

Ultraplankton distribution in surface waters of the Mozambique Channel — flow cytometry and satellite imagery

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ABSTRACT: The composition of ultraplankton (UP) in near-surface samples collected underway every 1 to 6 h from a ship sailing from Durban to the Seychelles was determined by flow cytometry, using both autofluorescence pigments and fluorescence DNA staining. *Prochlorococcus* (*Pro*) (17 to 160×10^3 cells ml^{-1}) numerically dominated the ultraphytoplankton (UPP), followed by *Synechococcus* (*Syn*) (4.5 to 57×10^3 cells ml^{-1}) and eukaryotic algae (EA) (0.6 to 4.2×10^3 cells ml^{-1}). The abundance of heterotrophic bacterioplankton (HB) was 0.4 to 1.3×10^6 cells ml^{-1} . A strong correlation ($r = 0.8$ to 0.97) was observed between sea-viewing wide field of view sensor (SeaWiFS) satellite estimates of total chlorophyll *a* (chl *a*) concentration and chl *a* concentration, abundance and biomass of EA as well as abundance and biomass of HB. This shows the potential for deducing spatial distributions of these 2 groups for ecosystem modelling using satellite data. Although the correlation between satellite chl *a* estimates and *Syn* chl *a* concentration was strong ($r = 0.83$ to 0.88), the correlation with its abundance and biomass was poor ($r < 0.6$) due to high variability (factor of 12) in cellular chl *a* content and to a lesser extent to diurnal cycles. The relationships were similar when either only daytime or all UP measurements were compared with the satellite data. No relationship was found between satellite data and *Pro* chl *a* concentration, abundance or biomass, even after correction for a pronounced diel cycle, suggesting that the SeaWiFS instrument might not detect *Pro* chl *a*.

KEY WORDS: Picoeukaryotic algae · *Synechococcus* · *Prochlorococcus* · Cyanobacteria · Bacterioplankton · Picoplankton · Remote sensing · Sea surface temperature · Sea surface colour · Chlorophyll

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INTRODUCTION

Monitoring of ocean colour using satellites provides high-resolution data about the spatio-temporal distribution of photosynthetic pigments in the surface waters (Garçon et al. 2001). A valuable addition to space-based studies of primary productivity (Behrenfeld & Falkowski 1997) would be the interpretation of these data in terms of the concentration of certain groups of abundant micro-organisms. Considering their concentrations (10^3 to 10^6 cells ml^{-1}) and size (0.2 to $5 \mu\text{m}$ in diameter) ultraplankton (UP) (Li 1995) should have the largest effect among organisms on light attenuation and reflection in the surface waters (Morel et al. 1993). In tropical and subtropical

oceanic waters, ultraphytoplankton (UPP) account for ~80% of total chlorophyll *a* (chl *a*) and phytoplankton biomass as well as for ~70% of total primary production, e.g. (Li & Harrison 2001, Maranon et al. 2001). Eukaryotic algae (EA) and cyanobacteria, the 2 main components of UPP, contribute equally to UPP biomass (Zubkov et al. 2000a). The carbon standing stock of heterotrophic bacterioplankton (HB), another component of UP, on average equals that of total phytoplankton (e.g. Li & Harrison 2001). However, despite the apparent UP domination in tropical to equatorial oceanic waters, according to our knowledge, there have been few comparative studies of satellite measurements and *in situ* concentrations of UP groups.

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Using flow cytometry (Olson et al. 1993), UP can be accurately enumerated and discriminated into 3 main cytometric groups: (1) Picoeukaryotic algae; (2) cyanobacteria, i.e. *Prochlorococcus* (*Pro*) (Chisholm et al. 1988) and *Synechococcus* (*Syn*) (Waterbury et al. 1979); and (3) other nonautofluorescent HB. The latter are usually visualised by staining cellular nucleic acids with fluorescence dyes (e.g. Marie et al. 1997). Another advantage of flow cytometry is the ability to generate large sets of high quality data necessary for studying large-scale distributions of UP (e.g. Buck et al. 1996, Li 1995, Li & Harrison 2001, Zubkov et al. 2000b). This ability is particularly useful for validating space-based biological measurements.

In the present study, we counted UP on a meridional transect along the Mozambique Channel with the aim of assessing the possibility of using satellite data for estimating the concentrations of UP groups in the surface waters.

MATERIALS AND METHODS

Sampling site. A total of 65 samples were collected from 23 to 29 August 2001 during a passage of the royal research ship RRS 'Charles Darwin' on a route from Durban (South Africa) to the Seychelle Islands (Fig. 1). The samples were collected from the ship's non-toxic seawater supply, drawn from a 5 m depth 10 or 11 times a day with hourly samples between 18:00 and 24:00 h local time and every 3 to 6 h during the rest

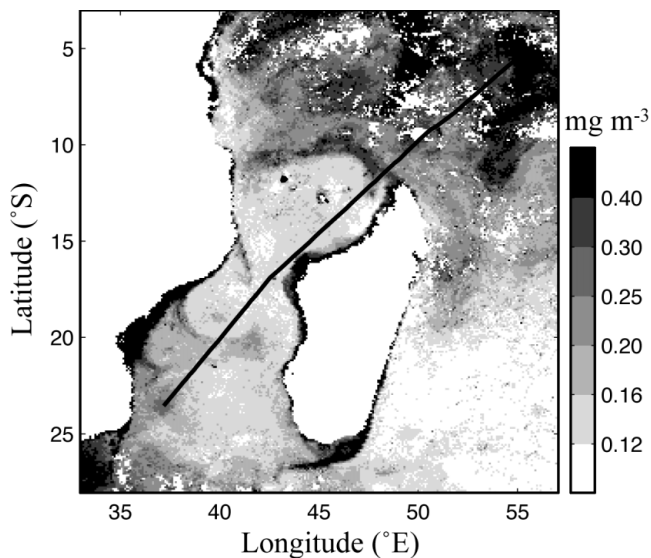


Fig. 1. Composite satellite image (SeaWiFS) of chlorophyll a spatial distribution in the Mozambique Channel from 23 to 29 August 2001, with the track of RRS 'Charles Darwin' superposed

of the day. The ship's coordinates and seawater temperature were recorded at the time of sampling.

Satellite data. Two sources of satellite data were used in this study. Sea surface temperature (SST) measurements were obtained from the Tropical Rainfall Measuring Mission (TRMM) microwave imager (TMI), processed by Remote Sensing Systems. TMI is a microwave instrument, enabling SST measurements through clouds. Ocean colour data were recorded by the sea-viewing wide field of view sensor (SeaWiFS), a visible light sensor on the SeaStar platform, launched by ORBIMAGE. The sea surface was observed at 1 km resolution in 8 frequency bands spanning 410 to 860 nm. Chl a concentration was determined from the radiances observed at 440, 490, 510 and 555 nm (O'Reilly et al. 1998), with the other channels being used in screening for clouds and correction for scattering from the atmosphere. We used the daily chl computations provided by the NASA Goddard Space Flight Centre at a resolution of 9 km. The flow cytometric measurements were compared with both the individual satellite passes and a weekly composite (the latter improves spatial coverage, see Fig. 1).

Flow cytometry. Replicated subsamples of 1.8 ml were taken from each collected sample, fixed with 1% paraformaldehyde (PFA), incubated at 2°C for 24 h and stored at -20°C before being analysed after their return to the laboratory. UP were analysed by flow cytometry (FACSort, Becton Dickinson), using a 15 mW 488 nm laser. UPP groups, namely EA, *Pro* and *Syn* were enumerated separately on the basis of the differences in their autofluorescence properties and light scattering (Olson et al. 1993, Zubkov et al. 1998) (Fig. 2). Nonautofluorescent, predominantly HB were enumerated after staining with the DNA dye, SYBR® Green I (Marie et al. 1997). Yellow-green beads, 0.5 µm in diameter (Polysciences), were used as an internal standard of 90° light side-scatter as well as of red fluorescence. The ratio of the mean side-scatter or red fluorescence intensity of a UP group to the respective bead intensity was used to normalise samples and to calculate relative values of side-scatter, as an index for mean cellular biomass (Bernard et al. 2000), and red fluorescence as a substitute for mean cellular chl a content (Li 1995).

The absolute concentration of beads in a standard stock suspension was determined by flow cytometric counting of beads in volumes dispensed with an automatic micro-injector (KD Scientific). The ratio of bead abundance to that of UP groups was used to compute the absolute concentration of the latter. The concentration of HB was calculated by subtracting the *Syn* and *Pro* concentrations, determined in unstained samples from the total bacterial concentration. An estimate of total UPP chl a or biomass was a sum of the estimated

group specific values. The latter were computed by multiplying the mean relative chl *a* or side-scatter values on group abundance.

Because of their high degree of synchronised division, cell cycle analysis of cyanobacteria was carried out to estimate their growth rates. The stained cyanobacterial cells were discriminated from other bacteria based on their higher red fluorescence due to the presence of chl *a*. Larger *Syn* cells scattered more light than smaller *Pro* cells. Cyanobacterial cells with a single copy of DNA (G_1 stage) were distinguished from those in which DNA was replicating (S stage) or had already replicated (G_2 stage), and the proportion of cells at S and G_2 stages was monitored during a diurnal cycle. The minimum growth rates of both *Syn* and *Pro* cyanobacteria were calculated according to Vaultot et al. (1995).

Data analysis. Acquisition and preliminary analysis of flow cytometric data were done using CellQuest software (Becton Dickinson). Correlation and regression analyses were used for comparison of the data sets at 99% confidence.

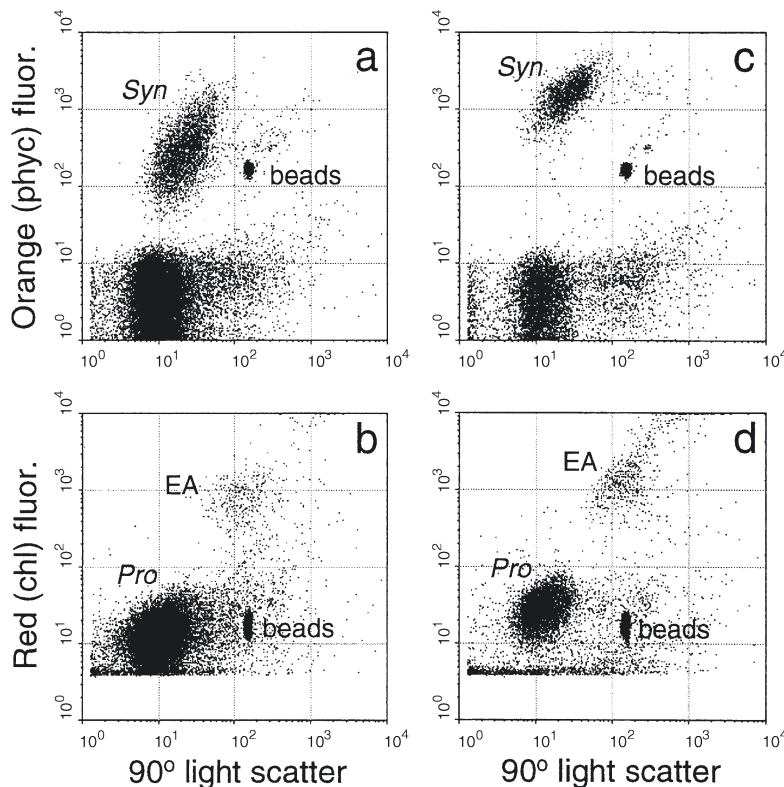


Fig. 2. Characteristic flow cytometric signatures of natural ultraplankton in the (a,b) Mozambique Channel and at (c,d) the approaches to the Seychelles. The *Synechococcus* (*Syn*) cluster (a,c) was revealed by its specific orange phycoerythrin (phyc) autofluorescence and excluded from the chlorophyll *a* (chl *a*) plots (b,d). The *Prochlorococcus* (*Pro*) (b,d) and eukaryotic algae (EA) (b,d) clusters were clearly resolved by their red autofluorescence of chl. Yellow-green beads, 0.5 μm in diameter, were used as an internal standard

RESULTS

The SeaWiFS data revealed the derived chl *a* concentrations to vary on quite small spatial scales (Fig. 1). High chl *a* concentrations, associated with upwelling, were found along many parts of the coast of Madagascar and Mozambique. In this region, eddies and currents play an important part in the spatial distribution of the chl *a* signature (Quartly & Srokosz 2003). To the south of Madagascar lies a region of high chl *a* waters, which have been upwelled near the coast and then entrained by the East Madagascar Current, which flows westward around the south of the island. At 20° S in the Mozambique Channel, there is a 100 km wide band of high chl *a* waters that have been advected into mid-channel from the Mozambican coast. De Ruijter et al. (2002) have noted 4 to 5 anticyclonic eddies per year heading south along the western edge of the channel, and Quartly & Srokosz (2003) have shown the effect these have on the chl *a* distribution. At the northern end of the channel, there is another sharp transition, presumably related to the westward-flowing North Madagascar Current entraining productive waters from the northern tip of the island.

The TMI data showed surface temperatures of around 25°C, with values higher in the shallower regions of the central Mozambique Channel and the approach to the Seychelles (55° E, 5° S). The ship and satellite records of temperature agreed closely (Fig. 3a), with a significant correlation at $p < 0.0001$, $r = 0.86$. The ship measurements, at a depth of 5 m, were systematically 0.8°C lower than the satellite measurements, which correspond to the top few millimetres of the sea surface.

Encouraged by the agreement of the temperature data sets, we compared SeaWiFS chl *a* measurements with chl *a* concentration, abundance and biomass of different UP groups. Although SeaWiFS was overflying the studied area daily at about 12:00 h local time, SeaWiFS does not cover the entire globe in 1 d, so there were only simultaneous satellite observations on 3 out of the 7 d of shipboard observations. To increase the number of comparisons with satellite measurements, we combined all satellite measurements along the ship track during the 7 d of on-board sampling. The mean values of chl *a* concentration are compared with individual observations on Fig. 3b. The coefficient of variance of mean values was rela-

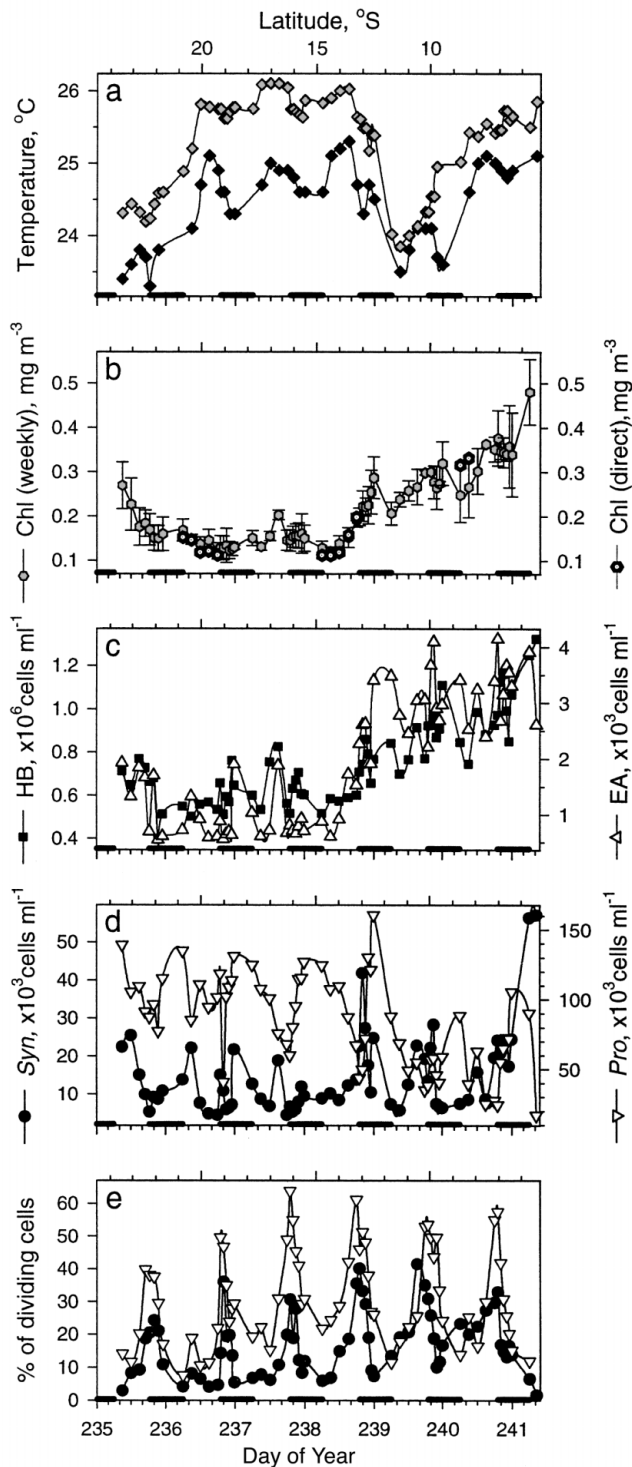


Fig. 3. (a) Latitudinal distribution of the shipboard (\blacklozenge) and satellite (\blacklozenge) sea surface temperature; (b) composite mean (\bullet) and direct (\bullet) concentrations of chlorophyll *a* (chl *a*) measured by SeaWiFS satellite; error bars show single SDs; (c) heterotrophic bacteria (HB) and eukaryotic algae (EA); (d) *Synechococcus* (*Syn*) and *Prochlorococcus* (*Pro*) abundance in surface waters; and (e) temporal variation of percentages of dividing cyanobacterial cells along the ship track (symbols as in d). Thick lines at the bottom show night periods

tively low (less than 30%). To reduce the possibility of artificial relationships caused by averaging satellite data, we correlated the weekly composite as well as individual satellite chl *a* measurements with flow cytometric measurements of UP groups (Fig. 3, Table 1).

The direct satellite measurements (Fig. 3b, black hexagons) were compared with matched daytime (from 06:00 to 18:00 h local time) measurements of UP parameters, assuming that the ocean colour did not change during the light period (Table 1, Direct). To minimise the effect of diel variability, a subset of UP measurements recorded during the daytime was correlated with composite satellite measurements (Table 1, Daytime). Finally, the whole set of onboard UP measurements was compared with the 7 d satellite composite (Table 1, All). Irrespective of the analysed data set the EA and UPP chl *a* correlated strongly with satellite chl *a* measurements. A similar relationship was found for the *Syn* chl *a*, except for the comparison of the direct data sets, most likely because of a small

Table 1. Pearson correlation coefficients and their significance (given by asterisks) for the various observed relationships between the SeaWiFS chl *a* concentrations and flow cytometric estimates of ultraplankton group measurements, such as chl *a* concentrations, abundance and biomass, done between 06:00 and 18:00 h on the day of satellite measurement (Direct), during the daylight period (Daytime), as well as with a complete data set of the transect measurements (All). Pearson coefficients of correlation higher than 0.8 marked in bold. Difference between 2 values insignificant: $p > 0.01$; significant: * $p < 0.01$, ** $p < 0.001$, *** $p < 0.0001$ (99% CI)

| Variables | SeaWiFS chl <i>a</i> | | |
|--------------------------------------|----------------------|---------------------|-----------------|
| | Direct (n = 12) | Daytime (n = 30) | All (n = 63) |
| Chl <i>a</i> | | | |
| Eukaryotic algae | 0.97*** | 0.83*** | 0.86*** |
| <i>Prochlorococcus</i> | 0.27 | 0.03 | 0.32* |
| <i>Synechococcus</i> | 0.57 | 0.88*** | 0.85*** |
| Total ultraphytoplankton | 0.81* | 0.93*** | 0.89*** |
| Averaged daytime chl <i>a</i> | | | |
| Eukaryotic algae | 0.94*** | 0.84*** | 0.89*** |
| <i>Prochlorococcus</i> | 0.05 | -0.03 | 0.02 |
| <i>Synechococcus</i> | 0.85** | 0.84*** | 0.83*** |
| Total ultraphytoplankton | 0.92*** | 0.89*** | 0.88*** |
| Abundance | | | |
| Eukaryotic algae | 0.93*** | 0.84*** | 0.87*** |
| <i>Prochlorococcus</i> | -0.7* | -0.56* | -0.48** |
| <i>Synechococcus</i> | -0.01 | 0.67* | 0.59*** |
| Heterotrophic bacteria | 0.91*** | 0.89*** | 0.87*** |
| Biomass | | | |
| Eukaryotic algae | 0.8* | 0.79** | 0.82*** |
| <i>Prochlorococcus</i> | -0.57 | -0.45 | -0.38* |
| <i>Synechococcus</i> | 0.11 | 0.67* | 0.57*** |
| Total ultraphytoplankton | -0.01 | 0.45 | 0.44** |
| Heterotrophic bacteria | 0.92*** | 0.86*** | 0.86*** |

data set combined with high variability of *Syn* cellular chl *a* content.

Each of the UPP groups comprised a substantial proportion of total UPP chl *a*: *Syn* $40 \pm 17\%$, EA $30 \pm 9\%$ and *Pro* $30 \pm 17\%$. Therefore, the very weak correlation between the *Pro* chl *a* and satellite chl *a* require an explanation. We plotted the satellite chl *a* versus the EA, EA+*Syn* and EA+*Syn*+*Pro* group sums, using the 3 data sets (Fig. 4). Unsurprisingly, our confidence in the slopes of the regression lines is dependent on the size of the data sets, emphasising the necessity of increasing the number of observations for more reliable regression approximations. A similar feature of all 3 plots was the parallel regression lines of the EA+*Syn* and EA+*Syn*+*Pro* group chl *a* sums that showed that the contribution of *Pro* chl *a* was similar at all stations, despite of the fact that *Pro* chl *a* concentration varied 6-fold along the transect. Most likely, the SeaWiFS sensor could not detect *Pro* chl *a* due to the physical properties of the *Pro* cells (e.g. small cell size, 0.5 to 0.6 μm), which is comparable to wavelengths of visible light.

One of the sources of variability of the UPP group chl *a* is diel change in cellular chl *a* content (Jacquet et al. 2002). We reduced the diel effect by using only daytime cellular chl *a* data. For each individual day, we compared mean chl *a* contents of the UPP groups using only measurements taken at 09:00, 12:00 and 15:00 h local time (as the relevant satellite overpass is near 12:00 h local time), and multiplied these mean values by the concentration of cell determined during the particular day, i.e. 'averaged daytime' chl *a*. The diel correction increased the correlations for the EA, *Syn* and UPP chl *a* (Table 1), confirming that diel cellular chl *a* variation made a moderate contribution towards spatio-temporal variability of the group chl *a* fields. However, although *Pro* had the most significant diel variation, the correction had no effect on its relationship with satellite chl *a*, supporting our speculation that the satellite instrument could not detect *Pro* chl *a*.

If one decides to employ real-time satellite colour data to parameterise marine ecosystem models, the strong empirical relationships between the satellite chl *a* and the EA and *Syn* chl *a* measurements would have limited usefulness, because additional conversion factors are required for converting the UPP group chl *a* estimates into the group abundance or biomass. To reduce the number of conversion steps, we looked for direct correlation between the latter 2 parameters and the satellite chl *a* (Table 1). Among the UP, only EA and HB group abundances have similarly strong relationship with the satellite chl *a*. Compared to the low variability of the EA and *Pro* cellular chl *a* contents (3 and 4.3 times, respectively), *Syn* cellular chl *a* varied 12 times, and consequently, the correlation between *Syn* abundance and satellite chl *a* was weaker than the

correlation between *Syn* chl *a* and satellite chl *a* measurements. As *Pro* has its greatest abundance in oligotrophic (chl *a* poor) waters, it has a weak negative correlation with the satellite chl *a* measurements.

The satellite data would be even more useful for modelling if one could use them for approximation of

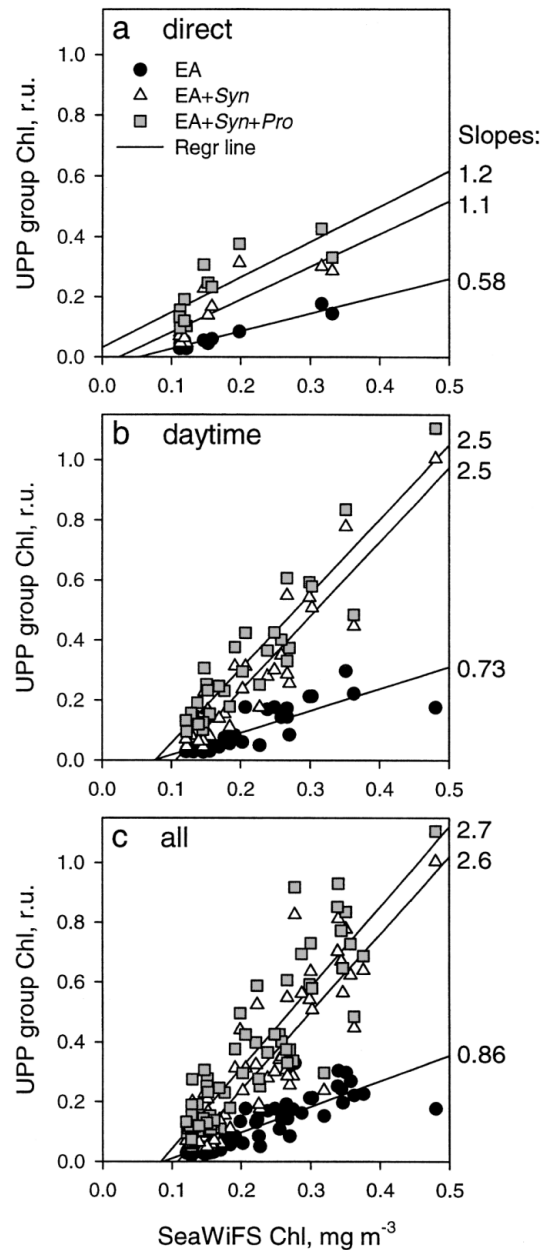


Fig. 4. Comparison of SeaWiFS chlorophyll *a* (chl *a*) measurements with (a) matched/direct, (b) daytime and (c) all measurements of chl *a* concentrations of ultraphytoplankton (UPP) groups: eukaryotic algae (EA) plus *Synechococcus* (*Syn*) and plus *Prochlorococcus* (*Pro*) in relative units (r.u.). Lines indicate linear regressions. Corresponding slope values are shown outside the plots. See details in the text and Table 1

the UP group biomass. In the present study, side-scatter of the UP groups was used as an index of cellular biomass. Only *Pro* cellular side-scatter correlated strongly with the *Pro* cellular chl *a* content ($r = 0.8$, $p < 0.0001$). However, because no significant relationship was found between satellite chl *a* measurements and *Pro* chl *a* concentration, there were also very weak correlation between *Pro* biomass and satellite chl *a*. Similarly to the abundance data, only the EA and HB group biomasses showed strong correlation with the satellite data.

DISCUSSION

The strong relationships found between the SeaWiFS chl *a* concentrations and abundance and biomass of EA and HB (Figs. 3b,c & 4) are very encouraging, because there is a potential for estimating concentrations of these 2 groups from satellite chl *a* data using linear regressions. For example: EA (cells ml^{-1}) = $-670 \pm 200 + 12\,000 \pm 850 \times \text{chl } a$ (ng ml^{-1}); HB (cells ml^{-1}) = $320\,000 \pm 32\,000 + 1\,900\,000 \pm 140\,000 \times \text{chl } a$ (ng ml^{-1}). The negative intercept of the EA regression can be explained by the fact that EA represent only part of a phytoplankton community, the total chl *a* amount of which is determined by the satellite.

On the other hand, the satellite data used in this comparison contained no values below 0.1 mg m^{-3} , whereas the cytometric measurements at a number of stations showed minimal concentrations of EA, *Syn* and *Pro* (see Fig. 4). In general, the SeaWiFS algorithm can yield chl *a* concentrations below 0.05 mg m^{-3} , so there would appear to be a bias in the particular satellite data set used here. This could be due to the satellite detecting chl *a* much deeper in the water column than the 5 m depth intake for the underway water supply system, or it could be due to thin undetected cloud increasing the atmospheric radiance, leading to a local bias in satellite-derived chl *a* values.

Higher concentration of total chl *a* would mean higher concentration of phytoplankton in general and EA in particular, and higher phytoplankton concentration leads to more organic nutrients for sustaining higher HB concentration. Using EA and HB groups (Fig. 3c) as indicators of productivity, one can speculate that the northern part of the Mozambique Channel, characterised by warmer waters (Fig. 3a), probably had lower productivity than the other traversed regions. The sharp decrease in temperature in the proximity of the northern tip of Madagascar indicated the water mass change and coincided with the increase in the EA and HB groups as well as total chl *a* concentrations.

Strong correlation between total chl *a* measurements and HB abundance is well documented for various

aquatic systems (e.g. Simon et al. 1992). An explanation of this relationship requires an assumption of steady state for the microbial planktonic community. HB populating the open oceanic waters are dependent on phytoplankton as the ultimate source of organic nutrients, and in the studied area, the microbial community was in homeostasis. There has been considerable doubt that concentrations of unpigmented HB are likely to be approximated by remotely sensed parameters (e.g. Zubkov et al. 2002), and before generalising, we would like to check the robustness of observed relationships in other oceanic regions and during other seasons.

The abundances of *Pro* and *Syn* cyanobacteria showed a weak correlation with satellite measurements (Table 1). In the case of *Syn*, it was most likely due to high variability of cellular chl *a* content. In the case of *Pro*, the relationship with chl *a* was negative, because usually the abundance of these cyanobacteria is greater in more oligotrophic and, consequently, chl *a*-depleted waters (Partensky et al. 1999). Our attempts to use other SeaWiFS wave band measurements (e.g. $555 \pm 10 \text{ nm}$) for correlating with *Syn* concentrations, exploiting their unique phycoerythrin orange fluorescence, were not successful.

For the required spatial coverage, the satellite images collected over a period of several days had to be composited, and consequently, the temporal resolution was sacrificed. Both EA and HB were comprised of many different species, and consequently, did not show pronounced diel cycles. Highly variable *Syn* and *Pro* abundance (Fig. 3d) were clearly controlled at an hourly scale not by nutrient sources, i.e. a 'bottom up' factor (Verity & Smetacek 1996), but more likely by a dynamic equilibrium of synchronised replication (Vaulot et al. 1995) and mortality – predatory pressure and/or viral infection, i.e. a 'top down' factor. The primary cause of the high variability of cyanobacterial abundance was a remarkable diel synchronisation of both *Syn* and *Pro* cell division along the track (Fig. 3e) with estimated minimum growth rates of 0.25 ± 0.037 and $0.35 \pm 0.042 \text{ d}^{-1}$ averaged for 6 full diel cycles, respectively. Up to 60% of *Syn* and *Pro* cells were generally going into division after dusk between 18:00 and 22:00 h. Similar division patterns of cyanobacteria were observed both in the Pacific (Vaulot et al. 1995) and Atlantic (Partensky et al. 1999, Zubkov et al. 2000b) Oceans, showing that this phenomenon appears to be global. The average minimum growth rate of *Pro* in the Mozambique Channel was about $\frac{1}{3}$ of *Pro* growth in the Equatorial Pacific and twice as high as estimated *Pro* growth in the oligotrophic Atlantic Ocean. The high diel variability seemed the most plausible explanation of the weak relationship between cyanobacterial abundance/biomass and satellite data;

however, we were able to determine that it was not the main cause.

Thus, the present study demonstrates the utility of a combination of flow cytometry and satellite remote sensing in surveying UP. It shows both the great potential and also certain limitations of current satellite remote sensing methodology. There is a possibility of using parameters like ocean colour for predicting distribution of UP organisms like EA and even HB. New spaceborne sensors with improved spatial and spectral resolution, such as MERIS on Envisat (Rast et al. 1999), may better enable the quantification of separate pigment concentrations, and thus, classification of the different phytoplanktonic groups present. However, monitoring of cyanobacteria from space will probably remain a challenge, until satellite instrument sensitivity is significantly improved and model corrections for diel cellular pigment adaptations are established and parameterised.

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