, red tides due to *Chattonella antiqua* (Raphidophyceae) have occurred sporadically during summer in the Seto Inland Sea, Japan (Ono & Takano 1980, Nakamura et al. 1989). Such blooms kill farmed fish, especially yellowtail, and the damage reached 30 million dollars during the red tide of 1972 (Iwasaki 1979). In order to assess the roles of chemical substances [such as N- and P-nutrients (N,P-nutrients), vitamin B12, and trace metals] in inducing these red tides, we have monitored the marine environment around the Ie-shima Islands (Seto Inland Sea) each summer since 1985 and conducted bioassay and laboratory culture experiments. The results from these field surveys and experiments (Nakamura et al. 1988, 1989, Nakamura 1990) revealed that: (1) N,P-nutrients in the surface layer (0 to 5 m) were insufficient to support rapid growth of *C. antiqua*, but those below the nutrient cline were sufficient; (2) vitamin B12, essential for the growth of *C. antiqua*, did not become a growth-rate-limiting factor; (3) Fe in the surface layer (0 m) was usually insufficient to support the maximum growth rate, but it was not as severely deficient as were N and/or P; (4) Cu, whose toxicity to phytoplankton species may regulate the structure of the phytoplankton community, was always detoxified by natural organic ligand(s); and (5) the presence of a shallow nutrient cline (5 to 10 m), coupled with the ability of *C. antiqua* to migrate vertically and take up nutrients at night, played an important role in the development of the red tide.
tides. Thus, we now have some understanding of the population dynamics of this organism, but little attention has been paid to life cycle aspects of the bloom. Life cycle changes that allow populations of some red-tide flagellates to alternate between a benthic and a planktonic existence clearly play important roles in the initiation and decline of blooms (e.g. Anderson & Wall 1978, Anderson et al. 1983). In the case of Chattonella antiqua, it has been believed that this organism overwinters as a benthic form, since incubation of the bottom sediments from the Seto Inland Sea with fresh medium led to recovery of planktonic populations (T. Akizuki unpubl.). However, the benthic form of C. antiqua (hereafter termed 'cysts') was not identified until 1988 and life cycle studies were rather restricted: Imai & Itoh (1985, 1987) used an indirect, extinction dilution method (i.e. sediments were incubated under a regime of serial dilution and the liberation of planktonic cells was monitored), examined the distribution of cysts in the Seto Inland Sea and found that cold treatment (11°C) for 4 mo was necessary for cyst maturation. Recently, cysts of C. antiqua were isolated from the bottom sediments of the Seto Inland Sea (Imai & Itoh 1988), and we began a study of this organism's life cycle. We succeeded in forming cysts in laboratory cultures by transferring the N- or P-depleted cultures (mixture of 2 strains) to continuous darkness. We also monitored the encystment process (Nakamura et al. 1990): under N- or P-depleted conditions, 'small' cells (gametes) were formed. These fused to become 'triangle'-shaped cells (planozygotes), which in turn changed into cysts (hypnozygotes). DNA measurement of the cysts and interclonal crosses supported the above observations that cysts are sexual products.

In order to assess the ecological roles of cysts in the population dynamics of Chattonella antiqua, there is a clear need to understand the encystment/excystment process as a function of environmental variables (e.g. Anderson & Wall 1978, Anderson et al. 1984, Binder & Anderson 1987). In this context, we examined the numerical yields of cysts under a variety of nutritional and environmental conditions. Furthermore, throughout the period of the C. antiqua red tide outbreak around the Ie-shima Islands in the summer 1989, the flux of cysts, together with environmental variables, were monitored using a sediment-trap method to clarify when, in what layer, and how many cysts were formed. In the present paper, the laboratory and field results are described and combined to interpret the encystment process of the natural population with reference to environmental conditions.

**MATERIALS AND METHODS**

**Laboratory experiments**

**Organism and culture conditions.** Two strains of Chattonella antiqua (A-0 and B-20), isolated from the Seto Inland Sea (Nakamura et al. 1990), were used throughout the experiments. The strains were clonal but not axenic. The composition of the media used in the

<table>
<thead>
<tr>
<th>Component</th>
<th>TH medium</th>
<th>P-encystment medium</th>
<th>N-encystment medium</th>
<th>Fe-encystment medium</th>
</tr>
</thead>
<tbody>
<tr>
<td>Filtered seawater (ml)</td>
<td>950</td>
<td>950</td>
<td>950</td>
<td>950</td>
</tr>
<tr>
<td>Distilled water (ml)</td>
<td>50</td>
<td>50</td>
<td>50</td>
<td>50</td>
</tr>
<tr>
<td>NO₃⁻ (µM)</td>
<td>450</td>
<td>30</td>
<td>10</td>
<td>150</td>
</tr>
<tr>
<td>PO₄³⁻ (µM)</td>
<td>30</td>
<td>1-1.5</td>
<td>3</td>
<td>10</td>
</tr>
<tr>
<td>Si(OH)₄ (µM)</td>
<td>35</td>
<td>20</td>
<td>20</td>
<td>20</td>
</tr>
<tr>
<td>N1-metals (ml)</td>
<td>30</td>
<td>30</td>
<td>30</td>
<td>not added</td>
</tr>
<tr>
<td>Vitamin B₁₂ (ng l⁻¹)</td>
<td>200</td>
<td>20</td>
<td>20</td>
<td>20</td>
</tr>
<tr>
<td>Thiamine (µg l⁻¹)</td>
<td>100</td>
<td>100</td>
<td>100</td>
<td>100</td>
</tr>
<tr>
<td>Biotin (ng l⁻¹)</td>
<td>100</td>
<td>100</td>
<td>100</td>
<td>100</td>
</tr>
<tr>
<td>Tris (g)</td>
<td>1.0</td>
<td>1.0</td>
<td>1.0</td>
<td>not added</td>
</tr>
<tr>
<td>EDTA (µM)</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>1</td>
</tr>
<tr>
<td>pH</td>
<td>8.0</td>
<td>8.0</td>
<td>8.0</td>
<td>not adjusted</td>
</tr>
</tbody>
</table>

*Surface seawater from the Kuroshio area (ca 35 °N. S) was used

*1000 ml of N1-metals contains: Na₂EDTA. 2H₂O, 1.0 g; FeCl₃. 6H₂O, 63 mg; CoSO₄. 7H₂O, 8.9 mg; ZnSO₄. 7H₂O, 12 mg; MnCl₂. 4H₂O, 32 mg; CuSO₄. 5H₂O, 0.2 mg; Na₂MoO₄. 2H₂O, 0.2 mg

*PH of the medium was adjusted by addition of HCl

*Fe-encystment medium was sterilized by membrane filtration (Nuclepore, 0.2 um) and others were sterilized by autoclaving (121 °C, 20 min)

*Comparisons of minimum cell quota of C. antiqua for N, P, and B₁₂ indicate that N and P limit the final cell yield in N- and P-encystment medium, respectively (C. antiqua cannot utilize Tris as N source, Nakamura & Watanabe 1983b)
present study is shown in Table 1. Maintenance and inoculum cultures were grown in TH medium (Table 1; Nakamura et al. 1990). The strains (cell conc. = ca
2 × 10^4 ml⁻¹) were inoculated into experimental medium in combination (0.5 ml l⁻¹, each strain) and cultured at 25°C with 150 μE m⁻² s⁻¹ illumination under a 12 h light:12 h dark photoperiod. In most cases, encystment experiments were started when the cultures reached the growth maximum (small cell (≤ 40 μm) fraction = 10 to 20% of motile cells) by dividing the culture into subcultures (100 ml each; see below).

**General method for encystment.** Since cysts of *Chattonella antiqua* have a tendency to adhere to the surface of solid materials (Imai & Itoh 1988), cysts were formed on the surface of microscope coverslips (Nakamura et al. 1990). Four coverslips (15 mm in diameter, 1.8 cm²; made of borosilicate glass) were placed into a 200 ml Erlenmeyer flask with a 100 ml of distilled water and autoclaved. The distilled water was then discarded and replaced with 100 ml of C. antiqua culture. The culture was incubated under a given experimental condition for a minimum of 6 d, since this represents the interval after which total cyst counts do not increase (Nakamura et al. 1990). At the end of this interval, each coverslip was taken from the culture and rinsed in filtered seawater to remove motile cells, and the surface was observed using a microscope. The number of cysts adhering to the 4 coverslips was counted and converted to the cyst yield (cysts cm⁻²).

Small-cell concentrations were also monitored by microscope just before the start of the encystment experiment and after the end of incubation.

**Attempts to form cysts under N- and P-sufficient conditions.** Before examining the cyst yield as a function of environmental variables, some attempts were made to form cysts in the absence of nutrient depletion. Three experiments were conducted:

1. A mixed culture of A-0 and B-20 (1 l) was grown in P-encystment medium (1.5 μM PO₄³⁻) under normal conditions (25°C, 150 μE m⁻² s⁻¹, 12 h light:12 h dark). When the culture was in exponential phase (no small cells), 5 subcultures (100 ml × 5) were established and placed in darkness (25°C) for 7 d. The remaining culture (500 ml) was further incubated under the normal conditions until growth reached a maximum (P-depleted). Then, the culture was divided into 5 subcultures and placed in darkness (25°C) for 7 d.

2. A mixed culture of A-0 and B-20 (1.5 l) was grown in TH medium under the normal culture conditions. When the cells were growing exponentially (ca. 2.5 x 10^3 cells ml⁻¹), the culture was divided into 3 parts. These parts were placed under high-light (450 μE
m⁻² s⁻¹, 25°C, 12 h light:12 h dark), high-temperature (150 μE m⁻² s⁻¹, 28°C, 12 h light:12 h dark), and normal conditions respectively, for 2 d. Then each part was divided into 5 subcultures and placed in darkness (25°C) for 6 d.

3. Precultured *Chattonella antiqua* (mixture of A-0 and B-20; cell conc. = 3.7 x 10^5 ml⁻¹) were inoculated in medium with a low Fe content (Ni-metals = 0.38 ml⁻¹; Tris omitted; other components at levels for TH medium; see Table 1) was inoculated (ca 1 ml) into Fe-encystment medium (1 l). When the culture reached growth maximum, it was divided into 10 subcultures. Five were placed in darkness (25°C) for 7 d to form cysts, and the other 5 were used to ensure that the encystment experiment was conducted under Fe-deficient conditions: 3 received Fe-enrichment (0.2 μM, as FeCl₃), while the others did not. These were then incubated under normal conditions, and growth was monitored.

**Effects of light, temperature and motile cell concentration.** When light was an experimental variable, a mixed culture of A-0 and B-20 [2 l; grown in P-encystment medium (1 μM PO₄³⁻); at growth maximum and P-depleted] was divided into 18 subcultures, and each part was further divided into 3 parts. When temperature was an experimental variable, a mixed culture of A-0 and B-20 [2.5 l; grown in P-encystment medium (1.5 μM PO₄³⁻); at growth maximum and P-depleted] was divided into 12 subcultures, and each part was further divided into 3 parts. When cell concentration was an experimental variable, 12 subcultures with 4 different concentrations (0.9 to 4.5 x 10^5 cells ml⁻¹) were established from a mixed culture of A-0 and B-20 [1 l; grown in P-encystment medium (1.5 μM PO₄³⁻); at growth maximum and P-depleted] by diluting the culture with TH medium from which N and P were omitted. These subcultures were then incubated in darkness (25°C) for 7 d.

When temperature was an experimental variable, a mixed culture of A-0 and B-20 [2.5 l; grown in P-encystment medium (1 μM PO₄³⁻); at growth maximum and P-depleted] was divided into 24 subcultures, wrapped with aluminum foil and placed in incubators (3 flasks at each of 8 temperatures, range 17.7 to 26.6°C) for 8 d.

When cell concentration was an experimental variable, 12 subcultures with 4 different concentrations (0.9 to 4.5 x 10^5 cells ml⁻¹) were established from a mixed culture of A-0 and B-20 [1 l; grown in N-encystment medium; at growth maximum and N-depleted] by diluting the culture with TH medium from which N and P were omitted. These subcultures were then incubated in darkness (25°C) for 7 d.

**Nutrient re-addition effects.** A mixed culture of A-0 and B-20 (1 l; grown in N-encystment medium; at growth maximum and N-depleted) was divided into 9 subcultures. The remaining culture was used for isolation of small cells (see below). Nutrients were then added to the subcultures. The modes of enrichment were: control (no addition), +N (10 μM NO₃⁻), and +N and P (10 μM NO₃⁻ and 1 μM PO₄³⁻). After enrichment, the subcultures were incubated at 25°C with 150 μE
m⁻² s⁻¹ illumination for 3 h and then placed in darkness (25°C) for 7 d. Nutrient re-addition effects were also examined using P-depleted cultures [1 l; grown in
P-encystment medium (1 μM PO₄³⁻); at growth maximum and P-depleted). The modes of enrichment were: control, +P (1 μM PO₄³⁻) and +N and P (10 μM NO₃⁻ and 1 μM PO₄³⁻). Other experimental procedures were the same as those described above.

Morphological changes in the small cells formed in the N- or P-depleted cultures were monitored under nutrient-sufficient conditions. From the remainder of N- or P-depleted cultures, small cells were isolated using a Pasteur pipette and placed into the wells of a tissue culture plate (Sumitomo Co., 96 wells) with 0.25 ml well⁻¹ of N,P-reduced TH medium (10 μM NO₃⁻ and 1 μM PO₄³⁻). Isolated cells were incubated at 25°C under darkness or 150 μE m⁻² s⁻¹ illumination (12 h light:12 h dark) for 4 d. Morphological changes in the cells were monitored daily using an inverted microscope.

**Vertical distribution of small cells.** A mixed culture of A-0 and B-20 [21], grown in P-encystment medium (1.5 μM PO₄³⁻); P-depleted; small cell fraction = ca 40 % of motile cells] was transferred to 2 cylindrical glass tubes (25 cm high by 8 cm i.d., sides and bottom blackened). One received nutrient enrichment (10 μM NO₃⁻ and 1 μM PO₄³⁻) and the other did not. After thorough mixing of the cultures, they were placed in the normal culture conditions (lights on at 08:00 h and off at 20:00 h) at 13:00 h. On the following day, water samples (ca 3 ml) were taken from the 'upper', 'middle' and 'bottom' layer (Watanabe et al. 1983) at 00:30 h and 11:30 h. Concentrations of small cells (≤ 40 μM) were measured by microscope counts using a 1 ml Sedgewick-Rafter chamber with appropriate dilutions if necessary. As a control, vertical distributions of exponentially growing cells in TH medium were also monitored.

Field survey

**Monitoring of marine environment.** In summer 1989 (20 July to 13 August), field surveys were conducted at Stn B (depth = 21 m) around the Ie-shima Islands in the Seto Inland Sea (see Nakamura et al. 1988). Water temperature and salinity were monitored daily at 2.5 m intervals using a Surveyor H (Hydrolab Co.). Transparency (Secchi depth) was monitored daily. Water samples for chemical analysis [NO₃⁻, NO₂⁻, PO₄³⁻, Si(OH)₄, chlorophyll a] and enumeration of *Chattonella antiqua* and *C. marina* cells were taken daily from depths of 0, 5, 10, 15 and 19 m using a 10 l Van Dorn type bottle. Samples for vitamin B₁₂ analysis were obtained using the same bottle from depths of 0, 10 and 19 m at 3 d intervals. Monitoring and sampling were conducted in the morning (08:30 to 09:30 h) and water samples were treated (filtration, enumeration of cells, etc.) immediately in our field laboratory (Nakamura et al. 1988). Analytical procedures for each chemical variable have been summarized in previous papers (Nakamura et al. 1988, 1989). Concentrations of *C. antiqua* and *C. marina* cells were measured by observing 1 ml of the intact seawater sample in a Sedgewick-Rafter chamber using a microscope.

**Cyst trap.** In order to monitor the flux of *Chattonella* spp. cysts (cysts of *C. antiqua* are morphologically indistinguishable from those of *C. marina*; Imai & Itoh 1988), sediment trap bottles (50 cm high by 10 cm i.d., made of acrylic acid resin) were immersed daily from 5 to 12 August (except on 6 August) at depths of 15 and 19 m in the afternoon (ca 14:00 h). At each depth, 2 bottles were placed in a cage suspended from the surface by nylon rope and a buoy. The following day, trap bottles were recovered (ca 13:30 h) and all but ca 650 ml of the supernatant was removed by siphoning on board. Two trap samples from the same depth were combined into a 2 l polypropylene bottle and carried to our field laboratory.

At the field laboratory, trap samples were sieved through 20 μm netting, washed with and suspended in filtered seawater, and poured into a graduated cylinder for volume measurement. Then the samples were introduced into 100 ml polyethylene bottles, stored at 10°C in the dark until the end of the survey period and carried to our institute (ca 10°C, in the dark).

Cysts were enumerated, using an epifluorescence microscope, by observing autofluorescence under blue-light excitation irrespective of degree of maturation (Imai & Itoh 1988). The samples were placed in a Palmer-Maloney counting chamber (0.18 ml) and enumeration was conducted at least 10 times for each sample.

**RESULTS**

**Encystment in laboratory cultures**

Attempts to form cysts under nutrient-replete conditions

Results of experimental attempts to form cysts under N- and P-sufficient conditions are summarized in Table 2. When an exponentially growing culture (growth rate = 0.7 d⁻¹) in P-encystment medium was transferred to the dark, no cysts were formed and essentially no cells survived. In contrast, following P-depletion in the same culture, transfer to the dark induced encystment and a significant fraction of the cells (especially small cells) survived after dark incubation (Table 2).

We hypothesized that a rapid increase in temperature or light intensity would cause formation of small cells (precursors of cysts) under nutrient-replete conditions. Although these treatments caused retardation
of the growth rate (temperature shift) or a slight swelling of the cells (light shift), no small cells were formed. Following these treatments, the cultures were transferred to the dark, but no cysts were formed (Table 2).

Another attempt to form cysts under N,P-replete conditions was conducted using an Fe-limited culture. Although Fe-enrichment to the culture revived the growth, indicating that the culture was Fe-depleted (data not shown), small cells were not observed before transfer to the dark and cysts were not formed (Table 2).

Cyst yield under environmental and nutritional stress

Cyst yield was strongly affected by light intensity (Fig. 1). The yield decreased steadily with light intensity. However, encystment was not inhibited completely even under the maximum light intensity examined (150 μE m⁻² s⁻¹; yield = 1 cyst cm⁻²). Although total motile cell concentrations after the incubation were 50 to 75% of the initial values (data not shown), concentrations of small cells after the incubation were higher than the initial value (250 cells ml⁻¹) at each irradiance level and did not show any systematic trend with irradiance (Fig. 1).

In the temperature range examined (17.6 to 26.6°C), cyst yield ranged from 6 to 67 cysts cm⁻², with the highest production at and above 21.6°C (Fig. 2). The concentration of small cells after the incubation appeared to vary in conjunction with cyst yield, except at 26.6°C, and was higher than the initial value (310 cells ml⁻¹) at all temperatures examined.
Cyst yield increased linearly with total motile cell concentration (Fig. 3). In other words, encystment efficiency was constant and no cell-concentration-dependent effects were apparent.

Nutrient re-addition to N- or P-depleted cultures did not affect the cyst yield significantly (Table 3). Irrespective of nutrient re-addition, small-cell concentrations after the incubation were higher than the initial value (760 and 800 cells ml\(^{-1}\) for N-depleted and P-depleted cultures, respectively). Furthermore, small cells isolated from N- or P-depleted cultures and placed into N- and P-replete medium did not become large (vegetative) cells again, irrespective of light conditions (12 h light:12 h dark, or darkness). Division of small cells was not observed.

**Vertical distribution of small cells**

In an attempt to explain the field observation that small cells were only found near the bottom layer (see below), the vertical distributions of small cells in a cylindrical tube were examined. Although an exponentially growing culture displayed vertical migration, small cells always accumulated at the bottom irrespective of nutrient enrichment (Table 4).

**Field observations**

In summer 1989, we conducted a field survey around the Ie-shima Islands (20 July to 13 August). Vertical profiles of environmental variables throughout the survey period are summarized in Table 5. A small red tide due to *Chattonella antiqua* was observed from 7 to 11 August (Fig. 4A; see also Fig. 5). The bloom occurred on 7 August and the vegetative cell concentration averaged over the water column reached a maximum on 8 August (47 cells ml\(^{-1}\)). Thereafter, the population decreased rather rapidly to 2 cells ml\(^{-1}\) on 12 August.

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### Table 3. *Chattonella antiqua*. Effects of nutrient re-addition to nutrient-depleted cultures on cyst yield (no. cm\(^{-2}\)) and small-cell concentrations (no. ml\(^{-1}\))

<table>
<thead>
<tr>
<th>Culture</th>
<th>Treatment</th>
<th>Small-cell conc. before dark incubation</th>
<th>Cyst yield(^a) (SE)</th>
<th>Small-cell conc. after dark incubation(^b)</th>
</tr>
</thead>
<tbody>
<tr>
<td>N-depleted</td>
<td>Control</td>
<td>760</td>
<td>46 (6)</td>
<td>1550</td>
</tr>
<tr>
<td></td>
<td>+N</td>
<td>760</td>
<td>45 (3)</td>
<td>2040</td>
</tr>
<tr>
<td></td>
<td>+N and P</td>
<td>760</td>
<td>47 (3)</td>
<td>2240</td>
</tr>
<tr>
<td>P-depleted</td>
<td>Control</td>
<td>800</td>
<td>76 (14)</td>
<td>1370</td>
</tr>
<tr>
<td></td>
<td>+P</td>
<td>800</td>
<td>76 (15)</td>
<td>1670</td>
</tr>
<tr>
<td></td>
<td>+N and P</td>
<td>800</td>
<td>69 (6)</td>
<td>2070</td>
</tr>
</tbody>
</table>

\(^a\) Mean of 3 replicates
\(^b\) Measured in 1 of 3 replicates
Table 4. *Chattonella antiqua*. Vertical distributions of small cells and vegetative cells (no. ml⁻¹) in cylindrical bottles under 12 h light/12 h dark. For each sample the number of hours into the light (L) or dark (D) period is shown (e.g. 5.0 L = 5 h after the light went on).

<table>
<thead>
<tr>
<th>Sampling period</th>
<th>Sampling layer</th>
<th>Small cells (P-depleted)</th>
<th>Small cells (nutrients re-added)</th>
<th>Vegetative cells (exponential phase)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Day 1, 13:00 h (5.0 L)</td>
<td>Mixed</td>
<td>1820</td>
<td>1820</td>
<td>940</td>
</tr>
<tr>
<td>Day 2, 00:30 h (4.5 D)</td>
<td>Upper</td>
<td>20</td>
<td>60</td>
<td>150</td>
</tr>
<tr>
<td></td>
<td>Middle</td>
<td>10</td>
<td>30</td>
<td>110</td>
</tr>
<tr>
<td></td>
<td>Lower</td>
<td>56900</td>
<td>38200</td>
<td>22300</td>
</tr>
<tr>
<td>Day 2, 11:30 h (3.5 L)</td>
<td>Upper</td>
<td>30</td>
<td>10</td>
<td>12850</td>
</tr>
<tr>
<td></td>
<td>Middle</td>
<td>90</td>
<td>10</td>
<td>120</td>
</tr>
<tr>
<td></td>
<td>Lower</td>
<td>147600</td>
<td>217600</td>
<td>400</td>
</tr>
<tr>
<td>Day 2, 14:00 h (6.0 L)</td>
<td>Mixed</td>
<td>1800</td>
<td>1410</td>
<td>1720</td>
</tr>
</tbody>
</table>

Table 5. Vertical profiles of environmental variables at Stn B from 20 July to 13 August 1989. Secchi depth during this period changed from 5.5 to 9 m (avg. = 7.4 m). Values in parentheses are averages of each parameter throughout the survey period.

<table>
<thead>
<tr>
<th>Depth (m)</th>
<th>Temp (°C)</th>
<th>S (%)</th>
<th>NO₃⁻ (µM)</th>
<th>NO₂⁻ (µM)</th>
<th>PO₄³⁻ (µM)</th>
<th>Si(OH)₄ (µM)</th>
<th>B₁₂ (ng 1⁻³)</th>
<th>Chl a (µg 1⁻³)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>24.0-26.4 (25.0)</td>
<td>31.0-32.2</td>
<td>&lt;0.1-1.9</td>
<td>&lt;0.1-0.2</td>
<td>&lt;0.01-0.32</td>
<td>2.7-10.3</td>
<td>2.2-4.1</td>
<td>0.7-5.8</td>
</tr>
<tr>
<td>5</td>
<td>23.5-26.3 (24.7)</td>
<td>31.0-32.2</td>
<td>&lt;0.1-1.6</td>
<td>&lt;0.1-0.3</td>
<td>&lt;0.01-0.14</td>
<td>2.6-11.6</td>
<td>2.1-4.6</td>
<td>1.0-6.5</td>
</tr>
<tr>
<td>10</td>
<td>22.5-26.1 (24.4)</td>
<td>31.2-32.3</td>
<td>&lt;0.1-3.7</td>
<td>&lt;0.1-0.8</td>
<td>0.01-0.25</td>
<td>3.1-15.6</td>
<td>2.8-3.7</td>
<td>0.9-4.6</td>
</tr>
<tr>
<td>15</td>
<td>21.2-25.0 (23.7)</td>
<td>31.6-32.5</td>
<td>0.5-9.0</td>
<td>0.1-1.3</td>
<td>0.03-0.43</td>
<td>5.3-27.5</td>
<td>1.2-3.7</td>
<td>0.9-2.9</td>
</tr>
<tr>
<td>19</td>
<td>21.1-24.2 (23.0)</td>
<td>31.7-32.5</td>
<td>1.1-9.3</td>
<td>0.3-1.4</td>
<td>0.29-2.64</td>
<td>6.6-38.6</td>
<td>1.9-3.7</td>
<td>1.4-3.3</td>
</tr>
<tr>
<td>Bottom</td>
<td>21.0-24.1 (22.8)</td>
<td>31.6-32.5</td>
<td>1.1 (4.7)</td>
<td>0.7 (0.7)</td>
<td>0.71 (22.9)</td>
<td>(2.7) (2.3)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

(see Fig. 5). During the blooming period, other phytoplankton species were rather scarce.

Changes in the concentration of *Chattonella antiqua* vegetative cells and in temperature and nutrients (NO₃⁻, PO₄³⁻) from just before until the end of the bloom (5 to 13 August) are shown in Fig. 4. Throughout this period, temperature was optimum for *C. antiqua* growth (Nakamura & Watanabe 1983a, K. Yamanaka unpubl.). In the period from pre-initiation to bloom development (5 to 8 August), concentrations of nitrate and phosphate in the surface layer (0 to 5 m) were relatively high and comparable to the half-saturation constants for growth of *C. antiqua* (K₉) for these nutrients (K₉NO₃ = 1 µM, K₉PO₄³⁻ = 0.11 µM; Nakamura et al. 1988). After this period, nutrients were exhausted in the surface layer. Nutrient concentrations at 10 m were higher than K₉'s from 5 to 8 August and decreased after 9 August (except or 10 August). Below 15 m, nutrients stayed at high levels irrespective of bloom development.

On 7 and 8 August, vegetative cells of *Chattonella antiqua* appeared healthy in terms of size (ca 100 µm) and color, and accumulated in the surface layer in the morning. However, in the period of bloom decline, cells became smaller (ca 80 µm) and looked pale, and their distribution was shifted downward (Fig. 4). Furthermore, small cells (ca 30 µm) were observed on 10 and 11 August at 19 m, at concentrations of 19 and 170 cells ml⁻¹, respectively. 'Fusing' pairs of small cells were also found. Except for this period and layer, no small cells or 'fusing' pairs were observed during the entire survey period. *C. marina*, whose cysts are morphologically indistinguishable from those of *C. antiqua* (Imai & Itoh 1988), were scarce (≤ 2 cells ml⁻¹) from 5 to 12 August. A colorless dinoflagellate, *Gyrodinium* sp., which contained debris of *C. antiqua*, was apparent after 10 August (cell concentrations = ca 10 to 100 cells ml⁻¹), but quantitative enumeration was not conducted.

Changes in *Chattonella* spp. cyst flux (5 to 13
Chattonella antiqua. Bloom outbreak time series at Stn B. (A) Vegetative cell concentration; (B) temperature; (C) nitrate concentration; (D) phosphate concentration. H: high, L: low.

August) are shown in Fig. 5. The number of cysts at 19 m showed small values (≤ 1.5 cysts cm⁻² d⁻¹) during the pre-bloom and bloom-development periods (5 to 8 August), increased rapidly to reach a sharp maximum between 9 and 10 August (bloom-declining period; 45 cysts cm⁻² d⁻¹) and then decreased. The total flux at 19 m for the period was 98 cm⁻², comparable to the cyst concentration in the bottom sediment of the Seto Inland Sea (Imai & Itoh 1985). Cyst flux at 15 m showed a slight maximum in the bloom-declining period (3 cysts cm⁻² d⁻¹), but was negligible in comparison with that at 19 m. These results indicate that cysts of Chattonella spp. were formed mainly below 15 m in the bloom-declining period of the 1989 red tide.

**DISCUSSION**

As shown previously (Nakamura et al. 1990), cysts of Chattonella antiqua are formed through the sexual fusion of small cells (gametes). In the present study, small cells were observed only in N- or P-depleted cultures, in spite of attempts to form them under nutrient-replete conditions. Once small cells were formed by nutrient depletion, total cyst yield was influenced by temperature and light conditions, but not by the re-addition of nutrients. In the field, the cysts were formed
mainly below 15 m in the bloom-declining period when nutrients were exhausted in the *C. antiqua* habitat (0 to 10 m). These observations seem to be consistent with the laboratory results.

**Laboratory experiments**

It is well known that a reduction in nutrient concentrations is often effective in inducing gamete formation and encystment (e.g. Pfiester 1975, Turpin et al. 1978, Anderson et al. 1984, 1985, Anderson & Lindquist 1985). Within the scope of the present study, cysts of *Chattonella antiqua* were obtained only in N- or P-depleted cultures. No small cells were observed in the cultures without nutritional stress (exponentially growing or having undergone a light/temperature shift), and transfer of these cultures to the dark resulted in mortality of vegetative cells (Table 2). On the other hand, small cells formed under N- or P-depleted conditions survived in the dark (Tables 2 & 3). These findings, together with the observation that small cells did not revert to the vegetative state in nutrient-replete conditions, indicate that small cells were physiologically differentiated from nutrient-replete vegetative cells.

Since we used non-axenic cultures of *Chattonella antiqua*, one might conjecture that bacteria could have broken down Tris buffer (see Table 1) to ammonium and that small cell formation in N-encystment medium (see Table 3) occurred under N-sufficient conditions. However, the results of our preliminary experiments showed that motile cell (= vegetative plus small cell) concentration at the growth maximum in N-encystment medium (5 to 20 µM NO₃⁻) increased almost linearly with initial nitrate concentrations (Umemori 1990). Thus, we do believe that small-cell formation in the N-encystment medium occurred under N-depleted conditions even if some ammonium liberated from Tris was utilized by *C. antiqua*.

Since iron in the surface layer in summer around the Ie-shima Islands is usually insufficient to support rapid growth of *Chattonella antiqua* (Nakamura 1990), and Fe limitation induces encystment in the dinoflagellate *Alexandrium tamarense* (Doucette et al. 1989), we tried to form cysts in Fe-limited cultures (Table 2). Although the total motile cell concentration before dark incubation (130 cells ml⁻¹) was much lower than that in P-encystment medium (4120 cells ml⁻¹; Table 2), small cells were not found and no cysts were detected in 5 replicate counts (i.e. 20 coverslips; 35 cm²). If we assume that the efficiency of encystment in Fe-depleted cultures is not very different from that in P-depleted cultures (cyst yield = 80 cysts cm⁻²; Table 2), the expected cyst yield in the former would be 80 x (130/4120) = 2.5 cysts cm⁻², much higher than the observed value (no cysts in a 35 cm² area). Although the possibility of *C. antiqua* forming cysts under Fe-depleted conditions cannot be ruled out completely due to the low population density used, we conclude that the efficiency of small-cell formation and encystment under Fe-limited conditions is much less than that in P- (and N-) depleted conditions.

In some phytoplankton species, sexual reproduction occurs under nutrient-replete conditions (e.g. Wall et al. 1970, Nozaki 1986, Montresor & Zingone 1988) and our attempts to form cysts without N,P-depletion were rather restricted. Thus, it is too early to conclude that small cells and cysts are formed only under N- or P-depleted conditions. However, it should be stressed that nutrient depletion is at least one important factor for small-cell formation by *C. antiqua*.

As shown in Fig. 1, the cyst yield was at a maximum under darkness and decreased with increasing light intensity. These trends have also been observed for *Chattonella marina* (Imai 1989) and explain the generally low cyst yields at high irradiance (15 m; Fig. 5). The concentration of small cells after incubation did not revert to the vegetative state in nutrient-replete conditions, indicate that small cells were physiologically differentiated from nutrient-replete vegetative cells. The concentration of small cells after incubation did not revert to the vegetative state in nutrient-replete conditions, indicating that small cells were physiologically differentiated from nutrient-replete vegetative cells.

Vegetative growth of *Chattonella antiqua* was positive above 15°C, with maximum growth rate occurring in the range of 24 to 27°C, and cell mortality was observed at 31°C (Nakamura & Watanabe 1983a, K. Yamanaka unpubl.). At 21.6°C, the lower temperature limit for the highest level of cyst production, the growth rate was about 60% of the maximum. Although the upper temperature limit for the highest level of cyst production was not determined and the strains used in the present study (A-0 and B-20) were different from that used for growth rate measurements (Ho-1), the range for the highest cyst production (21.6 to ≥ 26.6°C; Fig. 2) is apparently wider than that for growth.

One might expect encystment efficiency to be optimal at or above a specific cell concentration, with the lowest values at reduced cell concentrations, where interaction between small cells (gametes) would be less frequent. However, the linear relationship between cyst yield and motile cell concentration (Fig. 3) indicates that encystment efficiency was constant across a wide range of motile cell concentrations. Relatively constant encystment efficiency was also observed in N- or P-limited cultures of *Gonyaulax tamarensis* (Anderson et al. 1984). Unfortunately, we cannot presently make any conclusive statements about the constancy of encystment efficiency.

In the field, cysts of *Chattonella* spp. were formed mainly in the nutrient-replete layer (Figs. 4 & 5). In
contrast, cysts were not formed in laboratory cultures without nutrient depletion, within the scope of the present study. In order to explain this discrepancy, nutrient re-addition effects were examined (Table 3). The results indicated that once small cells have been formed under N- or P-depleted conditions, the process cannot be reversed and their ability to form cysts is maintained even if ambient nutrient concentrations increase again.

The concentrations of small cells in N- or P-depleted cultures (and nutrient re-addition cultures) after dark incubation were higher than those before dark incubation (Tables 2 & 3). This indicates that small cells continued to be formed in the dark, and that most of the small cells did not participate in encystment. Since small cells were tentatively defined as cells ≤ 40 μm in length, some of them might have been nutrient-depleted vegetative cells (which cannot become encysted). In addition, encystment of *Chattonella antiqua* is completed within 3 d after transfer to the dark (Nakamura et al. 1990). The formation process for small cells, duration of their fusing ability, and differentiation of gametes from small vegetative cells should be clarified in order to understand the encystment process on a biological basis.

### Field observations

In comparison with the *Chattonella antiqua* red tide in 1987 (Nakamura et al. 1989), the magnitude and distribution of the 1989 red tide were rather limited (S. Yoshimatsu pers. comm.). Furthermore, the occurrence of the 1989 red tide around the Ie-shima Islands was coupled with the decline of the bloom at the northern edge of the Seto Inland Sea (Futami fishing harbor, 30 km from the Ie-shima Islands; H. Kobayashi pers. comm.), suggesting that the sudden occurrence of the bloom (7 August 1989) was due (partly) to advection.

In spite of the great difference in the magnitude of red tides between these 2 years, changing patterns of environmental variables in 1989 from just before initiation to the termination of the bloom (5 to 13 August) resembled those in 1987. In both years, temperature and vitamin B₁₂ were optimal throughout the period (cf. Nakamura & Watanabe 1983a, Nakamura et al. 1988). N- and P-concentrations at or above 10 m were relatively high just before bloom initiation and during the bloom-developing period, and nutrients were exhausted in the *Chattonella antiqua* habitat (0 to 10 m; Hamamoto et al. 1979, Nakamura et al. 1989) during the bloom-declining period (Fig. 4). However, in contrast to the 1987 red tide, nutrients at or below 15 m were not exhausted and the decline of the bloom was apparently coupled with the grazing of *C. antiqua* by the colorless dinoflagellate *Gyrodinium* sp. The cyst flux data are discussed below with reference to the changes in environmental variables.

Turbid bottom water is widely present in the Seto Inland Sea and resuspension of the bottom sediments is considered to be the cause of the turbidity (Kawana & Ōmoto 1984). Thus, one might consider that cyst flux at 19 m (F₁₉) was strongly affected by the resuspension of the cysts in the bottom sediment. However, it should be noted that F₁₉ was very small (≤ 3 cysts cm⁻² d⁻¹) before the initiation of the bloom (5 to 6 August), during the development period (7 to 8 August) and after termination (12 to 13 August) (Fig. 5). In addition, since the weather was sunny and calm between 8 and 11 August (the period when F₁₉ showed high values), as well as during the other period of cyst-trap experiments (Ie-shima Weather Monitoring Station, Meteorological Agency, Japan), it is unlikely that resuspension of the cysts between 8 and 11 August was greater than in the other period. We conclude that the contribution of resuspension to F₁₉ was negligible.

Cysts of *Chattonella antiqua* are morphologically indistinguishable from those of *C. marina* (Imai & Itoh 1988) and the cyst flux observed was the total for the 2 species. However, vegetative cells of *C. marina* were scarce (≤ 2 cells ml⁻¹) during the period of cyst-trap experiments and its contribution to the flux was probably negligible.

Since encystment of *Chattonella antiqua* is affected markedly by irradiance levels (Fig. 1), comparison of the light intensity between 15 and 19 m (i.e. the depth where traps were immersed) is important for interpretation of the cyst flux data (Fig. 5). Although light intensities were not measured in the 1989 survey, light attenuation with depth was monitored once a day in 1990 (21 July to 13 August; no red tides; Nakamura unpubl.) and light intensities at 15 and 19 m in 1989 were roughly estimated as follows. Using the daily total irradiance data at Osaka (100 km from the Ie-shima Islands; Meteorological Agency, Japan) and light attenuation data, light intensities averaged over the daytime period at depths of 15 and 19 m (I₁₅ and I₁₉) in 1990 were calculated. Since light attenuation coefficients were correlated with Secchi depth (6 to 18.5 m; Nakamura et al. 1989), the formation process for small cells did not participate in encystment. Since encystment of *Chattonella antiqua* is completed within 3 d after transfer to the dark, it is unlikely that resuspension of the cysts between 8 and 11 August was greater than in the other period. We conclude that the contribution of resuspension to F₁₉ was negligible.
maximum (at 0 μE m⁻² s⁻¹). In other words, encystment efficiency at 19 m is 3.7 times higher than that at 15 m, and this difference partly explains the fact that cysts were mainly formed below 15 m (Fig. 5).

Based on the results obtained from the laboratory culture experiments and field observations, the cyst flux data (Fig. 5) seem to be explained consistently by the changes in the environmental variables, and a simple scenario for the encystment of *Chattonella antiqua* in the field can be drawn. Namely, following the development of the bloom, nutrients in the habitat of *C. antiqua* (0 to 10 m) were exhausted (Fig. 4) and small cells were formed due to N- and/or P-depletion (Tables 2 & 3). Since small cells have a tendency to sink (Table 4), they descended below 15 m depth. The observation that small cells were found only at 19 m in the bloom-declining period (10 and 11 August) confirms the above reasoning. Below a depth of 15 m, temperature was in the optimum range for encystment (Figs. 2 & 4) and replete nutrients (Fig. 4) did not affect the encystment ability of the small cells (Table 3). Furthermore, encystment efficiency increased with increasing depth (or decreasing light intensity) (Fig. 1), and light conditions above 15 m were somewhat inhibitory for encystment [see above]. Thus, as shown in the cyst flux data (Fig. 5), cysts were formed mainly below 15 m in the bloom-declining period.

Although the above scenario for the encystment process in the field is apparently reasonable, there are some uncertainties with respect to the time required for encystment. In our time-course measurements of encystment in batch culture (Nakamura et al. 1990), it took 2 or 3 d for encystment following the formation of small cells. If this is also the case for the natural populations, the maximum cyst flux observed between 9 and 10 August (Fig. 5) was due to the small cells formed on 7 and 8 August, when nutrients at or above 10 m were not completely exhausted and *Chattonella antiqua* populations were developing (Fig. 4). In addition, sinking rates of small cells were not determined and the time required for small cells to reach a depth below 15 m is still unknown. In this context, the above scenario is not conclusive.

Laboratory culture results and field observations involving the red tide dinoflagellate *Gonyaulax tamarensis* are inconsistent. Cysts were not formed under nutrient-replete conditions in batch culture (Anderson et al. 1984, Anderson & Lindquist 1985), whereas encystment in the field occurred under conditions seemingly favorable for growth (Anderson et al. 1983). Although the reason for the inconsistency observed in *Chattonella antiqua* and *G. tamarensis* has not yet been identified, one possible explanation is that it is due to the characteristics of batch culture. In batch culture, concentrations of limiting nutrient decrease rapidly to reach ‘zero’ (below the detection limit) with population growth, and it is difficult to examine small-cell (gamete) formation at nutrient levels in the growth-rate-limiting range (ca 0.1 μM PO₄³⁻ or 1 μM NO₃⁻ for *C. antiqua*; Nakamura et al. 1988). Continuous culture is needed in order to consider the possibility that *C. antiqua* may form small cells under conditions where ambient nutrient is not depleted but regulates growth.

Since cyst flux at 19 m changed drastically during the experimental period (5 to 13 August; Fig. 5), it would be interesting to monitor the cyst concentration in the bottom sediment throughout the blooming period. We are now planning to monitor the cyst concentrations in the bottom sediments by direct counting (total cysts) and by the extinction dilution method (germinative cysts) together with measurements of cyst flux.

In summary, encystment conditions of *Chattonella antiqua* obtained in laboratory cultures were used to explain the changes in *Chattonella* cyst flux. Although some uncertainties still remain with respect to the timing of small-cell formation, the observation that cysts were formed mainly below 15 m is explained well in terms of the conditions of irradiance, temperature and nutrients required for encystment.

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