The Influence of Dissolved Organic Carbon on the Microbial Community Associated with *Tetraselmis striata* for Bio-Diesel Production

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Featured Application: The following topics are assessed in this study: (1) microbial disturbance due to dissolved organic carbon secreted from *T. striata*; (2) risk of large-scale microalgae cultivation to marine environment; (3) relationship between bacteria and phytoplankton by dissolved organic carbon; (4) ecosystem impact assessment through mesocosm study.

Abstract: The green alga *Tetraselmis striata* is regarded as a suitable candidate microalga for bio-diesel production. Recently, *T. striata* was cultured near Yeonghueung Island, Korea, in a “marine culturing field”; however, its environmental impacts are not yet studied. We estimated the amount of dissolved organic carbon (DOC) released from *T. striata* cultivation in the marine culturing field, and we investigated the changes in bacterial composition. Then, we designed and installed a mesocosm for further understanding. From the mesocosm results, the DOC released from the cultivation of *T. striata* led to changes in bacterial communities, disturbance of the microbial food web structure, rapid depletion of nutrients, and a decrease in dissolved oxygen (DO) and pH. Our novel work demonstrates that large amounts of DOC secreted by large-scale microalgal cultures such as that of *T. striata* can potentially have a significant impact on the structure and function of the surrounding microbial ecosystem.

Keywords: *Tetraselmis*; DOC; bacterial community; microbial disturbance; food web structure; mesocosm

1. Introduction

Over the past two decades, the cultivation of microalgae as a source of sustainable biomass for the production of biofuels gained much attention [1,2]. Marine microalgae cultivation is relatively low cost, has a low space requirement, and can utilize a natural source of nutrients for algal growth [3]. The green alga *Tetraselmis* sp. (Chlorodendraceae) is regarded as a suitable candidate microalga for...
bio-diesel production due to its high lipid content and fast growth [1,4,5]. As a consequence, various studies were carried out to search for the optimal growth conditions of this species [6–8] and to increase the lipid content using genetic, biotic, and environmental manipulations [9–13].

More recently, *Tetraselmis striata* was cultured near Yeonghueung Island, Korea, in a “marine culturing field”—a floating marine system which uses semi-permeable membranes for cost-effective mass cultivation [14,15] (Figure 1). This system also uses minimal power for temperature control, as well as a minimal supply of nutrients, and it requires insignificant mixing compared to conventional, large-scale algal culturing systems [16–19]. This large culturing facility, however, may pose adverse effects on the surrounding marine environment, many of which are yet to be investigated. Firstly, this large-scale marine culturing field may cause detrimental effects to the underlying benthic ecosystem by obstructing light usually transmitted through the water column [20–22]. Secondly, any damage to the culturing marine culturing field due to natural disasters such as typhoons or tsunamis may cause large quantities of microalgae to be released into the natural ecosystem. Finally, large quantities of dissolved organic matter (DOM) are continually released into the environment from this facility due to the use of semi-permeable membranes [23,24]. While reduced light transmission and natural disasters are localized events, the release of DOM into the coastal environment can be significant, resulting in coastal eutrophication and the potential for other microalgal blooms to occur including those that are harmful. The DOM released by phytoplankton is also a ubiquitous process, often resulting in 2%–50% of the carbon fixed by photosynthesis leaving the cell [25]. This supply of carbon is thought to be the main autochthonous source of dissolved organic carbon (DOC) into marine ecosystems and a major source of carbon for heterotrophic prokaryotes [25]. A subsequent change in the release of DOC into the marine environment may also cause a change in bacterial composition within the microbial ecosystem [26–28]. As these bacterial communities form the basis of microbial food webs, playing a significant role in marine biogeochemical cycling and energy transfer, any change in their community structure is likely to also affect organisms at higher trophic levels [29,30]. Indeed, recent research by Wang et al. (2017) [31] showed that the DOC measured near Yeongheung Island appeared to be an important factor affecting the abundance and composition of bacteria and phytoplankton in this area.

At present, the size of the marine culturing field near Yeonghueung Island, Korea, gradually expanded to 1 ha (10,000 m²) with the capacity to cultivate 280 tons of *T. striata*. Whilst the size of the marine culturing field gradually expanded, the environmental impacts are yet to be studied. It is hypothesized that a large amount of DOC entering the region may change the composition of bacteria, and a sudden bloom of *T. striata* may inhibit the growth of other competitive organisms and/or affect organisms at higher or lower trophic levels. With this in mind, we firstly estimated amount of DOC released from *T. striata* culture when the marine culturing field was in operation, and we compared the bacterial composition inside and outside of the semi-permeable membrane in which *T. striata* was cultured. After confirming the effect of DOC on the bacterial community composition, a mesocosm experiment was then designed and carried out to gain a more in-depth understanding of the influence of DOC release from this *T. striata* cultivation on the surrounding water column physicochemical properties, bacterial community, and microbial food web structure.
Figure 1. National (A) and regional (B) locations where the “marine culturing field” is located, viewed on a Google Map (C) (http://map.google.com; 37°14′00.3″ north (N) 126°25′50.6″ east (E)). The view of the operating marine culturing field (D), and the installed cultivation pond made using semi-permeable membrane (E).

2. Materials and Methods

2.1. Tetraselmis striata Strain and Culturing

The Tetraselmis striata strain (KCTC12432BP) used in this study was obtained from the Department of Biological Engineering, Inha University, Incheon, South Korea, and maintained under axenic conditions [28] to exclude any phycosperic and/or symbiont bacterial effect. Prior to the experiment beginning, the strain of T. striata was checked for any bacterial contamination using 4′,6-diamidino-2-phenylindole (DAPI) staining and observation using a fluorescence microscope (BX51, Olympus, Japan) [28]. Ten liters of T. striata stock culture was maintained using f/2 media [32] at 20 °C exposed to cool-white fluorescent lamps of 50 mmol photons·m⁻²·s⁻¹ under a 12-h/12-h light/dark photoperiod cycles until the experiment began.

2.2. Sampling of Marine Culturing Field

Culturing of T. striata for bio-diesel biomass was carried out from May to November 2016 using 30 floating cultivations, each 3 m wide and 10 m in length, made using semi-permeable membranes with a total holding capacity of nine tons of seawater (Figure 1). The ponds were initially inoculated...
with *T. striata* at a cell density of $9.1 \times 10^4$ cells·mL$^{-1}$. From the surface of one randomly selected pond, a 1.0-L sample of *T. striata* was sampled on days 0, 1, 2, 3, 5, 7, and 9 to measure the changes in cell density of *T. striata* and the concentration of DOC over time (Section 2.4). From each of these samples, a 50-mL subsample was preserved with 5% Lugol’s solution [33] and returned to the laboratory for cell enumeration. A Sedgewick–Rafter counting chamber (Rigosha Co., Tokyo, Japan) was used to estimate the number of *T. striata* using light microscopy (Axioplan, Zeiss, Jena, Germany) at 400× magnification. Another set of 0.5-L samples were collected on days 0, 3, and 9 to analyze the bacterial composition of the cultivation pond (Section 2.5). Finally, a 1.0-L sample of seawater was collected from the surface approximately three meters away from the marine culturing field for comparison to the DOC and bacterial composition.

### 2.3. Mesocosm Design and Sampling

Four mesocosms (each one ton) were set up to evaluate the impacts of *T. striata* cultivation on the surrounding bacterial community and the microbial ecosystem—see below (Figure 2). Mesocosms were floated by buoys and tied to the pier using ropes (Figure 2B). Each mesocosm was filled with one ton of natural seawater and then left for two days for acclimation. A 20-L bioreactor containing *T. striata* enclosed by a semi-permeable membrane was then installed into two of the mesocosms and secured with ropes. These are termed “bioreactor installed mesocosms” (BMs) for the duration of the study. The other two adjacent mesocosms had no bioreactors installed, and they are termed “control mesocosms” (CMs) (Figure 2A,C,D). All mesocosms (BMs and CMs) were then spiked with 30 µM nitrate and 2 µM phosphate to induce sufficient growth of all biota living in mesocosms. On days 0, 2, 3, 4, 5, 6, 7, 9, 11, 13, and 15, four liters of seawater was subsampled from each mesocosm for DOC concentration, as well as various biotic factors (bacterial abundance and composition, abundances of heterotrophic nano-flagellates (HNF), phytoplankton, ciliates, and zooplankton, and the nutrients ammonium, nitrate, nitrite, phosphate and silicate) (see below). On each day, a 50-mL sample of *T. striata* was also collected using a suction flask through tubes which connected to the faucet of each bioreactor (Figure 2C,E). This water sample was subsequently preserved with 5% Lugol’s solution [33] and returned to the laboratory for cell enumeration. Water temperature (°C), salinity, pH, and dissolved oxygen (DO, mg·L$^{-1}$) were also measured within and adjacent to (3 m away) mesocosms using an YSI 6600 sonde (YSI inc., Marion, MA, USA).

### 2.4. DOC Analysis

To analyze the total concentration of DOC, seawater samples were collected by passing 50 mL of the filtrate through a pre-combusted (400 °C for 8 h) GF/F filter (Whatman, Middlesex, U.K.) in a pre-combusted (400 °C for 8 h) total organic carbon (TOC) vial (Shimadzu, Kyoto, Japan) and fixed with 2% H$_2$PO$_4$ (final concentration) to protect the total amount of DOC from any further biological activity. The samples were then stored at 4 °C until measured using a TOC Analyzer (Shimadzu, Kyoto, Japan). To characterize the DOC, 0.5-L subsamples were filtered with a 47-mm-diameter GF/F filter (Whatman, Middlesex, U.K.) and the filtrates were stored at −80 °C in the dark until further analysis. Synchronous fluorescence spectra of the DOC were estimated using a luminescence fluorometer (FS-55, Scinco, Korea) with a constant offset (∆λ = 30 nm), based on the protocol of Hur et al. (2008) [34]. Four regions were allocated to the integrated areas of the fluorescence intensities: 250–300 nm for protein-like fluorescence (PLF); 300–380 nm for fulvic-like fluorescence (FLF); 380–420 nm for humic-like fluorescence (HLF); 420–600 nm for terrestrial humic-like fluorescence (THLF) [35]. For discrete data, integration was made by the summation of the fluorescence intensities at the interval of 1 nm. Each relative fluorescence region was then calculated over the whole fluorescence intensity as %PLF, %FLF, %HLF, and %THLF, respectively.
2.5. Bacterial Abundance and Composition

To determine the abundance of bacteria in each water sample collected, a 10-mL subsample was preserved with 2% glutaraldehyde final concentration (Sigma-Aldrich, St. Louis, MO, USA) and stored in the dark at room temperature until further processing. Preserved samples were then filtered using a 0.2-µm-pore GTTP Millipore filter membrane (Millipore Filter Corporation, Cork, Ireland) and stored using the DAPI method [36]. The bacterial cells were enumerated using an Olympus epifluorescence microscope (XC10, Olympus, Tokyo, Japan) under 1000× magnification.

Next-generation sequencing (NGS) was used to analyze the bacterial community. For DNA extraction, a 0.5-L seawater sample obtained from each experiment was filtered onto a 0.2-µm-pore and 47-mm-diameter ISOPORE membrane filter (MILLIPORE, Cork, Ireland). Loaded filters were placed in a 2-mL micro-tube (Axygen Sciences, CA, USA) which contained 800 mL of extraction buffer (100 mM Tris-HCl, 100 mM Ethylenediaminetetraacetic Acid Disodium Salt (Na2-EDTA), 100 mM sodium phosphate, 1.5 M NaCl, and 1% Cetrimonium bromide (CTAB)) and stored at −80 °C until further processing. DNA extraction was performed following the EX DNA extraction protocol [37], and an exploration of the bacteria community was targeted using primers 27F (5′−AGA GTT TGA TAA GAC TCA TAA GTC TA−3′) and 519R (5′−GWA TTA CCG CGG CKG CTG−3′) [38,39], for the V3–V4 region of 16S ribosomal RNA (rRNA) [40]. High-throughput sequencing was performed using the Illumina Miseq platform (Molecular Research LP; Shallowater, TX, USA) following the manufacturer’s guidelines. Clustering analysis was performed on the sequence data using the Bray–Curtis distance and Ward’s method [41] with the cluster package in R (version 1.14.3, R-studio). The relationship between the bacterial communities and the DOC was then assessed using redundancy analysis (RDA) in the CANOCO 5.0 software package [42] in BMs.
2.6. Preservation and Enumeration of HNFs, Phytoplankton, Ciliates, and Zooplankton

For enumeration of the HNFs, a 10-mL sample was filtered onto a 1.0-µm-pore Nuclepore filter (Japan KK, Tokyo, Japan) stained with Nigrosin, and the HNFs were stained with fluorescein isothiocyanate (FITC). The filters were observed under blue excitation (resulting in green fluorescence). One hundred HNFs were enumerated per sample. For phytoplankton, ciliate, and *T. striata* enumeration, a 1.0-L subsample was preserved with 0.5% Lugol’s solution [43], concentrated to approximately 50 mL by decanting the supernatant [33] and counted using a Sedgewick–Rafter counting chamber (Rigosha Co., Tokyo, Japan) and light microscopy (Axioplan, Zeiss, Jena, Germany) at 400× magnification. Morphologically distinct phytoplankton cells were identified to species or genus level [44]. Zooplankton were sorted using a 60-µm-pore sieve from the 1-L subsample and collected into a 50-mL conical tube (SPL, Life Sciences, Pocheon, Korea) using filtered seawater. Zooplankton samples were subsequently preserved in 5% formalin, and individuals were counted using a zooplankton counting chamber under a dissecting microscope (Carl Zeiss, Jena, Germany) [45].

2.7. Nutrients Analyses

To analyze the concentration of inorganic nutrients, a 0.5-L subsample collected from each day was immediately filtered with a 47-mm-diameter GF/F filter (Whatman, Middlesex, U.K.) and placed in acid-cleaned polyethylene bottles. The filtered seawater was stored at −20 °C in the dark until laboratory analysis. Ammonia (NH$_4^+$), nitrate + nitrite (NO$_x^+$), phosphate (PO$_4^{3-}$), and silicate (Si$^-\) concentrations (µM) were determined in the laboratory using a flow injection autoanalyzer (QuikChem 8000; Lachat Instruments, Loveland, CO, USA). All nutrient concentrations were calibrated using standard brine solutions (CSK Standard Solutions; Wako Pure Chemical Industries, Osaka, Japan) with the detection limits of DIP (0.02 µM), DIN (0.05 µM), and DSi (0.1 µM).

3. Results

3.1. DOC and Bacteria in Cultivation Pond

The dynamics of *T. striata* cell density and DOC concentration in the cultivation pond sampled are shown in Figure 3. The cell density of *T. striata* increased after initial inoculation of *T. striata* with the maximum cell density (1.8 × 10$^5$ cells·mL$^{-1}$) measured on day 5, and then gradually decreased until day 9. The DOC concentration rapidly increased from day 2 with the maximum DOC concentration (428.7 mg·L$^{-1}$) measured on day 3 and then a gradual decrease until day 7. The DOC concentration within the natural seawater also increased and decreased from day 2 to day 7, but its maximum concentration (24.7 mg·L$^{-1}$) was significantly lower than that of the samples from the cultivation pond.

The bacterial community within the cultivation pond was substantially different to the natural seawater community (Figure 4). The Planctomycetaceae bacterial group dominated at day 0 and maintained its dominance (29.4%) in natural seawater until day 9. On the other hand, the Rhodobacteraceae and Saprospiraceae dominated the cultivation pond on day 3 and day 9. The maximum percentage contributions of the Rhodobacteraceae group and Saprospiraceae group were 37.1% and 26.5%, respectively, on day 3 in the cultivation pond.
The bacterial community within the cultivation pond was substantially different to the natural seawater community (Figure 4). The Planctomycetaceae bacterial group dominated at day 0 and maintained its dominance (29.4%) in natural seawater until day 9. On the other hand, the Rhodobacteraceae and Saprospiraceae dominated the cultivation pond on day 3 and day 9. The maximum percentage contributions of the Rhodobacteraceae group and Saprospiraceae group were 37.1% and 26.5%, respectively, on day 3 in the cultivation pond.

Figure 3. Dynamics of abundance of *Tetraselmis striata* cells cultured in cultivation pond, as well as concentrations of dissolved organic carbon (DOC) in cultivation pond and near (3 m away) seawater for nine days.

Figure 4. Changes in bacterial composition in cultivation pond and near (3 m away) seawater for nine days.
3.2. Environmental Factors during Mesocosm Experiment

The seawater temperature gradually decreased from 22.0 °C to 18.8 °C during the mesocosm experiment (Figure 5A). Salinity gradually increased from 29.2 to 29.9 in the BMs and CMs, while salinity reached 30.7 on day 15 in the natural seawater (Figure 5B). The concentration of DO rapidly decreased after installation of the *T. striata* bioreactor starting at 0.47 mg·L⁻¹ on day 2 and remained low until the end of experiment (Figure 5C). The DO in the CMs, however, varied from 3.32 to 5.86 mg·L⁻¹, while the natural seawater varied from 5.35 to 7.89 mg·L⁻¹ (Figure 5C). Fluctuating patterns of DO in the BMs and natural seawater were similar (Figure 5C). The pH of the BMs gradually decreased from 7.80 to 7.06 on day 9, although the low pH values of the CMs (7.40) and the natural seawater (7.50), as measured on day 9, were relatively high compared to the BM pH of 7.12 (Figure 5D).

![Figure 5](image-url)

*Figure 5.* Changes in environmental factors: water temperature (A), salinity (B), dissolved oxygen (C), and pH (D) of each mesocosm and near seawater during mesocosm experiment period (15 days). Each value of white and dark circles was averaged using two measured values of the two bioreactor installed mesocosms or control mesocosms, respectively, but values of white squares were measured one time.

3.3. Abundance of *T. striata* and DOC Concentration

The initial mean cell density of *T. striata* in each bioreactor was 6343 ± 696 cells·mL⁻¹ at day 0 (Figure 6). Cell density then gradually increased on day 4, then rapidly increased until day 6 (maximum cell density 149,550 ± 18,847 cells·mL⁻¹). Cell abundance then rapidly decreased to 11,670 ± 1232 cells·mL⁻¹ on day 9 (Figure 6). The lowest cell density (1250 cells·mL⁻¹) was measured at the end of the mesocosm experiment on day 15. The DOC concentration of natural seawater was 22.7 mg·L⁻¹ two days prior to the bioreactor installation, and it gradually decreased until day 0 for the CMs and day 1 for the BMs. The DOC concentration within the BMs gradually increased until day 4, when it noticeably increased until day 6 (Figure 6). DOC then rapidly decreased to 6.74 mg·L⁻¹ in one day and maintained its low concentration until the end of the mesocosm period (3.50 mg·L⁻¹). The maximum DOC concentration within the BMs was 74.1 mg·L⁻¹ on day 6. The DOC concentration with the CMs remained lower than that of BMs across the entire experiment, with its maximum concentration measured on day 13 as 5.38 mg·L⁻¹. The relative proportion of FLF was higher than the other fluorescence regions of DOC (PLF, HLF, and THLF) in both mesocosms (Figure 7). The proportion of FLF varied from 57.7% to 59.6% in the BMs and from 49.3% to 58.6% in the CMs. Increments and
The bacterial abundance within the BMs rapidly increased from day 0 (2.24 × 10^6 cells·L⁻¹) to day 2 (1.04 × 10^7 cells·L⁻¹) after the bioreactor installation. It then decreased until the end of the mesocosm experiment and finished on day 15 at 2.97 × 10^6 cells·L⁻¹ (Figure 8A). The bacterial abundance within the CMs increased from day 1 to day 4 reaching 9.58 × 10^6 cells·L⁻¹ (Figure 8A). At the end of the mesocosm experiment, the CMs contained a similar bacterial abundance (1.09 × 10^7 cells·L⁻¹) compared to the BMs, but the highest abundance (1.09 × 10^7 cells·L⁻¹) was reached on day 13. Cluster
analysis revealed five clades (A–E) of bacteria across all sampling days and both mesocosms (Figure 9). The bacteria grouped in the same clade (B) until day 0, separated into two clades (D and E) on day 4, and separated into another two clades (C and E) on day 9. By day 15, the bacterial communities of BMs were separated into clade A alone.

In BMs, an Alteromonadaceae group dominated on day 1, but its proportion gradually decreased throughout the experiment. Alternatively, the Flavobacteriaceae and Oceanospirillaceae groups increased from day 4 to 9, a Cellvibrionaceae group increased from day 9, and a Rhodobacteraceae group gradually increased throughout the experiment. In the CMs, Alteromonadaceae and Oceanospirillaceae groups dominated at day 1. The proportion of the Alteromonadaceae group decreased, but that of the Flavobacteriaceae group noticeably increased from days 4 to 6 (Figure 9). The proportions of ect. bacterial groups (<2%), which dominated two days prior to the beginning of the experiment and rapidly decreased in both mesocosms, but increased again in the CMs from days 9 to 15. At the end of mesocosm experiment, the Shannon diversity index decreased to 4.8 and 4.1 in each BM, but relatively higher diversities (6.0) were observed in the CMs compared to the BMs. ANOVA analysis revealed that the bacterial diversity of clade A (day 15 of BMs) was significantly ($p < 0.05$) lower than that of clades B and C.

Figure 7. Total concentrations of dissolved organic carbon (DOC) and relative ratios of four fractions of DOC (PLF, FLF, HLF and THLF), and its changes of each day in bio-reactor installed mesocosm and control mesocosm. Grey bars indicate the relative DOC concentrations, each ratio of orange, green, violet and yellow bar indicates the ratios of four fractions, respectively, and changed amounts of ratio of each four DOC fractions are indicated as blue (plus) and red (minus) bars, respectively.

Figure 8. Dynamics of abundance of biotic factors: bacteria (A), heterotrophic nano-flagellates (B), phytoplankton (C), ciliate (D), and zooplankton (E) during mesocosm experiment period (15 days). Each value of white and dark circles was averaged using two measured values of bioreactor installed mesocosms or control mesocosms, respectively.
3.5. Relationship between Bacterial Community and DOC

The RDA showed a significant relationship ($p < 0.05$) between the amount of DOC and relative amounts of the four DOC fractions, as well as the occurrence of family-level bacterial groups in the BMs (Figure 10). The concentration of DOC was strongly correlated to *T. striata* cell density and the *Colwelliacae* group. The concentration of PLF also showed a strong relationship with two distinct bacterial groups—the Rhodobacteraceae and the Flavobacteriaceae. RDA revealed that the majority of the variance was explained by axis 1 (66.9%) and axis 2 (22.2%).

3.6. Other Biotic Factors

The increasing and decreasing patterns of HNFs in both mesocosms were similar, but the CMs showed higher HNF abundances than those within the BMs (Figure 8B). The maximum abundance of HNF in the CMs was 3551 cells·mL$^{-1}$ on day 5 and 2630 cells·mL$^{-1}$ on day 4 for the BMs. The abundances of HNFs in both mesocosms rapidly decreased from day 7 until a final abundance of 21 (BM) and 22 (CM) cells·mL$^{-1}$ at the end of the mesocosm experiment (Figure 8B). The CM phytoplankton abundance gradually increased from day 1 until its highest abundance on day 11 (2185 cells·mL$^{-1}$) (Figure 8C). On the other hand, the phytoplankton within the BMs gradually decreased from day 1, and very few phytoplankton cells were observed from day 3 to the end of the mesocosm experiment (Figure 8C). The most abundant species throughout the experiment within the CMs was *Chaetoceros* spp. (data not shown). The abundance patterns of ciliates within both mesocosms were similar to...
those of phytoplankton (Figure 8D). The maximum ciliate abundance within the CMs was recorded as $1.21 \times 10^5$ cells·mL$^{-1}$ on day 11, but that in the BMs decreased from day 1, with most dead by day 3 (Figure 8D). Zooplankton abundance in both mesocosms was similar until day 2, whereby the abundance in the BMs rapidly decreased from day 2, with none observed by day 9 (Figure 8E). In the case of the CMs, zooplankton were observed until day 9 and then rapidly increased from then until day 13 (Figure 8E). The maximum zooplankton abundance within the CMs was $2.56 \times 10^5$ individual·ton$^{-1}$ on day 13.

Figure 10. The redundancy analysis to determine the relationship between dissolved organic carbon (DOC) including four different fractions (PLF, protein-like fluorescence; FLF, fulvic-like fluorescence; HLF, humic-like fluorescence; THLF, terrestrial humic-like fluorescence) and bacterial groups at the family level.

3.7. Nutrient Dynamics

Concentrations of inorganic nutrient in the surrounding seawater two days prior to the mesocosm experiment were 12.7 µM for NOx, 9.6 µM for ammonium, 1.1 µM for phosphate, and 32.8 µM for silicate (Figure 11). Concentrations of NOx and phosphate then increased on day 0 to 40.1 (BMs) and 41.7 µM (CMs) for NOx, and 2.5 (BMs) and 2.7 µM (CMs) for phosphate, due to the additional inoculations of NaNO$_3$ and NaH$_2$PO$_4$ (Figure 11A,C). Except for the silicate within the BMs,
the concentrations of nutrients gradually decreased during the mesocosm period, but decreased more rapidly in the BMs as compared to the CMs (Figure 11A,C). The patterns of ammonium concentrations of both mesocosms were similar throughout the experiment (Figure 11B). The concentration of silicate within the BMs remained steady, but rapidly decreased from day 7 within the CMs (Figure 11D).

Figure 11. Changes in inorganic nutrient concentrations: nitrate + nitrite (A), ammonium (B), phosphate (C), and silicate (D) during mesocosm experiment period (15 days). Each value of white and dark circles was averaged by two measured values of bioreactor installed mesocosms or control mesocosms, respectively.

4. Discussion

4.1. DOC Released from T. striata

Dissolved organic carbon (DOC) in the ocean is one of the largest carbon pools on Earth [46]. The extracellular release of recently fixed photosynthate is the major DOC production process in the marine ecosystem [47]. Our study revealed significant correlations between the cell density of T. striata and DOC concentration in both the cultivation pond and the mesocosm experiments. When the growth rate of T. striata was high, the amount of DOC secreted was high. The highest T. striata cell densities were observed on day 5 (cultivation pond) and day 6 (mesocosms). Although the high cell densities were still observed on day 7 in both the cultivation pond and the mesocosms, the DOC concentrations rapidly decreased. Phytoplankton usually sequester excess carbon in the form of storage polysaccharides, such as chrysolaminarin in diatoms [48,49] and glycogen in cyanobacteria [50]. These polymers also constitute the main fraction of phytoplankton exudates. According to Aluwihare and Repeta (1999), up to 37% of the DOC originating from phytoplankton consists of high-molecular-weight (>1 kDa) components, mainly hydrolyzable carbohydrates. The polysaccharides represent the largest fraction of high-molecular-weight DOC, with around 50% relative amount of the DOC in surface waters and around 25% in deep waters [51]. Various studies from the North Atlantic also support this large polysaccharide component with samples from the North Atlantic (<11% of total DOC) [52], the Norwegian Sea (<14.8%) [53], and the North Pacific (<10.1%) [54].

In aquatic systems, bottom-up (inorganic and organic nutrient supply) factors significantly influence bacterial succession. The DOC in marine ecosystems is almost completely cycled by bacterial groups over different timescales depending on different biochemical characteristics of the water column. The DOC pool constitutes the so-called “labile” fraction and turns over rapidly (hours to days).
The “semi-labile” fraction is cycled by bacteria on time scales of weeks to months [55], while the so called “refractory” DOC pool is biologically cycled on time scales ranging from centuries to millennia [55,56]. Therefore, in order to understand the close variation of the bacterial community, it is necessary to examine the differing fractions of DOC. Hur et al. (2011) [35] proposed that the methodology of DOC be divided into four fractions to characterize the DOC depending on the varying biological component, as well as temporal and differing regional sources. In addition, the proportion of PLF measured from phytoplankton was higher (73.9%) than the other organic matters (leaf litter, compost, and paddy water) in their experiment. In our study, a high concentration of DOC (74.1 mg·L$^{-1}$) and a significant correlation between the amount of DOC released from $T.\ striata$ and the proportion of PLF was observed in our mesocosm study; however, the PLF fraction was relatively lower than the other DOC fractions during the mesocosm experiment. Although the amount of PLF fraction was small, its changes were significantly correlated with the total amount of DOC and biomass of $T.\ striata$. This may be explained by the fact that the release of compounds from phytoplankton is susceptible to rapid uptake by heterotrophic bacteria [57]. This microbial degradation can induce the dramatic decrease of PLF and the enhancement of the FLF, the HLF, and the THLF fractions at the same DOC concentration [35].

4.2. Changes in Bacterial Community in Cultivation Pond and Mesocosms

The bacterial community within the cultivation pond was significantly different to that observed in the natural seawater during our experiment. A noticeable increase in the ratio of Rhodobacteraceae was recorded; however, these results do not consider the attached and/or free-living bacteria. It is, therefore, difficult to determine the direct effect of DOC released from $T.\ striata$ on the bacterial community as a whole. The $T.\ striata$ strain employed in our experiments was axenic, which may have introduced bacteria into the cultivation pond upon inoculation. It is difficult to be certain, therefore, whether a large proportion of the Rhodobacteraceae and Saprospiraceae bacterial groups was present at the time of inoculation or naturally increased, but the bacterial species diversity certainly decreased over the experimental duration. Despite the experimental set-up being a “semi-natural” condition, results from the mesocosm experiment suggest that there was an effect on the bacterial community from the DOC release. Evidence to support this include the following: the bacterial compositions of CMs and BMs were clearly differentiated at the end of the experiment, with a significantly decreased bacterial species diversity; the initial bacterial diversity in both mesocosms and the CMs on day 15 were high, but the diversity of the BMs on day 15 was significantly ($p < 0.05$) lower; in the BMs, the Cellvibrionaceae group occupied a large proportion, while the uncultured alpha and gamma Proteobacterium and etc. groups noticeably decreased at the end of mesocosm experiment (Figure 9). An increase in bacterial groups which favor the DOC artificially increased by the large cultivation of $T.\ striata$ would have resulted in a reduction of the natural bacterial group.

The Rhodobacteraceae group increased in the $T.\ striata$ cultivation pond. Interestingly, this bacterial group also gradually increased in the mesocosm experiment after the installation of the $T.\ striata$ bioreactor. The most relevant bacterial group was the Rhodobacteraceae for the PLF fraction in the analysis, followed by the Flavobacteriaceae (Figure 10). Many previous studies reported that the Rhodobacteraceae group is closely correlated with phytoplankton blooms, for example, blooms of Akashiwo sanguinea, Alexandrium spp., Gymnodinium catenatum, Karenia mikimotoi, Prorocentrum lima, and Leptocylindrus spp. [58–64]. Interestingly, most of the Rhodobacteraceae bacteria (90%) contain the vitamin B12 synthesis pathway, which is suggested to help in the growth of phytoplankton [65]. Furthermore, the Flavobacteriaceae are chemo-organotrophic [66,67], and strains were shown to synthesize proteins that actively bind and exploit algal polymeric substrates, such as carbohydrates, polypeptides, and lipids [68]. This bacterial group can be a dominant group in particle-attached communities [69] and are also closely associated with phytoplankton blooms [59,70]. Another bacterial group, the Colwelliaceae, although not strongly correlated with the PLF fraction in our study, is highly associated with the total amount of DOC. The genus Colwellia, the type genus in the Colwelliaceae,
only found in oceanic environments, is known as polysaccharide-degrading bacteria [71]. It is also a classic marine secondary producer and a key player in the global carbon cycle. Colwelliaceae are involved in decomposing organic material such as hydrocarbons, lipids, proteins, and polysaccharides within particulates of the pelagic zone and sea ice, and within algal, faunal, and floral associations, where it engaged in anaerobic respiration and/or fermenting [72]. The Rhodobacteraceae and Flavobacteriaceae groups, therefore, seem to be related to the growth of T. striata, and the Colwelliaceae group seems to be related to the degradation or termination of T. striata or the other phytoplankton cells in the BMs. Specific bacterial–algal associations are not cultural artificial effects, but relevant and important relationships in natural field populations. Further investigations into the specific functions of bacterial and algal metabolites, as well as transcriptomic studies, could provide a clue to the clarification of the relationship between these bacteria and phytoplankton groups.

4.3. Microbial Food Web Structure

Installed mesocosms maintained an environment similar to the natural condition during experiments. There was little difference between the surrounding water temperature and salinity compared to that measured within the mesocosms. The DO and pH, however, varied slightly between the natural seawater and the mesocosms, but the pattern of these variables over time remained the same in each. The abrupt dissipation of phytoplankton communities could lead to a decrease in DO and pH in the BMs. The major differences in biological factors between the CMs and BMs was the abrupt decrease in the abundance of phytoplankton, followed by the ciliate and zooplankton of the BMs. The abundance of phytoplankton began to decrease on day 1 of T. striata cultivation, while the ciliates began to decrease over the following two days and the zooplankton over the following three days. The initial decrease in phytoplankton coincided with a rapid decrease in n and p after T. striata inoculation. Especially, the mechanism of “luxury uptake” is known to occur when phytoplankton including Tetraselmis rapidly uptake and store excess phosphorus as inorganic polyphosphate granules [73–76]. In contrast, silicate rapidly decreased in the CMs after day 3 due to the dominance of the diatom Chaetoceros spp. Since inorganic nutrients are continuously supplied in natural marine environments, it is difficult to partition the rapid nutrient uptake of T. striata and its effect on the growth of other phytoplankton. Another possible influence on phytoplankton growth/succession patterns could be due to allelopathy and algicidal bacteria. Although this was not the focus of the present study, it is known that many phytoplankton species are able to secrete an allelopathic substance that can inhibit the growth of other organisms [77]. A rapid increase in DOC was associated with the growth of T. striata, and the possibility of the presence of a large amount of allelopathic material could be a possibility to explain this increase. On day 1 when growth of phytoplankton began to decrease, the proportion of the Alteromonadaceae bacterial group increased noticeably. The dominant genus group was Alteromonas. Many species of Alteromonas show an algicidal effect on various phytoplankton communities such as dinoflagellate, raphidophyte, and diatom, even working on phytoplankton species of freshwater [78–83].

The decrease in phytoplankton was followed by a decrease in the population of zooplankton and ciliates, which are higher trophic levels in nature. There may, however, be an alternate reason in our study. The cultivation of T. striata using semi-permeable membranes may have physically inhibited the transfer of energy from phytoplankton to higher trophic levels. Bacterial-sized organisms are able to migrate, but the semi-permeable membrane obstructed the direct interaction of phytoplankton (T. striata) with ciliates and zooplankton, such that activities including feeding and decomposition were disabled. This may have resulted in an imbalanced trophic structure within the BMs, whereby the higher trophic levels were removed (or extremely reduced).

5. Conclusions

Our novel work demonstrates that large amounts of DOC secreted by large-scale microalgal cultures such as that of T. striata can potentially have a significant impact on the structure and function of the surrounding microbial ecosystem. From our experimental results, DOC released from the
cultivation of *T. striata* led to (i) changes in bacterial communities, (ii) disturbance of the microbial food web and trophic structure, and (iii) a rapid depletion of nutrients and a decrease in DO and pH. An opened or semi-opened mesocosm is needed to better reflect the natural ecosystem, and a steady monitoring study around the marine culturing field should be done in parallel. In addition, more detailed molecular analysis is required to understand the interaction between specific phytoplankton with the surrounding bacterial community. Recent “omics” approaches allow for better insight into the mutual interactions between phytoplankton and heterotrophic bacteria, with metagenomes revealing partnerships and transcripts that resolve the different physiological processes executed by the different partners. Proteomes may further reveal species-specific exudates in multispecies assemblages, as well as the response to the presence of specific bacteria.

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