Gametogenesis in Liquid Cultures of *Chlamydomonas eugametos*

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At the end of the exponential phase of vegetative growth in liquid cultures, cells of *Chlamydomonas eugametos* develop into gametes. Gametogenesis can occur without nitrogen deficiency. In continuous light, mating competence lasts only for a few hours, resulting in a low percentage of gametes. Nevertheless, mating competence is considered to be a general property of newly born cells in late exponential phase, since cultivation of both mating-types in mixed cultures normally leads to a high yield of paired cells. No evidence was found for an influence of the presence of partner gametes on gametogenesis. During gametogenesis of the *mt*– mating-type, extractable biologically active agglutination factor appears in the cell body. Subsequently, after a gametogenic division, mating competence and agglutinability are expressed, both fluctuating synchronously with the light/dark regime.

**INTRODUCTION**

*Chlamydomonas eugametos* is a biflagellate unicellular green alga, reproducing vegetatively by sporulation. For sexual reproduction, vegetative cells have to differentiate into gametes. There are no morphological differences between gametes and vegetative cells, nor between the gametes of the two mating-types: *plus* (*mt)* and *minus* (*mt*–). Gametes are distinguished from vegetative cells by their ability to agglutinate and to form vis-à-vis pairs with gametes of the opposite mating-type. Plasmogamy and karyogamy of the paired cells lead eventually to the formation of zygotes. Agglutination takes place via the flagella. The glycoprotein responsible for agglutination (the *mt*– agglutination factor) has been isolated from flagella of *mt*– gametes (Musgrave et al., 1981; Homan et al., 1982), and has also been found in relatively large quantities in the cell body (Pijst et al., 1983).

In the related species *C. reinhardtii*, gametogenesis is induced by nitrogen starvation (Sager & Granick, 1954). Synchronous gametogenesis is accomplished by placing synchronously dividing vegetative cells in the mid G1-phase into nitrogen-free medium (Kates & Jones, 1964). Gametogenesis usually involves a mitotic division (Kates & Jones, 1964; Schmeisser et al., 1973). Nitrogen depletion has also been reported to induce sexual differentiation in *C. eugametos* (Bernstein & Jahn, 1955), but there are, as far as we know, no reports of synchronous gametogenesis. In nature, at relatively low nitrogen levels, asexual and sexual reproduction of *C. eugametos* seem to co-exist (Trainor, 1975), which prompted us to reinvestigate the role of nitrogen depletion in gametogenesis in this species.

In this work gametogenesis was studied using the following parameters: mating competence, flagellar agglutinability and the presence of extractable biologically active *mt*– agglutination factor.

**Abbreviations:** LD, 16 h light/8 h dark; LL, continuous light; CL-medium, conditioned medium from stationary-phase liquid cultures; CA-medium, conditioned medium from flooded agar cultures.
METHODS

Strains. Chlamydomonas eugametos strains UTEX 9 (mating-type plus, mt+) and UTEX 10 (mating-type minus, mt-) were used from the Algal Collection at the University of Texas at Austin, USA.

Plate cultures. Cells were cultivated in Petri dishes on agar as described by Mesland (1976). Standard gamete suspensions were obtained by flooding 2-week-old cultures with water.

Liquid cultures. Cells were cultivated in Fernbach flasks with 1000 ml medium containing 1/100 of the minerals used by Kates & Jones (1964), with trace elements according to Wiese (1965) and Fe-EDTA according to Jones (1962). The cultures were grown at 19 °C with illumination by white fluorescent tubes, 3500 lx (42 μE s⁻¹ m⁻²); a 16 h light/8 h dark (LD) regime was used unless indicated otherwise. The cultures were aerated with compressed air and gently agitated at 50 r.p.m. on a rotary shaker. Cells from 2- to 3-week-old agar plates, flooded with medium the previous day, were used as inoculum; the inoculation density was 1 × 10⁴ cells ml⁻¹. To determine cell densities, samples with glutaraldehyde (final concn 1:25%, v/v) were counted in a haemocytometer.

Mixed cultures. Mixed cultures were started with equal numbers of non-competent mt+ and mt- cells from late stationary phase cultures, to ensure that pairs did not form in the mixed inoculum. Samples, fixed with glutaraldehyde, were counted for single cells and vis-a-vis pairs in a haemocytometer to give both cell density and percentage of paired cells.

Mating-type analysis in mixed cultures. To determine the ratio of the two mating-types in a mixed culture, a mouse monoclonal antibody raised against mt- flagellar antigens (Homan et al., 1984) was used. This antibody specifically reacted with mt- flagella and, after removing the cell wall with NaOH, also with the plasma membrane of mt+ cells. Therefore, to label all mt- cells, including sporulating cells that had retracted their flagella, cells of a mixed culture were fixed with glutaraldehyde and treated with 1 m-NaOH for 30 min at 40 °C according to Pijst et al. (1983). After washing, the cells were spread on a slide and incubated for 30 min with the antibody. Bound antibodies were labelled with anti-mouse antibodies bound to fluorescein isothiocyanate (goat anti-mouse IgG-FITC, Tago Inc., Burlingame, Calif., USA). The samples were then scored for fluorescent mt+ cells and non-fluorescent mt- cells.

Conditioned medium. Cell suspensions were centrifuged and the supernatants filtered through a 0.8 μm Millipore filter to obtain conditioned cell-free culture medium.

Mating competence test. To determine the percentage of mating-competent cells in a sample, mt- cells were mixed in triplicate in test tubes with an excess of standard mt+ gametes for 1 h at 19 °C with illumination by white fluorescent tubes, 5500 lx (65 μE s⁻¹ m⁻²). After fixation with glutaraldehyde, individual cells and vis-a-vis pairs were counted in a haemocytometer. With the known ratio of the two mating types in the mixture, the mating competence of the mt+ cells (= percentage of mt- gametes) was calculated as follows: (no. of pairs ml⁻¹ × 100)/ (no. of mt+ cells ml⁻¹). This test, based on an experiment described by Lewin (1956), gives no information about the mating competence of the mt+ cells forming the majority of the mixture. For correct test conditions, the mating competence of the major partner had to be at least 20%, so the standard mt+ gametes were always cross-checked as the minor partner against standard mt- gametes in a control test. It was established that the optimal test density was 2-4 × 10⁶ cells ml⁻¹, the portion of the minor partner not exceeding 20% of the mixture.

For determining the mating competence of a mt+ suspension, the same procedure was followed with an excess of standard mt- gametes.

Test of flagellar agglutinability. mt- cells were fixed with glutaraldehyde for 30 min, washed, and mixed on a glass slide with standard mt+ gametes. The agglutinability was compared microscopically with the activity of an equivalent sample of fixed standard mt- gametes with a known high mating competence. The agglutinability of the test cells was expressed in relation to the agglutinability of the standard gametes, which was arbitrarily given a value of 5 points.

Extraction and bioassay of mt- agglutination factor. A constant amount of 10⁷ mt- cells was exhaustively extracted with 1% (w/v) SDS. The extract was assayed with mt+ gametes for biological activity by the charcoal technique and the activity was expressed as the titre of the highest dilution of a series (2¹, 2², 2³, etc.) still showing biological activity (Musgrave et al., 1981).

Determination of nitrate and phosphate in the culture medium. The concentration of nitrate in cell-free medium was determined according to Stainton et al. (1974) after quantitative reduction to nitrite in a Cd/Cu-column (Woods et al., 1967; Nydahl, 1976). Phosphate concentrations were calculated by determining the phosphorus contents of cell-free medium according to Murphy & Riley (1962).

RESULTS

Gametogenesis in liquid cultures under continuous light

The standard method for producing gametes in liquid cultures of C. reinhardtii and C. moewusii is to transfer vegetative cells from a complete medium to nitrogen-free medium (Kates & Jones, 1964). When this method was applied to C. eugametos, under the conditions of Kates &
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Fig. 1. Growth (●) and gametogenesis (percentage of mating-competent cells, ▲) of C. eugametos mt− grown under LL. Results for mt+ cells were similar, except for the early exponential growth phase (dashed line).

Fig. 2. Growth (○) and gametogenesis (percentage of paired cells, □) of C. eugametos grown under LL in a mixed culture (mt+ with mt−). Cultures were inoculated with non-competent cells from single cultures in the late stationary phase of growth, to prevent forming of pairs in the inoculum.

Table 1. Growth and nutrient status of C. eugametos mt− with and without additional KNO₃

<table>
<thead>
<tr>
<th>Nitrate concn (μM)</th>
<th>Phosphate concn (μM)</th>
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<tr>
<td>0·01 mM-KNO₃</td>
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<tr>
<td>initial</td>
<td>final</td>
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<td>1</td>
<td>28</td>
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<td>0·64</td>
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<td>140</td>
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<td>1100</td>
<td>950</td>
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<td>4·5</td>
<td>0·12</td>
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Jones (1964), cells kept their vitality and motility, but no gametes were found. Similar results were obtained with cells washed and resuspended in H₂O.

When the mineral content of the medium was decreased (see Methods), gametes were found during 1 or 2 d at the end of the period of vegetative growth in liquid cultures under continuous light (LL), where they accounted for up to 25% of the population (Fig. 1). To determine whether the gametogenesis had been due to nitrogen deficiency, the concentration of nitrate in the culture medium was raised from 0·01 mM to 1 mM (the concentration used by Kates & Jones, 1964). Under these conditions, cultures reached a much higher plateau density, indicating that nitrogen depletion caused growth arrest in the normal medium. Mating-competent cells accounted for up to 40% of the cultures with extra nitrate, which had by no means been exhausted at the end of the growth period (Table 1), indicating that gametogenesis can occur in cultures in which the nitrogen content is not growth limiting. In contrast, only 3% of the original phosphate was left at the end of growth. Presumably deficiency of some nutrient other than nitrogen had caused the growth arrest and induced gametogenesis.

Although under normal conditions the number of fusing gametes did not exceed 25%, it seemed possible that a higher percentage of the cells did participate in gametogenesis. This was suggested by the appearance and disappearance of the mating-competent cells, which closely corresponded with the production of new cells in the late exponential phase (Fig. 1). The implication is that gametes are mating competent for only a limited period. To test this possibility, mt+ and mt− cells were cultivated in mixed cultures. Assuming that growth and development of the two mating-types occur in parallel and that during the mating-competent period of each particular cell it is possible to find a partner, the prediction is that by
accumulation the percentage of paired cells should easily exceed 25%. In most experiments, the percentage of paired cells was indeed 40% or more, with a maximum of 60% (Fig. 2). This result supported the idea of a short mating-competent period per cell, but an alternative explanation for the high gamete yield in mixed cultures could be the mutual stimulation of gametogenesis in both mating-types. However, evidence against this was provided by two deviating cases when hardly any pairs were formed (Fig. 3). Determination of the ratio of the individual mating-types in mixed cultures, using a monoclonal antibody specifically reacting with the mt- strain, indicated that in one experiment pair formation failed because mt+ lagged behind mt- in the vegetative growth phase. As a result, there was almost no overlap in the gamete phases, so that few pairs could be formed (Fig. 3a). Another experiment involved mt- cells with a very low mating competence, as determined in the single control culture. In the presence of the opposite mating-type, the mt- cells maintained a low mating competence, while the mt+ strain produced a normal amount of gametes (Fig. 3b). Therefore, the mating-types seem to develop independently both vegetatively and sexually, without any mutual influence. This leads to the conclusion that mating competence is expressed for only a short period. From the experiment shown in Fig. 1 it was calculated that if all cells born in the late exponential phase underwent gametogenesis, they expressed their mating competence on average for 5 h.

Gametogenesis in liquid cultures under a light/dark regime

Under LD-conditions sporulation of C. eugametos mt- occurred in the dark period, new cells being born in daily bursts at the D−L transition (hatching). Under these conditions gametes were produced during the last cell burst before the culture entered stationary phase (Fig. 4), showing that a gametogenic division was involved. The gamete percentages, though rather variable, were consistently higher than in LL-cultures: up to 80%. Significantly, mating competence was expressed in peaks, in synchrony with the LD-rhythm, with maxima at the D−L transition. Fig. 4 shows three successive peaks of mating competence of 70, 60 and 45%, respectively, at constant cell density. This means that the oscillation of the mating competence concerns the same cells, i.e. they lose and recover their mating competence. Cells born during the gametogenic cell division were mating competent for 8 h on average. This also supports our conclusion that gametes born in continuous light are only temporarily able to fuse. However, since the mating competence returned in the dark period, darkness seems to extend the mating competence. Fig. 4 also demonstrates the relationship between mating competence, agglutinability of the flagella and the presence of extractable, biologically active, agglutination factor. The agglutination factor was first detected in cell extracts towards the end of exponential growth, before the cell burst. After hatching, the daughter cells were immediately agglutinable and mating competent. The flagellar agglutinability fluctuated, like the mating competence, with the LD-rhythm. Both disappeared 2 or 3 d after the gametogenic division, even though the cells maintained the same amount of extractable agglutination factor. In general, flagellar agglutinability closely followed mating competence, but occasionally strong agglutinability was accompanied by low mating competence, as can be seen on day 6 in Fig. 4. Therefore a complementary gametic property can be distinguished: fusion competence. Only cells with both flagellar agglutinability and fusion competence are mating competent.

Gametes derived from agar plates

As inoculum for liquid cultures 2-week-old agar plates were flooded with either H2O or complete nutrient medium. Both yielded suspensions with high mating competence. This implies that on 2-week-old agar plates induction to gametogenesis had already occurred. After flooding, the cells generated flagella and became mating competent. When flooded with H2O, the cells remained mating competent for at least 2 weeks. In complete medium, however, the gametes lost all sexual properties within 1–3 d. When this degametogenesis was complete, vegetative reproduction started (Fig. 4).

Gametes derived from agar plates (standard gametes) contained the same amount of biologically active agglutination factor per cell as gametes in liquid cultures (Fig. 4). However, the prolonged mating competence of standard gametes strongly contrasted with that of the
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Fig. 3. Growth and gametogenesis of *C. eugametos* during two experiments (a and b) similar to that of Fig. 2, but where forming of pairs was negligible (inset). In experiment (a) the development of *mt*+ lagged behind that of *mt*-, so that few pairs could be formed. In experiment (b) *mt*+ had a low mating competence compared to its partner, leaving most *mt*+ gametes unmated. The separate growth patterns of the *mt*+ and *mt*− cells in one culture were measured by determining the ratio of the mating-types using a *mt*−-specific monoclonal antibody (see Methods). The percentage of competent *mt*+ cells was determined by adding an excess of standard *mt*− gametes to culture samples and scoring the number of pairs formed. The percentage of competent *mt*− cells was determined in a similar way by adding standard *mt*+ gametes. ○, Growth of *mt*+; ●, growth of *mt*−; ○, growth of whole culture; ■, percentage of paired cells; △, percentage of mating-competent *mt*+ cells; ▲, percentage of mating-competent *mt*− cells.

Fig. 4. Growth and gametogenesis of *C. eugametos mt*− grown under a circadian LD-rhythm. Dark periods are represented as hatched rectangles. Cultures were inoculated with cells derived from agar plates. ●, Growth; ▲, percentage of mating-competent cells; ○, relative agglutinability; ■, titre of *mt*− agglutination factor in bioassay.

liquid-induced gametes. To investigate whether this difference was caused by the medium, standard *mt*− gametes were washed and resuspended in conditioned medium from liquid cultures in the stationary phase of growth (CL-medium) and from agar cultures flooded with water (CA-medium) and kept on a LD-regime. The mating competence was determined in both
the dark and the light periods. When gametes were resuspended at high density (about $1.5 \times 10^7$ cells ml$^{-1}$, i.e. the normal cell density of agar-derived gamete suspensions), the mating competence in CA-medium was preserved throughout the experiment and showed a LD-rhythm similar to that of liquid-induced gametes (Fig. 5). In CL-medium, however, mating competence soon dropped and was lost within 3 d. When gametes were resuspended at low density (about $2.5 \times 10^5$ cells ml$^{-1}$, i.e. the normal cell density of liquid-induced gamete suspensions), mating competence in CA-medium dropped rapidly and disappeared within 2 d. In CL-medium gametes lost their mating competence immediately and behaved like gametes induced in this medium. In all cases cell densities were constant throughout the experiment. These results indicate that conditioned medium from agar cultures contains substances that maintain the mating competence, but only when the cell density is high. In our liquid cultures the cell density is too low to produce enough of these substances to preserve the mating competence for a long time.

**DISCUSSION**

Gametogenesis in *C. eugametos* seems not to be triggered by the same events as gametogenesis in *C. reinhardtii* and *C. moewusii*. Cells could not be induced to undergo gametogenesis by transferring them to nitrogen-free medium. Furthermore, gametogenesis not only occurred when nitrate was at growth-limiting concentration, but also when nitrate was abundant and growth was prevented by deficiency of some other nutrient. These findings indicate that, as suggested by Trainor (1975), nitrogen does not play a key role in the gametogenesis of *C. eugametos*. Our conclusion is that nutrient stress in general is the factor that induces cells to undergo a final and gametogenic division. Wiese (1984) has also argued that gametogenesis is not induced by nitrogen depletion and that nitrate does not prevent gametogenesis, claiming that ammonium depletion rather than nitrate depletion triggers gametogenesis in *C. eugametos*. However, it is obvious that this was not the case in our experiments.

In the colonial Volvocales *Gonium* and *Pandorina*, gametogenesis only occurs after mixing the two mating types (Wiese, 1969). Our study did not provide any evidence for a mutual influence on vegetative or sexual development of the two mating-types.

In LD-cultures gametes appeared during the last cell burst. So, as in *C. reinhardtii*, gametogenesis in *C. eugametos* involves a gametogenic division (Kates & Jones, 1964;
Schmeisser et al., 1973). Agglutination factor could be detected in the daughter cells prior to hatching. Gametes in liquid cultures contained the same amount of agglutination factor as gametes derived from agar plates. This may mean that gametes contain a specific amount of agglutination factor per cell. Only a small part of the agglutination factor is extracted from the flagella, most of it being located on the cell body, as shown by Pijst et al. (1983). These authors proposed that agglutination factor could perhaps diffuse from the cell surface to the flagella and that the cell body therefore acted as a pool. However, the fact that many of our cells contained a full complement of agglutination factor, but were not agglutinable, suggests that there is no free diffusion between the membranes of the cell body and the flagella.

Flagellar agglutinability and mating competence of gametes oscillated with the LD-rhythm. Since the decline in flagellar agglutinability always occurred after the onset of the light period, a possible explanation is that light has a negative effect on this property and consequently on mating competence. This effect was only partly reversible and after 2 or 3 d no mating-competent cells were left. The short period of mating competence found in cultures grown under continuous light can also be seen as a result of the negative effect of light. This loss of mating competence must be distinguished from degametogenesis resulting in vegetative growth. Gametes induced in liquid cultures, though losing their mating competence, did not revert to the vegetative state. This is concluded from the absence of cell division and the presence of agglutination factor.

Under a LD-regime, liquid-induced gametes rhythmically expressed their mating competence for only a few days. In contrast, agar-derived gametes in medium from flooded agar cultures preserved this rhythmic mating competence, but only at a high cell density, indicating that substances secreted by the cells play some role. Taking this phenomenon together with the oscillation of flagellar agglutinability, an alternative explanation for the fluctuation in mating competence is that it is caused by a rhythm in the secretion of such substances, taking place only in the dark period, followed by a breakdown, not necessarily restricted to the light period as suggested above. An interesting question is to what extent transport of agglutination factor from the cellular pool via the flagella into the medium takes place. Agar-derived mt- gametes excrete large amounts of agglutinative membrane vesicles, so-called isoagglutinins (Homan, 1981), and these may be needed for the re-establishment of the flagellar agglutinability.

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