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Annual cycle of primary production in the Cariaco Basin: Response to upwelling and implications for vertical export

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Abstract. Monthly hydrographic, primary production, bacterial production, and settling particulate carbon flux observations were collected between November 1995 and December 1997 at 10.5°N, 64.67°W within the Cariaco Basin, off Venezuela. Upwelling of Subtropical Underwater (SUW) started around October and lasted through approximately May of the following year. Wind speeds >7 m s⁻¹ were observed between January and June, with weaker winds (<5 m s⁻¹) between July and December. The upwelling cycle was therefore out of phase with that of the trade winds by 2–3 months. A seasonal cycle punctuated by transient extremes associated with subsurface ventilation events was observed in primary production. High bacterial activity and organic carbon recycling rates were observed near the oxic-anoxic interface. Integrated primary production was 690 gC m⁻² yr⁻¹ in 1996 and 540 gC m⁻² yr⁻¹ in 1997. Settling carbon flux measured with sediment traps was about 5.6% of integrated primary production at 275 m and about 1.7% at 1225 m, with no seasonality in the proportion of vertical flux to primary production. In total, between 10 and 11 gC m⁻² yr⁻¹ were delivered to the bottom sediment of Cariaco, which suggests that between 4 × 10⁵ and 1 × 10⁶ t of C yr⁻¹ were delivered to sediments within the upwelling area of the Cariaco Basin. This represents permanent sequestration of carbon previously entrained in the North Atlantic gyre in the area of formation of SUW. Results suggest that upwelled inorganic nitrogen, rather than nitrogen fixation, is responsible for the large productivity and particulate carbon settling flux in the Cariaco Basin.

1. Introduction

Over geological timescales, geochemical imbalances at the Earth’s surface are closely tied to carbon sequestration by marine organisms and to the sinking flux of particulate organic material [Berner, 1992]. This variable flux serves as a key to interpreting past climate when preserved at depth. However, the relationship between fossil flux and oceanographic conditions near the ocean’s surface must be analyzed to understand how these records can be used to make inferences about climate change. There are few places where flux observations have been linked to climatic and oceanographic forcing conditions [Deuser et al., 1990; Karl et al., 1996; Thunell, 1998a]. Indeed, only in a few locations do the underlying sediments preserve variability spanning a wide range of timescales. This has made it difficult to establish the relationship between productivity in surface waters and biogeochemical feedbacks acting over decadal and century timescales [see Falkowski et al., 1998].

Many studies have focused on the present balance between inorganic and organic pools in the deep ocean [see, e.g., Hansell and Carlson., 1998, and references therein], but the contribution toward this balance by waters near continental margins remains unclear. All attempts at global assessments of marine primary production indicate that ocean margins support on the average 2–5 times the annual production of open ocean waters [Koblentz-Mishke et al., 1970; Field et al., 1998]. Indeed, the mean particle flux at 2300 m along continental margins is of the order of about 7.0 gC m⁻² yr⁻¹, compared to ∼0.8 gC m⁻² yr⁻¹ at 2300 m in the deep sea [Deuser et al., 1990; Walsh, 1991; Pilskaln et al., 1996; Thunell, 1998b]. Globally, over 65% of the particulate carbon that falls below 1000 m is derived from the slope and rise of continental margins [Jahnke, 1996], and this estimate does not include areas shallower than 1000 m [see also Honjo et al., 1982; Spencer, 1984; Walsh et al., 1992]. These simple statistics suggest that the biological pump [Volk and Liu, 1988] is much more efficient in the vicinity of continental margins than in the ocean’s interior.

To try to understand the connection between surface production and vertical carbon flux in a productive setting, we started a series of monthly hydrographic and productivity observations and deployed a set of sediment traps in the Cariaco Basin, off the coast of Venezuela. These observations form the basis of the Carbon Retention in a Colored Ocean (CARIACO) study. Here we discuss the first 2 years of observations at the CARIACO site.

The Cariaco Basin (Figure 1) offers a unique geographical setting that permits studies that are difficult or impossible to conduct at other sites. This is a large (~160 km long, 70 km
wide) and deep (~1400 m) basin within a continental shelf. It is bound to the north by a sill connecting Margarita Island to Cabo Codera at a mean depth of about 100 m. There are two channels breaching this sill (Figure 1): one in the northeast of 135 m depth (La Tortuga) and a narrower one in the northwest of 146 m depth (Centinela) [Richards, 1975; Lidz et al., 1969].

Current conceptual models attribute a marked cycle in upwelling and associated sea surface temperature (SST) changes in this region primarily to the seasonal intensification of the trades along an east-west coastline [Richards, 1960, 1975; Herrera and Febres-Ortega, 1975; Muller-Karger and Aparicio, 1994]. Similar and concurrent variations in upwelling occur along the entire south/central Caribbean Sea, but we show here that winds lag behind the onset of upwelling by up to 3 months. Within the Cariaco Basin the seasonal upwelling process provides a source of nutrients that leads to vigorous phytoplankton growth near the surface. Previous studies suggest that much of this remains ungrazed and sinks, allowing pigments to reach the sediment [Richards, 1975; Richards and Vaccaro, 1956].

In contrast to the situation at other continental margins, the Cariaco Basin has a muted advective regime at depth. The sill restricts water motion and the lateral flux of material, thus forming a natural sediment trap within a continental shelf. As the turnover of basin waters is slow [Deuser, 1973], the decomposition of the sinking material leads to permanent anoxia below about 275 m depth. The Cariaco Basin is well known to be the largest anoxic basin of truly oceanic character. Indeed, the hydrography, chemistry, and sediment deposition processes in the Cariaco Basin are sensitive to changes in climate [e.g., Holmen and Rooth, 1990; Scranton, 1988; Scranton et al., 1987; Deuser, 1973]. Varved sediments that accumulate within the bottom anoxic waters provide a detailed record of annual- to decadal-scale change [Peterson et al., 1991; Hughen et al., 1996, 1998; Haug et al., 1998; Black et al., 1999]. The evidence suggests that even weak disturbances with a muted expression in the deep ocean are recorded temporarily within the water column in Cariaco and permanently in the sediments at the bottom of the basin.

In addition to the recent importance of the Cariaco Basin as the site of an important paleo-oceanographic time series, the Cariaco Basin has served as a natural laboratory for biogeochemists for over 40 years. This basin has been key in constructing stoichiometric models of organic matter remineralization [Redfield et al., 1963; Richards, 1975], developing residence time and box models, studying metallic sulfides [Bacon et al., 1980], and numerous other studies. However, very little is known about the variability in primary production or about the actual processes that control the carbon and nutrient fluxes within the basin.

2. Methods

Twenty-six CARIACO cruises were conducted between November 1995 and December 1997 to examine the hydrography, primary productivity, and vertical flux of material at 10.5°N, 64.67°W. Figure 2 is a schematic of the CARIACO logistics. The base of operations is the Estacion de Investigaciones Marinas Isla Margarita (EDIMAR) of the Fundacion La Salle de Ciencias Naturales (FLASA), located on Punta de Piedras, Margarita Island, Venezuela, and cruises use the R/V Hermano Gines (FLASA).

Wind data were collected at Santiago Mariño Airport (10.9°N 63.96°W). Hourly records were averaged to daily values, and a five-point running mean was applied to generate the figures shown in this paper. Historical wind records from Punta de Piedras, Margarita Island (10.9°N, 64.2°W) were also exam-
ined to assess the quality of observations at Santiago Mariño Airport.

SST was estimated in situ at 1 m depth with our conductivity-temperature-depth (CTD) sensors and more frequently from imagery collected by the advanced very high resolution radiometer (AVHRR) sensors on the NOAA 11, 12, and 14 satellites. The satellite SST were derived using the split window techniques [Walton, 1988; Strong and McClain, 1984; McClain et al., 1983]. While nonlinear algorithms developed for AVHRR data could be used, there is no conclusive evidence that these algorithms are any better than the multichannel sea surface temperature algorithms. The nominal accuracy of AVHRR SST retrievals is in the range of ±0.3–±1.0 K [see also Brown et al., 1985; Minnett, 1991]. In Figure 3 we show a smoothed representation of the satellite-derived SST (100-point running mean) and the monthly 1 m CTD temperatures at CARIACO.

A minimum of five hydrocasts were performed during each monthly cruise to collect a suite of core observations. Additional hydrocasts were performed for specific process studies. Water was collected with a SeaBird™ rosette equipped with 12 (8 L) teflon-coated Niskin bottles with teflon-coated springs. The rosette housed the CTD, YSI oxygen probe, and Chelsea Instruments profiling fluorometer outfitted for chlorophyll a estimates. A SeaTec c beam transmissometer (660 nm) was added to the system on the eleventh month of the time series.

For the first 10 CARIACO cruises we used a series of SeaBird SBE-19 CTDs, but we had some difficulty with inconsistent and obviously erroneous salinity data below the oxic-anoxic interface with these devices. Through repetitive attempts, good profiles were obtained for each cruise. However, the SBE-19 was permanently replaced with an SBE-25 in September 1996. In addition to annual calibration by the manufacturer, each salinity profile was corrected with discrete salinity observations collected at 20 depths. The salinity measurements were conducted onshore on a Guildline™ Portasal 8410 salinometer calibrated with International Association for Physical Sciences of the Oceans (IAPSO) Standard Ocean Water.

Phytoplankton biomass was estimated with chlorophyll a extracted in methanol and read on a Turner Designs fluorometer using standard methods [Holm-Hansen et al., 1965; Falkowski and Kiefer, 1985]. During the high production season...
(approximately January–May) 250 mL seawater were filtered onto 25 mm glass fiber filters (GF/F) filters, while 500 mL were filtered during the rest of the year. Three replicates were taken per depth except where biomass was clearly at its minimum, when only two were collected. Methanol extractions permitted direct comparison of concentrations estimated separately using the Kishino et al. [1985] method for measuring the light absorption coefficient by particulate matter.

Suspended particulate organic carbon (POC) was estimated by filtering water samples of known volume (generally 2 L) onto precombusted GF/F. The filters were folded inside cleaned tin disks and combusted in a PE2400 Elemental Analyzer. Cystine standard was used for calibration, and blank filters were used to determine background counts.

Primary productivity observations were made using a modified Steeman Nielsen (1952) NaH14CO3 uptake assay. One hour before sunrise, water from 1, 7, 15, 25, 35, 55, 75, and 100 m was obtained. Water was poured directly from the Niskin bottle under low-light conditions into 250 mL clear polycarbonate bottles, which had been acid-washed, rinsed, and soaked in deionized water for over 48 hours. Bottles were rinsed three times before filling using a near-total fill. Four clear polycarbonate bottles were filled from each depth. One inoculated bottle from each depth was wrapped in aluminum.

Figure 3. Wind ((a) v, (b) u, and (c) scalar speed (m s⁻¹)) at Santiago Mariño Airport (Margarita Island). Broken line laid over the scalar wind represents the monthly mean scalar wind. Negative u and v values represent westward and southward winds, respectively. (d) AVHRR-SST in a 12 × 12 km² area centered at the Cariaco site. Heavy line (squares) overlay represents CARIACO (in situ) temperature at 1–7 m depth.
foil to obtain the dark $^{14}$C uptake rates. An extra bottle for 1, 15, 35, and 75 m was filled but not inoculated to provide time zero ($t_0$) filter and seawater blanks. The $t_0$ samples were kept in the dark in the laboratory and were filtered after deploying the floating incubation buoy.

Each sample was inoculated under low-light conditions with 1.0 mL (4 $\mu$Ci) of the $^{14}$C sodium bicarbonate working solution. A 200 $\mu$L aliquot for counting total added $^{14}$C activity was removed from one of the three bottles from each depth and placed in a 20 mL glass scintillation vial containing 250 $\mu$L ethanolamine and 10 mL of liquid scintillation cocktail (CytoScint $^\text{TM}$). The mixture was held at 5°C until subsequent liquid scintillation analysis onshore. A 50 $\mu$L aliquot of the $^{14}$C working solution in a vial with ethanolamine (250 $\mu$L) and scintillation cocktail was counted for reference.

A Licor photosynthetically active radiation (PAR) integrator was used to estimate photoperiod length. Between December 1995 and November 1996, samples were incubated from 0600 to 1000 LT. Starting December 1996, the protocol was changed to incubate between 0700 and 1100 LT, which more accurately represents one third of the daily photoperiod and one third of the total energy received in 1 day at 10$^\circ$30'$^\prime$N, as verified with a Licor PAR light sensor. Incubation times were relatively short because of the potentially high productivity (>1000 mg m$^{-2}$ d$^{-1}$) of this continental margin. Upon completion of the incubations, sample bottles were removed from the mooring and placed in labeled dark plastic bags. A 50 $\mu$L aliquot was withdrawn from each productivity bottle and was filtered onto a 25 mm Whatman GF/F, which was rinsed with 0.25 mL 0.5 N HCl, placed in a 20 mL glass scintillation vial, sealed, and held at 5°C until subsequent processing onshore. Immediately upon return to the shore lab and within 15 hours of sample collection, 10 mL of liquid scintillation cocktail were added to vials with filters. The vials were refrigerated until analysis at Empresas Polar in Caracas, Venezuela.

Carbon uptake calculations followed the standard formulation outlined in the Joint Global Ocean Flux Study (JGOFS) manual [United Nations Educational, Scientific, and Cultural Organization (UNESCO), 1994], taking into consideration a (very low) quenching curve. To obtain the daily productivity rate, a photoperiod scaling factor that varied between 1.89 and 3.90, according to the fraction of the energy received during the incubation period relative to the total energy received in a day, was used. This was determined by the shipboard PAR photometer. Depth-integrated and annual production estimates were derived by trapezoidal integration over the upper 100 m and then across months.

Total dissolved inorganic carbon (T$^{13}$CO$_2$, hereinafter referred to as DIC) was calculated from measurements of total alkalinity and of pH on the total hydrogen ion concentration scale at 25°C. These estimates are performed using the precise spectrophotometric dye methods developed by Robert-Baldo et al. [1985] and Byrne and Breland [1989] and modified from Clayton and Byrne [1993] and Breland and Byrne [1993]. A single-beam spectrophotometer (Ocean Optics) was used. The pH was calculated using the original equations plus an additive factor of 0.0047, which results from a correction to the buffers used in the original paper [DelWalls and Dickson, 1998]. These methods circumvent the problem that arises when potentiometric electrodes are transferred from dilute buffers to seawater samples because of the sample’s high ionic strength. Total DIC was calculated using the K1 and K2 equilibrium constants from Mehrbach et al. [1973]. Values for the other equilibrium constants were taken from the review of Millero [1995].

On 11 cruises, total bacterial production was measured using a modification of the method of Kirchman [1993]. Discrete samples from 12 to 18 depths were transferred (in triplicate), under a nitrogen atmosphere, to 40 mL vials with Teflon-lined butyl rubber septa. Each vial was allowed to overflow two to three vial volumes and was sealed without headspace to maintain ambient redox conditions. Hydrogen 3-leucine (10 nM final concentration) was introduced into each vial via syringe. Samples were incubated in the dark for 8–10 hours in water baths maintained at in situ temperatures. Incubations were terminated by transferring the contents of an entire vial to tubes containing trichloroacetic acid (5% final concentration), which lysed cells and precipitated macromolecules. Acidified and chilled samples were processed as described by Kirchman [1993] upon return to EDIMAR. Bacterial production was estimated by assuming constant proportions of leucine and protein in bacterial biomass [Kirchman, 1993]. Bacterial production on an areal basis (mg C m$^{-2}$ d$^{-1}$) was calculated from volumetric rates using trapezoidal interpolations between discrete depth intervals.

Acetate uptake rate constants were measured using the radiotracer method described by Wright and Hobbie [1966] and Lee [1992]. Water samples were collected as described above for the bacterial production assay. One hundred microliters of a nitrogen-purged solution of $^{14}$C-acetate (4.4 $\times$10$^6$ dpm mL$^{-1}$) were added to each vial using a gas tight syringe. Samples were incubated in the dark at in situ temperatures. At four time intervals (between 6 and 12 hours), vials were sacrificed by filtering duplicate 5 mL subsamples through 0.22 $\mu$m polycarbonate Nuclepore filters to determine incorporation. Filters were placed in 20 mL scintillation vials containing 5 mL Optiflor scintillation liquid. The remaining 30 mL sample was killed by addition of 0.5 mL 10 N KOH. The amount of acetate respired to carbon dioxide was measured later in the laboratory by acidifying the sample in a closed flask with filter soaked in 2 N KOH [Wright and Hobbie, 1966; Lee, 1992].

Settling carbon flux was measured with sediment traps. A mooring with four Mark-VII automated sediment traps [Honjo and Doherty, 1988] has been deployed at the CARIACO site in 1400 m of water since November 1995. The mooring has been retrieved and redeployed every 6 months, in May and November. Traps were positioned at approximately 275, 455, 930, and 1225 m within the quiescent flow regime below sill depth. The cone-shaped traps had an aperture of $\sim$0.5 m$^2$ cross section and had a baffle designed to minimize advective disturbances. Descending particles were funneled into a sample jar at its base. Each trap had 13 sampling jars filled with high-salinity water, 12 of which were poisoned with formalin. The jars were sequentially rotated beneath the trap by a microprocessor-controlled stepping motor, which allows for sample collection over specified time intervals. We synchronized the computers to collect samples for concurrent 2 week periods at each depth. Upon retrieval, samples were split, and subsamples were subjected to a number of routine geochemical analyses to estimate carbonate, organic carbon, nitrogen, and biogenic silica fluxes. To estimate organic carbon, samples were dried, homogenized, weighed, and placed in silver boats. To eliminate inorganic carbon, boats were placed in a dessicator alongside a beaker of
concentrated HCl and put under a vacuum overnight. The samples were then folded and combusted in a PE2400 Elemental Analyzer. Cystine standard was used for calibration, and blank boats were used for determining background counts. Another subsample was used for radioisotope and stable isotope determinations and organic geochemistry. Approximately half of each sample remained available for additional study.

3. Results

3.1. Trade Wind

Figure 3 shows variations in the wind experienced at Santiago Mariño Airport on Margarita Island, about 80 km to the northeast of the CARIACO station. The zonal component of the wind \( u \) was almost identical to that observed about 15 km to the west at Punta de Piedras. At both locations the zonal component dominated variations in the wind, showing marked seasonality with values ranging from \( 4 \) m s\(^{-1}\) between about August and January to \( 10 \) m s\(^{-1}\) between about February and June. Muller-Karger and Aparicio [1994] compared the winds at Margarita Island with those observed at Orchila Island 200 km to the northwest. They concluded that variations in the zonal component of wind at Margarita are representative of those observed over the region.

The meridional component of the wind \( v \) observed at the airport in Margarita was weaker, varying between about 0 and \( 1 \) m s\(^{-1}\). This component showed cyclic shifts with a period of about 3 months (Figure 3). Muller-Karger and Aparicio [1994] found that at both Margarita and Orchila Islands the meridional component of the wind stress was usually \(<10\%\) of the zonal component of stress, particularly if averaged over periods >1 month.

The trade winds were relatively weak (order of \( 4 \) m s\(^{-1}\)) when we initiated the CARIACO program in November 1995 (Figure 3). Winds increased in December and January, peaked at \( 9–10 \) m s\(^{-1}\) in May 1996, but decreased rapidly thereafter, dropping to \(<5 \) m s\(^{-1}\) again in June 1996. Zonal winds remained relatively weak (\( u \sim 5 \) m s\(^{-1}\)) in February–April 1997, but the meridional component showed substantial northward intensification (\( v \gg 2 \) m s\(^{-1}\)) at this time. Zonal winds increased to about \( 9 \) m s\(^{-1}\) in May–June, with a nil meridional component, but then \( u \) decreased and \( v \) increased after July. The trade winds remained weak (\(<5 \) m s\(^{-1}\)) through the rest of the year, but they exhibited an anomalous northward component over this extended period (\(<1–2 \) m s\(^{-1}\)).

3.2. Hydrography

Between November 1995 and March 1996, surface temperatures decreased from \( 27.5^{\circ}C \) to \( 23.0^{\circ}C \) (Figure 3). The \( 21^{\circ}C \) isotherm migrated from a depth of \( \sim 130 \) m in November 1995 to \( \sim 30 \) m by May 1996 (Figure 4a). In May 1996 the \( 22^{\circ}C \) isotherm reached the surface. A peak in SST of \( \sim 29.0^{\circ}C \) was observed in September 1996. In 1997 an SST minimum of \( 21.5^{\circ}C \) was observed in March as a result of the \( 21^{\circ}C \) isotherm shoaling as early as January–February (Figure 4a). In both years an SST decrease was observed in July and August (Figures 3 and 4a).

Features noted in the temperature data are evident in the other hydrographic parameters. From November 1995 through January 1996 the salinity maximum (36.9) rose from \( \sim 100 \) m to the surface, and high surface salinities were observed until May 1996 (Figure 4b). A relatively fresh surface layer (\( \sim 36.3 \)) appeared during August–October, while the salinity maximum retreated to 50–100 m. Salinity isopleths turned upward again after October 1996, and the salinity maximum (\( >36.75 \)) reached the surface again in January 1997. This situation persisted through March. Surface salinities between August and
October were fresher (<36.5), and temperatures were warmer, coincident with the onset of the rainy season.

Intrusions of Caribbean water, identified on the basis of oxygen data from the CTD profiler and the discrete (bottle) O₂ and DIC data (Figures 4c and 4d), were observed in January 1997 at 200–220 m, in June 1997 at 300 m, in August 1997 at 275 m, and in December 1997 at 220–240 m. The January 1997 event was not seen in the January bottle data but did appear in the February bottle data; it is possible that in January the bottle at 200 m missed this feature. In the months following an event the anomalous water mass could still be detected.

When an intrusion of water from outside the basin was detected, waters from depths above the intrusion were observed closer to the surface. This can be seen in Figures 4c and 4d, where isolines of oxygen and DIC are relatively flat in early 1997 below depths of 200 m, while above those depths, water containing low oxygen values and high DIC are raised toward the surface, tracing the effects of an intrusion in January. The patterns of intrusion and the effects on hydrography and geochemistry are presented in more detail by Sarnion et al. [2001] and Y. Astor et al. (manuscript in preparation, 2001).

A strong seasonal cycle was observed both for DIC (Figure 4d) and near-surface CO₂ fugacity. The near-surface fugacity values during July–October of both 1996 and 1997 were typically in the 410–415 μatm range. During January–May 1996 we observed surface values in the 385–390 μatm range. During January–May 1997, values were smaller, typically between 370 and 390 μatm, with one observation of ~335 μatm in March.

3.3. Chlorophyll and Primary Production

A subsurface chlorophyll a maximum (~0.4 mg m⁻³) was located near 55 m in November 1995, but it rose to 25–30 m in December (Figure 5a). Between January and March 1996, values of 1.5–3.0 mg m⁻³ chlorophyll a were observed in the upper 35–55 m. In April and May, values in the upper 25 m exceeded 6 mg m⁻³ and reached 8 mg m⁻³ at 15 m. Phytoplankton concentrations then decreased, reaching <0.2 mg m⁻³ by June at the surface. A small chlorophyll maximum (~0.4–0.8 mg m⁻³) remained at about 35–55 m through October.

This cycle was repeated in 1996–1997, but the bloom started earlier than in 1995–1996, with biomass values of ~4 mg m⁻³ seen at 25 m in November 1996 (Figure 5a). After a brief decline in chlorophyll a to <1 mg m⁻³ in the upper 20 m in December 1996 a bloom reformed in January 1997 with 3–5 mg m⁻³ in the upper 25 m and lasted through May.

Surface particulate organic matter also increased tenfold between November 1995 (40–50 mgC m⁻²), particulate organic nitrogen (PON) ~ 2.5 mgN m⁻³, and March 1996 (500–800 mgC m⁻³, PON > 60 mgN m⁻³). POC and PON reached a peak during May 1996, but unlike chlorophyll, which declined in June 1996, POC (and dissolved organic carbon (DOC)) remained high until July 1996. We also commonly observed a minimum in POC (10–20 mgC m⁻³) just above the oxic-anoxic interface throughout the year. Concentrations around the interface at 275 m were higher but variable (20–70 mgC m⁻³). Lower but also variable concentrations (25–50 mgC m⁻³) were observed in deeper waters (>275 m).

Primary productivity showed marked seasonality (Figure 5b). Between about July and December, rates were <2.5 mg m⁻³ h⁻¹ in the upper 30 m, with lowest values (<1.0 mg m⁻³ h⁻¹) in August and September of both 1996 and 1997. Deeper waters typically showed values of <0.3 mg m⁻³ h⁻¹. Rates increased tenfold during January through June of both years, with values reaching 15 mg m⁻³ h⁻¹ and on occasion 20–40 mg m⁻³ h⁻¹ in the upper 7 m. The change in incubation time, from 0600–1000 to 0700–1100 LT, starting in January 1996 caused a small decrease only in the surface (1 m) primary productivity estimates.

Depth-integrated (0–100 m) primary production changed substantially over the course of our 1996–1997 field effort (Figure 6), in part because the annual cycle was punctuated by strong events. The average depth-integrated production between January and May 1996 was about 2800 mg C m⁻² d⁻¹, which was heavily influenced by the May 1996 upwelling event. During the same period in 1997 it was about 2600 mg C m⁻² d⁻¹. Productivity during June–December was lower (~1000 mg C m⁻² d⁻¹ in 1996 and ~860 mg C m⁻² d⁻¹ in 1997) (Figure 6). Integrated production in 1996 was ~500 (not counting May) to 690 gC m⁻² yr⁻¹ (if May is included). In comparison, production in 1997 was between 480 (not counting January) and 540 gC m⁻² yr⁻¹ (counting January).

The assimilation number PB (amount of carbon fixed per unit chlorophyll) is shown in Figure 7. PB at 7 m was typically the highest throughout the series and was likely the closest to PB max. Average PB at 7 m during January–June was about 7 mg C mg Chl⁻¹ h⁻¹ (range 4.6–9.0 mg C mg Chl⁻¹ h⁻¹), and during July–November it was 12 mg C mg Chl⁻¹ h⁻¹ (range 7.5–14 mg C mg Chl⁻¹ h⁻¹). Surface (1 m) PB was generally 6–14 mg C mg Chl⁻¹ h⁻¹ prior to December 1996 and decreased to 4–9 mg C mg Chl⁻¹ h⁻¹ afterward (not shown). This is likely due to increased photoinhibition in the surface incubation bottles caused by the later incubation times. This effect was not apparent in deeper samples.

3.4. Vertical Organic Carbon Flux

The first 6 months of sediment trap samples were retrieved successfully in May 1996. Unfortunately, samples for the following 6 months were lost. The four traps clogged immediately after redeployment of the mooring, resulting in a data gap in the sediment flux series from May–November 1996. We attribute this to unusually high particle fluxes caused by the large plankton bloom of May 1996 (see Figures 5 and 6). Subsequent redeployments, in November 1996 and May 1997, were successful.

The vertical flux of organic carbon followed a regular pattern, with minima between September and January, maxima between February and May, and similar temporal variability at all four depths (Figure 8). In 1996, carbon flux showed a minimum of ~0.01 gC m⁻² d⁻¹ in January at all four trap depths (Figure 8). Flux increased in the subsequent 2 months to peak in March 1996 at 0.17 gC m⁻² d⁻¹ at 275 m and at about 0.06 gC m⁻² d⁻¹ at 1225 m. The largest organic carbon flux recorded in this 2 year period was seen in the 1225 m trap sample for early July 1997 and was associated with a turbidite generated by an earthquake that occurred along the Venezuelan coast [Thunell et al., 1999]. Otherwise, the highest flux (0.18 gC m⁻² d⁻¹) was observed in the shallowest trap in May 1997. In general, organic carbon flux decreased from the shallowest to the deepest trap. The 275 m trap usually exceeded the flux captured at 455 m by 20–200% during January–May and frequently by 200–400% during July–November periods. However, in 27% of collections the flux to the 455 m trap matched or slightly exceeded the flux observed in the shallowest (275 m) trap.
3.5. Bacterial Production

Bacterial abundance, bacterial productivity, acetate concentration and acetate uptake rate constants, and dark carbon fixation rates were measured on six cruises between November 1995 and December 1997, and bacterial production was measured on an additional five cruises. The distributions seen for May 1997 (Figures 9 and 10) were similar to those observed on other dates. The most noticeable feature in the profiles is the low bacterial activity throughout the water column except at the surface and at the oxic-anoxic interface. Rates of bacterial activity (leucine incorporation and acetate uptake) near the oxic-anoxic interface often are comparable to rates in the surface mixed layer.

4. Discussion

The Cariaco Basin serves as a unique laboratory to examine the sinking flux of material because it forms a natural sediment trap. Outside the Cariaco Basin, materials that sink from surface waters and reach the deep parts of the Caribbean Basin
may be trapped there for periods of at least 50–800 years [Redfield et al., 1963; Kinder et al., 1985]. However, sinking material is quickly dispersed because of the strong Caribbean Current that washes the South American continental margin. Because of the strong currents along this margin, this flux is hard to measure. Waters in the Cariaco Basin have comparable residence times [Deuser, 1973], but material sinks near its place of origin. In either case the important quantity is the amount of material that is incorporated into the sediments because this material may be sequestered for periods spanning multiple centuries or millenia [Haug et al., 1998; Haagen et al., 1998; Black et al., 1999].

The results from the CARIACO series site demonstrate that production along continental margins in the tropics can be substantial and, indeed, redefines earlier estimates as being minimum values. Our annual production estimates (500–690 gC m$^{-2}$ yr$^{-1}$) are higher than those reported previously from the vicinity of the Cariaco Basin (200–400 g C m$^{-2}$ yr$^{-1}$) [Ballester and Margalef, 1965; Curl, 1960; Richards, 1960]. These values are comparable to those estimated for Monterey Bay (7 year average of $\sim$460 gC m$^{-2}$ yr$^{-1}$) [see Olivieri and Chavez, 2000; Chavez, 1996] and considerably higher than those estimated for Georges Bank, the New York Shelf, and the Oregon Shelf (380, 300, and 190 gC m$^{-2}$ yr$^{-1}$, respectively) [Walsh, 1988]. Previous estimates of production in the Cariaco Basin may be lower either because they were obtained by indirect methods [Richards, 1960; Curl, 1960] or derived from sparse temporal data that missed production peaks [Ballester

Figure 6. Cariaco Basin daily primary production integrated over the upper 100 m (mg C m$^{-2}$ d$^{-1}$).

Figure 7. Time series of water temperature at 7 m measured with a CTD (solid line, circles (°C)) and of the assimilation number $P^B$ at 7 m (broken line, squares) at the CARIACO station between December 1995 and December 1997.
sometimes reaches.

graphic studies [e.g.,

used without rigorous testing in studies of modern processes

fines this continental margin. This conceptual model has been

down the (roughly) east-west oriented coast that de-

the first half of the year, it leads to elevated nutrients in the

water is brought to the surface in the Cariaco Basin during

seasonal strengthening of the trade winds (Figure 3). For ex-

ample, rapid cooling occurred between October 1995 and Jan-

uary 1996 and again between October 1996 and January 1997,

prior to any significant increase in zonal wind speed above 5–6

m s⁻¹. Similarly, every year, SST started increasing prior to, or

coincident with, the onset of the seasonal weakening of the

trades, while zonal wind speeds were still in excess of 7 m s⁻¹

(Figure 3). Even though there appeared to be small decreases

in SST associated with the cyclic increase in the northward

component of the trade wind (Figure 3), there was no clear

evidence that the larger seasonal changes seen in the hydrog-

raphy and SST were driven solely by the local wind.

At this time, there is no conclusive explanation for the out-

of-phase changes in hydrography relative to the wind. For the

Caribbean Sea a hypothesis may be that the vertical displace-

ment of the SUW in the southern Caribbean is controlled by seasonal changes in the geostrophic flow through the basin

[see, e.g., Morrison and Nowlin, 1982; Morrison and Smith,

1990]. Such changes in the meridional slope of the isopleths

are likely connected to remote processes. We may speculate

that the circulation through the Lesser Antilles is strongly

affected by the seasonal switch between strong combined in-

flow from the North Equatorial Current and the Guiana Cur-

rent between January and June and weaker flow between June

and about December, when the North Equatorial Countercur-

rent and the North Brazil Current Retroflection form [see

Muller-Karger et al., 1995, and references therein]. These

changes would provide the correct phase relative to the wind in

the southern Caribbean. In addition, the seasonal cycle in the

Cariaco Basin may be superimposed on larger interannual

variation in the flow through the various passages in the Ca-

riaco Sea [see Murphy et al., 1999].

At shorter timescales of 2–3 months, small variations in the

hydrography were apparent, particularly in 1996. These varia-

tions included relatively small oscillations in isotherms, isoha-

lines, and oxygen concentrations to a depth of about 150 m

(Figure 4). They were also reflected as roughly 1°C amplitudes

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raphy and SST were driven solely by the local wind.
at least in part, to intrusion of Caribbean water into the Carib-
aco Basin at or near sill depth.

The intrusion of deeper water (below SUW) from the Ca-
ribbean Sea into the Cariaco Basin is dependent on this water
being present at sill depth, a situation that occurs during the
first half of the year [see Morrison and Smith, 1990]. When the
influx is large, it may be classified as a ventilation event be-
cause it brings substantial oxygen into the deeper, oxygen-poor
waters in the basin. Intrusions are probably more likely during
times when the SUW layer tilts upward toward the southern
Caribbean margin [e.g., Morrison and Smith, 1990], i.e., be-
tween about December and the following June.

There is at this time no definitive explanation for the 2–3
month signal observed in the 1996 hydrography nor for the
repeated ventilations of 1997. There is a 2–3 month variation in
the meridional component of the wind (Figure 3), which even
though is of a small amplitude (≈1–2 m s\(^{-1}\)), may play a role
in these fluctuations (Figure 4). Another possible scenario is
that the hydrography of the shelf was affected by eddies mov-
ing west in the Caribbean Current. A synoptic view of these
eddies and their relation to the wider circulation in the Carib-
bean Sea has emerged from recent numerical models of the
Caribbean Basin and from satellite altimeter data [Murphy et
al., 1999; Carton and Chao, 1999]. Once an intrusion or venti-
lation event occurs, the local wind assists in further elevating
the SUW layer and in enhancing mixing, thereby lowering the
temperature further along the coastline. A detailed discussion
of the deep geochemical effects of intrusions is provided by
Scranton et al. [2001], and a more detailed description of the
hydrographic effects is presented by Y. Astor et al. (manuscript

4.2. Primary Production

Similar to the pattern observed in SST, seasonal changes in
primary productivity precede changes in the wind. Figure 5b
shows that higher surface productivity (≥2 mg m\(^{-3}\) h\(^{-1}\)) was
observed as early as December–January, 1–2 months prior to
substantial increases in the westward wind speed. Productivity
tracked temperature changes, which is a proxy for nutrient
availability. In 1997, production was sustained at a higher level
and for a longer period relative to 1996 because of waters with
higher nutrients being pushed toward the surface from below
by a strong ventilation event [see Scranton et al., 2001; Y. Astor
et al., manuscript in preparation, 2001].

The biomass specific carbon uptake rates \(P^B\) were close to
the physiological maximum, which suggests that phytoplankton
were growing at near-maximum specific growth rates through-
out the year and that there was no nutrient limitation in the
Cariaco Basin during the upwelling season. Nevertheless, there
was a distinct seasonal variation in \(P^B\) (Figure 7). While Be-
havenfeld and Falkowski [1997] suggest that \(P^B\) should decrease
above about 20°C, \(P^B\) at CARIACO increased with tempera-
ture over the seasonal temperature range of 21°C–29°C. This is
likely to be due in part to the temperature effect described by
Eppley [1972] and not to a nutrient effect. At CARIACO,
nutrients follow an inverse relationship to SST, and \(P^B\) should
decrease with a decrease in nutrient concentrations. There is
also an inverse relationship between \(P^B\) and biomass, suggest-
ing that succession favored organisms with higher specific
growth rates during periods of lower nutrient availability. \(P^B\) in
1997 showed increased variability relative to the 1996 obser-
vations possibly because of an interaction between increased
photoinhibition caused by the change in incubation time and
nutrient supply due to increased (but variable) upwelling.

4.3. Sinking Organic Carbon Flux

Because of their high productivity, surface waters are a
source of organic carbon to deeper water. While we have not
measured vertical velocities of the water being upwelled at the
CARIACO station, a numerical model developed by Walsh et
al. [1999] suggested upward vertical speeds of 1–2 m d\(^{-1}\) under
westward winds of 6–10 m s\(^{-1}\), compared to 5–10 m d\(^{-1}\) about
below the 140 m sill depth, are within the statistical cone of
driven by biological production (see Figures 6 and 8). The flux of opaline silica was
Basin followed an annual cycle similar to that of surface pro-
zontal losses within this statistical funnel. We assume that vertical settling of particles dominates hori-
[Deuser et al., 2000]. Therefore, even if the actual sinking rates for partic-
ulate matter at CARIACO were much less than 100 m d\(^{-1}\) [Deuser et al., 1990], it is likely that the sediment traps, located below the 140 m sill depth, are within the statistical cone of sinking flux [Deuser et al., 1990] that originates at the station. We assume that vertical settling of particles dominates horizontal losses within this statistical funnel.

As mentioned above, the settling POC flux in the Cariaco Basin followed an annual cycle similar to that of surface production (see Figures 6 and 8). The flux of opaline silica was tightly coupled with organic carbon flux, and the C/N and \(^{13}\)C\(_{org}\) values for the sediment trap samples mostly varied from 5.8 to 7.9 vol/vol and from ~18.0 to ~22.5%, respectively [Thunell et al., 2000]. This is consistent with the idea that the material reaching the bottom of Cariaco was diatomaceous in origin and that the flux of terrigenous material into the middle of the Cariaco Basin was small at all times of year [Thunell et al., 2000].

Vertical carbon flux was directly proportional to integrated production (Figure 11), with the magnitude of the flux usually decreasing with depth. Carbon flux at 275 m was on the average 5.6% of integrated primary production. This decreased to 5.1% at 455 m, although in February, March, and April 1996 it was essentially the same at both 275 and 455 m at 6–7%. The average proportion decreased to 2.8% at 930 m and to 1.7% at 1225 m. One of the effects of the decrease in the ratio of settling carbon flux to primary productivity with increasing depth is a marked change in the difference of the flux between 225 and 1225 m with time. Specifically October, November, and December (low flux months) feature the smallest vertical differences in the flux, while March–May (high flux months) show large differences (see Figure 8). This implies that during the spring bloom, remineralization of the sinking organic matter is more efficient than during the nonupwelling season.

Acetate uptake rate constants (respiration plus incorporation; Figure 10) also provide a measure of organic remineralization under low-oxygen conditions [Henrichs, 1993]. Rate constant profiles consistently showed a strong maximum at the oxic-anoxic interface. In addition, rate constants for upwelling seasons (March and May) were higher than rate constants measured during November and July (data for July not shown). The abrupt deepening of the acetate uptake rate constant maximum in November 1997 as compared to other dates was associated with a similar deepening of the oxic-anoxic interface caused by the repeated intrusions of 1997 described above [Scranton et al., 2001]. These data support the conclusions from the sediment traps that more organic matter is respired during upwelling periods than during quiescent periods at all depths.

On the basis of the rate of decrease of fluxes with depth it appears that almost 95% of the particulate carbon produced by phytoplankton near the surface is consumed or regenerated within the upper 275 m of the water column. This can be compared to about 86–98% in Monterey Bay [Pilskaln et al., 1996] and 84–93% in the Santa Barbara Basin [Thunell, 1998b]. Our observed carbon fluxes are in excellent agreement \((r^2 = 0.87)\) with predicted fluxes determined using the Pace et al. [1987] model developed for the eastern Pacific Ocean and suggest that organic matter degradation occurring in an anoxic water column is as efficient as that occurring in well-oxygenated waters [see Thunell et al., 2000].

Suspended POC also decreases sharply with depth even during the high production season, from values in excess of 200 mg m\(^{-3}\) in the upper 25 m to <50 mg m\(^{-3}\) below 50 m. The vertical distribution of bacteria and bacterial activity (Figure 9) suggests that an active biological population is located at the oxic-anoxic interface. While not associated with any apparent increase in bacterial biomass, a secondary maximum in bacteria productivity between 100 and 160 m suggests that intensive carbon cycling also takes place in this depth range (Figure 9). Peaks in bacterial activity are commonly found at the oxic-anoxic transition zone in many environments.

While most of the bacterial production occurs in the upper 275 m, a significant fraction occurs below this depth. Our data show that 4–37% of integrated bacterial production (mean = 17%; standard deviation = 11) can occur below the depth of

![Figure 11. Relationship between monthly values of daily depth-integrated primary production (mgC m\(^{-2}\) d\(^{-1}\)) and monthly means of the biweekly integrations of organic carbon flux (mgC m\(^{-2}\) d\(^{-1}\)) at 275 (diamonds), 455 (squares), 930 (triangles), and 1225 m (crosses) at the CARIACO site. Lines are least squares regressions. January 1996 and 1997 were not included in the regressions as these were considered to be possible outliers in production.](image)
the top sediment trap (275 m). This is in contrast with the sediment trap data, which imply that on average, 95% of the labile export production is consumed or regenerated at depths shallower than 275 m and therefore that <5% sinks below this depth. However, bulk carbon delivery to the 455 m trap exceeded that delivered to the 275 m trap in 27% of our observations. Furthermore, on average we observe as much, and frequently more, bacterial production between 275 and 450 m, below the oxic-anoxic interface (17% integrated bacterial production), as in the overlying 175 m (14% integrated bacterial production).

Clearly, the interface is a region of vigorous carbon cycling fueled by labile carbon. We propose that this carbon is not necessarily provided by surface-derived export production. While bulk carbon delivery to the deepest trap appeared to conform to open water predictions [Thunell et al., 2000], the composition and source terms for this material are not well defined. Rapid heterotrophic activity at the oxic-anoxic interface suggests introduction of fresh labile organic matter at depth either through vertical migrators or possibly through in situ production by chemooautotrophs. Biologically enhanced layers established by either accumulation of particles along isopycnals or lateral advection of more productive waters are less likely explanations in the Cariaco Basin because waters below 200 m have nearly constant density and waters across the interface consistently have a uniform temperature-salinity (T-S) signature. Furthermore, these biologically enhanced layers are persistent features, and flux anomalies do not appear to correspond to the intrusions reported above. If in situ production at the interface contributes significantly to the vertical flux, then total fluxes to the seafloor and the chemical and stable isotopic composition of this material may be decoupled from surface processes.

Eppley and Peterson [1979] found that in the deep ocean the fraction of new to total production grows as total production increases. One of our early goals in the CARIACO program was to test this hypothesis at our continental margin site. For this purpose we may assume that the vertical organic carbon flux observed at the CARIACO station represents new production (export production equals new production) and that our productivity estimates are an approximation of total production. While there was seasonal variation in the settling organic carbon flux, there was no noticeable seasonality in the proportion of vertical flux to primary production. Indeed, the e ratio was relatively constant to within a few percent at any of the trap depths (Figure 11). In Monterey Bay [Pilskaln et al., 1996] and in the Santa Barbara Basin [Thunell, 1998b] the e ratio decreased as primary productivity increased. At these sites off California this result was interpreted as an effect of lateral advection. In any event, this suggests that Eppley and Peterson’s [1979] relationship applies primarily to areas of lower productivity and not to continental margins.

The trap and primary productivity observations suggest then that between 10 and 11 gC m⁻² are delivered to the bottom sediment of Cariaco every year. This agrees well with the 7–8 gC m⁻² seen at 2300 m at other continental margin locations [cf. Walsh, 1991; Pilskaln et al., 1996; Etcheber et al., 1996; Thunell, 1998b]. Preservation of organic matter within the anoxic environment of the Cariaco Basin therefore does not seem to be either enhanced or diminished relative to that of other continental margins [Thunell et al., 2000].

If these annual rates of carbon delivery to the bottom (10–11 gC m⁻² yr⁻¹) apply over the upwelling plume that covers the Cariaco Basin and adjacent areas, which can be estimated to cover >4 × 10⁴ km² and sometimes >9 × 10⁴ km² from infrared satellite data, between 0.4 × 10¹² and 1.0 × 10¹² gC yr⁻¹ (4 × 10⁵ and 1 × 10⁶ t of C yr⁻¹) may be delivered to sediments of the southeastern Caribbean Sea.

The final question is What is the source of the CO₂ that provides the carbon that is deposited in sediments of the Cariaco Basin system? In open ocean surface water, carbon cycling tends to deplete near-surface CO₂, resulting in a net flux from the atmosphere into the ocean. However, in a system like the Cariaco Basin the upwelling process brings water enriched in DIC and with a high fugacity of CO₂ into the euphotic zone. Even though CO₂ fugacity in surface waters decreased during the highly productive upwelling season, aquatic CO₂ remained near or above the atmospheric partial pressure of ~360 μatm essentially at all times. Thus, counterintuitively, the southeastern Caribbean is a source of CO₂ on a year-round basis in spite of the substantial primary production occurring there. This is important since intensification of the “biological pump” is often considered to be a key mechanism for drawing down atmospheric CO₂.

This poses an interesting question regarding the role of continental margins in the global carbon cycle. Viewed as a purely vertical system, the Cariaco upwelling system is a source of CO₂ to the atmosphere on a year-round basis. In this respect the CARIACO basin, as an analog for continental margins, does not appear to provide a ready mechanism for sequestration of anthropogenic atmospheric CO₂ in spite of the high primary productivity and organic carbon sedimentation rates seen there. However, the significance of the downward flux of particulate carbon at this continental margin lies in its role as a sink for CO₂ captured within SUW as it is formed in the North Atlantic. Because SUW is advected into the Cariaco system, it is a source of new nutrients as well as of new CO₂. Some of this carbon sinks when incorporated into the particulate flux, and some is vented to the atmosphere as CO₂. The upwelling process along the southern Caribbean margin therefore represents a remote sink for CO₂ previously sequestered from the surface within the North Atlantic gyre in the area of formation of the SUW. We may extend this analogy to other continental margins and view upwelling areas as both nutrient traps and areas of CO₂ outgassing.

5. Conclusions

The CARIACO results show that primary productivity of this tropical ocean margin is extremely high, with the assimilation ratio at its temperature maximum all year round. Previous estimates of annual production for the Cariaco Basin were low by factors of 2–3 because of undersampling of the high production events driven by transient physical structures. Indeed, upwelling of SUW and ventilation of the Cariaco Basin lead to increased primary production in surface waters and to increased settling flux of particulate carbon. Surface CO₂ fugacity is lower during the upwelling season relative to nonupwelling periods, reflecting the lower temperature, complete nutrient utilization, and upwelling of water that is nutrient-enriched relative to carbon. This is further evidence that the Cariaco Basin serves as regional carbon sink for CO₂ entrained at remote locations, such as in midlatitude gyres where the SUW is formed.

Estimates of rates of carbon remineralization in the water column of the Cariaco Basin from sediment traps suggest remi-
eralization is comparable to that seen in the open ocean in spite of the suboxic-anoxic nature of much of the water column. Data from direct microbial measurements suggest that remineralization is largely localized in the surface mixed layer and near the suboxic-anoxic interface. These data have also shown that remineralization rates vary somewhat with season (and primary production rate). The export rates (normalized to total primary production) seem to vary little in the transition from upwelling to nonupwelling season. This is different from the *Eppley and Peterson* [1979] finding that the fraction of new to total production increases as total production for the deep ocean increases but is similar to what others have found more recently in assessing export production. An additional finding is that the oxic-anoxic boundary is a zone of both high microbial heterotrophic and autotrophic processes.

Results suggest that upwelling along the southeastern Caribbean Sea margin is not only responsive to seasonal trade wind intensification. It is possible that there is a more direct association with the meridional tilting of the thermocline in the Caribbean Sea, which reflects the remotely forced changes in geostrophic flow through the basin. Transient eddies in the Caribbean and fluctuations in the meridional component of the local wind are likely to play a role in ventilation events. This suggests that this site can be used to assess the impact of larger-scale phenomena, which then would also help to understand better the paleo-oceanographic record.

The CARIACO site is located above sediments that have been shown to retain a high-quality record of sediment deposition for at least the past 12,000 years [Hughen et al., 1996]. A recent study of these sediments [Haug et al., 1998] concluded that nitrogen fixation plays an important role in controlling the settling carbon flux in the modern Cariaco Basin and that this is a consequence of high denitrification rates. The CARIACO study found the oxic-anoxic interface at about 250 m, and nitrate data show an indication of denitrification as shallow as about 130 m during July–November, the nonupwelling period. However, during upwelling the source of nutrients for surface production is water associated with the SUW, which has an elevated nitrate content. Therefore it is likely that mechanisms other than local nitrogen fixation may control the δ15N signature detected by Haug et al. [1998] in the Cariaco Basin sediments.

The results suggest that the Cariaco Basin serves as a recorder of both ancient and modern changes occurring over regional and possibly basin scales. The question remains of how widely some of these findings can be extrapolated to other margins.

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