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Late Pleistocene paleoproductivity patterns during the last climatic cycle in the Guyana Basin as revealed by calcareous nannoplankton

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Abstract

Variations in the assemblages and abundances of calcareous nannoplankton have allowed us to interpret changes in oceanic and atmospheric dynamics in the Guyana Basin, mainly linked to the southeast trades over the last climatic cycle.

Records of the paleoproductivity index of coccolithophores (N ratio) allowed us to monitor the nutri-thermocline fluctuations. Additionally, nannofossil accumulation rates vary closely with the N ratio, indicating a strong correlation between these two paleoproductivity proxies. The dominance of upper (small Noelaerhabdaceae, Emiliania huxleyi and Gephyrocapsa oceanica), over lower photic zone dwellers (Florisphaera profunda) during Termination II and interglacial substages 5.1 and 5.3 is related to eutrophic conditions due to a shoaling of the nutri-thermocline as a consequence of enhanced southeast Trade Winds. This activated an upwelling at the continental margin of the Guyana Basin. Low N ratio values and the dominance of F. profunda over the glacial substages of MIS 5 and glacial MIS 2-4 are linked to a deep nutri-thermocline (deep stratification of the mixed layer), at times of low influence of the southeast Trade Winds, and a weak upwelling. However, the N ratio during MIS 2-4 was slightly higher than those seen for the MIS 4/5 boundary and glacial substages 5.2 and 5.4. These micropaleontological proxies follow the insolation at high northern latitude (65° N): the high N ratio and NAR data from the Guyana Basin during Termination II and interglacials 5.1. and 5.3 are correlated with high insolation values, and low values of the N ratio and NAR during the MIS 4/5 boundary, glacials 5.2, 5.4 and MIS 2-4 are correlated with low insolation at the same latitudes. This situation suggests a link between the ITCZ, the southeast Trade Wind dynamics and the Northern Hemisphere climate changes during the last climatic cycle.

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1 Introduction and background

Coccolithophorids are photosynthetic planktonic algae living in the photic layer of the ocean (Hibberd, 1976, Brand, 1994; Young, 1994; Winter et al., 1994) that play a significant role in carbonate precipitation (Siesser and Winter, 1994; Steinmetz, 1994). The temporal and spatial distribution of coccolithophores may respond to major environmental factors in the upper oceanic layer such as temperature, nutrients, the trophic regime and sunlight levels (Brand, 1994; Young, 1994). For example, F. profunda, a phytoplankton species, thrives in the lower photic layer (Okada and Honjo, 1973; Okada and McIntyre, 1977) at times of high nutrient concentration, pointing to a deep nutricline (Molfino and McIntyre, 1990a, b; de Menocal, 1995; Beaufort et al., 1997; Wells and Okada, 1997; Flores et al., 1999; 2000; Kinkel et al., 2000; Beaufort et al., 2001; Gibbs et al., 2004; among others). Thus, the relationship between dwellers from the upper photic zone and those from the lower photic zone has been used to reconstruct changes in nutricline depth caused by variations in the southeast Trade Winds in the 15 tropical Atlantic (Molfino and McIntyre, 1990a, b) and in the Western Tropical Atlantic (Kinkel et al., 2000), as well as in the other parts of the ocean (Okada and Honjo, 1973; Young, 1994; Okada and Wells, 1997; Wells and Okada, 1997; Beaufort et al., 1997, 1999, 2001, 2003; Beaufort and Buchet, 2003; Flores et al., 2000; Liu and Herbert, 2004; Baumann and Freitag, 2004).

The production and sedimentation of coccolith-derived calcite in the deep ocean plays an important role in the carbon cycle through photosynthesis and calcification (Westbroek et al., 1993; Marsh, 2003; Brand, 1994, Young, 1994; Steinmetz, 1994). These organisms are one of the most important contributors to carbonate sedimentation in the open ocean and in continental slopes (Milliman, 1993; Steinmetz, 1994), such as in the Guyana Basin (Shipboard Scientific Party, 2003), an area controlled by a setting of hemipelagic deposition. This region has been documented as the main route of interhemispheric heat flux exchange from the South to the North Atlantic.

The Guyana Basin is located in the area of annual shift of the Intertropical Conver-

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gence Zone (ITCZ) that moves between 10° N and 5° S (Müller-Karger, 1989). The northward or southward position of the ITCZ causes a significant impact on water discharge from the Amazon and Orinoco rivers, as well as variations in the direction and speed of the Guyana Current (GC) and the southeast Trade Winds (Müller-Karger, 1989; Wilson et al., 2002; Ffield, 2005; Stramma et al., 2005) (Fig. 1).

During the boreal summer, the southeast Trade Winds are stronger and, the South Equatorial Current (SEC) and the North Brazilian Undercurrent (NBUC, which brings salty water from the South Atlantic) are well developed (da Silveira et al., 1994; Stramma et al., 1995, and Masson and Delecluse, 2000). Over that season, the westward displacement of the SEC pileup surface waters along the eastern coast of South America deepens the nutri-thermocline in the Western Tropical Atlantic (WTA) (Hastenrath and Merle, 1987). When the SEC reaches South America, it splits into two branches: one turns southward, supplying the Brazilian current (BC), and the other one takes off towards the north and feeds the North Brazilian Current (NBC), which is called the Guyana Current (GC) when it reaches the Guyana Basin (Fig. 1). The Intertropical Convergence Zone (ITCZ) is displaced to the northernmost position (6° N-10° N), generating a wet season north of ~5°N (Müller-Karger and Aparicio-Castro, 1994). The NBC (or GC) is mostly retroflexed, contributing to the North Equatorial Countercurrent (NECC) between 5° N-10° N (Müller-Karger et al., 1988), allowing anticyclonic rings to enter the continental margin and pass northwestward to merge with the Orinoco River plume (Müller-Karger, 1989; Ffield, 2005). These current rings transport and advect the Amazon River plume and, together with the NBUC, develop a zone of minimum surface salinity, warm temperatures and a shallower and stronger halocline, where the river flow is dispersed (Masson and Delecluse, 2000). The other component of the GC is directed to the North Atlantic, transporting relatively warm and fresher waters via the Caribbean Sea (feeding Caribbean current, CC) and the Gulf of Mexico (where it supports the Gulf Stream, GS) (Showers and Bevis, 1988; Schmitz, 1995; Stramma and Schott, 1996; Lynch-Stieglitz et al., 1999).

During the boreal winter the northeast Trade Winds dominate and relocate the ITCZ

to the southermost position (0°-5° S), causing dry conditions in the north of South America. River plumes decrease in intensity, and together with the GC, are mostly driven into the coastal area, carrying relatively colder and saltier waters to the North Atlantic. During this season, the NBUC mainly feeds the Equatorial Undercurrent (EUC) at the equator and cannot cross the equator to the North Atlantic (Masson and Delecluse, 2000 and references therein). The actual mean position of the North Equatorial current (NEC) has been found north of 10° N (Lazar et al., 2002). This atmosphere-ocean dynamic in the WTA plays a major role in the cross-equatorial heat flux to the North Atlantic (Ffield, 2005), producing relatively cooler surface waters south of the equator, and warmer surface waters north of the equator (>24°C) in the WTA.

This study focuses on a high-resolution micropaleontological analysis of coccoliths over the last climatic cycle. The primary goal is to reconstruct the patterns of calcareous nannoplankton production and their relationship with fluctuations in the nutrithermocline and ocean dynamics in the Guyana Basin (Fig. 1).

2 Materials and methods

2.1 Core location and materials

Core MD03-2616 was recovered in the Guyana Basin in the Western Tropical Atlantic (WTA) (7°48.75′ N, 53°00.80′ W) at a water depth of 1233 m, during the PICASSO Cruise by the Marion Dufresne R/V (Fig. 1). This area is characterized by the influence of the Guyana current, at the continental margin of French Guyana. The core shows an undisturbed high-quality sedimentary section of 39 m spanning the Pleistocene. It mainly consists of olive green silt and clay, rich in foraminifera and calcareous nannoplankton and with slight bioturbation levels and high organic matter remains (Shipboard Scientific Party, 2003). Here we report the uppermost 14.75 m (a detailed description was reported by the Shipboard Scientific Party, 2003).

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2.2 Age model and biostratigraphy

The age-depth assignments are primarily based on a series of 16 oxygen isotope control points performed on tests of the benthonic foraminifer *Uvigerina peregrina*. Stable isotope analyses were performed using an automated carbonate preparation line coupled to a Finningan MAT 251 mass spectrometer at the Laboratoire des Sciences du Climat et de l'Environnement in Gif-sur-Yvette (France). Isotopic events were identified up to Marine Isotope Stage 6, MIS 6 (\sim 155 Ka), by comparison of the benthic δ^{18} O from the Core MD03-2616 and the benthic δ^{18} O stack of Lisiecki and Raymo (2005) (Fig. 2) (López-Otálvaro et al., 2008¹).

We analyzed the >150 μ m fraction to detect the qualitative abundance of planktonic foraminifer species typical of the WTA. Study of planktonic foraminifer assemblages allowed the identification of the Ericson Climatic Biozones W1, X, Y, Z of Ericson and Wollin (1956) and the $Y_{P.obliq.}$ of Kennett and Huddlestun (1972). In our record, Biozone W1 was identified by the consistent and lower occurrence of the Globorotalia menardii group from the bottom (155 Ka) of the section studied up to 133 Ka. This event was dated by Kenneth and Huddlestun (1972) from 150 Ka to Termination II (128 Ka); i.e., slightly above ours. Biozone X was dated from 133 Ka to the middle phase of substage 5.1 (84.3 Ka). Prell and Damuth (1978) reported the top of the Biozone X at 85 Ka; i.e. slightly below ours. The G. menardii complex is absent from Biozone Y (Ericson and Wollin, 1956; Kennett and Huddlestun, 1972), as we report here. The disappearance datum of *Pulleniatina obliquiloculata* ($Y_{P, oblig}$) was recognized at ~39 Ka, but this datum is diachronous across the tropical Atlantic (Prell and Damuth, 1978; Kennett and Huddlestun, 1972). Biozone Z was identified close to the end of MIS 2 (over the latest Wisconsin) in agreement with Kennett and Huddlestun (1972), although other authors have placed its bottom at the MIS 1/2 boundary (Ericson and Wollin, 1956; Maslin and Mikkelsen, 1997) (Fig. 2).

According to currently available data, the reversal in the abundance of the coccol-

¹López-Otálvaro et al., in preparation, 2008.

ithophore species *E. huxleyi* and *Gephyrocapsa muellerae* is not clear. Thierstein et al. (1977) placed this datum at around 73–85 Ka. The absence of *Pseudoemiliania lacunosa*, the relatively low abundance of *G. caribbeanica* species (less than 3%), and the occurrence of both *E. huxleyi* and *G. muellerae* along the interval studied suggest that our sediment sequence is younger than 268 Ka (Fig. 2).

2.3 Calcareous nannoplakton preparation and counting techniques

Slides were prepared following the decantation technique outlined by Flores and Sierro (1997) to obtain quantitative measurements (coccoliths per gram, accumulation rates of coccoliths and coccolith-derived carbonate). These slides were examined at a 4-cm spacing (representing approximately $\sim\!0.120\,\mathrm{Ka}$ to $\sim\!0.700\,\mathrm{Ka}$) with a polarized microscope at 1000X magnification. 380 samples were examined, and more than 500 coccoliths were counted on each slide in order to identify the coccolith assemblage quantitatively.

Variations in the production of calcareous nannoplankton, and hence fluctuations in the nutricline, were expressed as a function named the *N* ratio (modified from Flores et al., 2000; López-Otálvaro et al., 2008), based on the relative proportion of taxa usually living in the upper photic zone (small Noelaerhabdaceae, *E. huxleyi* and *G. oceanica;* Okada and Honjo, 1973; Okada and McIntyre, 1979; Giraudeau, 1992; Young, 1994; Okada and Wells, 1997; Wells and Okada, 1997; Flores et al., 1999, 2000, 2003; Bollman et al., 1998; Beaufort et al., 1999; Beaufort and Buchet, 2003; Hagino and Okada, 2004, among others) versus the lower photic zone dwellers (*F. profunda*; Okada and Honjo, 1973; Okada and McIntyre, 1977; Molfino and McIntyre, 1990a, b among others). High *N* ratio values (close to 1) indicate a high production of upper photic species against the production of lower photic species, as result of a shallow nutrithermocline. Low *N* ratio values (close to 0) reveal a proliferation of *F. profunda*, in agreement with a deep stratification and a deep nutri-thermocline (Okada and Honjo, 1973; Okada, 1980; Molfino and McIntyre, 1990a, b; Wells and Okada, 1997; Flores et al., 2000).

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A qualitative preservation index of coccoliths, CEX', a proxy of the coccolith lysocline position (Dittert et al., 1999; Boeckel and Baumann, 2004), was used based on the relationship of the small Noelaerhabdaceae and *E. huxleyi* as the thinnest coccolithophoral species versus *Calcidiscus leptoporus* as a highly calcified species.

The accumulation rate of coccoliths (NAR, liths*cm⁻²*Ka⁻¹) has been widely used by several authors as a reference for high paleoproductivity of coccolithophores and particle flux out of the mixed layer (Steinmetz, 1994 and references therein; Su, 1996; Baumann et al., 2004). This parameter was estimated following the standard method of Flores and Sierro (1997), and it involves knowledge of the absolute abundances of coccoliths, the sedimentation rate and the dry sediment bulk density.

3 Results

The small Noelaerhabdaceae, *G. oceanica* and *F. profunda* proved to be the dominant species in the calcareous nannoplankton assemblage. *Gephyrocapsa muellerae* and *Umbilicosphaera sibogae* species (Appendix B and C) were identified as important secondary components. Additionally, *Helicosphaera* spp. and *C. leptoporus* were considered subordinate species because they were present in lower proportion (less than 2%) along the record studied. *U. sibogae* was the most abundant species (up to 12%) of the group of warm taxa, accounting for less than 4% and include *Calciosolenia murrayi*, *Discosphaera tubifera, Neosphaera coccolithomorpha, Oolithothus* spp., *Pontosphaera* spp., *Rhabdosphaera clavigera*, *Syracosphaera* spp. and *Umbellosphaera* spp. *U. sibogae* was separated from this group to analyze its paleoecological behavior along the section studied (Appendix B). The preservation of coccoliths was good along the last climatic cycle, except during the cooler MIS 5.2 and MIS 5.4 and the end of MIS 5.5. Decreases in the CEX' index and marked increases in the relative abundances of the most resistant species in the assemblage (*G. oceanica* and *F. profunda*), at least during MIS 5.4 and later MIS 5.5, confirmed this situation.

Based on the temporal evolution of calcareous nannoplankton, the sedimentary se-

quence was separated in two scenarios: Glacial MIS 6, interglacial MIS 5 and Termination II, and glacial stages 2–4 and the Holocene (Figs. 3 and 4).

3.1 Glacial MIS 6, Termination II and interglacial MIS 5

Small Noelaerhabdaceae dominate the assemblage, showing high relative abundances during MIS 6, the interglacial substages 5.1 and 5.3 and Termination II. By contrast, the relative abundance of *F. profunda* indicates a predominance of this species in the assemblage throughout the glacial substages of MIS 5, following an inverse trend to small Noelaerhabdaceae and *G. oceanica* (Fig. 3). The proportion of *G. oceanica* has low values and this points to moderate fluctuations during MIS 5, except during the end of MIS 5.5 and MIS 5.4 (when the assemblage underwent dissolution, as seen by the low CEX' values, Fig. 4). Mild percentage peaks of *G. oceanica* are seen along the MIS 4/5 boundary and glacial substage 5.2. The percentages of *U. sibogae* vary between 2% and 6%, with maxima during interglacials 5.1, 5.3 and early 5.5 (Fig. 3).

Common high NAR values during MIS 5.1, 5.3 and early 5.5 are seen for small Noelaerhabdaceae and *U. sibogae*. The former species also display high NAR values along Termination II and MIS 6. The NAR peaks of *G. oceanica* only occur during MIS 5.1 and early MIS 5.5, while the highest NAR of *F. profunda* occurs along MIS 5.2 and MIS 5.3. Owing to dissolution, all species show minimum NAR values throughout MIS 5.4 and late MIS 5.5.

Stronger gradients of the main taxa are observed in the relative abundances of calcareous nannofossils (between 10% and 80%) and partial NAR values $(1\times10^{11}$ and 1×10^{9} liths*pg $^{-1}$ *cm $^{-2}$ Ka $^{-1}$ for small Noelaerhabdaceae; 8×10^{9} and 1×10^{8} liths*pg $^{-1}$ *cm $^{-2}$ Ka $^{-1}$ for *G. oceanica*; and $1*10^{9}$ and $3.5*10^{10}$ liths*pg $^{-1}$ *cm $^{-2}$ Ka $^{-1}$ for *F. profunda*) during MIS 5 (Figs. 3 and 4).

The variation in total NAR is very similar to that of the N ratio, showing stronger gradients between the interglacial and glacial substages of MIS 5, with values between 0.9 to 0.2 for the N ratio and, 1.2×10^{11} and 1×10^{10} liths*pg $^{-1}$ *cm $^{-2}$ Ka $^{-1}$ for the total NAR (Fig. 4).

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3.2 Isotope stages 2-4 and the Holocene

During MIS 2-4 and the Holocene, dominant species exhibit less amplitude (lower gradients) but more frequent variability in their relative abundances than along MIS 5. *F. profunda* is the most abundant species, showing stable values in its percentages (up to 45%). The proportions and NAR values of this species vary in the opposite sense to those of *G. oceanica* and small Noelaerhabdaceae (Fig. 3).

The relative abundances of *G. oceanica* (10–20%) are lower than the steady proportion of small Noelaerhabdaceae (20–38%), in spite of its slight increase in the former species during MIS 3. Higher NAR values of *G. oceanica* appear during early and late MIS 3 and the Holocene. On the other hand, the stronger drop in the NAR values of small Noelaerhabdaceae during MIS 2-4 and the Holocene is comparable to the values found for the glacial MIS 5.2.

The percentage abundance of *U. sibogae* is somewhat higher during glacial 2–4 (values comparable to those of MIS 6) than during MIS 1 and MIS 5. However, the variability in the NAR values of this species is similar between the Holocene, the boundary of MIS 5/4 and MIS 5.1.

Frequent and mild fluctuations in the N ratio and total NAR occur throughout MIS 2-4 and the Holocene. They are comparable to those observed at the end of MIS 5.2 (Fig. 4). These parameters are strongly influenced by the dominant species in the assemblage and in consequence display low gradients during glacial MIS 2-4 (Figs. 3 and 4).

4 Discussion

4.1 Productivity of calcareous nannoplankton

The evolution of the calcareous nannoplankton assemblage (revealed by the N ratio, the NAR of individual species and the total NAR values) is mainly a response to a

productivity pattern in the Guyana Basin (Figs. 3 and 4), but the production rate of calcareous nannoplankton is limited by the release of nutrients to the mixed layer (Young, 1994). The question thus arises as to which oceanic mechanism allowed the production of calcareous nannoplankton.

All proxies studied in Core MD03-2616 (N ratio, species percentages and NAR values) indicate a shallow nutri-thermocline and more nutrient-enriched conditions in the mixed layer during MIS 6, Termination II and the interglacial substages of MIS 5 (Figs. 3 and 4). This situation would have resulted in a higher primary production and eutrophic conditions, and such a scenario implies the influence of the southeast Trade Winds blowing parallel to the coast (Gibbs, 1980), favoring upwelling and eastward water transport through the well-developed NECC (a situation analogous to the hydrological conditions prevailing during the boreal summer, Fig. 1). Moreover, during the glacial substages of MIS 5, the data support a deeper nutri-thermocline and more nutrientdepleted surface waters, resulting in a lower primary production in a deep stratified 15 photic layer (Figs. 3 and 4). This scenario implies a weaker influence of the southeast Trade Winds in the Guyana Basin and the intensification of the northeast Trade Winds blowing perpendicularly to the coast (Gibbs, 1980). Enhanced northeast trades favor the piling up of surface waters dragged by the NEC and the disappearance of the NECC that transports water away from shore. Additionally, the ITCZ is shifted to a southward position (between 0° to 5° S, Müller-Karger, 1989) and the GC follows a northwest direction (Fig. 1). Bassinot et al. (1997) and Vink et al. (2001) have suggested a southward latitudinal migration of the atmospheric and hydrographic circulation in the WTA during the last glacial and Vink et al. (2001) proposed a displacement of 6° southward for the NEC during the same time. Similarly, Kinkel et al. (2000) recorded a shallow nutricline and high coccolithophore productivity along the interglacial times and a deeper nutrithermocline along the glacial stages and substages in the WTA during the last climatic cycle.

The glacial MIS 2-4 and the Holocene are associated with recurrent, steady and low amplitude variations in the shoaling/deepening of the nutri-thermocline, pointing to a

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weak upwelling. This latter situation reveals the entrance of a mesotrophic regimen in surface waters as a result of the periodic variability of Trade Wind dynamics in the WTA (Peterson et al., 2000; Vink et al., 2001) (Figs. 3 and 4).

Gibbs (1980), Pujos and Froidefond (1995, and references therein) indicated seasonal scenarios where southeast Trade Winds are enhanced and blow alongshore in French Guyana, favoring wind-induced upwelling in the basin. Their work was restricted to the equator and 5° N. The geographic position of the core studied falls within the areas studied by those authors, and hence the same hydrological conditions can be considered (wind-induced coastal upwelling). This atmospheric dynamic displaces the ITZC to a more northern position (6° N–10° N), and permits the NBC (or GC) retroflection (Müller-Karger, 1989) and the formation of the NECC between 5° N–8° N (Busalacchi and Picaut, 1983). Additionally, other authors (Ryther et al., 1967; Hulburt and Corwin, 1969; Müller-Karger, 1995) have examined the occurrence of upwelling at the continental margin of the Guyana Basin (7° N±1°, Müller-Karger, 1995; Wilson et al., 2002) associated with the origin of the retroflection of the NBC (GC), and with the enhancement of the southeast Trade Winds.

Furthermore, cool sea surface temperatures from the North Atlantic are in phase with changes in ice sheet volume, reflecting glacial-interglacial oscillations and controlling Trade Wind intensity (Manabe and Broccoli, 1985; Ruddiman and McIntyre, 1984; Ruddiman et al., 1989). Changes in the temperature/pressure gradients between the North and South Atlantic affect the meridional wind component and define the latitudinal position of the ITCZ (Hastenrath and Greischar, 1993; Hastenrath and Druyan, 1993). This association accounts for the relationship between cold/warm periods in high latitudes in the North Atlantic and drier/wetter conditions over northern South America (Haugh et al., 2001). Visual comparison of the *N* ratio and NAR records with a northern insolation curve (mean insolation, 1 June to 30 September, 65° N) reveals peaks in the production and accumulation rates of calcareous nannoplankton and a shallow nutrithermocline that follows the maxima in insolation at 65° N. This correlation suggests a close connection between the shifts of the nutri-thermocline, Trade Wind dynamics,

and the latitudinal migration of the ITCZ. Molfino and McIntyre (1990a, b) and Wolff et al. (1999) found that variations in the position of the nutri-thermocline are controlled by the Trade Winds in the eastern tropical Atlantic (but in contrast with the present findings in the WTA, they reported a shallow nutri-thermocline during glacial periods).

Variations in the position of the thermocline have been reported by Hüls and Zahn (2000) and Vink et al. (2001) in a nearby core taken from the Tobago Basin. These authors studied Core M35003-4, which was collected at 12°05′ N, 61°15′ W at a water depth of 1229 m (a similar water depth to Core MD03-2616; 1233 m). They found the shallowest thermocline for MIS 3 (~50 m depth) in comparison with the glacial MIS 2 (~80 m depth) and the Holocene (150 m depth). Furthermore, Pailler et al. (1999) reported that the main thermocline is currently 150 m deep in a section of the WTA located between 0° N–45° W and 8° 20′ N–41° W, which surrounds the core position of this study.

4.2 River runoff

Some authors have suggested that the main freshwater source that affects the WTA is the Amazon River, which is mostly responsible for the low sea surface salinity in the WTA (Masson and Delecluse, 2000) since the waters of the Orinoco flow downstream towards the Caribbean Sea (Ffield, 2005). Several authors have indicated that high abundances of phytoplankton are associated with higher seasonal river runoff and lower salinities in the northern Caribbean Sea (14° N and 18° N) than in the typical waters of the WTA (Müller-Karger et al., 1989 and references therein).

Likewise, but addressing paleoclimatic changes at orbital scales, other studies have confirmed the northward position of the ITCZ along the interglacial substages of MIS 5 (in the Caribbean Sea, Martínez et al., 2007), and interglacial substages of MIS 3 (in the Cariaco Basin, Peterson et al., 2000; in the WTA and in the Tobago Basin, Vink et al., 2001), favoring the influx of river nutrients to the ocean and the configuration of an oceanic system that supports a superficial nutri-thermocline and consequently increased biological productivity in those regions. Similarly, Showers and Bevis

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(1988) documented an increase in the Amazon River runoff during Termination I until 5-6 Ka B.P. Other authors have reported the southward geographic situation of the ITCZ over the last glacial and consequently drier conditions over northern South America (Kolla et al., 1979; Balsam et al., 1995), decreasing the influence of the river flow system, deepening the nutri-thermocline and reducing bioproductivity (Peterson et al., 2000; Vink et al., 2000, 2001, 2002). According with these suggestions and the chlorophyll concentration data (primary distribution) presented by SeaWIFS Project (Fig. 1), the influence of an enhanced nutrient supply by rivers in the production of calcareous nannoplankton cannot be completely ruled out. However, there is no clear indicator within the association of calcareous nannoplankton able to account for a greater discharge from the Amazon during the interglacials and neither are there any indicators within the association to suggest relationships with variations in the salinity of surface waters. U. sibogae has been directly related to surface water salinity in the North Pacific (Roth and Coulbourn, 1982), has been found in warm oligotrophic waters (Okada and McIntyre, 1979; Roth, 1994; Young, 1994; Böeckel and Baumann, 2004; Ziveri et al., 2004), but also in medium-to-high fertility waters (Roth and Berger, 1975; Roth and Coulbourn, 1982) with temperatures ranging from 18°C to 25°C (McIntyre and Bé, 1967; Okada and McIntyre, 1979). Additionally, G. oceanica has also been observed in relatively warm, highly saline and fertile waters (Winter, 1982, 1985; Mitchell-Innes and Winter, 1987). However, U. sibogae and G. oceanica do not show a systematic correspondence with temperature or salinity. This implies that G. oceanica and U. sibogae respond to a productivity pattern rather than to a pattern of temperature or salinity, especially significant in MIS 2-4 (Fig. 3).

4.3 Dissolution of calcareous nannoplankton

Coccolith dissolution episodes (Figs. 3 and 4) suggest either a shallowing of the lysocline or dissolution above the lysocline. Curry and Cullen (1997) have documented high carbonate dissolution during glacial periods in response to deep water mass changes at greater depths in the WTA (>3000 m water depth). Owing to the shallow depth po-

sition of Core MD03-2616, dissolution below the lysocline is less probable, because aragonite and calcite lysoclines have been reported at 2200 and 4000 m, respectively (in Vink et al., 2001). Dissolution above the lysocline may be a response to the acidification of interstitial waters due to the oxidation of organic matter within the sediments. However, this hypothesis is not supported by the available observations, and hence its confirmation requires detailed benthonic isotope δ^{13} C data, sedimentological studies, and further oceanographic reconstructions in the Guyana Basin.

5 Conclusions

Variations in the evolution of calcareous nannofossils have allowed us to elucidate changes in the oceanic and atmospheric dynamics in the Guyana Basin; such changes are primarily linked to the southeast Trade Winds.

The data on the accumulation rate of coccoliths (NAR) resemble those of the *N* ratio when dissolution is lacking, suggesting a close connection between the production of calcareous nannoplankton and the accumulation of coccoliths. Higher values of the *N* ratio and NAR during interglacials are linked to a persistent shallow nutrithermocline, in contrast to low *N* and NAR values during glacials, which indicate a deep nutri-thermocline. A shallow nutri-thermocline gives rise to more surface nutrient-enriched waters during Termination II and interglacial substages of MIS 5.1 and 5.3, involving more productive conditions. By contrast, a deep nutri-thermocline reveals deep stratified conditions during the glacial substages of MIS 5 and MIS 2-4.

The greater productivity of calcareous nannoplankton was due to the upwelling of cool waters from depth (termination II and interglacial substages of MIS 5), which is consistent with the enhanced southeast Trade Winds that blew parallel to the coast and displaced the ITCZ to a more northerly position, while deep stratified conditions would have resulted from weakened southeast Trade Winds and enhanced northeast Trade Winds. Northeast Trade Winds allowed the southward displacement of the ITCZ and blew perpendicularly to the coast, piling up waters transported by the NEC and re-

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ducing upwelling. However, the influence of continental-derived nutrients and a freshening of surface water masses in the ocean through Amazon River runoff cannot be discarded.

The production of calcareous nannoplankton and the shifting of the nutri-thermocline covary with insolation at high northern latitudes: a high *N* ratio and a high NAR occurred during periods of high northern insolation (interglacials) and viceversa. Such variations are in agreement with an intensification (weakening) of the southeast Trade Wind dynamics and with the northern (southern) latitudinal migration of the ITCZ that activated (reduced) the upwelling system at the continental margin of the Guyana Basin.

Appendix A

Taxonomic appendix

Calcidiscus leptoporus (Murray and Blackman, 1898) Loeblich and Tappan, 1978

Calciosolenia murrayi Gran, 1912

Coccolithus pelagicus (Wallich, 1877) Schiller, 1930

Emiliania huxleyi (Lohmann, 1902) Hay and Mohler in Hay et al., 1967

Florisphaera profunda
Gephyrocapsa aperta
Gephyrocapsa ericsonii
Gephyrocapsa caribbeanica

Okada and Honjo, 1973
Kamptner, 1963
McIntyre and Bé, 1967
Boudreaux and Hay, 1967

Gephyrocapsa muellerae Bréhéret, 1978 Gephyrocapsa oceanica Kamptner, 1943

Hayaster perplexus (Bramlette and Riedel 1954) Bukry 1973

Helicosphaera carteri (Wallich, 1877) Kamptner, 1954

Neosphaera coccolithomorpha Lecal-Schlauder, 1950

Oolithotus (Cohen, 1964) Reinhardt, in Cohen and Reinhardt, 1968

Oolithotus antillarum Reinhardt, in Cohen and Reinhardt, 1968

Pontosphaera Lohmann, 1902

Pseudoemiliania lacunosa (Kamptner 1963) Gartner 1969 Rhabdosphaera clavigera (Murray and Blackman, 1898) Reticulofenestra Hay, Mohler and Wade 1966

SyracosphaeraLonmann, 1902Syracosphaera laminaLecal-Schlauder 1951Syracosphaera pulchraLohmann, 1902

Umbellosphaera Paasche, in Markali and Paasche, 1955

Umbilicosphaera hulburtiana Gaarder, 1970

Umbilicosphaera sibogae var. foliosa (Kamptner, 1963) Okada and McIntyre, 1977 Umbilicosphaera sibogae var. sibogae (Weber-van Bosse, 1901) Gaarder, 1970

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Appendix B

Taxonomic information

The systematic classification of coccoliths is based on their morphology. The morphological differences used for the genus Gephyrocapsa were adopted from Flores et al. (2000) and are summarised in Appendix C. Small placoliths (such as Gephyrocapsa aperta and G. ericsonii, small Reticulofenestra and Emiliania huxleyi) with an open or closed central area and smaller than 3 µm were classified together in the group of small Noelaerhabdaceae. Gephyrocapsa spp. contain Gephyrocapsa species larger 10 than 3 μm such as G. oceanica, G. muellerae, and G. caribbeanica. Different sizes of G. oceanica were also considered for carbonate calculations (medium Gephyrocapsa 3-4 μ m and large Gephyrocapsa 4-5 μ m), but later on these sizes were lumped together as G. oceanica for paleoecological purposes. Additionally, Oolithotus spp., Pontosphaera spp., Rhabdosphaera clavigera, Syracosphaera spp., Umbellosphaera spp. and Umbilicosphaera spp. were included as the group of warm taxa (Hiramatsu and De Deckker, 1997; Boeckel and Baumann, 2004). The genus Umbilicosphaera includes U. hulburtiana, U. sibogae var . foliosa and U. sibogae var. sibogae (the latter species here designated *U. sibogae*). The marked dominance of *Umbilicosphaera* sibogae in the group of the warm taxa suggests that its paleoecological behaviour in Guyana Basin should be analyzed. Calcidiscus leptoporus and Helicosphaera spp. species are characterized by their lower abundances and poor consistence along the record studied. Helicosphaera spp. mainly include Helicosphaera carteri and in a lower proportion H. pavimentum (below 1%).

Appendix C

Taxonomic notes and morphological differences used for the genus *Gephyrocapsa* (mainly adopted from Flores et al., 2000) in this study.

This Study	Small Gephyrocapsa (G. ericsonii and G.aperta)	G. muellerae (it includes G. margereli)	G. caribbeanica	G. oceanica (G. oceanica and large Gephyrocapsa)
Coccolith length	< 3 mm	> 3 µm	> 3 µm	3-5 μm (3-4 μm and 4-5 μm)
Bridge angle		5°-40°	Central area closed	>50°
Author equivalence				
Thierstein et al., 1977		G. caribbeanica		
Raffi et al., 1993	Small Gephyrocapsa	Small Gephyrocapsa	Small Gephyrocapsa	Medium Gephyrocapsa
Bollmann, 1997	G. minute	G. cold	G. oligotrophic, G. transitional	G. large, G. equatorial
Flores et al., 2000	Small Gephyrocapsa	G. muellerae	G. caribbeanica	G. oceanica
Baumann and Freitag, 2004	G. ericsonii/G. aperta	G. muellerae/G. margereli	G. caribbeanica	G. oceanica

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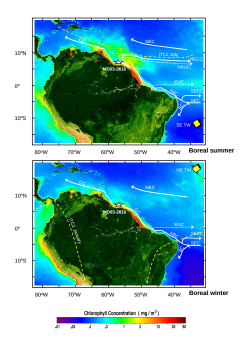


Fig. 1. Satellite images acquired by the SEAWIFS Program at the location of the Core MD03-2616. Hydrographic and atmospheric features in WTA: Surface currents are indicated by continuous lines and subsurface currents are indicated by dotted lines. SEC: South Equatorial Current, BC: Brazilian Current, NBC: North Brazilian Current, CC: Caribbean Current, NECC: North Equatorial Counter Current, SECC: South Equatorial Counter Current, EUC: Equatorial Undercurrent, NBCU: North Brazilian Countercurrent, ITCZ: Intertropical Convergence Zone. (Adopted from http://oceancolor.gsfc.nasa.gov/SeaWiFS; Richardson et al., 1994 and Haug et al., 2003.)



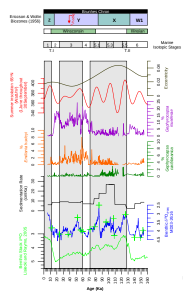


Fig. 2. Benthic δ^{18} O stack record (Lisiecki and Raymo, 2005), benthic δ^{18} O record from Core MD03-2616 (López-Otálvaro et al. 2008¹), sedimentation rates at Core MD03-2616 and the relative abundance of the specific index species from calcareous nannofossils at Core MD03-2616. The biozones X, Y, Z and W represent the biostratigraphic events from planktonic for aminifers identified at Core MD03-2616. The $Y_{P.obliq}$ biozone corresponds to the disappearance of *Pulleniatina obliquiloculata*. Las cruces verdes in the δ^{18} O record from Core MD03-2616 correspond to the control points used for constructing the chronostratigraphic framework at Core MD03-2616. Grey bars indicate interglacial marine isotope stages 1 throughout 5. MIS 5.1, 5.2, 5.3, 5.4, 5.5 correspond to the substages of MIS 5. Wisconsin and Illinoian episodes correspond to the Ice Ages between 12 Ka and 80 Ka in the former, and 128 Ka and 310 Ka in the later. T.I=Termination I, T.II=Termination II.

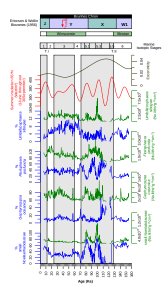


Fig. 3. Relative calcareous nannofossil abundances (%) and nannofossil accumulation rates (NAR, liths*gr⁻¹*cm⁻²) of the most significant species. Grey bars indicate interglacial marine isotope stages 1 throughout 5. MIS 5.1, 5.2, 5.3, 5.4, 5.5 correspond to the substages of MIS 5. Wisconsin and Illinoian episodes correspond to the Ice Ages between 12 Ka and 80 Ka in the former, and 128 Ka and 310 Ka in the later. T.I=Termination I, T.II=Termination II.

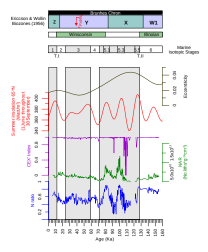


Fig. 4. Review of the bioproductivity proxies used for Core MD03-2616: N ratio vs. total nannofossil accumulation rates (NAR, liths*gr $^{-1}$ *cm $^{-2}$). Grey bars indicate interglacial marine isotope stages 1 throughout 5. MIS 5.1, 5.2, 5.3, 5.4, 5.5 correspond to the substages of MIS 5. Wisconsin and Illinoian episodes correspond to the Ice Ages between 12 Ka and 80 Ka in the former, and 128 Ka and 310 Ka in the later. T.I=Termination I, T.II=Termination II.