Springtime nutrient and phytoplankton dynamics on Georges Bank

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ABSTRACT: The dynamics of phytoplankton and nutrients before, during and after the winter-spring bloom on Georges Bank were studied on 6 monthly survey cruises from January to June 1999. We measured hydrography, phytoplankton cell densities, chlorophyll $a$, dissolved inorganic nutrients ($\text{NO}_3 + \text{NO}_2$, $\text{NH}_4$, $\text{Si(OH)}_4$, $\text{PO}_4$), dissolved organic nitrogen (DON) and phosphorus (DOP), particulate organic carbon (POC) and nitrogen (PON) and total particulate phosphorus (TPP). We present evidence that phytoplankton production may be significant year-round, and that the winter-spring bloom may have started in January. From January to April the phytoplankton was comprised almost exclusively of diatoms, reaching cell densities in March and April of ca. 450 cells ml$^{-1}$; chlorophyll $a$ concentrations exceeded 10 $\mu$g l$^{-1}$ in April. Diatoms decreased to relatively low levels in May (<50 $\times$ 10$^3$ cells l$^{-1}$) and increased again in June (>300 $\times$ 10$^3$ cells l$^{-1}$). Densities of dinoflagellates and nanoflagellates were low (<10 $\times$ 10$^3$ cells l$^{-1}$) from January to April, and increased in May and June to nearly 300 $\times$ 10$^3$ cells l$^{-1}$. Nitrate + nitrite concentrations in January were <3 $\mu$M in the shallow, central portion of the bank and decreased steadily each month. Silicate was also <3 $\mu$M over an even larger area of the central bank in January and declined to <1.5 $\mu$M over most of the Bank in April. The data suggest that silicate depletion, not DIN, contributed to the cessation of the diatom bloom. Regeneration of silicate occurred in May and June, presumably as a result of rising water temperatures in late spring which increased the dissolution rate of diatom frustules from the earlier diatom bloom. Dissolved organic nitrogen may have been utilized at the start of the winter-spring bloom; concentrations were ca. 14 $\mu$M in January, dropping to <6 $\mu$g l$^{-1}$ in February, after which DON concentrations steadily rose to >15 $\mu$g l$^{-1}$ in June. Overall micro- and nanoplankton biomass, measured as POC, PON and TPP, increased over the 6 mo period, as did nutritional quality of that biomass as indicated by declining C:N ratios. Our results suggest there may have been an increase in the heterotrophic component of the plankton in May and June which coincided with a second burst in diatom abundance. We discuss general features of planktonic production and nutrient dynamics with respect to year-round production on the Bank.

KEY WORDS: Nutrient Cycles · Phytoplankton · Primary production · Secondary production silicates · Nitrates · Georges Bank

INTRODUCTION

The biological productivity of Georges Bank, especially as it pertains to fisheries yields, is thought to be among the highest of any continental shelf sea (Backus 1987), with rates of primary production reported to exceed 400 g C m$^{-2}$ yr$^{-1}$ in the central portion of the Bank (O’Reilly et al. 1987). That production is known to be seasonal in nature and to exhibit a pronounced late winter-early spring phytoplankton bloom (Riley 1941, Cura 1987, Walsh et al. 1987, Townsend & Pettigrew 1997), but neither details of the seasonal cycle of primary production nor the dynamics of the spring bloom on Georges Bank are as yet well understood.
Much of what is known has been summarized by Backus (1987), and in recent years there have been renewed research activities as part of the US Globec Program (Global Ecosystems Dynamics, Wiebe & Beardsley 1996, Wiebe et al. 2001). New findings include those by Townsend & Thomas (2001), who reported that silicate is depleted before inorganic nitrogen and may be limiting to diatom production as early as February; regeneration of silicate may contribute to a second pulse in diatom production in late spring, concurrent with a developing nanoflagellate population (Kemper 2000). Inorganic nitrogen, on the other hand, appears to become depleted to levels that could limit phytoplankton growth by April (Townsend & Thomas 2001). For the remainder of the year, primary production is thought to be fueled largely by recycled nitrogen (Draxler et al. 1985, Horne et al. 1989, 1996). Fluxes of ‘new’ nitrogen (principally nitrate) delivered to the Bank from deeper waters around its edges appear to be too low to support significant levels of ‘new’ primary production (cf. Dugdale & Goering 1967, Eppley & Peterson 1979) across an entire submarine bank as large as Georges Bank, prompting Townsend & Pettigrew (1997) to argue that secondary production is likely to be nitrogen-limited. Consistent with that notion, Sherman et al. (1987) pointed out much earlier that the levels of zooplankton production on Georges Bank are anomalously low compared with rates of primary production; that is, total zooplankton production (microzooplankton plus macrozooplankton) on Georges Bank is about 18% of primary production, compared with the same ratio in nearby waters in the Gulf of Maine which is 26% (Cohen & Grosslein 1987). Thus, the nature of phytoplankton production, its dependence on nutrient dynamics, and their relationships to higher trophic level production on Georges Bank, a marine system renowned for its productive fisheries, remain puzzling in many respects.

This communication presents analyses of more complete data sets for Georges Bank, which encompass the presumed productive period from winter to the summer, with the hope of developing a better sense of how the basic biological oceanography of the Bank operates. Our results reinforce previous observations of the development of the winter-spring phytoplankton bloom on Georges Bank and suggestions of nitrogen and silicate limitation (Townsend & Pettigrew 1997, Townsend & Thomas 2001). In addition, we show that there was a species shift from nearly complete dominance of diatoms during the winter-spring bloom, to one of mixed diatoms and flagellates in May and June. During this 6 mo period there was a steady increase in overall plankton biomass and an increase in planktonic food quality (lower particulate C:N ratios). Much of this early summer plankton community depends upon recycled nutrients, especially nitrogen, and we argue that the heterotrophic component of plankton increases in May and June and facilitates the recycling of nitrogen, which drives the majority of planktonic primary production following the winter-spring bloom.

**MATERIALS AND METHODS**

Water samples were collected at stations sampled during survey cruises to Georges Bank in 1999 as part of the US Globec Program. The cruises were: 11 to 24 January (RV ‘Albatross IV’); 11 to 23 February (RV ‘Oceanus’); 10 to 23 March (RV ‘Oceanus’); 16 to 28 April (RV ‘Oceanus’); 19 to 27 May (RV ‘Albatross IV’); 14 to 24 June (RV ‘Albatross IV’). CTD casts for vertical profiles of temperature and salinity were made at 41 standard stations (Fig. 1); these hydrographic data were collected and processed by Dr. D. Mountain and M. Taylor of the Northeast Fisheries Science Center,
Woods Hole, Massachusetts. Water samples for analyses of nutrients and phytoplankton were collected during the same casts using Niskin bottles mounted on a rosette sampler. On all cruises we collected additional near-surface water samples (1 to 2 m depth) at positions halfway between the regular stations; these stations are numbered 42 to 81 (Fig. 1).

Samples for dissolved inorganic nutrients were filtered through 0.45 µm Millipore cellulose acetate membrane filters, after first flushing the filters with sample water. Samples were then frozen in 20 ml polyethylene scintillation vials by first placing the vials in a seawater-ice bath for about 10 min. Analyses for NO₃ + NO₂, NH₄, Si(OH)₄, and PO₄ were made on shore after the cruise, using a Technicon II AutoAnalyzer and standard techniques (Whitledge et al. 1986). Particulate organic carbon (POC) and nitrogen (PON) concentrations were determined at standard stations (1 to 41; Fig. 1) at 2 and 20 m depth, selected arbitrarily. Samples were collected by filtering 500 ml onto a preashed GF/F filter; the sample was frozen, and later fumed with HCl to remove inorganic carbon before analysis with a Control Equipment Model 240-XA CHN analyzer (Parsons et al. 1984). Samples for total particulate phosphorus (TPP) were collected as for POC and PON (but 200 ml were filtered) and the filters were frozen at sea and later analyzed using a modification of the method of Solórzano & Sharp (1980b). The filter was oxidized at 530°C for 2 h, digested in 1 ml 10% HCL for 24 h; 19 ml deionized water were then added; the sample was subsequently analyzed as for orthophosphate on the AutoAnalyzer. Concentrations of dissolved organic nitrogen (DON) and phosphorus (DOP) were determined by measuring total N and total P by the methods of Solórzano & Sharp (1980a,b) and subtracting the total concentrations of dissolved inorganic nitrogen and phosphorus; these samples were also collected at 2 and 20 m at the 41 standard stations. Phytoplankton chlorophyll a was measured fluorometrically on discrete water samples (Parsons et al. 1984). Samples of 100 ml were filtered onto GF/F filters, extracted in 90% acetone in a freezer (–18°C) for at least 6 h, and then analyzed at sea using a Turner Model 10 fluorometer.

Surface-water samples of 100 ml for phytoplankton cell densities and taxonomic composition were collected at 7 stations and preserved in Lugol iodine solution. The 7 stations were selected to represent the phytoplankton community on the Bank (Stns 3, 9, 11, 12, 20, 30 and 32; Fig. 1). A 50 ml subsample was allowed to settle in a graduated cylinder for 24 to 48 h, after which the top 40 ml were drawn off leaving a 5x concentrated sample. A 1.3 ml subsample of the concentrated sample was placed in a counting chamber and enumerated at either 100 or 400× magnification using a Nikon TMS inverted microscope. A minimum of 100 cells was enumerated for each station sample; on average, 200 to 700 cells were enumerated. The cells were identified as diatoms, dinoflagellates or other flagellates (nanoflagellates); lower taxonomic identifications were made only to record the dominant genera and species in each sample.

**RESULTS**

Seasonal distributions of near-surface temperature, salinity, chlorophyll a, nitrate + nitrite, silicate, phosphate and ammonium are presented as contour plots in Figs. 2 to 8 respectively.

Coldest surface water temperatures were not reached until March 1999, when temperatures were generally 4.5 to 5.5°C over much of the top of the Bank (Fig. 2). Slightly colder water could be seen moving onto the Bank as part of an intrusion of Scotian shelf water, but that intrusion was confined to the easternmost edge of the Bank. The characteristic lower salinity of the Scotian shelf water is also evident in Fig. 3. Slight warming of the top of the Bank only became evident during April, when surface temperatures were on average >5.5°C; however, during March and April much warmer (and saltier) surface waters could be seen along the southern flank of the Bank in association with slope water. The rate of springtime warming on the Bank accelerated between April and May, when temperatures rose by an average of 3 to 4°C. Temperatures rose another 3 to 4°C between May and June, reaching a bank-wide average of about 12°C in June. Vertical mixing by tides helped maintain a fairly uniform distribution of water properties throughout the water column over the entire Bank until May, when surface warming over deeper waters began to develop approximately along the 100 m isobath; this was most apparent in the southernmost portions of our sampling domain. Because of the nearly uniform vertical distributions of water properties, we refer here primarily to surface samples; cases of non-uniform vertical distributions are noted when appropriate.

Concentrations of chlorophyll a in the surface waters in January were greater than 1 µg l⁻¹ at most stations on the top of the Bank and >2 µg l⁻¹ at some (Fig. 4). By February, chlorophyll concentrations over the central portions of the Bank were >3 µg l⁻¹ (and >7 µg l⁻¹ at Stns 52 and 53) and increased to >5 µg l⁻¹ over much of the Bank in March and April, with overall highest chlorophyll concentrations appearing in April (>14 µg l⁻¹ at Stn 70).

The concentrations of nitrate + nitrite over the central portions of the Bank became steadily depleted between January and June, in keeping with the general
pattern of phytoplankton growth as revealed by chlorophyll distributions (Fig. 5). Concentrations of nitrate + nitrite in January were lowest (<1 µM) at Stn 10 on the central part of the Bank (Fig. 1), but were higher over most of the remainder of the Bank. By February, the patch of lowest nitrate + nitrite concentrations had become slightly larger in areal extent (Fig. 5), but the area encompassed by the 5.0 µM nitrate + nitrite concentration did not appear to change appreciably between January and February. Overall, dissolved inorganic nitrogen (nitrate, nitrite and ammonium) did not appear to be contributing significantly to the developing phytoplankton population during this initial period. By April, the shallowest central portions of the Bank were becoming depleted in nitrate + nitrite when much of the Bank had surface concentrations <2 µM. In May, only the northeast peak of the Bank and waters to the southwest (downstream) exhibited detectable levels of nitrate + nitrite; by June, only 1 station exhibited detectable levels. The shallower stations (inside the 60 m isobath) were generally vertically well-mixed, as indicated by isothermal temperature and isohaline salinity profiles, and consequently they did not exhibit vertical gradients in nutrient concentrations throughout the 6 mo period.

Silicate became limiting to the winter-spring diatom bloom before nitrogen (Fig. 6). In January, silicate levels were <3 µM over most of the top of the Bank, and a large area of the Bank exhibited concentrations <2 µM. The pattern of silicate depletion continued through February and March, and by April concentrations were undetectable at all but a few stations which were located along the southern flank. Perhaps by May, and certainly by June, there was clear evidence of a new flux of silicate over parts of the northern Bank; these increases were not associated with concomitant
and coherent increases in nitrate + nitrite, nor with colder and saltier water, as would be the case if the flux were the result of upwelling. Instead, these fluxes were probably associated with a regeneration of silicate on the Bank, as has been observed earlier (Kemper 2000, Townsend & Thomas 2001).

Concentrations of phosphate in surface waters in January were mostly in the range of ca. 0.5 to 1.0 µM (Fig. 7), and showed a steady decrease from January to June. The decrease was most apparent for the central portions of the Bank, similar to that of nitrate + nitrite and silicate. In May and June there was evidence of slightly higher concentrations in the vicinity of the northeast peak, consistent with upwelling and spreading of Gulf of Maine water into that area.

Surface concentrations of ammonium rarely exceeded 2.0 µM, and then only in isolated patches. Apart from these patches, the concentrations of ammonium reached a Bank-wide maximum in June, with average concentrations of 0.3 to 0.4 µM (Fig. 8).

Densities of diatoms, dinoflagellates and ‘other’ phytoplankton groups (mainly nanoflagellates) at 7 stations, selected to be representative of the Bank, are given in Fig. 9. Four stations are located on the top of the Bank, inside the 60 m isobath (Stns 11, 12, 30 and 32), and 3 are located along the eastern and southern flank between the 60 and 100 m isobaths (Stns 3, 9 and 20). The stations on the top of the Bank exhibited the highest densities of diatoms during the January to April period; densities were >450 × 10^3 cells l⁻¹ in February and March at Stn 12. The dominant taxa in January were the diatoms *Thalassiosira* spp. On average, diatom densities on the top of the Bank declined to their lowest densities in May, and subsequently increased again in June. Cell densities at the 3 deeper-water, flank stations (Stns 3, 9 and 20) did not show a pronounced winter-spring bloom compared with the top of the Bank, which most probably resulted from light-limitation at those stations because of tidal mixing and greater bottom depths. On average, highest

Fig. 3. Areal contour plots of surface salinity during 1999 for the January to June cruises.
diatom densities at these stations occurred during the May and June period. The dominant diatoms throughout the region were *Chaetoceros* spp., *Guinardia* spp., *Thalassionema* spp., and a variety of *Nitschia*-like pennates. Cell densities of dinoflagellates and other groups were generally very low from January to April (<10^3 cells l⁻¹), especially at the 4 shallower stations, and began to increase in May and June in a temporal pattern similar to what one would expect in terms of phytoplankton species succession. Dominant dinoflagellates were primarily *Ceratium* spp., plus *Peridinium* spp. and *Prorocentrum* spp. at somewhat lower densities. The ‘other’ group comprised almost exclusively nanoflagellates, especially the genus *Pyramimonas* (6 to 10 µm), which dominated the plankton in May at Stn 9 (ca. 300 × 10³ cells l⁻¹). Primarily because of the Lugol-stained samples, we were unable to determine reliably which cells might have possessed chloroplasts and which were more likely to be heterotrophic.

Concentrations of dissolved organic nitrogen (Fig. 10) showed initially high values in January, averaging 14.0 µM nitrogen (average of 2 and 20 m samples at all 41 standard stations), but dropped to an average of 5.5 µM the next month. The DON concentrations then increased each month to an average concentration of 15.3 µM in June. The concentrations of dissolved organic phosphorus (not shown) were extremely low from January to April, when they were not significantly different from zero. In May the concentrations averaged only 0.2 µM, and dropped to half that value in June.

The areal distributions of particulate organic carbon and nitrogen, and total particulate phosphorus were similar to that of chlorophyll, with elevated concentrations centered mostly on the top of the Bank through
April. The distributions were more patchy in May and June, with stations both on the top of the Bank and along the flanks and edges exhibiting elevated concentrations. Concentrations of POC, PON and TPP (Fig. 11) increased from January to June. This increase in particulate biomass measures stands in contrast to the trend for chlorophyll (Fig. 4), which reached a peak in April, dropped to lower concentrations in May, and then increased again in June. Concentrations of particulate organic nitrogen and carbon, and total particulate phosphorus taken together would represent total micro- and nanoplankton biomass (Fig. 11). These biomass measures did not undergo great changes in their proportions to each other over the 6 mo sampling period; i.e. they can be plotted on scales that do not need to be adjusted for each month. In general, the ratios of POC:PON:TPP are approximately in the proportions of 77C:18N:1P by weight (using the y-axes in Fig. 11 as overall 6 mo averages), which is a departure from the Redfield ratio of 40C:7N:1P (106C:16N:1P by atoms). Nitrogen is thus enriched relative to carbon and phosphorus in these samples, while phosphorus is low relative to nitrogen and carbon. These departures would be expected if our samples included heterotrophic organisms; for example, Elser et al. (1996) showed that protozoans can have atomic N:P ratios of 30 to 50 (13 to 22 by weight); this is similar to our results.

Ratios of average POC:PON from January to May (Fig. 12) show values close to, but slightly greater than the Redfield ratio, with values in June dropping below the Redfield ratio. The negative slope (−0.26) of the C:N ratios over the 6 mo period is significant (p < 0.05; Fig. 12). The lowest C:N ratios, which indicate nitrogen-enrichment in the particulate organic material, occurred in June, at the time of overall greatest particulate biomass (Fig. 11).

Linear regression lines fitted to POC versus chlorophyll a for each month (2 and 20 m samples for Stns 1
to 41) have non-zero \( y \)-intercepts of between 71 and 187 \( \mu g \; l^{-1} \) POC (Fig. 13). These \( y \)-intercepts are assumed to represent the non-photosynthetic component of the particulate biomass, which includes both living (heterotrophic) and non-living detrital particulate material; this biomass fraction generally increased from January to June, with the exception of a dip in May (Fig. 14). While there is a great deal of scatter in the data (Fig. 13), Table 1 shows that significant differences \((p < 0.05)\) in the \( y \)-intercepts plotted in Fig. 14 exist between 3 monthly pairings: January and March, January and April, and May and June.

The slopes of the POC versus chlorophyll \( a \) regression lines (Fig. 13) provide estimates of the carbon-to-chlorophyll ratio, ranging between 34 and 66 for all months except May, which had a slope of 100 (Fig. 14). Because these ratios are the slopes of the regression lines in Fig. 13, they do not include heterotrophic nor detrital organic carbon; thus, we have estimates of phytoplankton-specific carbon-to-chlorophyll ratios. Changes in these ratios between months (Fig. 14) can be interpreted to indicate changes in the relative phytoplankton growth rates, with low ratios corresponding to high growth rates, such as we see in the February to April period that encompasses the diatom bloom. Table 2 indicates that the slopes of the regression lines in Fig. 13 (phytoplankton carbon-to-chlorophyll ratios) for January, May and June, which represents a low growth period, are statistically different \((p < 0.05)\) from the February, March and April lines, which represent a high growth period.

**DISCUSSION**

Recent studies have demonstrated that the winter-spring phytoplankton bloom on Georges Bank starts as early as February and depletes inorganic nutrients by April (Townsend & Pettigrew 1997, Townsend & Thomas 2001). Our results indicate that the bloom may start even earlier, in that we observed a patch of elevated chlorophyll concentrations on the top of the
Bank in January of 1999, at which time the nutrient distributions, especially silicate, showed evidence of already having been drawn down. Diatom cell densities in January ranged between 50 and $>180 \times 10^3$ cells \text{l}^{-1} (at the stations where chlorophyll concentrations were high, Stns 11 & 12). This indication of relatively high diatom production rates as early as January may not represent the actual initiation of the winter-spring bloom, however. We cannot rule out the possibility that our survey in January was in fact describing some combination of incomplete wintertime nutrient replenishment on Georges Bank, driven by winter mixing and \textit{in situ} regeneration, and a continuing low—but nonetheless significant—level of winter primary production. Together, these could explain the relatively low nutrient concentrations over much of the Bank prior to the onset of the winter-spring bloom. With respect to nutrient replenishment, it is possible that the central portions of the Bank are too far away from the deeper, nutrient-rich waters beyond the Bank’s edges for complete replenishment to occur over the late fall and winter. Because much of Georges Bank is very shallow (the average depth inside the 60 m isobath illustrated in Fig. 1 is $>25$ m) we would expect that primary production would not be light-limited. That is, vertical thermal stratification and water-column stability (\textit{sensu} Townsend et al. 1994) are not requisites; the shallow bottom serves as the base of the upper mixed layer. If the water transparency is clear enough, and the ambient light levels high enough, then net positive photosynthesis is possible. Using values of the diffuse attenuation coefficient collected in February 1997 (0.18 m$^{-1}$; Townsend & Xu 1997) and calculations given in Townsend & Spinrad (1986), we see that the critical depth for net positive phytoplankton production is always deeper than the bottom, even on the winter solstice. Thus, it would appear possible that phytoplankton production on the shallower, central portions of Georges Bank may be maintained at a low but significant level year-round, and at a rate sufficient to keep
wintertime inorganic nutrient levels low. On the other hand, presumably because of tidal mixing and light limitation, waters between the 60 and 100 m isobaths did not exhibit such a pronounced winter-spring diatom bloom (Figs. 4 & 9).

Regardless of wintertime levels of primary production on Georges Bank, we clearly observed a pronounced winter-spring phytoplankton bloom, especially in waters on the top of the Bank and inside the 60 m isobath. At its height, chlorophyll concentrations exceeded 10 µg l⁻¹, which was almost exclusively attributable to diatoms. Following the bloom, we observed a shift in phytoplankton species composition, from a winter-spring dominance of diatoms (again, especially on the top of the Bank), to diminishing densities of diatoms in May (except at Stn 3), followed by increasing densities of diatoms in June; other flagellates began to increase in May. Cura (1987) noted the increasing importance of dinoflagellates in early summer, but based on his own samples and a review of earlier work in the area, he made no mention of the May period of very low diatom densities. This phenomenon of decreasing diatom densities reaching a minimum in May is an apparent result of silicate becoming depleted over the top of the Bank prior to nitrogen, and therefore limiting diatom growth. The reason is that source waters available for mixing with the waters on top of Georges Bank contain approximately 4 to 6 µM greater concentrations of inorganic nitrogen than silicate (Townsend & Thomas 2001). This is shown in Fig. 15 for all data collected on all 6 cruises in this study. From Fig. 15 we see even in January, near-zero silicate values with corresponding DIN values that are ca. 5 µM greater. Silicate values continued to decline through April, after which the DIN concentrations also decreased such that in May plots of both nutrients tend to converge at the origin. Also in May, but especially in June, the silicate levels began to increase to between 0 and 6 µM, which probably corresponds to an increase in silicate regeneration. Thus, as diatoms take up both

Fig. 8. Areal contour plots of surface ammonium during 1999 for the January to June cruises. (●) stations
Fig. 9. Bar graphs of cell densities of 3 phytoplankton groups (diatoms, dinoflagellates and other flagellates [nanoflagellates]) at 7 stations (Fig. 1) on Georges Bank from January to June 1999.

Fig. 10. Areal contour plots of dissolved organic nitrogen, presented as the average of 2 and 20 m depths at the standard stations, from January to June 1999. (●) stations.
inorganic nitrogen and silicate in approximately equal proportions (Brzezinski 1985), we would expect to see silicate becoming limiting before DIN. The areal distributions show that in February, silicate concentrations over much of the central portions of the Bank were reduced to < 2 µM (Fig. 6). This is at or below the half saturation constant for diatoms (2 to 4 µM; Paasch 1973), commonly interpreted as the concentration below which that particular nutrient becomes limiting. At the same time, nitrate + nitrite concentrations (Fig. 5) were in the range of 2 to 6 µM, which is well above the half-saturation constants reported for most species of phytoplankton. The drawdown of silicate concentrations on the Bank continued through March, which was when we observed the highest concentrations of diatoms, and subsequently diatom cell densities decreased. By April, most of the Bank had become depleted in silicate. Curiously, however, we did not see an immediate and concomitant increase in the abundance of flagellates, which did not exhibit relatively high cell densities until May, suggesting relatively slow growth rates compared with diatoms, a need for higher light levels and/or significantly warmer water temperatures which are not reached until May.

![Fig. 11. Seasonal trends (January to June 1999) in monthly mean particulate organic carbon and nitrogen and total particulate phosphorus (TPP), presented as an average of 2 and 20 m depths at standard stations excluding those in deeper waters off Georges Bank (Stns 38, 34, 40, 29, 25, 39, 16 and 7).](image1)

![Fig. 12. Monthly (January to June 1999) mean (±1 SD) ratios of particulate organic carbon to particulate organic nitrogen for samples collected at 2 and 20 m at standard stations (Stns 1 to 41), excluding those in deeper waters off Georges Bank (Stns 38, 34, 40, 29, 25, 39, 16 and 7). Dashed lines Redfield ratio of 5.7 (by weight), continuous line: linear regression of all data (n = 361). Regression results: C:N ratio = –0.26 (month) + 6.74. Slope is significant at 0.05 level.](image2)

![Fig. 13. Scatter plots of particulate organic carbon versus chlorophyll a each month (January to June 1999) for 2 and 20 m depth samples at all standard stations (Stns 1 to 41); stations in deeper waters off Georges Bank (Stns 38, 34, 40, 29, 25, 39, 16 and 7) were excluded. Linear regression lines are plotted for each month, statistics are given in Tables 1 and 2. The May, June and January fitted regression lines are dashed, while the April, March and February lines are continuous, to indicate statistical differences of slopes between the 2 sets of regressions (Table 2).](image3)
Localized patches of increased silicate concentrations in the surface waters were observed in April at stations along the southern flank of the Bank (Fig. 6); these could either have been silicate pools remaining after the height of the winter-spring diatom bloom, or patches of regenerated silicate. By June, there were relatively broad areas of the northern part of the Bank in which silicate concentrations were elevated to >2 µM. These results are similar to those recorded in 1997 (Townsend & Thomas 2001) and strongly suggest re-dissolution of biogenic silica. That is, we presume that diatom frustules from the winter-spring bloom a few months earlier were beginning to dissolve as a direct result of warmer water temperatures in May and June. Because of the shallow depths and the vigorous tidal mixing on the Bank, the silica is not lost by sinking as might be the case in deeper waters. There are 2 steps involved in the re-dissolution process, both of which are temperature-dependent. It is first necessary for the coating of organic material protecting the cell surface to be removed from the frustules by way of bacterial attack (Hecky et al. 1973), which would probably be a function of water temperature. Once the organic coating is removed, the dissolution of the silica is a direct function of water temperature (Kamatani 1982). The

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<th>Month</th>
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<th>Apr</th>
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Fig. 14. Bar graphs of monthly averages of particulate organic carbon concentrations in 2 fractions: the non-living detritus and the heterotrophic plankton (left graph; values determined from the y-intercepts of each of the regression lines in Fig. 13), and monthly average ratios of phytoplankton particulate organic carbon to chlorophyll a, given as the slopes of the regression lines in Fig. 13 (right graph; relationship to phytoplankton growth rate is given)
waters over most of Georges Bank remained at or below ca. 6°C from January to April, after which they warmed at a rate of about 3 to 4°C per month, supporting our assumption of temperature-dependent re-dissolution. We do not believe that the reappearance of dissolved silicate in May was the result of upwelling of deep-water nutrients into surface waters and subsequently onto the Bank. Although such a process could potentially renew silicate and other nutrients to the concentration levels seen in winter, and although it is certainly possible that phytoplankton could have rapidly taken up the new nitrogen making it difficult to detect, we did not observe the concomitant signatures in temperature and salinity that would be expected. Stations that exhibited elevated surface (and subsurface) silicate...
concentrations in May and June most often had nitrate + nitrite concentrations of <1 µM (cf. Figs. 5, 6 & 15), while potential deep-water sources beyond the Bank’s edges had an excess of about 4 to 6 µM more dissolved inorganic nitrogen than silicate (Fig. 15).

The depletion of DIN concentrations followed that of silicate, reaching concentrations of <1 µM over broad areas of the Bank in April (Fig. 5) at which point we presume that phytoplankton production as a whole became nitrogen-limited. Ammonium levels were only patchily distributed in April, and did not reach bank-wide concentrations higher than about 0.3 µM until June (Fig. 8). Phosphate concentrations declined throughout the entire 6 mo period, but never fell to levels below our detection limits (0.1 to 0.2 µM; Fig. 7). Fig. 16 shows a plot of DIN versus phosphate for all data from all stations and sampling depths, indicating that of the two nutrients, DIN, not phosphate, was the limiting nutrient from January to June. That is, nearly all the data points in Fig. 16 fall below the continuous line, indicating a N:P Redfield ratio of 16:1. However, in June, when DIN concentrations were at or near zero, we observed that phosphate levels also began to be depleted and crept toward zero. These dissolved inorganic nutrient analyses therefore strongly support the notion that primary production on Georges Bank becomes nutrient-limited very early in the year—by silicate in February and by DIN in April. Clearly, then, much of the primary production on Georges Bank is recycled production.

It is interesting to speculate here on the possible significance of the winter-spring diatom bloom on Georges Bank, especially in view of recent reports in the literature of the apparent toxic effects of diatoms on the reproductive success of copepods. Ban et al. (1997) reviewed earlier evidence and presented additional experimental results which showed that copepods fed diets of only diatoms produced eggs with low hatching success and nauplii with deformities. If fed a diet mixture that included flagellates, the diminished reproductive capacity disappeared. This subject is one of intense controversy (e.g. see Jonasdottir et al. 1998, Ianora et al. 1999) and as yet there is no consensus of opinion, but if the results of Ban et al. prove valid, it would imply that copepods on Georges Bank—inside the 60 m isobath, at least, where diatoms most clearly dominate—are unable to proliferate until flagellates become important components of the phytoplankton community in May and June. This indeed appears to be the case. Rather than exhibiting a springtime peak in zooplankton biomass concurrent with and shortly after the winter-spring bloom, the annual cycle of copepods on Georges Bank is reported to exhibit a biomass peak on about Calendar Day 151 (end of May/beginning of June), to decline quickly over the next 2 mo (June and July), and then to decline more slowly to minimum biomass levels in December and January (Sherman et al. 1987). The peak in zooplankton biomass, then, corresponds to the May-June period when we saw a second burst in diatom production and increasing abundances of flagellates (Fig. 9), and this is the time in our study when nutrient limitation was most pronounced.

In recent studies on Georges Bank, Gifford and Sieracki (pers. comm.) have shown that one of the dominant copepods, *Calanus finmarchicus*, feeds mostly on heterotrophic protozoans rather than phytoplankton throughout the winter-spring period. How well *C. finmarchicus* fares before May and June, e.g. before we would expect protozoans to become abundant, or if there is any difference in vital rates between populations residing on the top of the Bank versus its flanks, where flagellates are more prominent, has yet to be demonstrated. In addition to the potential deleterious effects of a relatively pure diatom food source, the diatom bloom occurs during the time of coldest water temperatures, while the flagellates dominate at warmer temperatures after May. Zooplankton population growth rates in general would be expected to experience this additional negative effect brought about by cold water temperatures; an extreme example of this potential low-temperature effect has been discussed earlier with respect to the winter-spring diatom bloom in the Gulf of Maine (Townsend & Cammen 1988).

Finally, with respect to zooplankton, we observed in this study that overall particulate biomass, in units of organic carbon, nitrogen and total phosphorus, increased fairly steadily from January to June, reflecting an overall steady increase in micro- and nanoplancktonic food supply for a grazing zooplankton population. The lowest observed C:N ratios (Fig. 12), the greatest particulate biomass (Fig. 11), and a mixed assemblage of diatoms and flagellates in May and June (Fig. 9) would seem to support the notion that this early summer period is most important to secondary production in this shelf ecosystem. Almost paradoxically, this is also the period of nitrogen limitation of new primary production.

Despite questions about the nutritional significance of the winter-spring diatom bloom on Georges Bank, this prominent oceanographic event is normally thought to be very important to secondary production in other temperate shelf ecosystems. If our results from Georges Bank suggest that the diatom bloom production may not be directly channeled through a copepod-dominated zooplankton population until May and June, then might not that carbon be consumed by the benthos? Unfortunately, this is an area of research that has been neglected in recent years, and we simply do not know. Regardless of the ultimate fate of a diatom
bloom, the contribution of the 1999 winter-spring bloom to total annual primary production on Georges Bank would at first appear to be very significant. We can calculate this contribution using the average decrease in total dissolved inorganic nitrogen on Georges Bank (approximated here by the area inside the 100 m isobath) from January to April. This decrease was approximately 5 µM (= 70 µg N l⁻¹), which, using the Redfield ratio, converts to a potential ‘new’ primary production (discussed below) of 400 µg C l⁻¹ over the 5 mo period from January to June. Ignoring any exports and imports of carbon or nitrogen across the bank’s boundaries, and assuming an average depth of 60 m for the bank as a whole, this converts to about 24 g C m⁻² new primary production over the 5 mo period encompassing the spring bloom. This simple calculation does not include dissolved organic nitrogen, because our results suggest that although there is evidence of its uptake between January and February, by June the DON was back to very near the January levels (ca. 15 µM N), resulting in no net change.

A new primary production of approximately 24 g C m⁻² over the winter-to-summer period on Georges Bank is surprisingly small considering that the total annual primary production is estimated to be ca. 400 g C m⁻² yr⁻¹ (O’Reilly et al. 1987). In addition to the wintertime accumulated nutrients, the remaining nutrient requirement for total primary production is met by a combination of upwelling fluxes of new nutrients (cf. Dugdale & Goering 1967, Eppley & Peterson 1979) as well as recycling in the well-mixed regions over most of the top of the Bank (Loder & Platt 1985, O’Reilly et al. 1987, Walsh et al. 1987, Horne et al. 1996, Townsend & Pettigrew 1997). Cross-isobath mixing and nutrient injections are thought to be most important along the northern flank of Georges Bank, where the topography is steepest and the hydrographic fronts are most pronounced (Loder et al. 1982, Loder & Platt 1985, Loder & Greenberg 1986, Horne et al. 1989, Townsend & Pettigrew 1997). This provides a greater flux of nutrients to the northern edge, which is at the upstream end of the residual clockwise circulation pattern around the bank. These higher nutrient concentrations on the northern flank of Georges Bank were first observed by Pastuszak et al. (1982), based on data from a limited number of stations. Cross-isobath fluxes of nitrogen-rich deeper waters have been calculated to explain new primary production of 50 to 100 g C m⁻² yr⁻¹ (or 12 to 25% of the total estimated primary production; Townsend & Pettigrew 1997). Thus, considering these continuous nutrient fluxes during the January to June period, which encompasses the winter-spring phytoplankton bloom, new production is more likely somewhere between 44 and 65 g C m⁻² yr⁻¹ (e.g. 24 g C m⁻² yr⁻¹ plus five-twelfths of 50 to 100 g C m⁻² yr⁻¹). As we have already argued, this exercise demonstrates that in order to explain high rates of primary production on Georges Bank we must conclude that, in summer, nutrient recycling is important in the bank’s central portions, with new primary production occurring along its edges. Such a tentative conclusion is consistent with the change we observed from spring to summer in the taxonomic composition of the phytoplankton.

The summertime planktonic ecosystem on Georges Bank would seem to be one of greater nutrient recycling on the Bank’s central portions, and more new primary production along the edges, as just discussed. The micro- and nanoheterotrophic components of the plankton community in this scenario are therefore important in nutrient cycling, but their abundances are not immediately evident from our cell counts, which were based on Lugol-preserved samples, thus making pigments difficult to identify. However, we can deduce some qualitative indications of their relative contributions to total biomass based on our analyses of particulate carbon, nitrogen and phosphorus (Fig. 11), the particulate organic C:N ratios (Fig. 12) and ratios of POC to chlorophyll (Fig. 13).

Assuming that the y-intercept of the POC:chlorophyll ratio in Fig. 13 corresponds to the standing stock of organic carbon in the detritus and heterotrophic plankton, we see that there is a general increase in that particulate fraction of the seston from January to June (Fig. 14). The generally-decreasing trend in the C:N ratios (Fig. 12) would suggest that the increases shown in Fig. 14 are more the result of living heterotrophs than detritus. It is probable, therefore, that a significant fraction of the flagellates we observed in May and June were heterotrophic, and may have been important in nutrient recycling. Quantitative estimates of the relative proportions of autotrophs and heterotrophs are lacking, however. Finally, the carbon-to-chlorophyll ratios of the phytoplankton in May and June were significantly higher than in February, March and April, indicating a lower phytoplankton growth rate in May and June. Whether a lower phytoplankton growth rate in early summer is offset by an increased biomass also remains unknown. Additional work in this area is clearly required before these interesting and potentially important questions can be addressed.

In summary, we observed that the winter-spring phytoplankton bloom in the Georges Bank region may begin as early as January. Likewise, there may be low but significant year-round primary production on the bank at a rate sufficient to impede the complete replenishment and buildup of new nutrients via winter mixing and local regeneration. The winter-spring bloom was well pronounced over the central portions of the Bank, and became silicate-limited as early as February; later in the spring, nitrate + nitrite became
limiting. Regeneration of silicate became apparent in May and June, and appeared to promote a second burst of diatoms along with increasing abundances of other non-diatom phytoplankton groups. Total micro- and nanoplanckton biomass and nutritional quality of that biomass increased from January to June. Zooplankton production would seem to be dependent on increases in the mixed planktonic assemblage forage during the May-June period.

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