Proteorhodopsin (PR) is a photoprotein that functions as a light-driven proton pump in diverse marine Bacteria and Archaea. Recent studies have suggested that PR may enhance both growth rate and yield in some flavobacteria when grown under nutrient limiting conditions in the light. The direct involvement of PR, and the metabolic details enabling light-stimulated growth, however, remain uncertain.

Here, we surveyed transcriptional and growth responses of a PR-containing marine flavobacterium during carbon-limited growth in the light and the dark. As previously reported (1), Dokdonia strain MED134 exhibited light-enhanced growth rates and cell yields under low carbon growth conditions. Inhibition of retinal biosynthesis abolished the light-stimulated growth response, supporting a direct role for retinal-bound PR in light enhanced growth. Among protein-coding transcripts, both PR and retinal biosynthetic enzymes showed significant upregulation in the light. Other light-associated proteins, including bacterial cryptochrome and DNA photolyase, were also expressed at significantly higher levels in the light. Membrane transporters for Na+/phosphate and Na+/alanine symporters, and the Na+-translocating NADH-quinone oxidoreductase (NQR) linked electron transport chain, were also significantly upregulated in the light (2).

Culture experiments using a specific inhibitor of Na+-translocating NQR indicated that sodium pumping via NQR is a critical metabolic process in the light-stimulated growth of MED134. In total, the results suggested the importance of both the PR-enabled, light-driven proton gradient, as well as the generation of a Na+ ion gradient, as essential components for light-enhanced growth in these flavobacteria (2).


Keywords: marine bacteria, proteorhodopsin, proton pump, photoheterotrophy, transcriptomics
Photoheterotrophy of marine aerobic anoxygenic phototrophic bacteria

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Aerobic anoxygenic phototrophic bacteria (AAnPB) containing the photosynthetic pigment bacteriochlorophyll (BChl) \textit{a} can grow phototrophically and/or heterotrophically. Therefore, their metabolic performance is called photoheterotrophy. Recently, AAnPB and other photoheterotrophs including proteorhodopsin-containing bacteria and cyanobacterium \textit{Prochlorococcus} have been classified into a new functional group in terms of energy acquisition. Although it has become clear the ubiquitous distributions of AAnPB in the upper oceans with their high spatiotemporal variations, what controls their population dynamics is still an open question. Based on the intriguing AAnPB ecophysiological characteristics, there is a hypothesis that their photoheterotrophy could be beneficial in nutrient-poor environments such as oligotrophic oceanic waters. However, this hypothesis has not yet been experimentally verified well. Therefore, we investigated the photosynthetic responses of a coastal \textit{Roseobacter} strain OBYS 0001 of marine AAnPB to an organic substrate limitation. In the batch cultures at 20°C, the growth curve and cellular BChl \textit{a} concentration of the substrate-limited strain grown in 1/100 ZoBell 2216E medium kept constant, while those in the undiluted medium increased with time. Variable BChl \textit{a} fluorescence measurements revealed that changes over time in the functional absorption cross-section (sigma) of the photosystem for the strain were little distinct between the two conditions. However, the maximum photochemical quantum efficiencies (\(F_{v}/F_{m}\)) of the photosystem under the substrate-limited condition were significantly higher than those in the substrate-rich circumstance. These results suggested that AAnPB can enhance their photosynthetic activity with increasing the photochemical conversion efficiency without changing their antenna size under organic substrate limitations. In this presentation, we would like to emphasize the significance of photoheterotrophy for AAnPB in the oceans using our latest results and the past literatures.

Keywords: aerobic anoxygenic phototrophic bacteria, bacteriochlorophyll, photoheterotrophy, variable fluorescence
Characterization of photosynthetic apparatuses from a new aerobic chlorophototroph discovered in microbial mats.

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Most anoxygenic chlorophototrophs are strict anaerobes and can only grow and perform light-energy capture and conversion under anoxic conditions. The recently discovered thermophilic acidobacterium “Candidatus Chloracidobacterium thermophilum” is the first aerobic chlorophototrophic bacterium that has a type-I, homodimeric reaction center (RC). This organism and its type-I RCs were initially detected by the occurrence of pscA gene sequences, which encode the core subunit of the RC complex, in metagenomic sequence data derived from hot spring microbial mats.

Here, we report the isolation and biochemical characterization of the type-I RC and light-harvesting complexes from Ca. C. thermophilum. The RC complex comprised two polypeptides: the reaction center core protein, PscA, and a 22-kDa carotenoid-binding protein. The light-induced difference spectra of the isolated RC showed maximal bleaching at 840 nm, which is attributed to the special pair and which we denote as P840. The RC was photoactive even in the presence of oxygen. In combination with the spectroscopic measurements, HPLC and MS analyses revealed that the RC complex contained bacteriochlorophyll (BChl)-a, chlorophyll-a and Zn-containing BChl-a molecules. The possible functions of the Zn-BChl-a molecules and the carotenoid-binding protein will be discussed. Light-harvesting complexes, chlorosomes, were also isolated from Ca. C. thermophilum. Although Ca. C. thermophilum is an aerobe, energy transfer among the BChls in these chlorosomes was very strongly quenched in the presence of oxygen, as measured by quenching of fluorescence emission. Spectroscopic, biochemical, and structural analyses showed that the chlorosomes of Ca. C. thermophilum possess a number of unique features but also share some properties with the chlorosomes found in anaerobic photosynthetic bacteria of other phyla.
Distribution and diversity of chlorophyll \(d\) containing cyanobacteria

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Almost all oxygenic photosynthetic organisms (land plants, algae and cyanobacteria) contain chlorophyll (Chl) \(a\) as their major pigment, and utilize visible light (400-700 nm in wavelength) for photosynthesis. This spectral range of radiation is called as photosynthetically active radiation (PAR), which has been thought to be necessary for oxygenic photosynthesis. *Acaryochloris* is a genus in the Cyanobacteria, which contains chlorophyll (Chl) \(d\) as the predominant pigment. They can utilize far-red light (700-750 nm) in addition to photosynthetically active radiation (PAR: 400-700 nm) for oxygenic photosynthesis. To reveal the ecological significances of this organism and the Chl \(d\)-based photosynthesis, the basic ecological studies on *Acaryochloris* were required. In this study, we aimed to investigate the distribution, diversity and amounts of *Acaryochloris* spp. using molecular biological methods.

The polymerase chain reaction-denaturing gradient gel electrophoresis (PCR-DGGE) was used to detect epiphytic *Acaryochloris* cells from macroalgae. However, *Acaryochloris* cells were hardly detected from most samples of didemnid ascidian and sponge which were collected from the coasts of the Republic of Palau, because of predominant cyanobacterial symbionts existing in these invertebrates. Then, I designed and used the *Acaryochloris*-selective primer set to detect *Acaryochloris* cells and analyzed phylogenetic diversity of them. By using this selective detection method, many phylotypes of *Acaryochloris* were detected in invertebrate samples collected from Palau and macroalgae collected from the coast of Japan and South Africa. Moreover, *Acaryochloris* phylotypes were also detected from pebbles, sands and seawater samples collected from Japanese coast. These results revealed that *Acaryochloris* spp. could randomly attach to various substrates, and they were widely distributed from tropical to subarctic region. It was also revealed that many and diverged phylotypes of *Acaryochloris* were distributed in environments. Phylogenetic analyses demonstrated that these phylotypes were diverged into three phylogenetic subgroups which were different one another at least in species level.

The detection and quantification methods for Chl \(d\) from environmental samples were also developed by employing high performance liquid chromatography, and I determined the amounts of Chl \(d\) extracted from macroalgal samples. As a result, certain amounts of Chl \(d\) were detected from most macroalgae, and the ratio of Chl \(d\) to Chl \(a\) was approximately 1% on average.

In conclusion, it was revealed that certain amounts of *Acaryochloris* spp. were globally distributed around the coastal environments, and the ratio of Chl \(d\)/Chl \(a\) reached up to approximately 1% in the seaweed beds. It suggested that Chl \(d\) could contribute about 1% of the primary production in seaweed bed area, which has been neglected from the estimation of carbon cycle at the coastal environments.

Keywords: chlorophyll \(d\), cyanobacteria
Isolation and characterization of Parmales and its dynamics in the Oyashio region, western North Pacific

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The order Parmales (Heterokonta) is a group of marine phytoplankton species with small solitary cells that are generally 2 to 5 micro-meter in diameter and surrounded by 5 to 8 silica plates. Since diatoms, which are the most successful groups of marine phytoplankton, are also composed of silica cell walls, Parmales has a possible close association with diatoms. Parmales may play a key role in the diatom evolution, to acquire silica synthesis and unique life cycle of diatoms. However, the relationship between Parmales and diatoms is unknown because there is no phylogenetic study due to lack of Parmales culture.

We successfully isolated and characterized a Parmales species, *Triparma laevis*, in the Oyashio region, the western North Pacific for the first time with the aid of a fluorescent silicon tracer. Molecular phylogenetic analyses of SSU rDNA and *rbcL* gene showed that our strain was within the bolidophycean clade of autotrophic naked flagellates and a sister group of diatoms. This indicates a close evolutionary relationship between Parmales, Bolidophyceae and diatoms. We can hypothesize that parmalean and bolidophycean algae or their common ancestor have a life cycle switching between silicified non-flagellated and naked flagellate stages. This hypothetical life cycle is similar to centric diatoms. Centric diatom species also have a silicified vegetative stage and produce naked flagellated male gametes (sperms).

It is important to elucidate similarity and difference between Parmales and diatoms for understanding the evolutionary relationship between these two algae. However, there is little ecological information about Parmales, such as seasonal variations and life cycles, while it is known that Parmales mainly distributes in polar and sub-polar waters. We investigated the seasonal variations of abundance and species composition of the Parmales assemblages in the Oyashio regions.

Samplings were conducted in the Oyashio region on March, May, July and October 2009. The samples for quantifying abundance of the Parmales assemblage were collected from seven depths at 0-100 m. On March and May, the water column was vertically mixed and water temperature was less than 10 degree centigrade at almost stations. Abundance of the Parmales assemblage was high at the stations where water temperature was less than 10 degree centigrade. On the other hands, the vertical stratification was evident between 10-40 m on July and October. Parmales was absent above pycnocline, being abundant in the subsurface layer of 30-50 m where water temperature was lower than 10 degree centigrade. Abundance of the Parmales assemblage on July and October was lower than that on March and May, as the decline of the abundance over pycnocline. Among the Parmales assemblage, *T. laevis* was almost dominant and the seasonal variation of the species composition was small. This characterized the Parmales assemblage compared to the diatom assemblage with clear seasonal succession. Our results showed that the Parmales assemblage, mainly composed of *T. laevis*, would grow in the surface layer in winter-spring and maintain its population under pycnocline in summer-winter. The population under pycnocline would play a role as the initial cells, so-called seed population, for the vegetative growth.

Keywords: Parmales, diatoms, bolidophytes, pico-phytoplankton, Oyashio region
Solar energy and life: the diversity of phototrophic processes in the environments

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The evolution and recruitment of phototrophic processes that convert solar radiation into biochemical energy are among the most critical issues for the life on Earth. Recent works has revealed mechanisms of the phototrophy that contribute to ecosystem and geochemical cycles are rather diverse beyond the well-known oxygenic photosynthesis is. In addition, recent works also revealed diversity and quantitative significance of pico-phytoplanktons in the aquatic environment, hence the microbial energy/material flow founding on them. Understanding of microbial ecology is now refreshing a basis for biogeochemical processes in the ocean.

Keywords: phototrophy, photosynthesis, chlorophyll, aquatic ecosystem, biogeochemical cycles
Examination for lipid biomarker compositions in culture samples of Parmales.

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Parmales is very small marine microalgae, in which cell size is 2-5 um, and is classified as picoplankton. It is pointed out that this alga is one of main primary producer in restricted subarctic regions. Parmales has siliceous tests, and may be closely related to diatom, which is a main important primary producer in the Cenozoic ocean. In 2008, Kuwata’s research group can succeed in isolation of the Parmales collected from the Oyashio region. In the present study, we try to search lipid biomarkers of the Parmales, and to give understanding for first appearance and first processes of evolution of diatom. There have been no reports for siliceous fossil of Parmales. It is known to well preserve siliceous diatom fossil in ancient sediment, and however, such fossil is frequently lost through its dissolution by diagenesis during postdeposition. Therefore, very small siliceous tests of Parmales must be easily dissolved by diagenesis, and it cannot evaluate the timing of first appearance and reconstruct productivity of Parmales by using its siliceous fossil. Thus, we clarified the Parmales biomarkers and their compositions, and these biomarkers are used as molecular fossils for giving understanding evolution processes and historical variations of productivity of this alga.

We use a cultural strain NIES-2565(TOY-0807) of Parmales Triparma sp. (Triparma laevis) for analysis of lipid biomarker. We can identify unsaturated alkene, unsaturated alkenoic acids, C27-C29 sterols as Parmales biomarkers, which have detected from culture samples of diatoms. In particular, this strain is found to be characterized by overwhelmingly abundance of C29 beta-sitosterol. In addition, we can detect a number of unknown polar lipids with higher molecular weight. In the present study, we report preliminary results for Parmales biomarker study and discuss its geoscientific significance as molecular fossil.

Keywords: Parmales, lipid biomarker, evolution of diatom, chemotaxonomy, culture, steroid
Photosynthesis of fern species having different habitats and frond morphologies

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Introduction
Ferns are known to live in various habitats. Many species live in the shady and humid sites under tree canopy but some species live in the open or xeric sites. Their sporophyte fronds have quite different morphologies between species. The morphology of fern frond is considered to reflect the adaptation strategies to their environments. To understand the evolution of ferns and their adaptation to the various habitats, the information of morphological and ecophysiological trait of ferns are needed. Recent studies showed the relationship between habitats and morphological traits or photosynthetic traits separately. The integration of frond morphological traits, photosynthetic traits and their habitats or life forms are essential, since the frond morphology of ferns reflects the adaptation for effective photosynthesis and water relations.

The aim of this study is to integrate the frond morphological traits and photosynthesis for understanding the adaptation strategy of ferns to the various habitats.

Materials and Methods
4 species of temperate fern such as shady terrestrial deciduous, Adiantum pedatum (kujyakushida), Open terrestrial deciduous, Pteridium aquilinum (warabi), shady terrestrial evergreen, Dryopteris erythrosora (benishida) and Epiphyte evergreen, Lepisorus thunbergianus (nokishinobu) in sporophyte stage were grown in a glasshouse. Leaf mass per area (LMA), stomatal density (SD), leaf water content (LWC), relative water content (RWC) and photosynthetic potentials ($V_{max}$ and $J_{max}$) were measured before stress treatment, and then they were subjected to drought stress to investigate the photosynthetic response.

Result
Stomata were on the abaxial side of fronds only in the all the ferns investigated. SD of the Open site fern, $P.$ aquilinum was twice of the other ferns. Evergreen fern had significantly higher LMA than the deciduous ferns. Epiphytic fern, L. thunbergianus showed the highest LMA among the ferns. Photosynthetic potentials ($V_{max}$ and $J_{max}$) of $D.$ erythrosora, L. thunbergianus and $P.$ aquilinum were similar. However, their photosynthetic rate ($A_{max}$) and stomatal conductance ($g_s$) were lower than $P.$ aquilinum.

Some fronds of the deciduous ferns under drought stress were dead. Tip of fronds were withered partly in $D.$ erythrosora. In L. thunbergianus, all of the fronds were withered, but they were recovered after reirrigation. In all ferns, $A_{max}$ and $g_s$ decreased concurrently under drought stress. There were positive correlations between $A_{max}$ and $g_s$. Epiphyte fern, L. thunbergianus showed negative transpiration in severe drought stress. The rubisco content of each ferns were not affected by drought stress.

Discussion
In well watered condition, open site fern, $P.$ aquilinum, have similar photosynthetic potentials ($V_{max}$ and $J_{max}$) to $D.$ erythrosora and L. thunbergianus, but showed the highest $A_{max}$ among the fern species. High SD and $g_s$ of $P.$ aquilinum may cause their high $A_{max}$ in the open sites.

The evergreen ferns showed higher WC than the deciduous ferns. The higher construction cost may be higher for water storage structure in the evergreen ferns. In stress conditions, some fronds of deciduous ferns were dead, but evergreen ferns maintained the frond.

All ferns decreased photosynthetic rate and $g_s$ concurrently but maintained the similar rubisco content in drought stress compared to the control. These results suggest that low stomatal conductivity may be a major factor reducing the photosynthetic rate in drought stress.

L. thunbergianus did not dead below 40% of RWC and showed negative transpiration in drought stress. These results suggest that L. thunbergianus is poikilohydry. Additionally, L. thunbergianus has the extremely higher LMA and WC than the other ferns, indicating that L. thunbergianus has succulent. L. thunbergianus live on the tree, where water resources may be limited. In such a prolonged drought environment, poikilohydry as well as succulent type morphology may be favorable.

Keywords: fern, pteridophyte, photosynthesis, adaptation, evolution, habitat
Acquisition of kleptoplast in *Planoglabratella opercularis* (foraminifer) and its putative function

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A rocky-shore benthic foraminifier, *Planoglabratella opercularis*, constructs specific host-symbiont relationships that has chloroplast as kleptoplast. The "kleptoplast" means "stolen chloroplast" or "symbiotic chloroplast", and refers to a chloroplast originating from a separate organism. Host organisms may have some benefit from kleptoplast, such as organic matters, or amino acids. To understand the functions of kleptoplast, we conducted molecular phylogenetic analyses of kleptoplasts, culture experiment and stable isotope analyses. The trophic position of individuals with or without kleptoplast, we measured stable isotopic composition of amino acid to understand whether their nutritional requirements come from kleptoplast or not. As a result, trophic position (TP) of the individual with kleptoplast shows 1.2. In contrast, TP of cultured individual specimens that digested kleptoplast shows 2.0. It is possible that *P. opercularis* behave as a primary producer, "phyto-benthos", in nature.

Keywords: Kleptoplast, benthic foraminifera, nitrogen isotope of amino acid
Intra- and interspecies variations in intercellular concentrations and compositions of alkene and alkenone in Haptophyte

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Haptophyte algae are one of the major primary producers in the ocean. Long-chain alkenones, unsaturated linear methyl and ethyl C_{37}-C_{40} ketones, are synthesized by few species of haptophyte algae (Emiliania huxleyi, Gephylocapsa oceanica, Isochrysis garbana and Chrysotila lamellosa). Alkenones have frequently been used for estimating the paleotemperature in geological samples, since the number of double bonds change in response to the growing temperature. Along with alkenones, these haptophyte species also produce polyunsaturated long-chain alkenes. Long-chain alkenes are more susceptible to diagenetic process (i.e. photochemical and bacterial degradations), hence are regarded less important to apply in geological past. Therefore, compositions and distributions of long-chain alkenes among its producers have not been systematically examined. Recently, algal biomass is expected to be a new energy resource. We look for the use of haptophyte algae for biorefinery, based on their high rate of reproduction, high content of long-chain lipids, and some other positive features. Alkenes gain importance in this context because of its chemical nature as hydrocarbon. In this study, we analyzed alkene and alkenones in the haptophyte algae in order to gain suite of concentration and compositional data covering wide range of Haptophyte algae.

More than 50 strains of haptophyte algae were obtained from stock culture of Shiraiwa’s laboratory and the other culture collections (e.g. NIES, NCMA, RCC). These strain were grown at 17°C & 20°C for 10-21days. Extraction and separation of lipids were performed based on Sawada and Shiraiwa (2004). After extraction, the lipids were separated by silica gel column, and fraction 1, 2 and 3 (hexane, hexane / ethyl acetate (95/5 v/v), hexane / ethyl acetate (9/1 v/v)) were analyzed by gas chromatography with flame ion detector (GC-FID) and gas chromatography / mass spectrometer (GC/MS).

Alkenes (C_{29}-C_{38}) and alkenones with alkenoates (C_{37}-C_{40}) were detected from strains of four species: E. huxleyi, G. oceanica, I. garbana and C. lamellosa. Other species lack all of these compounds. The total concentrations of alkenones and alkenes were 0.02-1.96 pg/cell (0.09-11.1 ug/ml) and 0.001-0.57 pg/cell (0.01-1.58 ug/ml), respectively. Both intercellular concentrations and compositions of these compounds showed significant differences between strains. A majority of strains mainly contain C_{31} and C_{33} alkenes, while some others contain C_{37} and C_{38} alkenes in significant proportion. Furthermore, some strains of E. huxleyi contained significant amount of C_{29} alkadienes. Rieley et al. (1998, Lipids 33, 617-625) reported that C_{37} and C_{38} alkenes have trans double bonds resemble to those of C_{37} and C_{38} alkenones while C_{31} and C_{33} alkenes have cis double bonds, suggesting distinct biosynthetic pathway for these two groups of alkenes. However, consistent occurrence of alkenes and alkenones highlights close biochemical relationship between these two groups of compounds, as well as importance of the four alkenone producing species as potential hydrocarbon resource.

Keywords: Haptophyte, alkenone, alkene, biorefinery
Microbial eukaryotes living in the coastal marine of Tokyo bay insights from the metagenome

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Progresses in sequencing technique makes it possible to reveal the existence and to detect the diversity of pico-planktonic organisms, and also decide the taxonomic position of uncultured organisms, rapidly and easy way. Metagenomic approaches show that the huge amounts of pico-planktonic organisms are living in various environments. It means that eukaryotic microorganisms play important roles as a producer, consumer or reducer in the biosphere such as microbial loop or biogeochemical cycle. There are so many taxonomically unknown organisms, even though their phylogenetic positions have been determined. It would not be an exaggeration to say that we just reach at the start line to reveal the eukaryotic diversity. Furthermore, there are less physiological or ecological information about them. In this study, we determined the 18SrDNA sequences using filter captured microbial eukaryotes collected every month from the sea water in Tokyo Bay.

Throughout the investigation period, the sequences belonging to Opisthokonta, Stramenopiles and Alveolata, were respectively obtained in each quarters of the total. The residual one quarter were the mixture of Rhizaria, Cryptophyta, Chlorophyta, Rhodophyta, Katablepharid, Telonemia and Haptophyta. When Heterosigma akashiwo made the bloom, the half of the determined sequence comes from the surface bloom water were originated from this alga, and the one from the lower layer under the bloom was mainly composed by Ciliophora and/or parasitic Dinophytes. In summer, centric diatoms, and the Opisthokonta in sometime, became dominant taxa. Most of the Opisthokonta sequences were originated from Metazoa. Notably, Chytridiomycetes were the dominant taxa in August. The sequences identified as Picobiliphyta or MAST(Marine Stramenopiles) were infrequently detected. These findings indicates the existence of many unidentified or missing link organisms in the universal aquatic ecosystems, and the importance of our studies. For the further understandings of the unknown organisms, we are developing the technique to get the both profiles of the phylogenetic position and morphological features together.

Keywords: Metagenome, Environmental sequences, Algae, Protist, aquatic ecosystem
Coping with toxicity of chlorophylls: a biochemical strategy

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Chlorophylls in aquatic environments have been known to be transformed into various derivatives; a portion of chlorophylls are degraded via a so-called Type I process in which tetrapyrrole macrocycles are preserved intact with a variety of defunctionalization, hence surviving into sediments. These survived chlorophyll derivatives could be further altered chemically to be fossil porphyrins and red pigments extracted from sedimentary rock as old as the Proterozoic. We improved HPLC methods that carefully excluded analytical artifacts. We thus revealed that a major component of the chlorophyll derivatives is derived from eukaryotic microbe, which is produced along chlorophyll detoxification catabolisms.

Keywords: chlorophyll, eukaryotic microbes, detoxification, chlorophyll catabolism
Vertical distribution and abundance of *Acaryochloris* spp. at the deepest area of Lake Biwa

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The genus *Acaryochloris* is the only organism which contains Chlorophyll *d* (Chl *d*) as their predominant light harvesting pigments. This cyanobacteria can absorb and use far-red light (FR), ranging from 700 to 750 nm, as well as visible light (400-700 nm) for oxygenic photosynthesis. Therefore, the ecological and the critical role of Chl *d* have been thought to utilize FR effectively, avoiding the competition for visible light with other chlorophyll *a* (Chl *a*) containing algae and providing niche to the genus *Acaryochloris*.

In our successive works, we have detected 16S rRNA gene of *Acaryochloris* spp. from the water samples collected at more than 30 m depth in Lake Biwa. At the depths, no FR was thought to be remaining. This suggested that the detected planktonic *Acaryochloris* spp. could not utilize FR for photosynthesis.

The purposes of this study were to reveal the niche of planktonic *Acaryochloris* spp. and the adaptive significances of Chl *d* in that depth. For these purposes, we established a real-time PCR method to quantify the copy of *Acaryochloris*-16S rRNA gene and investigated the vertical distribution of planktonic *Acaryochloris* spp. at the deepest area of Lake Biwa. Additionally, we compared the solar radiation spectra at the depth where *Acaryochloris* spp. were detected and the excitation spectrum of *Acaryochloris* cells. We would like to present the results of our investigations and discuss about the adaptive significances of Chl *d* for the planktonic *Acaryochloris* spp. in Lake Biwa.

Keywords: *Acaryochloris* spp., chlorophyll *d*, vertical distribution, real time PCR, Lake Biwa