ENVILAB: Measuring phytoplankton in-vivo absorption and scattering properties under tunable environmental conditions

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Abstract: Optical remote sensing of phytoplankton draws on distinctive spectral features which can vary with both species and environmental conditions. Here, we present a set-up (ENVILAB) for growing phytoplankton under well-defined light, temperature and nutrient conditions. The custom-built light source enables creation of light with spectral composition similar to natural aquatic environments. Spectral tuning allows for light quality studies. Attenuation is monitored with a spectrometer in transmission mode. In combination with automated spectrophotometer and fluorimeter measurements, absorption and excitation-emission-fluorescence spectra are recorded. The set-up opens the door for systematic studies on phytoplankton optical properties and physiology.

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References and links
1. Introduction

Phytoplankton forms the basis of the aquatic food web and plays a fundamental role in biogeochemical cycles in the ocean [1–3]. Certain phytoplankton species such as *Microcystis* are potential toxin producers and their monitoring is of interest to many stakeholders such as environmental agencies, aquaculture or water resource managers [4,5].

Due to its pigmentation and its specific scattering properties, phytoplankton can be detected in remotely sensed data from satellite, air- and shipborne as well as from ground based measurements. Optical signatures such as absorption and fluorescence by chlorophyll-*a* or specific backscattering properties serve for identification and separation from other water constituents in reflectance signals. These signatures are used to infer phytoplankton concentration and its spatial distribution on both local and global scales, e.g. [6–8]. In case of cyanobacteria, remote sensing algorithms often exploit spectral features originating from additional pigments in light harvesting complexes such as from phycocyanin [9–11]. Several studies on phytoplankton suggest that pigment composition can change not only from species to species but also with environmental conditions [12–16]. For remote sensing modelling, it is crucial to understand this variability in the inherent optical properties (IOPs), since it affects accuracy and sensitivity of retrieval algorithms. Systematic laboratory studies on phytoplankton *in-vivo* absorption and scattering characteristics under controlled environmental conditions are rare. Further, they are usually performed under so called day-light conditions (e.g. [17,18]). For this purpose, a light source (often a fluorescent tube) is used that provides continuous radiation in the 400 nm to 700 nm wavelengths range (photosynthetically active radiation – PAR). Assuring a certain PAR emission with day-light lamps helps to create stable and reproducible culturing conditions. However, the actual spectral composition (e.g., uneven spectral distribution and high distinctive emission peaks for fluorescent lamps), and how it compares to light that phytoplankton is exposed to in its aquatic environments is usually neglected. A few studies focus on the impact of spectral composition of light on phytoplankton and algae, but for this purpose usually lamps with color filters or a few individual light emitting diodes (LEDs) are chosen [19–21].

Recently, especially in the context of bio-fuel and biotechnology applications, there have been many studies on light influencing growth behavior of bacteria and algae. The motivation for these studies often is to increase growth rates and lipid production. Light sources for culturing are optimized towards yielding highest growth rates, lipid and large-scale production rather than to resemble natural conditions [22–24]. Therefore, culturing assemblies designed for bio-fuel applications usually do not allow for an investigation of phytoplankton growth under natural light conditions.
In the present contribution, a novel set-up for optical studies on phytoplankton is presented. It allows for controlled phytoplankton growth and serves as a stage for experimental investigation of changes in phytoplankton physiology in response to varying light field, nutrient or temperature conditions. With the help of a tunable multispectral light source, phytoplankton can be grown under a broad variety of artificial and natural light conditions with respect to intensity and spectral composition. Automated spectrometric measurements allow to analyze absorption and scattering characteristics.

After an introduction to the set-up design with focus on its individual components and the spectrometric data that is recorded, the functional range is demonstrated by growth and dilution of a cyanobacteria culture.

2. Design and Realization of the ENVILAB Set-up

2.1 System Concept

Phytoplankton is grown in a photo-bioreactor under light that is provided by a custom-built multispectral LED light source tunable in both spectral composition and intensity. Growth is monitored by a spectrometer that measures spectral transmission and thus attenuation of light through the reactor chamber. Via an automated peristaltic pumping system, the growth chamber is linked with a benchtop fluorimeter and spectrophotometer with an integrating sphere for recording high precision fluorescence and absorption spectra. An estimate on scattering properties can be obtained by subtracting absorption from attenuation spectra. Stirring and automated dilution of the culture with nutrient medium allows long term experiments and studies at fixed concentration ranges. Tubes for sample transportation can be cleaned with a cleansing liquid in between or after measurements. Specifically designed software allows for live monitoring of growth parameters such as the optical depth. A sketch of the experimental set-up is depicted in Fig. 1.

2.2 Tunable LED-light Source for Spectral Light Field Simulation

Tunable LED-based light simulators, as described in the literature, are realized in a variety of designs for applications such as color rendering [26,27], solar spectrum simulation [28–32] and responsivity measurements [33,34]. The respective light sources differ in number, types and combination of LEDs, as well as in their electronic control and additional optical components. Accordingly, they exhibit different characteristics in terms of spectral matching, illumination area and irradiance intensities. In photovoltaics (PV), commercial set-ups exist that are optimized towards accurate and stable reproduction of solar irradiance for characterization of PV cells [35–37]. For this purpose, assemblies with LEDs of typically around 20 different peak wavelengths are used to provide light in the visible and in parts of the ultraviolet and infrared. Illumination areas are designed to cover standard PV cell sizes. For biological studies, Fujiwara and Yano present a set-up consisting of 547 LEDs with 32 different peak wavelengths [28]. This light source provides high spectral flexibility in the 385...
- 910 nm wavelength range illuminating an area of about 7 cm². Linden et al. report on a tunable solar simulator with modular design for illumination of different sizes and shapes, but with less spectral matching flexibility in the visible (LEDs with 23 different peak wavelengths for cover the 350 to 1100 nm wavelength range) [31].

Here, we present a tunable, multispectral LED light source which is specifically designed for phytoplankton studies in a photobioreactor [38]. Special attention was paid to spectral flexibility and the resemblance of the obtained irradiance spectra to spectra found in aquatic environments. In addition to sun position and metrological conditions, spectral composition of underwater irradiance changes with depth, bottom type and vegetation, and water constituent concentrations. In order to simulate a variety of irradiance scenarios in the photosynthetically active radiation range of 400 nm to 700 nm, multiple LEDs were selected according to their high intensities, spectral peak position and bandwidths: Intensities are high enough to match (in sum) maximum spectral irradiance below the water surface. The latter was simulated using the Water Color Simulator (WASI) software (at a sun zenith angle of 0° and a sensor depth of 0 m) [26,27]. Actual bandwidths of LEDs span around 13 to 38 nm (FWHM), with the exception of two types of phosphor coated broad band LEDs which were added to fill the gap between 530 nm and 600 nm at which LEDs are difficult to find. Position of emission peaks are chosen to assure both spectral continuity and flexibility which is needed to match target spectra [39,40].

All LEDs (72 with 24 different peak wavelengths) are placed on Aluminium plates which together are arranged in arrays (see Fig. 2(a)). These arrays are mounted on heat dissipaters in such a way, that individual LED plates remain replaceable. The temperature of the heat sink is monitored; fans attached provide additional cooling in case the temperature rises above a given threshold (typically 25 °C).

In order to assure homogeneity of radiance across the illuminated area, the LED arrays are mounted into a mirror foil covered tube of rectangular shape (see Fig. 2(b)). Reflection at surface walls together with a diffuser plate (PLEXIGLAS® Satinice) at the tube’s exit, restrict inhomogeneity to less than 10% within the illuminated area of 10 cm × 20 cm.

Spectral characteristics of each LED (mounted in module) were determined at lab conditions with a spectrometer (CARL ZEISS MCS 500, 350-1021 nm, 0.8° field-of-view) at LED current intensity of 350 mA. The calibrated spectra are stored in a database. This database forms the spectral library used to determine pulse width modulation (PWM) duties for individual LEDs in designated control software. Via a graphical user interface, target spectra can be loaded. These target spectra are either artificially composed, measured or simulated irradiance spectra that cover the 400-800 nm wavelength region and which are to be reproduced by the light source. PWM duties of the corresponding LEDs are set by the control software using a least-square-fit between the target spectrum and the sum of individual LED spectra. In a cycling mode, user defined target spectra can be varied over time. This way, day and night cycles, spectral shifts or sudden changes in illumination can be simulated. A summary on technical details of the light source can be found in Table 1.
Table 1. Technical specifications of the LED light source.

<table>
<thead>
<tr>
<th>Specifications</th>
<th>Details</th>
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<tbody>
<tr>
<td>LEDs (total number)</td>
<td>72 LEDs of 24 different peak wavelengths.</td>
</tr>
<tr>
<td>Max. power/irradiance</td>
<td>~8 W on an area of 10 cm × 20 cm.</td>
</tr>
<tr>
<td>Hardware control</td>
<td>Custom designed circuit board with 32 independent PWM-controlled LED drivers (350 mA LED string current); setting of PWM duty via an I2C interface (12 bits resolution, frequencies up to 1526 Hz); temperature monitoring ICs and PWM-controlled fans.</td>
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2.3 Photobioreactor

Individual components of the photobioreactor [25] were designed while having both the microbiological and optical requirements in mind: In order to minimize contamination for microbiological studies on phytoplankton samples, all parts can be disassembled, cleaned and sterilized in an autoclave. In addition, electro-polished stainless steel of 2 mm thickness as frame material hinders deposition of phytoplankton on the inner walls. To fulfill optical demands, Borosilicate windows were chosen that provide over 90% transmission for light in the 390 to 750 nm wavelength range.

Figure 2(b) shows a sketch of the photobioreactor assembly in explosion view. The base area was matched to a magnetic stirrer plate of 12.5 cm diameter. This way, velocity controlled mixing within the reactor is possible by inserting a Polytetrafluoroethylene (PTFE) coated stirrer. The reactor’s inner dimensions comprise a volume of about 3.5 l of sample liquid which can be inserted and extracted via glass tubes that are connected to flexible tubes. Liquids in the tubing connection are transported with a four-channel peristaltic pump. Fluid pathways are controlled by magnetic pinch valves which open when voltage is applied. These valves are triggered by a microcontroller. Inside the incubation chamber, temperature is monitored with a digital temperature sensor in a steel casing. On the two sides of the chamber’s frame, custom-built copper plates provide heat-exchange to a liquid that can be heated or cooled by a thermostat. Insulation of the chamber body from ambient conditions assures additional temperature stability.

Software with Graphical User Interface allows creation of a timeline that includes all relevant pumping and measurement steps. A cycling function can be activated for repetitive experiments. Once the involved volumes are defined, the software visualizes current liquid levels and allows for live temperature and optical growth monitoring.
2.4 Spectrometric Monitoring

The ENVILAB set-up provides a two-step spectral monitoring making use of a miniaturized spectrometer (*FREEDOM VIS FSV-305*, Ibsen Photonics Inc., Denmark) for transmission measurements, and of two bench-top instruments for recording absorption and fluorescence spectra (*LAMBDA®1050*, PerkinElmer Inc., U.S., and *FLUOROMAX®4*, HORIBA Scientific, Japan). The transmission of light through the sample liquid is measured regularly using the *FREEDOM VIS* spectrometer. This way, the decrease in transmission is monitored and thus, by application of Beer–Lambert’s law, the growth rate is recorded. For high precision measurements with minimal scattering influence, samples are pumped into a flow-through quartz cuvette, which is mounted inside of the 150 mm diameter integrating sphere of the double beam spectrophotometer. In parallel, the fluorescence spectrum at a selected excitation wavelength or a full excitation and emission spectrum is recorded. Emitted light is detected under 90 degrees to the excitation light beam. Fluorescence measurements are normalized to excitation light intensities as measured with the instrument’s reference detector, and manufacturer’s correction factors are applied. After each measurement cycle, the respective cuvettes are automatically emptied into a waste collection flask and both tubes and cuvettes are flushed with a cleansing liquid.

3. Functional Tests and Discussion

3.1 Irradiance Simulation and Temperature Control

Motivation for designing the LED light source was to reproduce both artificial and natural irradiance spectra. Figure 3 illustrates target spectra resembling for three different irradiance scenarios: (a) at water surface, (b) in water rich of colored dissolved organic matter (CDOM) at 1 m depth and (c) at 3 m water depth. All spectra were simulated with WASI (for a sun zenith angle of 40 °, a total suspended matter concentration of 1.5 mg l⁻¹, an absorption of CDOM at 440 nm of 0.2 m⁻¹ with a slope of S_{CDOM} = 0.014 nm⁻¹ and a phytoplankton concentration of 3 µg l⁻¹). Black lines represent simulated spectra and colored lines individual LEDs, while grey diamonds reflect the total LED irradiance.

In Fig. 3, maximum relative deviation (per nm) from target irradiance of 54% in case (a) and 34-35% in case (b) and (c) can be observed, with an average deviation of 14%, 6% and 8%, respectively. This is caused by spectral overlap of the emission peaks. Especially if lower water depths are to be simulated, higher average deviation from target spectra will be observed. The reason for this is that, in larger depth, more light gets absorbed by the water itself, leading to pronounced spectral features around 600 nm. The coverage of the corresponding wavelength range with broad band LEDs does not allow for proper reproduction of these spectral features in the irradiance spectra. At the time when the light source was built, no suitable LEDs were available to cover the 550 to 600 nm wavelength regions.

Despite these drawbacks, the light source provides much more flexibility in resembling natural irradiance spectra than conventional light sources. No additional filters are necessary to avoid unwanted peaks. In addition, relative intensities of “overshoots” are much smaller compared to fluorescent tubes, where distinctive peaks of chemical compounds affect the emission spectrum.
Temperature stability inside the photobioreactor was tested in the laboratory at room temperature. For this purpose, 1.2 l and 3 l of purified water was cooled and heated via the heat exchanger using the thermostat which was to set values in the range of 1 to 40 °C. Tests confirmed stable temperature conditions in the range of 2.6 to 39.1 ± 0.1 °C with maximum deviation of 3°C for most extreme temperatures and depending on the filling level. Tests confirmed that the ENVILAB enables a variety of studies at temperatures typically found in natural water bodies [25].

3.2 Automated Spectral Measurements and Nutrient Control

First phytoplankton growth tests were performed with *Synechocystis sp. PCC 6803* [41]. This cyanobacterium is widely studied [42], making it an interesting model organism.

For bioreactor functionality validation, the culture was kept under light with fixed spectral composition. In order to produce an intensity similar to previous culturing conditions, the irradiance spectrum (at 2 m water depth; simulated with WASI default parameters) was additionally attenuated by a factor of 10. Light was switched on and off in a day-and-night cycle (14 h/10 h). A baseline measurement with pure nutrient medium was performed before adding the culture.

To validate the growth monitoring ability, transmission of a culture of PCC 6803 bacteria was measured with the FREEDOM VIS spectrometer during a period of 6.3 days with a series of 10 spectra every 15 minutes. Dark current measurements were recorded directly before start of the growth experiment and during night phases. The corresponding 3540 extinction spectra are shown in Fig. 4(a). They illustrate the continuous growth during day times which result in an increase in attenuation. The attenuation peaks correspond to phytoplankton pigments. The peak around 436 nm and the peak at around 673 nm can be associated with absorption by chlorophyll pigments. The peak around 620 – 630 nm relates to phycocyanin absorption [43,44].

In order to test automated measurement cycles and nutrient supply, a series of dilution measurements with PCC 6803 (grown under traditional day-light) was recorded. Over 24 hours, a dense culture inside the photobioreactor was stepwise diluted with nutrient medium (BG11 modified from [45]) until noise contribution became too high to acquire meaningful spectra [25]. Figure 4(b) shows the corresponding absorption spectra. The decrease in absorption corresponds to a reduction of cyanobacteria concentration. The absorption decreases linearly over the wavelength range of 300 nm to 700 nm as shown in the inset of Fig. 3(b).
At each dilution step, excitation-emission spectra were recorded. In order to reduce acquisition time, the wavelength range of interest was split in two: a) with an excitation ($\lambda_{\text{ex}}$) from 200 – 700 nm and an emission ($\lambda_{\text{em}}$) from 575 – 800 nm, b) with $\lambda_{\text{ex}}$ from 200 – 540 nm and $\lambda_{\text{em}}$ from 250 – 575 nm. The two EEM (excitation-emission matrices) spectra at every dilution step were automatically recorded right after each other (5 nm wavelength steps, 2 nm bandpass front and exit slit, 0.1 s integration time). After adding the spectra to combined EEMs, the pure nutrient EEM was subtracted. Figure 5(a) illustrates an example of such a blank corrected EEM. High intensities at diagonal values ($\lambda_{\text{ex}} = \lambda_{\text{em}}$) are due to elastic scattering (Rayleigh) and instrument effects. Also second order effects at doubled emission wavelength ($\lambda_{\text{em}} = 2\lambda_{\text{ex}}$) can be observed. For further analysis, Rayleigh scattering was masked out, leading to blanks in diagonal matrix values which occur as gaps in the excitation spectrum at 680 nm emission as shown in Fig. 5(b).

EEM spectra show that fluorescence emission maxima were shifted towards longer wavelengths as compared to absorption spectra, which is to be expected for this inelastic scattering process. Fluorescence intensity at 620 nm excitation and 680 nm emission ($f_{430/685}$) decreases linearly with dilution (see inset of Fig. 5(b)). This proves that measurements were not impacted by non-linear effects such as the inner filter effect (i.e. reabsorption of fluorescence photons by the sample) [25,46].

Fig. 4. (a) Growth of *Synechocystis* sp. (PCC 6803) monitored through the reactor windows; transmission was measured ten times during day every 15 minutes over a period of ~6.3 days leading to the depicted 3540 spectra. (b) Absorption spectra during automated dilution of a PCC 6803 culture; inset: corresponding concentration factor vs. absorption at 673 nm.

Fig. 5. (a) Excitation-mission spectrum of *Synechocystis* sp. (PCC6803) after blank subtraction. The dashed line indicates the excitation spectrum at 680 nm emission. (b) Excitation spectrum at 680 nm for PCC 6803 at different dilution steps. The inset indicates linear decrease of fluorescence intensity at 620 nm excitation and 680 nm emission with dilution.
4. Conclusion and Outlook

In this paper, a novel automated spectrometric set-up (ENVILAB) was presented, which enables controlled phytoplankton growth and at the same time allows for the characterization of its bulk attenuation, absorption and fluorescence characteristics. The custom-built, tunable, multispectral LED light source permits systematic studies on phytoplankton under varying light conditions. Consisting of 72 LEDs of 24 different peak wavelengths, irradiance intensities and spectral compositions of the light source are tunable in a wavelength range of about 380 to 700 nm. Maximum irradiance matches that of natural sun close to the water surface (with a zenith angle of 0°). The flexibility of the LED light source for natural light field simulation has been demonstrated on three examples at different underwater conditions. Thus, together with under-water irradiance simulation not only artificial but also under-water light scenarios similar to natural conditions (in terms of spectral composition) can be reproduced. In combination with the time line function, both sudden and slow changes in spectral composition and its impact on phytoplankton growth and optical properties can be studied. Further improvement of the light source should address finer reproduction of spectral feature, especially in the 550 to 600 nm wavelength range.

In the photobioreactor of the ENVILAB, cultures of phytoplankton can be grown under controlled environmental conditions. Temperature stabilization is realized via a heat exchanger which in combination with a thermostat assures temperatures in the range of 2.6 to 39.1 ± 0.1 °C. Currently, no gas exchange is included but additional openings in the steel frame of the bioreactor provide access if future implementation (e.g. of a bubbling system) is desired.

For validation of growth monitoring, *Synechocystis* sp. (PCC 6803) was grown in a stirred culture. Monitoring the LED light through the flat transmission windows using an external miniaturized spectrometer has been proven to be suitable for monitoring of changes in the attenuation of the culture.

Nutrient supply and automated measurement procedures were tested by continuously diluting a *Synechocystis* PCC 6803 culture with nutrient medium. With every automated dilution step, samples were programmed to be withdrawn from the bioreactor via the automated peristaltic pumping system, transported into the photometer and fluorimeter and spectra were recorded. This way, *in-vivo* absorption and fluorescence measurements could be performed externally with minimally invasive sample extraction. Absorption was determined inside an integrating sphere of the photometer, while fluorescence was recorded in a compact steady state fluorimeter which allows for excitation-emission measurements. For fixed concentration range studies, current attenuation of the phytoplankton culture inside the photobioreactor can be measured and the corresponding optical depth values are fed into the automated nutrient dilution programming loop. This loop controls concentration by mixing fresh nutrient medium into the bioreactor until the optical density has reached the desired value.

In summary, ENVILAB provides a novel laboratory environment for systematic studies on phytoplankton physiology and optical properties. Next to attenuation measurements, which give insight into scattering properties, absorption and fluorescence measurements on *in-vivo* samples are collected. This spectral information is highly valuable as it builds the basis for water constituent and pigment discrimination in many applications ranging from remote sensing to optical *in situ* devices [47–49]. Reduction of uncertainties in the IOPs and understanding its natural variability may therefore improve water quality monitoring in the long term [7]. In addition, control and adjustability of environmental factors during phytoplankton growth (such as light and its spectral composition, temperature and nutrient supply), enable experiments in various fields of phytoplankton research such as chromatic adaptation, day-night cycle or niche behavior [13,50].
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