NOTE

Variation in photosynthetic rates among individual cells of a marine dinoflagellate

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ABSTRACT: Rates of photosynthesis for individual cells from a field population of the dinoflagellate Ceratium tripos were measured by incubating them in situ with NaH\(^{14}\)CO\(_3\) then assaying them, 1 cell per scintillation vial. The mean rate per cell was a maximum of 0.51 to 0.71 ng C cell\(^{-1}\) h\(^{-1}\) (estimated doubling time 5.9 to 4.2 d) at the surface and an order of magnitude less at greater depths or on cloudy days. For each of 4 sampling dates the estimate of carbon specific growth obtained for C. tripos using this single cell assay was correlated significantly with the estimate obtained for the entire phytoplankton community using conventional filter techniques. We used cultures of the related Ceratium longipes to prove the single cell assay against a direct measurement (absolute increase in carbon per cell) and obtained agreement within 5%. Individual cells from the same water samples differed in their absolute rates of tracer uptake, a previously unknown dimension of population structure in phytoplankton. Factors that may contribute to this variation are discussed.

The reliability of conventional \(^{14}\)C estimates of phytoplankton production in the sea continues to be disputed (Williams et al. 1983, Smith et al. 1984, Jenkins & Goldman 1985). Interpretation of the tracer uptake is equivocal because the experiments are done in entire microbial communities, including heterotrophs as well as autotrophs (Peterson 1980). We have overcome the ambiguity by measuring the assimilation of tracer in individual phytoplankton cells and by verifying the result against a direct measurement that is independent of both tracer methodology and food web structure.

Of the techniques available for estimating the production of individual phytoplankton species, only the single cell \(^{14}\)C assay (Rivkin & Seliger 1981) is easy to learn, requires little equipment and can be used for most phytoplankton species (>15 \(\mu\)m). The results from grain density autoradiography may not be quantitative and the technique of track autoradiography is difficult to master (Knoechel & Kalff 1976). Swift & Durbin's (1972) technique of estimating \(\text{in situ}\) division rates seems applicable only for dinoflagellates with synchronized division periods and gives average growth rates (Rivkin et al. 1984) which are not useful if the relation between carbon fixation and light is of primary interest. Post-incubation sorting of \(^{14}\)C-labelled cells can be done automatically with a flow cytometer but the procedure requires considerable capital outlay and operator training, is restricted in the size range of cells it can handle (approximately 1 to 50 \(\mu\)m) and cannot separate species if they are of the same size and have the same pigment complement (W. K. W. Li, pers. comm.).

To measure the uptake of \(^{14}\)C we used a modification of the method previously described by Rivkin & Seliger (1981). The principal changes were (1) individual cells were picked hydraulically using a drawn-glass capillary tube (Throndsen 1978) in a micromanipulator; (2) the washed cells were acidified directly in the scintillation vials to remove residual inorganic \(^{14}\)C; and (3) the radioactivity of individual cells was assayed, 1 cell per scintillation vial, until the 95% confidence limits were within 5% of the mean counts per minute.

The procedure was tested on a culture population of Ceratium longipes maintained in F/10 medium (Guillard 1975) at 15 \(^\circ\)C under 400 \(\mu\)Em\(^{-2}\)s\(^{-1}\) with a 14:10 light:dark cycle. A subsample of the culture was incubated under the same conditions for the first 7 h of the light period with NaH\(^{14}\)CO\(_3\) at a final activity of 1.0 \(\mu\)Ci cm\(^{-3}\). Some 100 cells were isolated for radioactive assay and the results used to calculate average uptake rates (2.27 ng C cell\(^{-1}\) d\(^{-1}\), standard deviation, \(s = 0.041\); daily rate = twice 7 h rate). Three aliquots of the main culture containing an estimated 46,000 cells were removed at the beginning and the end of the light period and filtered onto silver filters with a pore size of 0.8 \(\mu\)m. The carbon retained on the filters was measured on a Perkin-Elmer Elemental Analyser (model 240). The change in the amount of carbon per cell was
2.16 ng C cell$^{-1}$ d$^{-1}$ ($s = 0.356$), which was within 5% of the single cell estimate.

The species chosen for our field experiments was *Ceratium tripos*, a large dinoflagellate commonly occurring in blooms of extremely low species diversity (Elbrächter 1973, Falkowski et al. 1980). These blooms can result in anoxic conditions which cause extensive fish and shellfish kills (Falkowski et al. 1980) and are common in near-shore areas on both sides of the Atlantic ocean (Elbrächter 1973) including Bedford Basin (44°32’ N, 63°39’ W). During November 1984, the phytoplankton was sampled on 4 occasions from 3 depths (0.5, 5.0, and either 10.0 or 15.0 m) and incubated in situ for 2 to 4 h with NaH$^{14}$CO$_3$ at a final activity of 1.0 μCi cm$^{-3}$. Two light bottles (volume = 250 cm$^3$) and 1 dark bottle were used at each depth. Temperature and salinity were measured with a Guildline SCTD, and water samples for the determination of nutrients, chlorophyll, and carbon were collected in Niskin bottles. At the end of the incubation period three 25 cm$^3$ samples were taken from each bottle and filtered through Whatman GF/F filters. These filters were fumed over concentrated HCl then counted with a Beckman 3100 liquid scintillation counter. Dark respiration was corrected for by subtracting the mean cpm for the filters from the appropriate dark bottle. The remainder of the sample was filtered through 76 μm Nitex, rinsed well with filtered seawater, backwashed and resuspended. Some 25 cells were then processed using the single cell procedure described above.

For individual cells radiocarbon was assimilated to a mean activity of 1.2 to 74.0 dpm above background, depending on light intensity (incubation depth). Estimates of total phytoplankton carbon per m$^2$ were obtained from a regression of total particulate carbon on chlorophyll [slope = 62.6, $r^2 = 0.681$, $p < 0.01$, $N = 9$]. A mean value of 34.7 ng C cell$^{-1}$ was obtained for an average *Ceratium tripos* by picking out 500 cells, filtering them onto a precombusted GF/F filter, and analysing this on the elemental analyser as above. For each of the 4 sampling dates the estimate of carbon specific growth rate $\left(\frac{\Delta C}{\Delta t} \cdot \frac{1}{C}\right)$ obtained for *C. tripos* using the single cell procedure correlated significantly with that obtained for the entire phytoplankton community sampled from the same light bottle using the conventional GF/F filter procedure (Fig. 1e to h). The ratio of the single cell estimate to the conventional filter estimate was 0.76 at the beginning of November but had decreased to 0.2 by the end of November perhaps because faster growing phytoplankton became more abundant.

Downwelling irradiance at depth was calculated from surface light levels attenuated by the extinction coefficient. Surface light levels were calculated by multiplying the values obtained from the KIPP CM6 pyranometer located at Halifax Citadel (44°39’ N, 63°35’ W) by 0.45 to obtain the photosynthetically active radiation (P.A.R.) that penetrated the air-water interface. The extinction coefficient was measured with a LICOR LI-185A quantum meter.

The mean productivity per cell decreased with the decrease in downwelling irradiance at greater depths for all 4 sampling dates (Fig. 1a to d). It is inappropriate to make a composite plot of all the data because the time between sampling dates (minimum of 1 wk) is too long. The maximum rates of photosynthesis obtained for *Ceratium tripos*, 0.51 to 0.71 ng C cell$^{-1}$ h$^{-1}$ (Fig. 1a to d) at approximately 10°C, give doubling times of 5.9 to 4.2 d, assuming a daylength of 8 h and not correcting for dark respiration (which was not distinguishable from background even when 5 cells were assayed together). The rates of photosynthesis at greater depths were of course lower, 0.30 to 0.049 ng C cell$^{-1}$ h$^{-1}$ at 5.0 m (Fig. 1a to d), and gave doubling times of 10 to 61 d.

Given that nutrients were always measurable (nitrate > 1.5 mg-at m$^{-3}$), but that average light levels during incubation periods were relatively low, these estimates agree with those previously published for *Ceratium tripos*. Data from the conventional filter technique, carbon estimates and direct cell counts suggested a doubling time of 20 to 30 d for large *C. tripos* (25 ng C cell$^{-1}$) during a bloom in the New York Bight (Falkowski et al. 1980). And doubling times estimated by observing the maximum frequency of dividing cells in Kiel Bay gave values ranging from 2.5 d at 16°C to 22 days at 10°C (Elbrächter 1973). *C. tripos* from Kiel Bay were observed to be capable of surviving up to 41 d without dividing (Elbrächter 1973), so even the low rates we obtained for the cells from 10.0 m may enable survival until the cells are again higher in the euphotic zone.

Even though any 2 replicate light bottles had very similar means (Fig. 1 & 2) there was considerable variation among cells within any 1 bottle, with productivity of some cells being twice that of others (Fig. 2). The coefficients of variation (s/σ) range from 28.7 to 73% (N = 25 cells) for the *Ceratium tripos* incubated in the field and from 28.8 to 39.7% (N = 100 cells) for the *C. longipes* incubated in the laboratory. Less than 1.5% of the variance in the means is attributable to the 2-sigma counting error of 5% ([N(1-s$^2$) + Nx$^2$]/ (1600 N x$^2$)). The most probable causes of this variation are: (1) variation in cell age, (2) variation in cell size, and (3) variation in the cells’ environmental history.

Variation in productivity associated with cell age is confounded with variation associated with cell size
Fig. 1. Primary production in Bedford Basin on 4 sampling dates from in situ incubations at 3 sampling depths: (●) 0.5 m; (■) 5.0 m; (▲) 10.0 m or (▲) 15.0 m. (a to d) Productivity of *Ceratium tripos* as a function of light intensity. Each point represents the mean of some 25 cells isolated from the same incubation bottle (see text). (e to h) Correlation between the estimate of carbon specific growth rate \( \frac{1}{C} \frac{\Delta C}{\Delta t} \) obtained for *Ceratium tripos* and the estimate obtained for the entire phytoplankton community from the same incubation bottle (see text). The correlations between the 2 estimates are: 0.976, 0.995, 0.978, 0.994 for Nov 1, 13, 20, & 29, respectively.

since recently-divided cells are also smaller. *Ceratium longipes* and *C. tripos* divide synchronously during the dark period. Recently divided cells can be identified by the relative length of their apical horn to their antapical horns. We used the single cell uptake procedure to measure the average uptake over 6 h of recently divided *C. longipes* and compared this with that of 2 to 3 d old cells from the same culture flask. The mean production per cell was significantly higher for the older cells (Mann-Whitney U, p < 0.001). The lower production of the recently-divided cells does not result from an inhibition of photosynthesis by the division process. In a subsequent experiment involving three 1 h incubations begun at 06:00 h (the beginning of the light period), 08:00 h, and 14:00 h, the production of the recently-divided cells was always significantly less than that of older cells from the same culture (ANOVA, p < 0.0001) but even at 06:00 h the recently-divided cells took up \(^{14}\text{C}\) at 79 % the rate of the older cells. In our field experiments we used a large mesh size and avoided picking-out recently divided *C. tripos*. Variance in cell size may be important however: we found cell body diameters to vary from 60 to 75 μm from the same water sample.

Variation in the cells' environmental history occurs even among cells collected in the same water sample because of the stochastic nature of local turbulent mixing, and it can lead to variation in physiological properties (Lewis et al. 1984).

An additional contributing factor could be genetic differences among cells. The importance of this could be assessed by initiating cultures with single cells isolated from natural populations, growing at least 2 replicate flasks of each of these original cultures.
under the same environmental conditions for several generations, then using ANOVA to compare the productivity within and between the different cell lines.

The variation that we report here in assimilation rates among individual cells cannot be seen unless cells are assayed 1 cell per vial and incubated under conditions that result in the uptake of at least 10 dpm above background. Our procedure has the power to detect differences both between species and within populations, both features of merit for investigation of phytoplankton ecology in an evolutionary context. By avoiding complications that arise from food-web structure, we have shown that the 14C method is able to give an estimate of the rate of carbon fixation in phytoplankton that is both accurate and precise.

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LITERATURE CITED


Fig. 2. *Ceratium tripos*. Distribution in productivity of individual cells. Data from November 20, 1984. Data from the 2 replicate bottles at the same depth are combined into the same histogram but the mean is shown for each bottle. All 3 histograms are plotted at the same scale which might give the illusion that the data for 10.0 m are less variable than the data for 0.5 m. Comparison of the coefficients of variation (s/x) (0.5 m: 73.4, 44.8; 5.0 m: 61.2, 70.0; 10.0 m: 66.1, 33.7) shows that the variance in productivity is roughly the same at all 3 depths, as it was on the other sampling dates.


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