Productivity patterns and N-fixation associated with Pliocene-Holocene sapropels: paleoceanographic and paleoecological significance


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Abstract. We have studied a suite of 35 sapropel sequences from a transect of four ODP sites across the Eastern Mediterranean to explore for paleoproductivity patterns and provide new insights on ecological changes during their deposition. Paleoproductivity variations were identified using TOC and Ba_bio mass accumulation rates and δ15N_total and δ13C_org values. Elevated Ba/Al and TOC mass accumulation rates record periods of basin-wide amplified productivity. Our data further support that sapropels were formed by cyclic increases in primary production of marine organic matter largely sustained by N-fixing bacteria. This productivity increase was triggered by climate factors leading to increased fluvial discharge and amplified nutrient input that also favored the establishment of N-fixing bacteria. Enhanced productivity led to depletion of deepwater dissolved oxygen and consequently improved organic matter preservation. Primary production was more intense during the middle to Late Pliocene compared to Pliocene equivalents, coinciding with increasing total sedimentation rates. δ15N values are dramatically lower in the sapropels than in TOC-poor background sediments, indicating a major contribution from nitrogen-fixing bacteria to the higher productivity during sapropel deposition. Different degrees of denitrification occurred as a consequence of water column oxygenation which in turns evolved from stagnant anoxic bottom waters during Pliocene sapropels to oxygen depleted and sluggish circulation in late Quaternary layers. These differences between sapropel layers provide new evidences for the general evolution of the Eastern Mediterranean basin during the last 3 Mys in terms of paleoceanographic conditions and the intensity of climate variability leading to sapropel deposition.

1 Introduction

The multiple sapropel layers that were deposited in the Neogene-Quaternary Mediterranean basin are of particular interest to the topic of enhanced organic carbon accumulation in marine sediment. Sapropels, as defined by Kidd (1978), are cyclically deposited dark colored sediment layers that are more than 2 cm in thickness and contain more than 2% total organic carbon (TOC). A debate about whether improved preservation or elevated productivity is the major cause for the enhanced accumulation of organic carbon in sediments has persisted in paleoceanographic studies for decades (e.g., Casford et al., 2003; de Lange et al., 2008; Emeis and Weissett, 2009; Jenkins and Williams, 1984; Mangini and Schlosser, 1986). The traditional interpretation of sedimentary layers rich in organic matter as representative of anoxic events has been challenged by Pedersen and Calver, (1990; see also Calvert et al., 1996; Calvert and Pederssen, 1993), who argued that increased productivity, rather than absence of oxygen, was the primary factor responsible for the enhanced accumulation of organic carbon in sediments. Indeed, sapropel deposition is usually considered to be related to an increase in marine export productivity (e.g., Calvert et
al., 1992; Diester-Haass et al., 1998; Lourens et al., 1992; Martínez-Ruiz et al., 2000, 2003; Meyers and Arnaboldi, 2005; Weldeab et al., 2003a, b). However, how marine productivity increased in a presently oligotrophic, nutrient deficient basin like the Mediterranean raises questions regarding the sources of nutrient that have fueled this productivity increase (e.g., Casford et al., 2003; Filippelli et al., 2003; Menzel et al., 2003; Sachs and Repeta, 1999; Tanaka et al., 2010).

Development of a number of geochemical paleoproductivity proxies that are independent of oxygen abundance has allowed reconstruction of productivity and improved interpretation of the causal mechanisms of organic matter accumulation. Among these proxies, barium excess (\(B_{\text{bio}}\)), and biogenic barite accumulation (e.g., Dehairs et al., 1987; Dymond and Collier, 1996; Dymond et al., 1992; McManus et al., 1998; Paytan and Kastner, 1996), and nitrogen content and isotopic composition (e.g., Altabet and Francois, 1994; Arnaboldi and Meyers, 2006; Calvert et al., 1992; Meyers and Bernasconi, 2005; Peters et al., 1978) have been found especially useful in a variety of paleoceanographic settings. In recent years several papers have employed some of these paleoproductivity proxies to describe spatial and temporal patterns of sapropel deposition (e.g., Arnaboldi and Meyers, 2006; Meyers and Bernasconi, 2005; Rinna et al., 2002; Struck et al., 2001). However, further comparisons of the isotopic and elemental proxies and \(B_{\text{bio}}\) paleoproductivity proxy most widely used in the Mediterranean, and additional documentation of any differences of these signals during the paleoceanographic evolution of the basin are important to resolve existing questions about the conditions under which sapropels were deposited.

In this paper, we describe the results of our analyses of a suite of 35 sapropels of different ages that span the Pliocene through the Holocene and that originate from a four-site transect across the Eastern Mediterranean Basin. The description of the spatial and temporal patterns of export productivity in sapropel deposition is interpreted in terms of their possible paleoceanographic significance. We correlate both organic C and total N content and isotope signatures (\(\delta^{15}N\) and \(\delta^{13}C\)) to complement the information given by Ba/Al, and we employ TOC and \(B_{\text{bio}}\) mass accumulation rates (MAR’s) to refine paleoproductivity patterns. The effect of diagenesis on sediment N and C contents is also considered, inasmuch as organic matter is susceptible to post-depositional oxidation and alteration (e.g., Martínez-Ruiz et al., 2000; Thomson et al., 1995; Van Santvoort et al., 1996).

2 Materials and methods

2.1 Sample settings

Sapropel sequences were sampled in cores recovered at four ODP Leg 160 sites (Fig. 1) that represent different oceanographic regimes within the Eastern Mediterranean Sea. To summarize, the sedimentary sequence deposited at Site 964, located in a deep marine setting (3658 m.b.s.l.) on the Pisano Plateau (Ionian Basin), is influenced by the Adriatic Sea and the water masses coming from the Western Mediterranean basin through the Strait of Sicily. Site 969, also in a relatively deep, open marine setting (2200 m.b.s.l.) on the Mediterranean Ridge, represents the centremost location in the Eastern Mediterranean. Cores recovered at Site 967 in the Levantine Basin, although containing a sequence of deep pelagic sediments (2555 m.b.s.l.), are influenced by detrital input from the Nile River, which drains the central African craton. Finally, Site 966 is situated on a pelagic high, the Eratosthenes Seamount, at the relatively shallow water depth of 926 m.b.s.l. At each site, high resolution sampling was carried out on selected sapropel-containing core intervals. These sections represent late Pleistocene and Holocene periods of sapropel deposition at all four sites, Lower Pleistocene layers at sites 969 and 967 and Pliocene layers for sites 964 and 969.

The sediments in these cores are composed mostly of variably bioturbated nanofossil clay, clayey nanofossil ooze and nanofossil ooze with some intervals of clay and foraminifera sand, variably bioturbated (Emeis et al., 1996). Dark colored to black sapropel layers appear periodically throughout the pelagic sequences. Some of these TOC-enriched sediments and sections of the overlying and underlying sediment from these cores were sampled at 2 cm intervals. Where lamination allowed, samples were collected at a resolution as fine as 1 mm.
2.2 Analysis

Samples were dried, homogenized in an agate mortar, and then subdivided for different analyses. TOC measurements for some samples were carried out using a Perkin-Elmer Elemental analyzer at the Stable Isotope Laboratory at Stanford University (Mucciarone, 2003). The rest of the samples were analyzed for total carbon (TC) and TOC separately at Bremen University using a TOC/TIC analyzer. TOC was measured on the TOC/TIC analyzer after acidification of sediment with 1N HCl to remove carbonates followed by heating to dryness. At Stanford University, parallel to TOC measurements, total N (TN), δ¹³C and δ¹⁵N isotopic composition were also carried out using a Finnigan MAT isotope ratio mass spectrometer (IRMS) connected to a Carlo Erba (now CE Elantech, Inc.) NA1500 Series II Elemental Analyzer. TOC/TN ratios are expressed on an atom/atom basis. δ¹⁵N and δ¹³C values are respectively expressed relative to atmospheric dinitrogen and Vienna PeeDee Belemnite (VPDB) standards. These samples were repeatedly acidified with HSO₃⁻ to eliminate all inorganic carbon prior to isotopic analysis.

Barium content was determined using an ICP-MS Perkin-Elmer Sciex Elan 5000 spectrometer (CIC; Analytical Facilities of the University of Granada), using Re and Rh as internal standards. Coefficients of variation calculated by dissolution and subsequent analyses of 10 replicates of powdered samples were better than 3% and 8% for analyte concentrations of 50 and 5 ppm, respectively (Bea, 1996). Aluminium content was analyzed by atomic absorption spectrometry at the Analytical Facilities of the University of Granada. These two analyses were carried out after HNO₃ and HF total digestion of the homogenized sample. Biogenic Ba (Ba₃bio) was calculated following the equation used by Wehausen and Brumsack (1999) in order to separate detrital barite from barium precipitated as biogenic barite.

Ages of individual sapropel layers and site to site age correlations are based on precessional age scales in Emeis et al. (2000a), Lourens (2004), and Sakamoto et al. (1998). Each sampled layer was directly assigned to the corresponding depth, assigned insolation cycle, age (mid-point) and calculated Linear Sedimentation Rate (LSR) for each interval.

<table>
<thead>
<tr>
<th>Sample Location</th>
<th>I-Cycle</th>
<th>*Age sapropel midpoint (ka)</th>
<th>Depth (mbsf)</th>
<th>Depth (mcd)</th>
<th>LSR (cm/ky)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Site 964, Ionian Basin (3658 m b.s.l.)</td>
<td>Hole 964A</td>
<td>i-c 2</td>
<td>7.75</td>
<td>0.73</td>
<td>0.73</td>
</tr>
<tr>
<td>Site 967, Levantine Basin (2555 m b.s.l.)</td>
<td>Hole 967C</td>
<td>i-c 10</td>
<td>102</td>
<td>6.55</td>
<td>6.58</td>
</tr>
<tr>
<td>Site 969, Mediterranean Ridge (2200 m b.s.l.)</td>
<td>Hole 969A</td>
<td>i-c 10</td>
<td>102</td>
<td>2.64</td>
<td>2.42</td>
</tr>
<tr>
<td>Site 966, Eratosthenes Seamount (926 m b.s.l.)</td>
<td>Hole 966B</td>
<td>i-c 8</td>
<td>81</td>
<td>4.32</td>
<td>4.28</td>
</tr>
</tbody>
</table>

* Ages are from Lourens et al. (1996) and de Kaenel et al. (1999) and are based on the orbital calculations. S1 age is based on De Lange et al. (2008).
assumptions of sapropel synchronicity considered by other authors (e.g., Cane et al., 2002; Capotondi et al., 2006; de Lange et al., 2008; Emeis et al., 2003). The LSRs used for MAR calculations are summarized in Table 1.

3 Results

3.1 TOC concentrations and MAR’s

TOC concentrations in the samples range from less than 0.1% in the background sediment up to 28.4% in one sapropel layer (Figs. 2 to 6), with the highest values detected in the Pliocene sapropels at sites 964 and 969. The general temporal trend is a progressive increase in sapropel TOC concentrations from the Holocene through the Late Pleistocene (Figs. 2 to 5) and into the Early Pleistocene and Pliocene (Fig. 6). At each site, TOC-MARs show maximum values in Pliocene sapropels. For sapropels deposited during the same insolation cycle, those at Site 966 have the lowest TOC concentrations and lowest TOC MARs (see Fig. 2 to 5).

3.2 Organic carbon isotopic compositions

The \( C_{\text{org}} \)-isotopic compositions of the sapropel sequences differ spatially and temporally (see Figs. 7 to 11). As a whole, \( \delta^{13}C \) values range between \(-19.0\%e \) and \(-26.5\%e \), and both the highest and lowest values are found within the Late Pleistocene sapropels sampled from Site 967. Although a general trend of increasing \( \delta^{13}C \) values appears at sites 964 and 966, this tendency is not observed for Site 967 nor in the Pliocene S53 at Site 969. An apparent shift towards higher \( C \)-isotopic compositions occurs at the base of the Quaternary sapropels or just before the increase in their TOC content.

3.3 Nitrogen isotopic compositions

The \( \delta^{15}N \) values are strikingly similar at all four locations. They are significantly lower in the TOC rich sections compared to the carbonate oozes. The \( \delta^{15}N \) values range from \( \approx 1.0 \) to \( 0.0 \) for the sapropel 1 (i-cycle 2) (Fig. 7) to between \( 0\%e \) to \(-3.0\%e \) for Pleistocene sapropels (see Figs. 8 to 10) as low as \(-3.1\%e \) in the Pliocene sapropels (Fig. 11). When same i-cycles are compared between sites, \( \delta^{15}N \) values are very similar; only those values at Site 966 tend to be slightly higher.

3.4 Ba concentrations and MAR’s

Barium concentrations vary in parallel with TOC concentrations (Figs. 2 to 5), although elevated Ba concentrations frequently extend above the organic enriched layer in late Quaternary sapropels (Fig. 2), a feature especially visible in sapropels from Site 966. The sapropel layers at this site show a distinct offset between organic carbon accumulation and the productivity signal as represented by Ba/Al. In contrast to the patterns common to the late Quaternary sapropels, Ba\(_{\text{bio}}\) values decrease more than TOC MARs near the tops of Pliocene sapropels at sites 964 and 969 (Figs. 6 and 11). Because of the different patterns in the two proxies, we estimated durations of the sapropel formation events by measuring the Ba peak width in the Holocene and Late Pleistocene sapropels and the TOC MAR peak in the Early Pleistocene and Pliocene sapropels.

Ba\(_{\text{bio}}\)-MARs are largest in Late Pleistocene sapropels, particularly during i-cycle 12 (S5, see Fig. 3). The highest Ba\(_{\text{bio}}\)-MARs values are found at Site 964, the deepest studied site. For same-age sapropels, the increases in Ba\(_{\text{bio}}\)-MAR are similar for the three sites located in the deeper parts of the basin, but they are notably lower at shallower Site 966 (Figs. 2 to 5).

3.5 TOC/TN ratios

The atomic TOC/TN ratio varies similarly through time at the four locations, showing values mostly between 5 and 10 for background “normal” pelagic sediment (carbonate ooze), and increasing up to 25 in the sapropels (see Figs. 6 to 11). The highest sapropel TOC/TN ratios are found at Site 967, whereas the lowest are found at the bathymetrically elevated Site 966.

4 Discussion

4.1 Paleo productivity proxies

4.1.1 \( \delta^{13}C \) patterns and paleoceanographic significance

Organic \( \delta^{13}C \) values in analyzed sediment samples mostly correspond to organic matter of marine origin, although some terrestrial influence might be argued. However, \( \delta^{13}C \) values lower than \(-23\%e \) are common across the Eastern Mediterranean, especially in low TOC sections. The more negative values are consistent with an oligotrophic setting in which C availability does not limit algal vital fractionation. In contrast, periods of higher productivity result in depletion of \( ^{12}C \) from increased uptake, increasing \( \delta^{13}C_{\text{org}} \) values (e.g., Meyers, 1997). This slight increase in \( \delta^{13}C_{\text{org}} \) values is the general response found in analyzed sapropels and is particularly evident at the base of the \( C_{\text{org}} \)-rich layers. This classic high-productivity isotopic trend is not evident at Site 967. Instead, cores recovered at this deep Levantine sub-basin location exhibit generally lower \( \delta^{13}C_{\text{org}} \) values and the most erratic trends in the sample suite. A possible explanation is that extensive organic matter recycling in the surface ocean limited the \( ^{13}C \) enrichment associated with net uptake (e.g., Arnaboldi and Meyers, 2006; Menzel et al., 2003; Meyers and Arnaboldi, 2008; Struck et al., 2001) which can also apply to slight decreases in the upper part of some sapropel layers (e.g., S49, Site 969). Increased continental runoff associated
with periods of sapropel formation (e.g., Gallego-Torres et al., 2010; Rossignol-Strick, 1985) lowered the salinity of the surface ocean and created a strongly stratified water column that impeded sinking of organic matter and discouraged vertical mixing. Oxidation of the isotopically light organic carbon could then occur within the lower part of the photic zone, where it could be re-assimilated by photosynthesizers. The near-surface recycling of marine organic carbon during times of fluvial dilution of the surface ocean would potentially be augmented by delivery of low $^{13}$C terrestrial organic matter, this site being under the influence of the distal plume of the Nile River, adding to the low isotope signal. In any case, a shift toward higher isotopic composition is visible at the base of the TOC-enriched layers at Site 967, which is a classical indication of a general increase in $^{12}$C removal in the form of marine organic matter during initiation of sapropel deposition.

4.1.2 Paleoproductivity evidence from nitrogen concentrations

Total nitrogen in sediment indicates organic matter production, provided N adsorbed to clay minerals is insignificant (e.g., Calvert, 2004; de Lange, 1992; Freudenthal et al., 2001). This was verified using the correlation method described by Nijenhuis and de Lange (2000), Calvert (2004), and Arnaboldi and Meyers (2006) in which the concentrations of TOC are plotted against those of TN. The correlation is extremely good ($R > 0.9$ for all cores, see Fig. 12) with an
intersection essentially at the origin of the $\gamma$ axis. Although this intersection point is not exactly zero, the amount of inorganic N is negligible, as C:N ratios of low TOC samples are within the range of previous studies (e.g., Meyers and Arnaboldi, 2005) and in agreement with expected Redfield ratios.

Because organic N is also susceptible to post-depositional oxidation similar to organic carbon, its use for paleoproductivity reconstruction is subject to the same restrictions. Nonetheless, the correlation between TN and TOC gives us confidence that the entire N in the sediment is associated with or derived from organic matter, and so our $\delta^{15}N$ analyses will reflect purely $N_{\text{org}}$.

**4.1.3 $\delta^{15}N$ patterns in sapropel sequences**

The $N_{\text{org}}$ isotopic composition can be used as an indicator for the origin of organic matter (e.g., Knicker and Hatcher, 2001; Meyers, 1997; Rinna et al., 2002; Schubert and Calvert, 2001) and, more importantly, for evaluating nutrient cycles in the water column (e.g., Altabet et al., 1999; Altabet and Francois, 1994; Freudenthal et al., 2001; Karl et al., 2002; Pantoja et al., 2002; Voss et al., 1996). The Mediterranean is presently an oligotrophic sea, characterized by low concentrations of macro-nutrients such as P and N in surface waters (e.g., Astraldi et al., 2002; Bethoux, 1989; Struck et al., 2001). This shortage of nutrients, as a whole and of P specifically, is particularly dramatic in the Eastern Basin. An important regional nutrient source to the Eastern Mediterranean is the Nile River (e.g., Diester-Haass et al., 1998; Jenkins
Organic matter in all the sapropels exhibits a shift to δ15N values lower than in the background marls at the four locations (Figs. 6 to 11). This signature is unlikely a result of diagenesis as under suboxic-anoxic water column conditions the expected trend of organic matter degradation is an increase in δ15N due to remineralization that is associated with an increase in TOC/TN ratios (e.g., Karl et al., 2002; Lehmann et al., 2002; Meyers and Arnaboldi, 2005; Nakatsuka et al., 1997). Moreover, Higgins et al. (2010) recently rejected the possible diagenetic overprinting by measuring chlorin δ15N in sapropels and marls. Instead, the possible explanations for the dramatic decrease in 15N values in the sapropels are: (a) incomplete nutrient utilization under nutrient excess conditions (e.g., Calvert et al., 1992), (b) an external source of light N such as river discharge or atmospheric deposition (e.g., Krom et al., 2004; Mara et al., 2009), and (c) an ecosystem change in which a bloom of nitrogen-fixing primary producers occurs (e.g., Arnaboldi and Meyers, 2006; Milder et al., 1999; Pantoja et al., 2002; Sachs and Repeta, 1999; Struck et al., 2001).

The evidence for increased productivity based on the high TOC and Ba concentrations of the sapropels is strong so we may reject the hypothesis of incomplete nitrogen utilization. Fluvial delivery is also unlikely as an extraordinary excess of nutrients would have to be delivered to lower the δ13N values, particularly considering the amount of organic carbon accumulated in the sapropels (Sachs and Repeta, 1999). Increase in atmospheric deposition of nitrogen is also unlikely as it is expected that less dust would be mobilized during the wetter climate associated with precessional minima. In addition much of the atmospheric nitrogen sources today are anthropogenic and thus atmospheric deposition was not as important in pre-anthropogenic times. Accordingly we believe that nitrogen fixation is the main source of the bioavailable nitrogen with low isotope values in surface waters (e.g., Altabet and Francois, 1994; Kuypers et al., 2004; Milder et al., 1999). This process would incorporate dissolved atmospheric dinitrogen (δ15N = 0‰) into the marine system, lowering the isotopic composition (e.g., Altabet et al., 1999; Karl et al., 2002; Meyers, 1997; Pantoja et al., 2002). The extremely light N is isotopic composition suggests a change in the photoautotrophic community to a primarily N-fixing biota, probably associated with a decrease in eukaryote phytoplankton activity during sapropel deposition times (e.g., Milder et al., 1999; Pantoja et al., 2002; Sachs and Repeta, 1999). Nitrogen fixing cyanobacteria, cyanobacterial mats and Williams, 1984; Rossignol-Strick, 1985; Weldeab et al., 2003a). This riverine input of nutrients has fluctuated over time as cyclic intensification in the monsoon system during insolation maxima increased runoff from its catchment (e.g., Hilgen, 1991; Rohling and Hilgen, 1991; Rossignol-Strick, 1985). With respect to N, nitrogen fixation and possibly atmospheric deposition are currently important sources for bioavailable N to the oligotrophic Mediterranean, although their significance is also variable over space and time.

Fig. 6. Geochemical signal and TOC and Ba_bio Mass Accumulation Rates, C:N, and C-N isotopic composition for representative Lower Pleistocene from Site 967 and Pliocene sapropels from Site 964. Note different scales for TOC and Ba-MAR’s. Shaded areas correspond to sapropel layers.
Nitrogen fixation is typically limited by Fe, Mo and/or P availability (Karl et al., 2002). However, although river runoff would not be able to supply enough N to support the observed increase in productivity, it may have been capable of supplying oligoelements, such as Fe, in quantities sufficient to induce higher productivity and N-fixation by cyanobacteria and archaea. The extra freshwater input occurring across the basin (e.g., Gallego-Torres et al., 2010; Osborne et al., 2010) decreased salinity in the upper part of the water column. Lower salinity enhances Mo reactivity and availability, in the form of MoO$_4^{2-}$, whereas P can readily be recycled by efficient scavenging in the water column (Karl et al., 2002) or reductive dissolution at the sediment-water interface (e.g., Slomp et al., 2004). These multiple factors, along with an increase in sea surface temperature (e.g., Emeis et al., 2000b; Lourens et al., 1992), probably allowed the maintenance of blooms of N-fixing organisms and associated biota.

The shift towards low $\delta^{15}$N values during sapropel events is evident across the whole region and parallels or precedes the increase in export production indicated by Ba$_{bio}$ MARs (e.g., Dehairs et al., 1987; Gallego-Torres et al., 2007; Weldeab et al., 2003a). $\delta^{15}$N values exhibit a progressive decrease from the onset of sapropel formation upwards, although not always parallel to Ba/Al enrichments, and generally preceding the productivity maxima, indicating that maximum N fixation was followed by an increase in total export production. As a whole, low $\delta^{15}$N values coincide well with increased organic carbon concentrations, in agreement with other sapropel studies (e.g., Calvert et al., 1992; Meyers and Arnaboldi, 2005; Milder et al., 1999; Struck et al., 2001). We thus infer that blooms of nitrogen-fixing biota were a central factor in creating the higher surface productivity that led to sapropel formation.
4.1.4 Paleoproductivity patterns based on Ba and \( C_{\text{org}} \) concentration.

Greater TOC concentration in sapropel sequences is consistently linked to an increase in \( B_{\text{a_bio}} \). Although \( B_{\text{a_bio}} \) cannot be used to quantitatively determine export productivity (e.g., Averyt and Paytan, 2004; Gingele and Dahmke, 1994; Kasten et al., 2001; McManus et al., 1998, 1994), it has been widely and successfully applied as a relative indicator for enhanced marine productivity in the Mediterranean (e.g., Dehairs et al., 1987; Diester-Haass et al., 1998; Emeis et al., 2000a; Weldeab et al., 2003a). Furthermore, it has been shown that Ba content generally permits a better reconstruction of the duration of enhanced productivity and thus, true sapropel duration, than TOC, because barite is less sensitive to postdepositional oxidative destruction (e.g., Paytan and Kastner, 1996; Thomson et al., 1995). In Mediterranean sapropels, TOC destruction is evidenced by oxidation fronts in the form of high Fe and Mn concentrations above sapropel layers coinciding with low TOC concentrations, whereas high Ba concentrations typically remain preserved (e.g., Gallego-Torres et al., 2010, 2007; Larraisoaña et al., 2003b; Martínez-Ruiz et al., 2000; Thomson et al., 1995, 1999).

TOC and \( B_{\text{a_bio}} \) MAR’s are calculated assuming constant LSR’s for each i-cycle (see Table 1). This assumption can conflict with the accepted “isochronous sapropel model” (e.g., de Lange et al., 2008; Rohling et al., 2006). However, assuming isochronous sapropel boundaries implies shifts in LSR’s of up to 90% higher or up to 45% lower from background to sapropel sediments. Nijenhuis and de Lange (2000) proved the inconsistency of such shifts and suggested that MAR’s should be calculated based on approximately linear sedimentation rate throughout each i-cycle. Owing to the necessity of selecting one hypothesis, constant LSR’s were assumed.

Although the amounts of \( B_{\text{a_bio}} \) increase may vary among the layers, they always accompany the shift to higher TOC-MAR at the base of the sapropels. As a whole, the Late Pleistocene sapropels, ranging in age from 81 ky to 217 ky.

Fig. 9. TOC, C and N ratio, \( \delta^{15}N, \delta^{13}C \) for S6 at all studied sites. Shaded area corresponds to organic enriched sediments.

Fig. 10. TOC, C and N ratio, \( \delta^{15}N, \delta^{13}C \) for S7 at three studied sites. Shaded area corresponds to organic enriched sediments.
Fig. 11. Temporal evolution of the depositional conditions from Pliocene to Pleistocene, represented by TOC, $\text{Ba}_{\text{bio}}$-MAR, N and C isotopic composition and C:N ratio on the Mediterranean Ridge (Site 969). For Holocene sapropel, see Figs. 2 and 6. Maximum off-scale values are numerically indicated in the graphs.

(see supplementary table) exhibit the highest export production levels of the studied sections, as represented by the highest $\text{Ba}_{\text{bio}}$-MAR, although TOC and TOC-MAR’s are higher during the Pliocene and Early Pleistocene. Thus, there is a poor relation between TOC accumulation and productivity ($\text{Ba}_{\text{bio}}$), as presented on Fig. 13 (see also TOC/$\text{Ba}_{\text{bio}}$ on supplementary table), which is particularly evident comparing Pliocene and Late Pleistocene layers. Factors other than export productivity appear to have influenced sapropel formation during the Pliocene.

A disconnect between concentrations of Ba (unaltered signal for increased productivity) and TOC at the top of the Holocene S1 sapropel across the basin (different gray shadings, Fig. 2) and, to a lesser extent, in some Late Pleistocene sapropels, (Figs. 3 to 5; see also Gallego-Torres et al., 2010), indicates post-depositional oxidation of the organic matter at the top of the sapropel (e.g., Martinez-Ruiz et al., 2000; Thomson et al., 1995; Van Santvoort et al., 1996) probably due to relatively easier and faster reventilation of bottom waters. Opposite to this trend, the Early Pleistocene and Pliocene sapropels (Figs. 6 and 11) generally lack oxidation.
4.2 Sapropel expression across the eastern basin through time

Our results reveal basin-wide patterns, temporal variations, and spatial differences in the history of sapropel deposition in the eastern Mediterranean Sea over the past 3 My. The general trend towards lighter N-isotopic composition in the sapropel layers is clear but not particularly strong during the deposition of the Holocene S1 (i-cycle 2). The most plausible explanation for this weak signal is a minor increase in nitrogen-fixing cyanobacteria population and in overall marine productivity. The remaining Pleistocene and Pliocene sapropels show similar and essentially concordant trends across the basin: a progressive decrease of $\delta^{15}N$ values from the onset of sapropel formation upwards, frequently preceding Ba/Al enrichment. At the oxidized top of the sapropel layers, loss of material does not permit us to assess whether N-fixation continued during the whole high productivity phase.

The Ba-MARs during sapropel deposition are equivalent for deep water sites (964, 969, and 967) whereas it is systematically lower at Site 966. The same pattern is seen in TOC-MARs. Sites 966 and 967 are geographically proximate to each other, but their water depths are very different (926 m.b.s.l. vs. 2555 m.b.s.l). Export production, which is a surface ocean process, is expected to be similar at both sites. The difference in Ba MARs may be instead related to the degree of barite saturation and thus its preservation in the water column, which is depth dependent (e.g., Paytan and Griffith, 2007; Van Beek et al., 2006). In any case, for
the same insolation cycle, the relatively shallow Site 966 exhibits the lowest TOC and Ba MARs of the four sites (Figs. 2 to 5). Therefore, it appears that greater water depth favors the burial and preservation of organic matter as proposed by Murat and Got (2000) and refined by de Lange et al. (2008), although it is not a factor influencing the primary process of sapropel formation itself. Sapropels on the top of Eratosthenes Seamount, although possessing lower TOC values, are characterized by all of the same features as those in the deep basin (increase in Ba/Al and Ba-MAR, lower δ15N, higher TOC/TN). These proxies point toward sapropel formation being induced by basin-wide increased surface productivity, at least during the period between i-cycles 20 and 2, where we are able to compare similar layers across the basin.

In terms of time evolution of sapropel events, productivity maxima were reached during i-cycle 12 (S5), coinciding with maximum monsoon index (Kraal et al., 2010; Rossignol-Strick and Paterne, 1999). This pattern allows us to conclude that the same climatic factors (warm/wet periods during summer insolation maxima and enhanced monsoon activity) that triggered increased fluvial discharge and amplified nutrient input also favored the establishment of N-fixing bacteria. This combination of factors ultimately induced sapropel formation.

From Figs. 3, 4 and 5, we observe that the duration of some sapropels (e.g., S5 to S7) appears to become longer from west to east. This pattern may indicate either a progressively shorter increased productivity event towards the west or an increase in sedimentation rates associated with sapropel forming conditions. A similar problem was addressed by Nijenhuis and de Lange (2000) and recently Katsouras et al. (2010) showed that S1 developed later in the Aegean Sea compared to the Lybian coast, by an offset of up to 0.9 ky. We do not have sufficient information (particularly, absolute dating) to resolve this issue. However, we are inclined to believe that this pattern results from differences in sedimentation rates inasmuch as the climate-driven surface paleoceanographic conditions that favored increased productivity were approximately stable and concordant across the basin.
4.3 Diagenesis

The combination of TOC/TN ratios and $\delta^{13}C$ and $\delta^{15}N$ values has been used to determine the origin of the organic matter in marine sediments (e.g., Bouloubassi et al., 1999; Meyers, 1997; Oldenburg et al., 2000; Rullkötter, 2000) by distinguishing between contributions from marine algae, marine cyanobacteria, lacustrine algae, and continental C$_3$-C$_4$ plants, as summarized by Meyers (1997). These parameters are plotted in Fig. 14 for the 35 analyzed sapropels. Holocene sediments plot close to the marine algae end member, with minor deviations coinciding with the highest TOC values. Pleistocene and Pliocene samples show a nearly bimodal distribution on the TOC/TN-$\delta^{13}C$ plot. The great majority of low TOC samples fall in the marine domain, the exception being a few samples from Site 967. Samples from within the sapropel shift towards higher C:N fields. In contrast, the $\delta^{15}N$ vs. $\delta^{13}C$ plot systematically shows a marine algae composition for non-sapropel samples and a typical marine cyanobacteria composition for sapropel samples.

Although the TOC/TN vs. $\delta^{13}C$ relation (Fig. 14) could be misinterpreted as resulting from an influence of continental input (i.e., C$_3$ plants), the low $\delta^{15}N$ and $\delta^{15}N$ vs. $\delta^{13}C$ plots suggest that the sapropels are dominated by marine organic matter, and independent evidence such as large values of the Rock-Eval Hydrogen Index (Bouloubassi et al., 1999; Emeis et al., 1996) and an abundance of marine biomarker molecules (Rinna et al., 2002) supports this interpretation. Nitrogen isotopic compositions are only slightly or not at all affected in preserved organic matter (Higgins et al., 2010), whereas under conditions of high surface productivity, the typical TOC/TN value of the exported organic matter is higher than the typical algal signal, partly due to selective remineralization of N and N-rich molecules below the euphotic zone and/or at the sediment surface (e.g., Twichell et al., 2002). According to Freudenthal et al. (2001; see also Macko et al., 1994), TOC/TN ratios would increase when remineralization and preferential degradation of amino acids (isotopically higher) occurs along the water column. Similar results were obtained in sediment traps studies done by Van Mooy et al. (2002). Selective remineralization of amino acids simultaneously yields lower $\delta^{15}N$ values and higher TOC/TN values of deposited organic matter. The increase in TOC/TN can also partly be caused by denitrification under a suboxic environment (Karl et al., 2002) that is easily achieved under highly productive waters. These diagenetic changes support the conclusions made by Arnaboldi and Meyers (2006) on similar sapropel-bearing sediments in the eastern Mediterranean. It is evident that this increase in TOC/TN ratio is more extreme during the Pliocene, in agreement with a highly restricted water column oxygenation for this period as postulated by Passier et al. (1999), Böttcher et al. (2003), Warning and Brumsack (2000), Larrasoña et al. (2003a), and Weldeab et al. (2003a, b).

Lower N-isotopic composition of organic matter mirrors the organic-rich levels, but it does not extend above oxidized sapropels to parallel the increase in productivity. Because nitrogen fixation seems to be crucial for sapropel formation, changes in export productivity (B$_{bio}$) should run parallel to N-fixing activity ($\delta^{15}N$), inasmuch as both reflect primary production. However, the post burial burn-down oxidation may affect this signal. Specifically, oxidation of deposited organic matter preferentially consumes $^{14}N$, so that the remaining organic matter in the sediment becomes enriched in $^{15}N$ thus altering the $\delta^{15}N$ signal. Higgins et al. (2010) postulate that the N isotope signal is not affected by oxidation of decaying organic matter, although the resolution of their analyses is not high enough to discriminate oxidation fronts. However, if we accept their hypothesis, we may infer that the increase of $\delta^{15}N$ values immediately above sapropels S1 or S7 (light shading in Figs. 2 and 5) is due to better oxygenation of the overlying water column while high productivity conditions still prevailed.

5 Summary and conclusions

The remarkably lighter N isotopic compositions of the sapropel layers indicate periods of nitrogen fixation and thus, amplified cyanobacterial productivity (probably associated with bacterial mats) in the water column. Although N-fixation is frequently associated with oligotrophic basins, in the case of Eastern Mediterranean sapropels it created a highly productive environment. This increase in N-fixing biota and bacterial primary productivity increased export production and is reflected in major increases in TOC and B$_{bio}$ MARs. The change in mode of productivity is fueled by nutrient input (such as P, Fe or Mo), most likely through riverine discharge and by a warming environment with higher SST and lower SSS (Emeis and Weissert, 2009; Emeis et al., 2000b). These climatically driven productivity events affected the entire Eastern Mediterranean, independent of water depth. Differences in sapropel expression depend on water depth, related to winnowing, deep water circulation and/or SO$_4^{2-}$ saturation.

Our results allow us to distinguish between the importance of organic matter production and its preservation in forming the sapropel layers. TOC-MAR and B$_{bio}$ MAR do not show a direct correlation with C$_{org}$ concentration and Ba/Al ratio, respectively. Some Late Pleistocene sapropels, such as S5 (i-cycle 12) have lower TOC concentrations but higher TOC and B$_{bio}$ MAR’s than older equivalents, indicating that both marine productivity and detrital sedimentation dramatically increased during the Late Pleistocene. Opposite to this trend, Early Pleistocene and Pliocene sapropels accumulated under markedly lower sedimentation rates and therefore exhibit higher Ba concentrations but lower MARs than Late Pleistocene equivalents. In other words, greater productivity does not directly result in higher TOC accumulation (Fig. 13). The Pliocene and Early Pleistocene relatively smaller increases
in productivity correspond to limited water mass circulation than in Late Pleistocene times; water mass circulation may have even reversed (estuarine circulation, e.g., Böttcher et al., 2003; Wehausen and Brumsack, 1999) favouring bottom water stagnation. This interpretation implies that organic carbon concentration in these layers is equally influenced by both preservation and productivity. Variations in deep water circulation are also responsible for different diagenetic imprints on TOC, TOC/TN ratio and $\delta^{15}$N signals.

Humidity maxima also contribute to the intensity of productivity increase by magnifying fluvial nutrient input into the basin. Higher humidity signifies stronger river runoff, higher nutrient supply from the continent and, at the same time, increased detrital input that would increase sedimentation rates. If we consider the increase in African derived nutrient input as climatically controlled (in turn defined by insolation cycles), the maximum increase in fluvial input should correspond to i-cycles 12 to 20, which show the greatest maximum summer insolation (500 to 520 W/m$^2$) according to Emeis et al. (1996) and Lourens et al. (2004). Sampled sapropels from i-cycles 152 to 182 correspond to insolation peaks that remain around 500 W m$^2$ (Larrasoana et al., 2003a) or below, for i-cycles 270 to 286. Thus, detrital and nutrient input would be the highest at maximum insolation (i.e., Late Pleistocene). This prediction is supported by our results that reveal relatively low TOC concentration compared to older equivalents and, at the same time, the very high TOC and $B_{\text{bar}}$-MARs found at all four sites, for example during i-cycle 12. The relative sizes of the increases in sedimentation rates and productivity rates are evidently related to the amplitudes of the precessional insolation and humidity maxima (monsoon index).

A remarkable increase in TOC/TN ratios, related to denitrification processes, suggests intense oxygen consumption below the euphotic zone. The $\delta^{15}$N signal, although mostly representing a primary signal, might be shifted to somewhat larger values by diagenetic oxidation that is evident in diminished TOC concentrations at the tops of sapropel layers because of oxidative burndown.

Supplementary material related to this article is available online at: http://www.biogeosciences.net/8/415-2011-supplement.pdf.

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