Blue-green light effects on light-limited rates of photosynthesis: relationship to pigmentation and productivity estimates for *Synechococcus* populations from the Sargasso Sea

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ABSTRACT: The impact of blue-green light incubation on short-term diurnal, daily, and integrated water column estimates of whole water (> 0.2 μm) and *Synechococcus*-specific photosynthesis was assessed throughout the euphotic zone at 2 stations in the Sargasso Sea. Replicate samples were incubated under both tungsten white light and broad band blue-green light, where the latter simulated light quality within the upper water column of the open sea. Diurnal variations in size-fractioned (0.2-0.6 μm, 0.6-1 μm, and 1-5 μm) blue-green vs white light photosynthesis-irradiance (P-I) curves, chlorophyll (Chl) and phycoerythrin (PE) concentrations, and cell abundance of PE-rich cyanobacterial *Synechococcus* spp. and Chl-fluorescing algae, were measured within samples from the surface, PE maximum, Chl maximum, and the base of the euphotic zone. *Synechococcus* spp. dominated ultra-phytoplankton communities down to the light depths of the PE maximum (3 to 7 % surface illumination, Iₜ), with maxima in cell abundance routinely located at light depths ≥ 50 % Iₜ. Blue-green and white light incubation conditions generally did not affect light-saturated rates of photosynthesis (Pₘₐₓ) but blue-green light routinely did provide much higher estimates of light-limited rates of photosynthesis (α). For size-fractioned subpopulations dominated by *Synechococcus* spp., blue-green light values of α were 25-fold greater than white light estimates. Compared to white light estimates, blue-green light estimates of total (> 0.2 μm) daily integrated water column primary productivity were 6 to 13 % higher, while the contribution of *Synechococcus* spp. to overall primary production rose from between 57 and 61 % to between 73 and 84 %. From the surface down to about 5 % Iₜ, the PE content of *Synechococcus* cells increased with decreasing light and/or increasing inorganic nitrogen availability. Increases in *Synechococcus* PE/cell occurred in direct proportion to blue-green light measurements of photosynthetic quantum efficiency, further indicating that these cyanobacteria are physiologically well suited to harvest photosynthetically utilizeable light throughout a large portion of the euphotic zone.

INTRODUCTION

Estimates of primary production in oceanic water masses have been revised upward as knowledge of the abundance and activity of very small (<3 μm) ultra-phytoplankton has advanced during the last decade and field approaches to measuring photosynthetic rates have improved (Glover 1985, Joint 1986, Li 1986, Waterbury et al. 1986, Marra & Heinemann 1987, Smith et al. 1987). As a result, the primary productivity of oceanic central gyres now appears to be based largely on the photosynthetic activity of ultra-phytoplankton communities (Glover 1985, Fogg 1986, Joint 1986, Li 1986, Shapiro & Guillard 1986, Waterbury et al. 1986, Li & Platt 1987). Ultra-phytoplankton are more abundant than net-phytoplankton in these oligotrophic environments (Murphy & Haugen 1985, Glover et al. 1988a, b, Itrirriaga & Marra 1998), perhaps in part because photosynthetic activity of smaller phytoplankters is favored by warmer temperatures found in tropical
water masses (Malone 1980), they have negligible sinking rates (Malone 1980, Takahashi & Bienfang 1983) and surface communities may be sustained by nanomolar levels of nitrate (Glover et al. 1988a). Furthermore, oceanic ultraphytoplankton can have near maximal rates of in situ growth (CAMPBELL & CARPENTER 1986, PREZELIN ET AL. 1987c, ITURRIAGA & MARRA 1988), which could lead to vertical stratification and accumulation of distinct ultraphytoplankton communities at different euphotic zone depths of relatively stable water columns.


During summer, communities of Prochloron-like and eukaryotic ultraphytoplankton are generally more abundant at and below the deep chlorophyll maximum (Chl max) (GLOVER 1985, MURPHY & HAUGEN 1985, GLOVER ET AL. 1986, CHISHOLM ET AL. 1988, GLOVER ET AL. 1988b), where they are thought to be photoadapted to low intensity blue-violet light (WOOD 1985, GLOVER ET AL. 1987) and/or exploiting the upward advection of nutrients across the thermocline (GLOVER 1985). From the Chl max down, a decline in both cell abundance and cellular rates of photosynthesis by Synechococcus spp. generally are observed (GLOVER ET AL. 1986, 1988b, PREZELIN ET AL. 1986). The decline may be linked to the very low photosynthetic quantum efficiency these cyanobacteria have for absorbed light at wavebands in the blue-violet portion of the visible spectrum, where their cell absorption is dominated by nonphotosynthetic carotenoids (LEWIS ET AL. 1986, BOUCHER ET AL. UNPUBL.). In contrast, Synechococcus spp. are often of order of magnitude more frequent in the upper euphotic layer of the mid-Atlantic than all size categories of Chl-fluorescing cells combined (MURPHY & HAUGEN 1985, GLOVER ET AL. 1986, 1988a, b, LI & WOOD 1988). Possible explanations for the predominance of Synechococcus over eukaryotic algae in the upper part of the euphotic zone include lower selective grazing pressure on the cyanobacteria, a higher efficiency in utilizing nanomolar levels of inorganic nitrate (GLOVER ET AL. 1988a), and/or higher efficiency in utilizing the blue-green spectral bands (500 to 550 nm) dominating the underwater light field of the upper mixed layer (WOOD 1985, LEWIS ET AL. 1986, GLOVER ET AL. 1987, CAMPBELL & ITURRIAGA 1988, LI & WOOD 1988, BOUCHER ET AL. UNPUBL.).

Further improvements in predicting oceanic primary production will come, in part, from a better understanding of the bio-optical interactions that influence the temporal/spatial distribution and activities of the major photosynthetic components within the ultraphytoplankton communities. Given that mixed community assemblages are routinely sampled in the field, shipboard determinations of metabolic rates for specific phytoplankton subpopulations can be very difficult. However, methodological advances have enabled us to combine selective size fractionation, epifluorescence microscopy, Chl and PE quantification, and small volume radiolabelling techniques in the present study, in order to detail the photophysiological characteristics and photosynthetic activities of Synechococcus subpopulations in a manner usually restricted to laboratory studies of clonal isolates. As part of a larger effort to determine the environmental variables that influence the natural distribution of Synechococcus, the present field study was carried out to assess the impact blue-green light incubation conditions can have on short-term diurnal, daily, and integrated water column estimates of whole water (>0.2 pm) and on Synechococcus-specific primary productivity. Blue-green light conditions simulated light fields in the upper euphotic zone and included the absorption waveband of phycoerythrin (PE), which has been shown in carbon action spectra measurements to drive a major fraction of photosynthetic activity in Synechococcus spp. (LEWIS ET AL. 1986, BOUCHER ET AL. UNPUBL.). For temporal and spatial replication, measurements of photophysiological variables were repeated at 3 intervals between dawn and dusk throughout the euphotic zone at 2 stations in the Sargasso Sea. By epifluorescence microscopic enumerations of Chl- and PE-fluorescing components within size-fractioned samples, we were able to compare the effect that blue-green and white light incubations have on estimates of water column primary productivity and the increased contribution that Synechococcus spp. make to that production. Furthermore, it was possible to relate the relative blue-green photosynthetic light quantum effi-
ciency of *Synechococcus* spp. to cellular concentrations of phycoerythrin in field populations sampled throughout the euphotic zone of the Sargasso Sea.

**METHODS**

The study was conducted at 2 stations in the Sargasso Sea during summer 1986, working from the RV ‘Endeavor’. Stn 1 was occupied between 21 and 28 July and was located 140 km east of Bermuda. Stn 2 was occupied between 2 and 9 August and was located 165 km northeast of Bermuda (Table 1). An 180 m submersible pumping system was constructed to collect discrete water samples and to simultaneously profile the vertical distribution of temperature, in vivo chlorophyll (Chl) fluorescence and in situ scalar irradiance (Biospherical Design). Intake hose diameter was 2” (5 cm) (ID), minimizing shear effects on phytoplankton communities and giving radiolabel uptake rates equivalent to those measured when replicate samples were collected from GoFlo bottles (Prézelin & Glover unpubl.). Pump flow rate approximated 30 gal (1131) min⁻¹, with a hose transit time of about 3 min from intake to outlet into a darkened container (10 gal [381], turnover time of 20 s) on deck where samples were debubbled. A portion of the debubbled pump effluent was directed via a submersible pump (Little Giant 2E-38N) through a secondary debubbler (1 l) and into an adapted Turner designs fluorometer to measure in vivo Chl fluorescence. Analog data from the fluorometer, in situ light, temperature and depth probes, as well as from a surface quantum irradiance sensor (Biospherical Design) were recorded on strip charts.

Just prior to passage through the fluorometer, 125 ml unfiltered samples were collected from a line pump directly into prewashed 125 ml polyethylene bottles and frozen for subsequent onshore analyses of inorganic nutrient concentrations. Nitrate concentrations in the upper 60 m were measured using a chemiluminescence technique with a precision of ± 2 nM (Garside 1982). At deeper depths, nitrate, nitrite and ammonia concentrations were determined by standard colorimetric methods with precisions of ± 0.08, 0.01 and 0.02 nM respectively (Strickland & Parsons 1972).

Larger-volume samples were collected directly from debubbled pump effluent in the darkened deck container and placed in darkened 20 l polyethylene carboys that were immediately transported inside to the laboratory for analyses. Shipboard enumeration of PE-fluorescing *Synechococcus* spp. and Chl-fluorescing algae were carried out by direct count epifluorescence microscopy, following procedures identical to those detailed by Glover et al. (1986).

For determination of Chl a concentration, duplicate water samples were filtered, with the chosen volume (50 ml to 1 l) depending upon the in vivo Chl fluorescence signal from the pump profile and on the filter pore size. Replicate samples were filtered through

<table>
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<th>Location</th>
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<th>% I₀</th>
<th>Temp (°C)</th>
<th>NO₃ (nM)</th>
<th>NH₄ (nM)</th>
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Nucleopore filters of 5, 1, 0.6 and 0.2 μm pore sizes. Methods of filtration and extraction were previously described (Glover et al. 1986). Chl concentrations were determined directly for > 5, > 1, > 0.2 μm communities, and by mathematical subtraction for 0.2–0.6 μm, 0.6–1 μm, and 1–5 μm fractions. Standard errors for chlorophylls were routinely less than 10% of the mean. Cell-specific Chl concentrations in pure θ-6–1 μm *Synechococcus* populations (95 to 100% of all auto-fluorescent cells), were used in conjunction with cell numbers in the 1–5 and 0.2–0.6 μm fractions, to estimate *Synechococcus* Chl in these other size ranges. The quantity of other algal Chl in 1–5 and 0.2–0.6 μm fractions could then be calculated by subtraction of the amount of *Synechococcus* Chl from the total concentration in each size category. It was only at the Chl max and the 0.5% light depth that the 0.6–1 μm fraction could not be considered a pure population of *Synechococcus*, since they represented < 50% of auto-fluorescing cells in this size range. At these depths, we could only estimate cellular Chl concentrations of *Synechococcus* spp. by proportion, using relative numbers of *Synechococcus*: algal cells within the 0.6–1 μm fraction. *Synechococcus* Chl may therefore have been overestimated at the base of the euphotic zone, since algae in this size range are believed to contain more Chl per cell than *Synechococcus* spp. (Glover et al. 1987).

Phycoerythrin (PE) concentrations were determined using a modification of procedures outlined by Wyman et al. (1985). *Synechococcus* cells were collected on 0.6 μm Nucleopore filters and resuspended in 4 to 5 ml of 50% glycerol, which uncouples energy transfer between phycoerythrin and phycocyanin. Cell numbers in resuspensions were determined by shipboard epifluorescence microscopy as previously described (Glover et al. 1986). PE was excited at 520 nm and the in vivo fluorescence emission of the phycoerythrobilin (PEB) chromophores detected at 577 nm (half bandwidth of 4 nm) in a Turner 111 fluorometer equipped with enhancer gates to increase sensitivity. The fluorescence intensity of the uncoupled PE in the natural communities was calibrated at sea against similarly treated whole cells of oceanic *Synechococcus* clone WH7803 of known PE content and against isolated phycobilisomes and purified phycoerythrin from the same organism. The purpose of using phycobilisomes and PE is to check for the amount of quenching resulting from package effects in whole cells. Since the effective concentration of PE localized in cells is high, there will always be a loss of emission intensity owing to quenching and reabsorption. In vivo fluorescence intensity in suspensions of uncoupled cells was linear over the range 0 to 550 ng PE ml⁻¹.

Diurnal measurements of photosynthesis-irradiance (P-I) relationships were conducted at 4 depths at both stations. Following vertical profiling of the water column, samples were collected at dawn, midday and dusk from the surface, the depth of the uncoupled PE m⁻³ fluorescence maximum, the depth of the in vivo Chl fluorescence maximum and the base of the euphotic zone (0.3 to 0.5% I$_{S}$) (Table 1). Unfiltered whole water samples were inoculated with sodium $^{14}$C-bicarbonate (3 to 5 μC ml⁻¹ final activity) and 15 to 17 ml aliquots dispensed into acid-washed glass scintillation vials, following procedures detailed in Prezelin et al. (1986, 1987b). Each P–I curve represented 14 light samples (measured at irradiances up to 550 μE m⁻² s⁻¹ in white light; up to 200 μE m⁻² s⁻¹ in blue-green light), duplicate darks and duplicate time zero controls. Identical samples were incubated over 'white' and 'blue-green' light, provided by a fan-cooled 500 W quartz-halogen lamp (GTE Sylvania 500T3/C/L-120V) with or without a Lee polyester #118 blue-green light filter.

Fig. 1. Spectral output of a tungsten lamp configured in a photosynthetic (see text) in the absence (solid line) and presence (dashed line) of a Lee polyester #118 blue light filter. The wavelength ranges for light absorption by major pigments present in oceanic phytoplankton are indicated by horizontal bars and include Chlorophyll (Chl) a, accessory Chl b and c, xanthophylls and phycoerythrin (PE). Absorption peaks for phycoerythrobilin (PUB) and phycoerythrobilin (PEB) chromophores within PE are indicated by
screen (Fig. 1). The spectral output of the transmitted light was measured at 5 nm intervals over the visible spectrum, using a Jarrel-Ash model 82-440 Ebert monochromator configured to reduce scattering to 0.0001 %. The monochromator was placed 10 cm from the light source, the path length through the monochromator was 50 cm, and a Li-cor collector was placed at the monochromator exit slit. Emerging light from the monochromator completely covered the irradiance collector, and measurements are reported as percent of maximum spectral output. Data were corrected for the spectral sensitivity of the Li-cor sensor.

Following incubation (2 to 3 h) in one of 4 white or blue-green light 'photosyntheticrons' (Lewis & Smith 1983) water-cooled to in situ temperature (± 2°C), both blue-green and white light samples were filtered onto 25 mm Nuclepore filters of 0.2, 0.6, 1.0 and 5.0 μm pore size. Methods of filtration, washing and addition of scintillation cocktail were based on methods described in Prézelin et al. (1986, 1987b). While at sea, isotope incorporation was determined on a portable LKB 1217 scintillation counter which was interfaced with an Apple II microcomputer for data storage and manipulation. Photosynthetic parameters were derived from P-I curves, using procedures detailed in Prézelin et al. (1987b). This approach does not allow an independent assessment of the standard deviation of individual parameters. However, independent determinations of the mean P_{max} from averages of data points on the light-saturated portion of the P-I curve, indicated that one standard deviation of P_{max} was routinely less than 15 %, and often less than 10 %, of the mean (n = 4 to 10). The lower limit of detection for I_{e} (the minimum irradiance required to light-saturate photosynthesis) was 3 μE m^{-2} s^{-1}. Alpha was usually determined as the slope of the light-limited region of the P-I curve. On occasions where I_{e} values were very low (< 3 μE m^{-2} s^{-1}) and insufficient data points were available in the light-limited region of a P-I curve, maximum alpha values were estimated as the quotient P_{max}/I_{e}. Rates of photosynthesis were determined directly for > 0.2, > 0.6, > 1.0, and > 5.0 μm fraction and by mathematical deduction for the 0.2–0.6, 0.6–1.0 and 1–5 μm fractions.

Estimates of water column productivity rates were derived from calculations of in situ photosynthetic performance, P_{r}, given the relation

\[ P_{r} = P_{max} \tanh (I/I_{e}) \]

which requires knowledge of both the diurnal (dawn-to-dusk) variations in underwater irradiance (I) and the diurnal patterns in size-fractioned P-I parameters as a function of depth in the water-column (Smith et al. 1987). By combining these estimates of in situ diurnal productivity patterns for each size fraction with measurements of midday vertical profiles of in situ Chl and primary productivity (Prézelin unpubl.), as well as with data on Synechococcus chlorophyll distribution within different size classes as a function of depth and time of day, it was then possible to sum Synechococcus primary production for all size categories and determine the relative contribution that Synechococcus makes to total primary productivity throughout the day at any depth.

**RESULTS**

The physical and chemical characteristics of critical depths at the time of sampling in the present study are summarized in Table 1. Details of the temporal variability in water column characteristics during 5 d at each of the 2 stations are presented elsewhere (Glover et al. 1988a, b). Most notable was the difference in nitrate concentrations in surface waters at the 2 stations. At Stn 1 there was a transient increase in nanomolar (nM) levels of nitrate (to between 21 and 27 nM NO_{3} in the upper 25 m) which selectively stimulated a unispecific bloom of Synechococcus spp. (Glover et al. 1988a); at Stn 2 there was negligible temporal variation in phytoplankton biomass and in the surface isothermal layer the nitrate concentrations were less than 10 nM (Glover et al. 1988b).

At both stations, the seasonal thermocline occurred at 25 to 30 m (24 to 25°C) and the depths of the PE maxima, the Chl maxima and the base of the euphotic zones occurred on similar isotherms. PE maxima were located at 80 to 85 m, at 3 to 7 % I_{e}, and on the 21.5°C isotherm. While PE maxima were within the upper region of the nitracline, ammonium was the major form of inorganic nitrogen assayed at these depths (Table 1). Broad Chl maxima occurred 15 to 30 m below the PE maxima and across the 1 to 3 μm fractions. The depth of the Chl maximum was shallower at Stn 1, where the downwelling attenuation coefficient (K_{par}) was 54 % higher than at Stn 2 (0.056 and 0.037 respectively). Both Chl maxima were located near the 21°C isotherm at nitrate concentrations around 600 nM, although there was twice as much ammonia measured at Stn 1 in comparison to Stn 2. The base of the euphotic zone at the 0.3 to 0.5 % I_{e} light depths were at near identical temperatures and were enriched to μM levels of nitrate at both stations. Ammonia levels continued to increase with depth at Stn 1 but decreased at Stn 2, resulting in relative proportions of NH_{4}:NO_{3}:NO_{2} of 1.6:1.0:0.07 and 0.2:1.0:0.14 at the base of the euphotic zone at Stns 1 and 2 respectively. More than 88 % of the integrated standing crop of Chl a at both stations occurred in the < 5 μm fraction (Glover et al. 1988b), indicating ultraphytoplankton dominated the photosynthetic communities. Within the ultraphytoplankton, the depth distribution of PE-
fluorescing Synechococcus spp. and Chl-fluorescing algae differed distinctly (Fig. 2). At both stations, maximum Synechococcus cell abundance maxima were at light depths ≥ 50% \( I_c \), where these cyanobacteria numerically dominated the ultraphytoplankton (note difference in scales used in Fig. 2). Given that Synechococcus cell abundance maxima in surface waters were not coincident with depths of the cyanobacteria-specific PE maxima found deeper in the euphotic zone, it was apparent that the deeper populations of Synechococcus were comparatively dilute in number but were enriched in cellular light-harvesting phycocerythrin (for greater detail, see Glover et al. 1988b). Conversely, abundances of Chl-fluorescing ultraplankton were low in surface waters but increased within the nitracline where both the PE and Chl maxima occurred. Near and below the 2% \( I_c \) depth, abundances of ultraplankton algae and Synechococcus cells are approximately equal within the same community.

A comparison of blue-green versus white light effects was made for Chl-specific rates of photosynthesis in whole water communities (>0.2 \( \mu m \)) under light-saturating \( (P_{max}) \) and light-limiting (alpha) conditions. The values of these parameters as well as the related variable \( I_k (P_{max}:alpha) \) are displayed in Fig. 3 to 5. Diurnal

![Graphical representation of data](image-url)
patterns of daytime changes in \( P_{\text{max}} \) and alpha were generally the same when measured under blue-green or white light incubation conditions (an exception being the alpha values for \( P_{\text{max}} \) at Stn 2, Fig. 4d). Midday maxima in photosynthetic rates were not routinely observed, the highest rates of light-saturated and light-limited photosynthesis being measured during early morning hours. The magnitude of daytime changes in Chl-specific \( P_{\text{max}} \) damped with depth, being greatest in surface waters (about 3-fold) where assimilation rates were also highest (Fig. 3). The magnitude of daytime changes in Chl-specific alpha were as high as 8-fold but did not show a depth-dependent relationship (Fig. 4). Highest alpha values were observed at the base of the euphotic zone and within the PE max at Stn 2. Since changes in \( P_{\text{max}} \) and alpha did not covary, diurnal patterns in \( I_k \) were evident and were distinctly different at individual depths at the 2 stations (Fig. 5). With the exception of the base of the euphotic zone, highest \( I_k \) values at all other sampling depths were observed at midday and were up to 7-fold higher than either dawn or dusk measurements. Diurnal variations in \( I_k \) dampened with depth and mean daily \( I_k \) values declined with depth.

For most samples, blue-green and white light incubation conditions did not influence the measured rates of Chl-specific \( P_{\text{max}} \). One exception was evident during morning within the PE max at Stn 1 (Fig. 3c), when blue-green light measurements of \( P_{\text{max}} \) (\( P_{\text{max blue}} \)) were twice as great as white light measurements of \( P_{\text{max}} \) (\( P_{\text{max white}} \)). Conversely, a second exception was detected in surface waters at Stn 2 where the value of \( P_{\text{max blue}} \) measured throughout the day were only about half of those measured for \( P_{\text{max white}} \) (Fig. 3b). Blue-green light measurements of alpha (\( \alpha_{\text{blue}} \)) were routinely equal or higher than white light alpha (\( \alpha_{\text{white}} \)) (Fig. 4). There was one exception at the base of the euphotic zone, where estimates of alpha were particularly difficult to make owing to the very low biological activity of the samples. The magnitude of \( \alpha_{\text{blue}}/\alpha_{\text{white}} \) varied with time of day and was greatest at Stn 2 in PE max populations.
Fig. 5. Comparison of blue-green (●) and white (∅) tungsten light measures of $I_k$ ($= P_{max} \alpha$) for whole water phytoplankton communities ($> 0.2 \mu m$) sampled over the day from (a, b) the surface (2 m); (c, d) the PE max; (e, f) the Chl max; and (g, h) the base of the euphotic zone ($< 0.5 \% I_o$) at 2 stations in the Sargasso Sea.

dominated by PE-enriched Synechococcus populations (Fig. 3). The resultant combined effect of blue-green light incubation conditions on estimates of both $P_{max}$ and alpha was that whole water samples required much lower irradiances of blue-green light than white light to saturate in situ rates of community photosynthesis. These observations are evident in the much lower $I_k$ values measured under blue-green light ($I_k$blue) than white light ($I_k$white) for all samples except the PE max at Stn 1 (Fig. 5).

In situ photosynthetic performance ($P_i$) was estimated from combined knowledge of P-I parameters and the underwater light field. A comparison of $P_i$ estimates derived from white light P-I parameters and blue-green light P-I parameters for whole water ($> 0.2 \mu m$) communities and the 0.6–1 $\mu m$ size fraction is given in Table 2. Note that in some instances the differences between blue-green and white light estimates were independent of time of day, while in other instances blue-green vs white light $P_i$ estimates were much greater at one time of day than another. For samples taken from below surface waters, blue-green light incubations generally gave rise to higher estimates of in situ productivity. The difference between white and blue-green light estimates of $P_i$ showed an increasing trend with depth for whole water communities. In surface waters, blue-green light estimates were similar to or were significantly less than white light estimates of photosynthetic performance. Interestingly, the magnitude of the blue-green vs white light increases were similar for whole water communities sampled at similar light depths at the 2 stations. For the 0.6–1 $\mu m$ size fraction, blue-green light incubation led to 3-fold higher estimates of photosynthetic performance at Stn 1 and 3-fold lower estimates of photosynthetic performance at Stn 2. Within the PE max at both stations, blue-green light estimates were about 40% higher than white light estimates. Within the Chl max at both stations, blue-green light estimates for the 0.6–1 $\mu m$ fraction were 2 to 3-fold higher than white light estimates.

To further define the differential effect blue-green light incubations might have on Synechococcus and algal subpopulations of ultraphytoplankton within the
Table 2. Blue-green:white ratio of estimates of in situ rates of photosynthetic performance ($P_I$) in subsurface whole water (> 0.2 μm) and 0.6–1.0 μm size-fractioned communities from 2 stations in the Sargasso Sea in July–August 1986. Samples were collected at different times of the day and replicates incubated under equal fluences of blue-green or white light. Estimates of $P_I$ were derived from knowledge of $P_I$ parameters and the underwater light field $PE_{max}$ and $Chl_{max}$ as in Table 1.

<table>
<thead>
<tr>
<th>Sample depth</th>
<th>Blue-green $P_I$: White $P_I$</th>
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</thead>
<tbody>
<tr>
<td></td>
<td>Dawn</td>
<td>Noon</td>
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<tr>
<td>&gt; 0.2 μm</td>
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<td></td>
</tr>
<tr>
<td>Surface (2m)</td>
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<td>1.03</td>
</tr>
<tr>
<td>Stn 1</td>
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<td></td>
</tr>
<tr>
<td>Stn 2</td>
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<td>0.54</td>
</tr>
<tr>
<td>PE max</td>
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<td></td>
</tr>
<tr>
<td>Stn 1</td>
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<td>1.37</td>
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<tr>
<td>Stn 2</td>
<td>2.75</td>
<td>0.87</td>
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<tr>
<td>Chl max</td>
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<td>Stn 1</td>
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<td>1.98</td>
</tr>
<tr>
<td>Stn 2</td>
<td>1.76</td>
<td>0.99</td>
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<tr>
<td>Base of euphotic zone</td>
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<td></td>
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<tr>
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<td>3.88</td>
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<td>Surface (2m)</td>
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<tr>
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<td>Base of euphotic zone</td>
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<tr>
<td>Stn 1</td>
<td>&gt; 6.00</td>
<td>–</td>
</tr>
<tr>
<td>Stn 2</td>
<td>–</td>
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</table>

-- Insufficient data to complete calculations

Fig. 6. Comparison of percentage of total Chl biomass attributable to *Synechococcus* spp. within different size fractions sampled at the surface, within the $PE_{max}$ and $Chl_{max}$, and the base of the euphotic zone (< 0.5 % $I_{s}$) at 2 stations in the Sargasso Sea.

Fig. 7 and 8 display the ratios of blue-green vs white estimates of Chl-specific $P_{max}$ and alpha within different phytoplankton size fractions as a function of depth at the 2 stations. Surface values were sorted by time of day, while values for other depths represent the means of daily averages. For Stn 1, whole community (> 0.2 μm) measures of $P_{max}$, blue: $P_{max}$, white approximated 1:1 and most subfractions within the same sample reflected whole community values. Notable exceptions were at midday within surface 0.6–1 and 1–5 μm fractions, exclusively comprised of *Synechococcus* Chl biomass, where blue-green light measures of $P_{max}$ were ≥ 4 times greater than white light measures of $P_{max}$. Similar blue-green light enrichment of $P_{max}$ values was observed at various times of day for *Synechococcus*-dominated populations within the PE max, but were less evident in size fractions where mixed populations of cyanobacteria and eukaryotic algae were found. At Stn 2, whole communities sampled from surface waters over the day...
and incubated under blue-green light actually showed only half the photosynthetic potential of white light samples. With increasing depth, whole water values for $P_{\text{max, blue}}:P_{\text{max, white}}$ steadily increased toward a value of 1.5 at the base of the euphotic zone. Size fractions within whole water communities showed similar $P_{\text{max, blue}}:P_{\text{max, white}}$ values, with the one exception of the 0.6–1 μm mixed community of the Chl maximum.

Blue-green light enrichment of measured rates of light-limited photosynthesis was common at all depths at both stations, with greatest alpha$_{\text{blue}}$:alpha$_{\text{white}}$ ratios associated with size fractions in which Synechococcus Chl biomass dominated phytoplankton samples (Figs. 6 and 8). Greater than 5-fold increases in Chl-specific alpha were evident when alpha$_{\text{blue}}$:alpha$_{\text{white}}$ ratios were compared for surface 0.2–0.6 and 0.6–1 μm fractions at Stn 1 (95% Synechococcus Chl) and 0.2–0.6 μm fraction at Stn 2 (72% Synechococcus Chl), and Chl max 0.6–1 μm fraction at Stn 1 (60% Synechococcus Chl). Working with only surface and PE max 0.6–1 μm size fractions, in which Synechococcus spp. accounted for ≥ 95% of Chl biomass (Fig. 6), it was evident that Chl-specific blue-green light alpha values increased in direct proportion ($r = 0.94$) to the PE concentration within these cyanobacteria (Fig. 9).

Photosynthetic parameters derived from white light measurements of P-I relations were used to estimate in situ rates of photosynthetic performance ($P_I$) for whole water (> 0.2 μm) communities and Synechococcus subcomponents and these were compared to estimates based upon blue-green light incubations. At Stn 1, the impact of blue-green light incubations was to increase total water column production estimates by less than 6% (from 360 to 380 mg C m$^{-2}$ d$^{-1}$) but to increase estimates of Synechococcus contribution to that production from 57 to 73% (Fig. 10). Similarly at Stn 2, the use of blue-green light incubation had much less effect on estimates of whole community production (which increased 13% under blue-green light conditions) than on estimates of Synechococcus spp. contribution to water column productivity (Fig. 11). Due primarily to increases in alpha estimates under blue-green light conditions for PE max populations of Synechococcus, estimates of Synechococcus spp. contribution to total water column productivity rose from 61% under white light conditions to 84% under blue-green light conditions at Stn 2 (Fig. 11).

**DISCUSSION**

The determination of photosynthesis-irradiance (P-I) parameters in phytoplankton communities is important because they can be used to discern temporal/spatial patterns of in situ productivity, quantum yield, photosynthetic potential and/or susceptibility to photoinhibi-

![Fig. 7. Comparison of ratio of blue-green vs white tungsten light measures of $P_{\text{max}}$ in different size fractions of phytoplankton sampled over the day from the surface (2 m). PE max, Chl max and the base of the euphotic zone (< 0.5% $L_0$) at 2 stations in the Sargasso Sea. Ratios of $P_{\text{max, blue}}:P_{\text{max, white}}$ for whole water communities (> 0.2 μm) are shown as black bars; ratios for size-fractioned (0.2–0.6, 0.6–1, 1–5, > 1 > 5 μm) communities are shown as hatched bars. In surface samples, noon time data was sorted out and compared to the mean of replicate measurements made at dawn and/or dusk. At PE max, Chl max and base of euphotic zone, the mean of all daytime measurements ($n = 1$ to 3) of blue-green:white $P_{\text{max}}$ ratio is presented. Vertical bars indicate 1 standard deviation of means](image-url)
features that may reduce their susceptibility to spectral bias by the tungsten incubation lamp source. First, the high abundance and species diversity within coastal assemblages may result in a spectral averaging of light absorption by the whole community that overcomes the spectral bias of the light field. Second, photosynthetic absorption and action spectra suggest that red and blue-violet light are functionally equivalent in light-harvesting capabilities by the accessory Chls that drive much of the photosynthetic activity of chromophytic and Chl b-containing organisms that dominate coastal waters (Prézelin & Boczar 1986). Lastly, there is less spectral dependency in the distribution of absorbed photosynthetic excitation energy between photosystems I and II in Chl c- and Chl b-containing algae than in the phycobilin-rich systems of cyanobacteria (Prézelin & Boczar 1986), which represent only a small fraction of the phytoplankton biomass in coastal regions.

The present study was prompted by questioning whether the spectral bias of ‘white’ light might be inappropriate for estimating oceanic primary productivity and in defining the photophysiological characteristics of cyanobacterial populations of PE-rich Synechococcus spp. One reason for concern arose from the knowledge that these cyanobacteria routinely dominate phytoplankton species abundance above the Chl max in the open ocean (Glover et al. 1988a, b) and blue-green light absorbed by PE alone can account for a large majority of the photosynthetically fixed carbon (Lewis et al. 1987, Boucher et al. unpubl.). The use of conventional tungsten ‘white’ light sources in photosynthetrons would therefore minimize the light-harvesting capabilities of PE and lower its contribution to photosynthetic carbon fixation. Some of the spectral bias in ‘white’ light sources can be overcome by placing colored filters over the tungsten lamps, such that the spectral output of the incubation illumination both simulates the spectral quality of light fields within the upper water-column of the open sea (Jerlov 1976) and overlaps the broad 485 to 565 nm maximum in spectral quantum yield for PE-driven photosynthesis that is characteristic of PE-rich cyanobacteria (Lewis et al. 1986, Boucher et al. unpubl.). The aim of the present study was to assess the impact of blue-green versus ‘white’ light incubation on primary productivity estimates of Sargasso Sea communities dominated by Synechococcus and to determine if any observed differences could be linked to other photophysiological characteristics of these cyanobacterial populations.

Blue-green versus ‘white’ light effects on P-I parameters

Diurnal (daytime) periodicity of $P_{\text{max}}$ and alpha have been documented for ultraphytoplankton communities
Fig. 10. Depth profiles of daily integrated rates of in situ primary production at Stn 1 in the Sargasso Sea, comparing estimates derived from (A) white light measures and (B) blue-green light measures of diurnal carbon fixation. Summation of water-column daily integrated rates of production are shown as a pie-diagram beneath each depth profile. Fractional contribution by Synechococcus spp. subpopulations to whole communities production (> 0.2 μm, length of bars) is shown by the black bars on the depth profile and the black portion of the pie-diagrams dominated by PE-rich Synechococcus (Puit & Prézelin 1985, Prézelin et al. 1986, 1987b, c). These studies demonstrated depth-dependent and water mass-dependent changes in the timing and amplitude of daytime variations in photosynthesis, which were independent of Chl concentrations and dark rates of carbon fixation. Generally, the timing of peak photosynthetic potential occurred between dawn and midday and were or were not coincident with diurnal variations in alpha. The amplitude of diurnal variations in P_max and alpha tended to dampen with depth. Laboratory experiments have shown that diel periodicity in Synechococcus photosynthesis is not driven by a biological clock (Sweeney & Borgese 1988), but is linked to changes in cell cycle photobiology (Boucher et al. unpubl.). In the present study, diurnal patterns of P_max and alpha in whole water samples at both stations were characterized by low amplitudes and when evident at all, peak activities occurred at times other than midday. While the possible mechanisms to account for wavelength-dependence of P_max and alpha are described below, it can be concluded that time of day can influence the degree of spectral dependency evident in the photosynthetic activity of an oceanic phytoplankton community.

The present study generally demonstrated P_max to be a wavelength-independent parameter for oceanic ultraphytoplankton (< 5 μm) communities (Fig. 7). However, there were 2 clear exceptions, both involving surface ultraphytoplankton that were almost exclusively comprised of Synechococcus cells. First, at Stn 1 there were 2 surface subpopulations of Synechococcus in 0.6-1 and 1-5 μm fractions, which showed a greater than 4-fold increase in P_max when incubated under blue-green instead of white light at midday (Fig. 7). Concurrent measurements determined that there was
no discernable change in the PE/Chl ratio within these size fractions at this time (unpubl.). This $P_{\text{max blue-green}}$ increase was not evident at all in independent whole water estimates, which combined ultraphytoplankton communities with the $>5 \mu m$ fraction (Fig. 3). One explanation for the blue-green light increases in *Synechococcus* $P_{\text{max}}$ might be a midday 'white' light-driven photoinhibition, whereby the unrealistic biasing in incubation illumination away from blue-green to red wave bands could have altered the cellular balance between the various deexcitation mechanisms which normally protect the photosynthetic components from photoinhibitory effects (cf. Powles 1984).

The second example of spectral effects on estimates of $P_{\text{max}}$ occurred in surface waters at Stn 2, where throughout the day all size fractions of ultraphytoplankton communities consistently gave blue-green light $P_{\text{max}}$ estimates that were only half of those measured under 'white' light (Fig. 7). Like Stn 1, *Synechococcus* populations accounted for $\geq 90\%$ of all ultraphytoplankton Chl in surface waters (Fig. 6). It is not clear whether these results indicate a blue-green light induced decrease or 'white' light induced increase in $P_{\text{max}}$, but it was apparent that the difference occurred independently of cyanobacterial size or time of day. One possible explanation is that nitrogen limitation in the Stn 2 *Synechococcus* population could be linked to a general mobilization of PE protein reserves to such an extent that the light-harvesting phycobilins were effectively decoupled from photosynthesis. Nitrate concentrations in surface waters at Stn 2 were less than half of
those measured at Stn 1, where a recent input appeared to have induced a *Synechococcus* bloom that was characterized by cells with a high PE content (Glover et al. 1988a, b). In contrast, surface *Synechococcus* cells at Stn 2 had only half the PE content as those at Stn 1, but contained the same amount of Chl. These observations are consistent with laboratory studies, which have shown nutrient deficient cyanobacterial cells contain normal Chl levels, while phycobilins are degraded and has led to the conclusion that light-harvesting phycobilins such as PE can also function as a nitrogen reserve (Boubissa & Richmond 1980, Wyman et al. 1983, Carr & Wyman 1986). Furthermore, the spectral quantum yield for photosynthesis, which is driven by PE light absorption, is markedly reduced in nutrient-depleted *Synechococcus* cells, while that of Chl remains about the same (Lewis et al. 1986, Boucher et al. unpibl.). If nutrient stress resulted in a decoupling of PE from the Chl-containing phototrops of *Synechococcus*, then photosynthesis could have been driven primarily by blue and red light absorption of cyanobacterial Chl. This explanation would account for the lack of spectral dependency on Chl-specific alphas in these surface samples (Fig. 8), whereby blue-green light absorbed by the blue Soret region of *Synechococcus* Chl is about as effective in driving carbon fixation as the tungsten ‘white’ light absorbed by the red Soret region of *Synechococcus* Chl (Boucher et al. unpubl.). Under such conditions, a spectral dependency in P max could result from an imbalance in the partitioning of absorbed light energy to Chls within photosystem I and II.

*Synechococcus* spp. numerically dominated ultraphytoplankton communities down to between 3 and 7 % I o, with maxima in cell abundance routinely located at ≥ 50 % I o (Fig. 2). It was within these populations that blue-green light routinely provided much higher (≥ 5-fold) estimates of light-limited rates of photosynthesis (alpha) than white light (Fig. 8). Concurrent with these observations, the PE content of *Synechococcus* cells increased with decreasing light and/or increasing nitrogen availability (Glover et al. 1988b), cells had a high ratio of phycourobilin to phycoerythrobilin chromophores (Campbell & Iturriaga 1989) and cellular rates of *Synechococcus* photosynthetic performance were equally high from surface waters down to the maximum in PE concentrations (3 to 7 % I o) (Prézelin & Glover unpubl.). These observations suggest that photoadaptive increases in the content of blue-green light absorbing PE was central to the maintenance of high photosynthetic rates by *Synechococcus* populations throughout a large portion of the euphotic zone. The present study also documents that the observed increases in PE/cell occurred in direct proportion to blue-green light measurements of photosynthetic quantum efficiency (Fig. 9), further indicating that these cyanobacteria are well suited to harvest the available light throughout a large portion of the euphotic zone. Hence, the color of the incubation light may be an important consideration in future studies aimed at determining the physiological bases of photosynthetic regulation in natural *Synechococcus* populations.

**Spectral effects on photosynthetic performance and contribution to primary productivity**

The present study emphasizes the comparison of photosynthetic rates measured in size-fractions of communities incubated under tungsten ‘white’ or blue-green spectral bands. However, we recognize that the base of the euphotic layer is dominated by blue-violet light (400 to 465 nm), which enhances light absorption and photosynthetic quantum efficiency in a variety of ultraplankton algae (Prézelin & Boczar 1986, Lewis et al. 1986, Glover et al. 1987). As a consequence, the present study may underestimate the productivity of eukaryotic algal communities and prokaryotic Prochlorophytes (Chisholm et al. 1987, 1988, Olson et al. 1988). Thus we are emphasizing *Synechococcus* spp., which dominated ultraphytoplankton communities from the surface down to depths just above the deep Chl maximum.

If the argument regarding the distinct nature of blue-green light photophysiology of *Synechococcus* is accepted, then blue-green light or in situ estimates of primary productivity may be more accurate than ‘white’ light estimates and the former may provide a better basis to determine their contribution to total primary productivity in the open ocean. Since blue-green light measurements tended to raise alpha and lower I o values, one impact on primary production estimates would probably come at depths where ‘white’ light measurements suggested light-limitation or photoinhibition. We observed that blue-green light incubations had the greatest impact on in situ productivity estimates just below the seasonal thermocline to the depth of the PE maximum (Figs. 10 and 11). Wavelength dependent effects on photosynthetic parameters were less evident within ultraphytoplankton communities at and below the Chl max, where phycobilin-rich organisms did not dominate and little difference in wavelength-dependent calculations of primary productivity were therefore found.

Since the blue-green light effects on individual size fractions tended to be greater than those expressed by the whole water sample, the percent contribution to total primary production by any fraction was also affected by the spectral quality of the incubation. A
comparison of the estimated contribution of the 0.6–1 μm Synechococcus dominated fraction to whole water rates of in situ photosynthetic performance is summarized in Table 2. With increasing depth, there is a clear trend toward increased estimates of total primary production when blue-green rather than ‘white’ light conditions were used.

In conclusion, the color of the incubation light did not alter the timing of daytime variations in Pₜ, although there were occasions when color of the incubation light did affect the magnitude of Pₜ measured throughout the day. In contrast, alpha was often wavelength-dependent and the ratio of blue-green vs ‘white’ light alpha measurements could depend on the time of day that the measurement was made. In general, small daytime variations in Pₜ combined with larger non-coincident changes in alpha gave rise to distinct diurnal patterns in Iₚ. The spectral dependency in Iₚ estimates were clearly evident, with greatest changes in amplitude occurring in surface waters. It was the spectral dependency on alpha and Iₚ that had the greatest impact on blue-green vs ‘white’ light estimates of in situ productivity, especially for Synechococcus-dominated size fraction. However, the difference between blue-green and white light estimates of daily integrated rates of primary productivity were small since Synechococcus populations were localized in the upper part of the euphotic zone where photosynthesis was light-saturated throughout most of the day. While spectral dependency of Synechococcus photosynthesis did not impact greatly on the estimates of water column productivity, it was an important photophysiological consideration that linked the relative quantum efficiency of photosynthesis to the cellular concentrations of PE in natural populations of Synechococcus. The results showed that increases in the cellular content of PE in Synechococcus populations were central to maintaining high photosynthetic rates throughout most of the euphotic zone.

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