Effect of Naturally Collected Bacteria on Growth of *Uroglena americana*,
a Freshwater Red Tide Chrysophyceae

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It was demonstrated that the presence of bacteria was essential for the growth of *U. americana*, a freshwater red tide Chrysophyceae, in a laboratory culture system. The effect on the algal growth of bacteria collected from Lake Biwa during the bloom was studied. All of the bacteria tested supported the algal growth and the growth extent of the alga was dependent on the species of associated bacteria. The algal growth was also promoted with the addition of nonsterilized natural lake water. The possible role of bacteria for the algal growth in nature was discussed.

In order to elucidate a mechanism of outbrake of a freshwater red tide of *U. americana* in Lake Biwa, Ishida *et al.* conducted an algal assay for analysis of major nutrients limiting the algal growth in the lake. However, the culture of *U. americana* used in this study was not an axenic, but a monoxenic one which was accompanied by a strain of bacteria. The alga, first isolated in 1978 from Lake Biwa during its bloom, was successfully grown in a bacterized unialgal culture using chemically defined medium (Ur-1), but failed to grow in bacteria-free condition.

The present paper describes that the algal growth is dependent on the presence of an associated bacteria, and that all of bacterial strains collected from Lake Biwa during the algal bloom are effective on the algal growth.

**Materials and Methods**

*Isolation and Cultivation of *U. americana***

*U. americana* strain 78 was isolated in 1978 from Lake Biwa. The monoxenic culture of *U. americana*, which was accompanied by a bacterial strain NA-1 (*Vibrio-Aeromonas sp.*), was obtained by washing the algal colonies 10 to 15 times in petri-dishes containing 0.5 ml of sterilized Ur-1 medium with the help of capillary pipettes. Stock cultures of *U. americana* were maintained in sterile paper-plugged (New Steri-Plug; W. Germany) 100-ml Erlenmeyer flasks containing 40 ml of chemically defined medium Ur-1, and transferred at intervals of 2 weeks. The composition of Ur-1 medium is as follows (per l): NH₄NO₃, 5 mg; NaH₂PO₄, 1.7 mg; MgSO₄, 10 mg; CaCl₂, 10 mg; KCl, 1 mg; vitamin B₁, 10 μg; vitamin B₁₂, 0.1 μg; biotin, 0.1 μg; Provasoli’s PIV metal solution, 1 ml; pH 7.0.

**Growth Conditions and Measurements of *U. americana***

For all growth experiments, 0.1 ml of the subcultures incubated for one week was inoculated into triplicate sterile Pyrex test-tubes plugged with P.P caps containing 4 ml of Ur-1 medium. Conditions for incubation were as follows: 15°C and 6000 lx cool white or daylight fluorescent on a 14-h light/10-h dark cycle. The growth of the cultures was monitored by measuring *in vivo* chlorophyll fluorescence of a 4 ml sample tube. Measurements were made with a Turner Model 111 Fluorometer, modified for chlorophyll analysis.

**Experimental Procedure for Bacterial Effect on the Growth of *U. americana***

(1) Preparation of a temporary axenic culture of *U. americana*

To make *U. americana* culture axenic, 1.5 μg/ml of filter-sterilized erythromycin was added to Ur-1 medium. A bacterial strain NA-1 associated with *U. americana* did not grow in the presence of more than 0.19 μg (per ml) of erythromycin.
After inoculating the alga, viability of a bacterial strain NA-1 was examined by use of LT10\(^{-1}\) medium\(^3\) containing 0.5 g of trypticase peptone (BBL), 0.05 g of yeast extract (Difco), and 1 l of aged lake water.

(2) Stability test of the antibiotic in the algal culture

Stability of the antibiotic (erythromycin) in the culture was assayed by the agar diffusion method. *Bacillus subtilis* (erythromycin-sensitive) as a test organism was spread on agar plates and metal cups containing samples were placed on the surface. After 24 h of incubation at 30°C, the clear zones on the plates were measured.

(3) A culture system for a bacterial effect on the growth of *U. americana*.

To examine bacterial effects on the algal growth, one hundred seventy-seven bacterial strains were isolated from Lake Biwa during April 16 to June 18, 1981, and among them 68 erythromycin-resistant strains were selected for an experiment employing a temporary axenic culture treated with erythromycin.

However, there is a limitation that erythromycin-sensitive bacteria could not be tested in this system. We, therefore, prepared a new monoaxenic culture system, in which the bacterial strain NA-1 associated with *U. americana* in the stock culture was replaced by an extremely slower growing bacterium strain 2B-11. To this culture system, each 24 strains of bacteria isolated from Lake Biwa on Apr. 27, May 9, 13 and 25, and Jun. 23 (1983) were inoculated and the algal growth was assayed after 14 days of incubation.

(4) The effects of sterilized and nonsterilized lake waters on the growth of *U. americana*.

Lake water samples were collected from Lake Biwa during the period of initial phase (Apr. 27, 1983) to final phase (Jun. 23, 1983) of the algal bloom. The samples were filtered through a combusted glass fiber filter immediately after sampling water. A portion of filtrates was sterilized by autoclaving at 121°C for 15 min. For the growth response test, 0.1 ml of either samples was added to the test tubes containing 4 ml of Ur-1 medium. The cultures were incubated for 14 days at light.

(5) In situ populations of *U. americana* and bacteria.

Population density of *U. americana* in situ was determined by counting its colony number in 0.5 ml of each water samples using hemacytometers immediately after sampling waters.

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**Fig. 1.** Growth of *U. americana* in the presence and absence of associated bacteria. *U. americana* culture accompanied by an erythromycin-sensitive bacterium NA-1 (○) was added with 1.5 µg/ml of erythromycin (●) or erythromycin plus erythromycin-resistant bacterium NbA-1 (●).

**Fig. 2.** Stability of erythromycin added to *U. americana* culture after 12 days of incubation. *Bacillus subtilis* as a test organism was spread on an agar plate, and metal cups placed in the agar were filled with the algal cultures which had been incubated for 12 days with no addition (A), with an erythromycin-resistant bacterium NbA-1 (B), with 1.5 µg/ml of erythromycin (C), and with 1.5 µg/ml of erythromycin plus the bacterium NbA-1 (D). Note the same inhibition zone size in (D) as in (C).
The number of bacteria in samples was counted by the spread plate method with LT $10^{-1}$ medium after three weeks of incubation at 20°C.

Result

Growth Response of {	extit{U. americana}} to Bacteria in the Presence of Erythromycin

Chemically defined medium Ur-1 supported good growth of the monoxenic culture of {	extit{U. americana}} which was accompanied by an erythromycin-sensitive bacterial strain NA-1 (Vibrio-Aeromonas sp.) (Fig. 1). However, after being made axenic by erythromycin, the alga failed to grow (Fig. 1). The growth in the erythromycin-added culture was recovered when inoculated with an erythromycin-resistant bacterial strain NbA-1 (Fig. 1). The erythromycin-stability test employing the agar diffusion method demonstrated that erythromycin was not decomposed by the bacteria in this culture during the course of the experiments (Fig. 2). These results present evidence that the presence of bacteria is an essential factor for the growth of {	extit{U. americana}}.

Sixty-six strains among 68 erythromycin-resistant bacteria isolated from Lake Biwa in 1981 supported the growth of {	extit{U. americana}} when examined in a similar system with that shown in Fig. 1, and the extent of the algal growth was variable, depending on the species of associated bacteria. Fig. 3A shows two typical examples. The growth of {	extit{U. americana}} in a mixed culture with bacterial strain 3A-3 showed distinct increase over 7–10 days, whereas the alga in a mixed culture with bacterial strain 2B-11 increased very slowly over a period of 40 days and subsequently reached only 40–60% of the yield of the cultures accompanied by bacterial strain 3A-3. During

Fig. 3. (A) Growth of {	extit{U. americana}} and two erythromycin-resistant bacteria in mixed cultures. Conditions for inoculum and incubation were same as described in the legend to Fig. 1. Symbols: ○, {	extit{U. americana}} with a rapid growing bacterium 3A-3; △, the bacterium 3A-3 with with {	extit{U. americana}}; ●, {	extit{U. americana}} with a slow growing bacterium 2B-11; ▲, the bacterium 2B-11 with {	extit{U. americana}}. (B) Effect of the supplemental inoculation of a bacterium 3A-3 on the growth of {	extit{U. americana}} accompanied by the bacterium 2B-11. Arrow indicates the addition of the bacterium 3A-3. Symbols: ○, {	extit{U. americana}} with the bacterium 2B-11; ●, {	extit{U. americana}} with the bacterium 2B-11 and 3A-3.
The course of the experiments the number of bacterial strain 2B-11 remained remarkably low as compared with that of a bacterial strain 3A-3 (Fig. 3A). Subsequent transfer of these cultures yielded virtually identical results. The growth of *U. americana* accompanied by bacterium 2B-11 was enhanced by additionally inoculating a bacterium 3A-3 to the culture (Fig. 3B). The growth of *U. americana* accompanied by a bacterium 2B-11 was also made much more enhanced by adding a small amount of peptone to promote the bacterial growth (Fig. 4B) or by making the initial number of bacteria higher (ca. $10^6$ per ml) (Fig. 4C).

The Effect on the Algal Growth of Various Bacterial Strains Isolated from Lake Biwa during the Bloom (1983)

To examine the effects of various bacterial strains on the algal growth, it is disadvantageous to employ the temporary axenic culture with erythromycin as an assay system, because erythromycin-sensitive bacteria can not be tested in this system. One hundred twenty strains of bacteria isolated from Lake Biwa during the period of initial phase to final phase of the bloom of *U. americana* (1983) were therefore inoculated each to the algal culture accompanied by a slow growing bacterium 2B-11 without erythromycin and the algal growth was assayed. Results were summerized in Table 1. All of the bacteria except one enhanced the growth of *U. americana*. In preliminary experiments several bacteria inhibited the algal growth. These inhibitory effects appeared mostly due to excessive growth of bacteria, but by making inoculum size lower, most of these inhibitory effects disappeared.

**In Situ Relationships between U. americana and Bacteria**

Fig. 5 shows the population density of *U. americana* and bacteria during the algal bloom in Lake Biwa in 1983. Before and during the algal bloom the number of bacteria was less than $10^5$ per ml. At the end of the algal bloom, the number of bacteria rose to $10^6$ per ml.

Fig. 6 shows the results of the growth of *U. americana* accompanied by a bacterial strain 2B-11 in the addition of heat-sterilized and nonsterilized lake water to Ur-1 medium. In all instances the growth of *U. americana* was enhanced with the

![Fig. 4. Growth of *U. americana* (○) and bacterium 2B-11 (●) in the mixed culture system in Ur-1 medium. (A) no addition. (B) with 50 mg/l tritycise peptone. (C) with additional supplement of a washed suspension of the bacterium (the initial number of the bacterium $10^6$/ml).](image)

### Table 1. Growth response of *U. americana* in addition of various living bacteria. Bacterial strains collected from Lake Biwa during Apr. 27 and Jun. 23, 1983 were inoculated to *U. americana* culture accompanied by a slow growing bacterium 2B-11

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<th>Growth nonpromoted</th>
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* Growth of experimental cultures was compared with that of a control culture (no addition) after 14 days of incubation. Positive growth effects were considered to be those which gave more than 200% growth of the control culture (taken as 100%).
addition of nonsterilized lake water but not with sterilized lake water. In some instances, these experiments were repeated with filter sterilization, but virtually identical results were obtained.

Discussion

The inability of *U. americana* to grow without the presence of bacteria in culture at first seemed surprising. Although numerous workers have found that algae release many substances that could be potential nutrients for bacteria, there is little information that algal growth is dependent on the presence of bacterial associates in cultures. Lange observed that under condition of carbon dioxide limitation the addition of organic carbon stimulated the growth of blue green algae via carbon dioxide production by bacteria. Berland *et al.* observed that several species grew much better in bacterized than axenic cultures and suggested release of vitamin substances by bacteria as a likely explanation of algal stimulation. Several other reports have been made as to bacterial commitment for algal growth and physi...
the number of bacteria has never exceeded $10^6$ per ml in Lake Biwa (Fig. 5) suggests that the algal growth in this period might be controlled by an interaction with bacteria similar to that observed in our mixed cultures. However, it is not possible to estimate the precise role of bacteria in initiating the bloom because of the ubiquitousness of bacteria in the environment to support the algal growth (Fig. 6, Table 1). The problem, in near future, will be better elucidated by the study on the mechanism undergoing the algal-bacterial association. We will describe it in a following paper.

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