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MIDDLE EOCENE-LOWER OLIGOCENE CALCAREOUS NANNOFOSSIL  
BIOSTRATIGRAPHY AND PALEOCEANOGRAPHIC IMPLICATIONS FROM  
SITE 711(EQUATORIAL INDIAN OCEAN)

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Calcareous nannofossils, biostratigraphy, Eocene, Oligocene, paleoceanography

**Abstract**

Nannofossil data from ODP Site 711 (equatorial Indian Ocean) yield a set of consistent, reliable biohorizons that form the basis of a revised calcareous nannofossil biostratigraphy for the low-latitude Eocene-Oligocene. We discuss 31 biohorizons occurring over an 11 myr time interval which we correlate to previous magnetostratigraphic data. Calcareous nannofossils from the middle Eocene through the lower Oligocene of the studied section are characterized by moderately well preserved assemblages consisting largely of low latitude and cosmopolitan species. A significant nannofossil dissolution interval is evidenced at the middle Eocene Climatic Optimum (MECO). We document a significant increase in late Eocene nannoplankton exhibiting a eutrophic preference. Analysis of the assemblage suggests important changes in the equatorial oceanic regime just before the onset of

the Eocene-Oligocene transition (EOT), that foreshadow the more dramatic climatic shift of the early Oligocene.

## 1. Introduction

Ocean Drilling Program (ODP) Leg 115 took place in the Indian Ocean along a south - north bathymetric transect in order to investigate the Reunion volcanic system and Paleogene to Quaternary stratigraphy and paleoceanography, focusing on the carbonate dissolution history of the Indian Ocean (Duncan et al., 1990).

Paleomagnetic data from the Initial Report (Backman and Duncan, 1988) provided a poorly constrained magnetostratigraphic interpretation for part of the late Oligocene and late Miocene - Pleistocene interval, thus the lower part of the cored succession was dated only by means of biostratigraphy.

Subsequently, the shipboard nannofossil specialists (H. Okada and D. Rio) produced several papers based on both semi-quantitative and quantitative analyses from the Paleogene to the Quaternary (Rio et al., 1990a; Fornaciari et al., 1990; Matsuoka and Okada, 1990; Okada, 1990). However, the “standard” zonations of Martini (1971) and Okada and Bukry (1980) have been proven to be of limited use for the Eocene-Oligocene because of the low reliability or scarcity of some markers. In the last decade, calcareous nannofossil biostratigraphy of the lower Eocene-Oligocene sediments has shown great potential for improvement, through identification of many new nannofossil species (e.g., Fornaciari et al., 2010; Bown and Dunkley

Jones, 2012; Toffanin et al., 2013). These studies formed the basis for the revised nannofossil biozonation of Agnini et al. (2014).

In this study we present a quantitative record of nannofossil assemblages through the middle Eocene –lower Oligocene succession in Hole 711A (ODP Leg 115). The abundance pattern and the stratigraphic range of nannofossil markers allowed us to increase the number of bioevents and to compare their reliability and synchronicity in other areas. Site 711 has been the focus of a recent magnetostratigraphic study (Savian et al., 2013) that spans the interval between Chrons C20n (middle Eocene) and C12n (lower Oligocene). Although at times the magnetic stratigraphy is of low resolution (Fig. 2) because of the high number of samples with no reliable magnetic directions (Savian et al., 2013), we have achieved a biochronology by means of higher resolution sampling and by using new biostratigraphic and taxonomic updates.

A further aim of this work is to determine if abundance variations of selected nannofossil taxa reflect the global paleoclimatic changes during this crucial time interval of the Earth climatic evolution. In fact, there is general agreement on the usefulness of calcareous nannofossils as proxies for Paleogene paleoenvironmental reconstructions (e.g., Wei et al., 1992; Bralower, 2002; Dunkley Jones et al., 2008; Pearson et al., 2008; Villa et al., 2008).

Site 711 has a paleodepth of greater than 3000 mbsl and, along with sites 1218 and 1219 (Lyle et al., 2005) and Integrated Ocean Drilling Program (IODP) sites U1331- U1334 (Pälike et al., 2010), is one of the equatorial sites that represents an opportunity to study the carbonate accumulation history and the large fluctuations

of the carbonate compensation depth (CCD) during the Eocene (e.g., Coxall et al., 2005; Pälike et al., 2012). The investigated interval encompasses the Middle Eocene Climatic Optimum (MECO), whose top is correlated with a dissolution event (Bohaty et al., 2009), and the long cooling trend that leads to the Oligocene glacial state (Oi-1, Miller et al., 1991). The relationship between CCD fluctuations, ocean acidity variations,  $p\text{CO}_2$  concentrations, and ultimately climate are matters of intense discussion (e.g., Pearson et al., 2008; Coxall and Wilson, 2011) and the study of calcareous nannofossils has been shown to be an excellent tool to add information on paleoclimate history (Villa et al., 2014).

## 2. Material and methods

ODP Leg 115 drilled five sites along a depth transect in the western equatorial Indian Ocean of which Site 711 represents the deepest end member. Hole 711A is located on Madingley Rise, at  $2^{\circ}44.56'S$  and  $61^{\circ}09.78'E$  at a water depth of 4428.2 mbsl (Fig. 1). For the studied interval the paleodepth has been estimated to have been between 3450 and 3750 mbsl (Peterson and Backman, 1990).

The studied samples were collected from Core 711A-14 to Core 711A-25, spanning the middle Eocene- early Oligocene interval. The mean sampling resolution is  $\sim 90$  cm, and in key time intervals, it is as high as 37 cm, allowing an average biostratigraphic resolution of 0.42 kyr, that reaches a value of 0.34 kyr in the Eocene. The studied cores in Hole 711A are composed of nannofossil oozes or clay-bearing nannofossil oozes with a consistent occurrence of radiolarians and

comprise part of the lithologic units III and IV of Backman and Duncan (1988) providing a continuous middle Eocene - lower Oligocene stratigraphic sequence. To better constrain the position of some bioevents of taxa with rare and scattered distribution (i.e., the topmost stratigraphic presence of *Nannotetrina* spp., *Chiasmolithus gigas*, and *Chiasmolithus solitus*) we supplemented our dataset with data from Wei et al. (1992). The smear slides were prepared from unprocessed samples using standard techniques (Bown and Young, 1998), and quantitative analyses were conducted in the light microscope at a magnification of 1250X, counting at least 300 specimens per sample. Two additional long traverses, corresponding to 12.56 mm<sup>2</sup>, were scanned in order to identify rare taxa. In order to create a clearer picture of assemblage changes over time the absolute abundance of each taxon was calculated by converting the number of observed specimens to the total number of specimens in an area of 1mm<sup>2</sup>. In addition, the same data plotted versus age, representing the assemblage variations, may indicate the paleoecological response to paleoclimatic events previously highlighted (Coxall et al., 2005; Bohaty et al., 2009; Villa et al., 2014). In this study we did not use percentage values because of the dissolution affecting some intervals, that could overestimate the dissolution-resistant taxa (e.g., *Discoaster* spp. and *Cyclicargolithus floridanus*). Only *Clausicoccus* spp. and *Cribozentrum erbae* were plotted as percentage values because their definition as bioevents is strictly related to their increase in abundance *versus* the total assemblage (Fornaciari et al., 2010). In the studied material, nannofossils are moderate to well-preserved, dissolution seems to affect moderately the assemblage,

whereas the overgrowth is weak but persistent along the studied interval. We observed at times a slight to moderate etching and a slight to moderate overgrowth in the same sample, which could be indicative of secondary overgrowth on more robust nannofossils at the expense of more delicate ones. At times this makes it difficult to distinguish some taxa consistently at specific levels (e.g., overgrowth of *Nannotetrina* and *Discoaster* nannoliths, and etching of the central area of *Reticulofenestra*). The scarcity of holococcoliths, represented here only by *Zygrhablithus bijugatus*, could be explained by a moderate dissolution and/or by low nutrient availability, a feature of open ocean environments. Most of the identified marker species are illustrated in microphotographs (Plates 1 and 2). In a short interval of the lower part of the studied section, we document specimens of *Sphenolithus* cf. *S. kempii* with four extremely elongated apical spines that could be related either to overgrowth or represent a morphotype of *S. kempii* (Plate 1, Figs 21-24).

The age model for the studied section is based on the magnetic polarity stratigraphy of Savian et al. (2013), correlated with the geomagnetic polarity time scale (GPTS) of Gee and Kent (2007), assuming constant sedimentation rates between the magnetostratigraphic tie points. The obtained biochronology provides a tool for comparison with other sites from the Pacific and Atlantic oceans (Tab. 1).

### **3. Middle Eocene-lower Oligocene biostratigraphy**



The traditional zonations of Martini (1971) and Okada and Bukry (1980) were developed respectively in mid latitude land and low latitude oceanic sections, but due to the lack of some zonal markers (e.g., *Blackites gladius* and *Chiasmolithus oamaruensis*) or scarcity of others (e.g., *Nannotetrina fulgens*, *Isthmolithus recurvus*), particularly some of the upper Eocene zones of both schemes are not recognized and correlated over wide areas. For this reason, we use additional bioevents, beside the aforementioned zonal schemes, in order to increase the biostratigraphic resolution (Fig. 2) for the middle Eocene - early Oligocene interval.

A reliable biohorizon should be easily reproducible, that means consistently correlated over wide distances, maintaining its position relative to other biohorizons and lastly, maintaining its position within the geomagnetic polarity time scale (Bralower et al., 1989; Rio et al., 1990b; Aubry, 1995; Berggren et al., 1995). Discrepancies may emerge in biomagnetostratigraphic correlations worldwide, related to the rate of dispersal over different biogeographic provinces.

The biohorizons recognized herein are summarized in Table 1 and, following Backman et al. (2012) and Agnini et al. (2014), have been labeled as Base (B), Base common (Bc), Top (T), and Top common (Tc) (Figs 3a, 3b and 3c).

The mean depths of some bioevents coincide (e.g., Base common of *R. daviesii*, Base of *S. akropodus* and Base common of *S. predistentus*) and this could be explained by a low sedimentation rate or by the large sample spacing, but the presence of a hiatus of short duration in Chron C13r cannot be ruled out.

The reliability and reproducibility of detected events from the base of the studied interval is discussed below.

### **Top of *Sphenolithus kempii***

*Sphenolithus kempii* (Plate 1, Figs 19-20) was first detected at Sites U1331-1334, in the equatorial Pacific (Bown and Dunkley Jones, 2012) restricted to the lower part of the NP15 Zone. The species is present from the base of the studied section and its Top was detected at 232.05 mbsf, (Fig. 3a), below the Top of *Chiasmolithus gigas* (base of CP13b), in a similar position to that recorded in the Pacific Ocean; further investigation in low to middle latitude areas may confirm the usefulness of this species over a wide geographic area.

### **Top of *Chiasmolithus gigas***

Okada and Bukry (1980) utilized this bioevent within their Zone CP13. As *Chiasmolithus* prefers temperate-cool waters, this species is quite rare at equatorial latitudes (Perch-Nielsen, 1985; Wei and Wise, 1989). At mid-latitudes, the Top of *C. gigas* is placed at the top of Chron C20r (Berggren et al., 1995; Jovane et al., 2007; Agnini et al., 2014). In Hole 711A, this event is recorded at 219.41 mbsf, in an interval without magnetostratigraphic data, below Chron C20n (Fig. 3a).

Likewise, in previous works this species may have been merged with *Coccolithus mutatus* misrepresenting its true distribution (Bown, 2005). Further investigations through a biometric analysis of the two species could point out the presence of transitional forms representative of an evolutionary trend.

**Base of *Sphenolithus runus* and *Sphenolithus strigosus***

*Sphenolithus runus* and *Sphenolithus strigosus* (Plate 2, Figs 1-2) were described from the onshore drilling sites of Tanzania (Bown and Dunkley Jones, 2006) and from the equatorial Pacific (Bown and Dunkley Jones, 2012) as first occurring in NP 16 Zone. Our data extend the range of these taxa to a lower stratigraphic level within Zone NP15, in agreement with the data from Demerara Rise (Lupi and Wise, 2006). Because of the distinctive morphology and distribution pattern of these species, both Bases are easily identifiable and are potential good markers for the middle Eocene.

**Base of *Reticulofenestra umbilicus***

This event defines the base of the Subzone CP14a, and though the initial range of *Reticulofenestra umbilicus* (Plate 1, Figs 15-16) is characterized by low percentages, its morphometric features ( $\geq 14 \mu\text{m}$ ) allow us to identify with confidence the Base of this taxon at 202.15 mbsf (Fig. 3a). The discrepancy with the data of Okada (1990), who detected the species about four meters above, is probably due to different sampling resolution, combined with the low number of this taxon in its initial range. This event has been recognized in Chron C19r at high and low latitudes (Wei and Wise, 1989; Fioroni et al., 2012; Toffanin et al., 2013) or in Chron C20n (Fornaciari et al., 2010; Agnini et al., 2014). At ODP Hole 711A, a lack of magnetostratigraphic data in this interval does not allow a direct

correlation of the event with the GPTS, however a correlation with Chron C20n is most plausible.

#### **Top of *Nannotetrina* spp.**

This bioevent has been largely used to approximate the base of Zone NP16 when *B. gladius* is missing (Perch-Nielsen, 1985), as in our material and in other equatorial sites (Bown and Dunkley Jones, 2012). In the studied section, *Nannotetrina* is difficult to identify to species level because of strongly overgrown specimens, therefore here we use the extinction of the genus (194.1 mbsf; 41.739 Ma) occurring in C19r (Fig. 3a). However, Agnini et al. (2014) observed this event at the top of Chron C20n, so the different results could be due to the scarcity of this genus.

#### **Top of *Sphenolithus runus***

This event has been detected at 191.92 mbsf (Fig. 3a), with an age of 41.335 Ma, in Zones NP16/CP14a and, as for the Base of this taxon, it is a promising biohorizon and further investigations in other areas are needed to test its reliability at low – mid latitudes.

#### **Base and Base common of *Cribozentrum reticulatum***

In Hole 711A, we record the Base of *Cribozentrum reticulatum* (Plate 1, Figs 1-2, 5) at 191.92 mbsf in Chron C19n (Fig. 3b), with an age of 41.335 Ma, whereas data from higher latitudes, i.e., at Site 1172 (Persico et al., 2012) and Sites 689 and 690

(Wei and Wise, 1990) report it in Chron C18n.2n. In the initial range it is very rare and sporadic, and therefore the Base of *C. reticulatum* has a low reproducibility (Fornaciari et al., 2010). The Base common (Bc) of the species is recorded at 181.36 mbsf (38.215 Ma), at the top of Chron C17r. Data from the Alano section and Site 1052, show these events in Chron C18r and in C18r/19n respectively (Fornaciari et al., 2010); Agnini et al. (2014) utilized the Base common (Bc) of *C. reticulatum* for the base of the CNE14 Zone, at the onset of Chron C19r. The high diachroneity of this bioevent casts doubt on its reliability over long distances, but it may be useful on a regional scale.

#### **Base and Base common of *Sphenolithus predistentus***

The first occurrence of *Sphenolithus predistentus* (Plate 2, Figs 17-21) occurs in upper Chron C18r with a temporary absence up to Chron C18n.2n in the mid-latitude record at Site 1052, and up to the base of Chron C17r in the Alano section (Fornaciari et al., 2010). We detect the first specimens at 191.92 mbsf (41.335 Ma), at the top of Chron C19n, at the same stratigraphic level as the Top of *S. runus* and the Base of *C. reticulatum* (Fig. 3b), suggesting condensed sedimentation, in agreement with the low sedimentation rate (0.833cm/kyr) reported by Savian et al. (2013) for this interval. The species becomes continuous and abundant (Bc) only from Chron C13r (157.12 mbsf; 34.476 Ma), as reported from Site 1218 (Blaj et al., 2009). The different distribution patterns of this sphenolith over different geographical areas could be related to biogeographical control, or to taxonomic problems tied to the evolutionary transition to *Sphenolithus akropodus* and

*Sphenolithus distentus* (e.g., Bown and Dunkley Jones, 2012). Considering that in the equatorial Indian Ocean the genus is abundant and well preserved through the Eocene-Miocene interval (Fornaciari et al., 1990; Wei et al., 1992), we tend to regard this Base common (Bc) as reliable.

### **Tops of *Sphenolithus furcatolithoides*, *Sphenolithus spiniger* and *Sphenolithus strigosus***

In Hole 711A the Tops of *Sphenolithus furcatolithoides* (Plate 2, Figs 7-10), *Sphenolithus spiniger* (Plate 2, Figs 13-14) and *Sphenolithus strigosus* (Plate 2, Figs 1-2) fall in the NP16/CP14a Zone at the same stratigraphic level (40.174 Ma), near the top of Chron C18r (Fig. 3b), which corresponds to the beginning of a carbonate dissolution interval marked both by the absence of nannofossils and by the mass accumulation rate (MAR) data (Peterson and Backman, 1990).

In the Pacific and Atlantic low to mid latitude sites (Fornaciari et al., 2010; Bown and Dunkley Jones, 2012; Toffanin et al., 2013) these events occur with minor age differences (Table 1) that can be ascribed to the dissolution interval in our material responsible for masking the real Tops. However, the absence of these sphenoliths above the dissolution interval suggests that they are older than the top of Chron C18r.

### **Base and Top of *Sphenolithus obtusus***

The short stratigraphic distribution of *Sphenolithus obtusus* (Plate 2, Figs 3-6) has a well constrained range between the upper NP16 and lower NP17 Zones (Agnini et

al., 2011; Bown and Dunkley Jones, 2012) and has been used for the new zonal scheme at mid latitudes (Fornaciari et al., 2010) and for the base of Zone CNE16 (Agnini et al., 2014). In Hole 711A, the Base of this species is recorded just above the middle Eocene dissolution event in Chron C18n (184.27 mbsf; 39.007 Ma) (Fig. 3b) and thus could occur in a lower sample. The Top is placed in the lower part of Chron C17n (178.56 mbsf; 36.925 Ma), above the Bc of *Criboecium erbae*, with a range apparently more extended compared to the biozonation of Agnini et al. (2014), where the Top is reported in Chron C18n.1n. This discrepancy could be due to different biogeographical distribution or, alternatively, to reworked specimens of *S. obtusus*.

#### **Base common of *Dictyococcites bisectus***

Following the original description (Hay et al., 1966), we include in this taxon specimens  $<10\ \mu\text{m}$  (Bown and Dunkley Jones, 2012; Persico et al., 2012), labeling the specimens  $>10\ \mu\text{m}$  as *Dictyococcites stavensis* (Plate 1, Fig. 11). In Hole 711A, *D. bisectus* (Plate 1, Fig. 12) occurs with low abundances in its initial range (from 191.71 mbsf, Fig. 3b), while the Base common (184.27 mbsf) is recorded just above the dissolution event of the middle Eocene (top of NP16 /CP14a Zones), in the middle-upper part of Chron C18n (39.007 Ma). Yet, the dissolution event may mask the true Base common of this taxon and we cannot exclude that this bioevent could be slightly older than assumed. Data from the Southern Ocean (SO) record the Base of *D. bisectus* in Chron C17n.1n (Persico et al., 2012) confirming this taxon as time transgressive, from middle to high latitudes (Wei and Wise, 1990).

**Top of *Chiasmolithus solitus***

Chiasmoliths are considered to belong to cool to temperate-water taxa (e.g., Wei et al., 1992; Persico and Villa, 2004) and are generally rare in tropical environments. The Top of *C. solitus* is reported as diachronous at different latitudes, disappearing at high southern latitudes later than at low latitudes (Villa et al., 2008; Fioroni et al., 2012). In Hole 711A, despite the scarce and scattered occurrence of the species, especially in its final range, this event is tentatively placed in the upper part of Chron C18n (38.744 Ma), in agreement with data from low-middle latitudes (Wei and Wise, 1989; Fornaciari et al., 2010; Agnini et al., 2011; Toffanin et al., 2013).

**Base common and Top common of *Cribrocentrum erbae***

This species (Plate 1, Figs 3-4) was first described in Fornaciari et al. (2010) and was also recorded from the equatorial Pacific (Bown and Dunkley Jones, 2012). In Hole 711A the species shows the same distribution pattern described at Alano and at Site 1052 (Fornaciari et al., 2010), with a noticeable acme reaching percentages higher than 12% of the total assemblage (Fig. 3b). The acme interval begins with a distinct increase (Bc; 38.03 Ma) and is followed by a sharp decrease in abundance (Tc; 36.925 Ma), with very rare and scattered specimens present before and after the acme interval. The acme is correlated with Chron C17n, and seems to be a promising biohorizon, at least at low to mid-latitudes. Agnini et al. (2011; 2014) proposed the Base common of *C. erbae* as an alternative biohorizon for the base of Zone NP18, usually defined by the Base of *Chiasmolithus oamaruensis*. This latter



species is very rare in low to middle latitudes, thus the *C. erbae* Base common could be useful for the definition of the GSSP for the base of the Priabonian Stage.

### **Base common of *Sphenolithus intercalaris***

Toffanin et al. (2013) have recently identified the Base common of *Sphenolithus intercalaris* at Pacific equatorial latitudes in the late Eocene (Chron C16n.2n) (Plate 2, Figs 15-16). In the studied section, we note an increase in abundance of this taxon in Chron C17n (37.632 Ma) (Fig. 3c), and further investigations are needed to test the reliability of this bioevent for correlations between low latitude sites.

### **Top of *Chiasmolithus grandis***

This event defines the top of the CP14b Zone of Okada and Bukry (1980). In Hole 711A the species (Plate 1, Figs 6-8) is present from the base of the studied section (Fig. 3c). The Top is detected at 177.11 mbsf in Chron C17n with an age of 37.57 Ma. Data from Site 523 indicate this bioevent in Chron C17n.1n (Backman, 1987) and in Chron C17n at Site 516 (Wei and Wise, 1989). Toffanin et al. (2013) identify this extinction in Hole 1333C in the upper part of Chron C18n.1n. At mid-latitudes Fornaciari et al. (2010) identify both the Top common and the Top of *C. grandis* in Chron C18n.1n and in the middle part of C17n.2n, respectively. In Hole 711A the distribution pattern of this taxon allows us to identify a neat Top, but not a Top common. Based on these data the *C. grandis* Top has to be considered diachronous and an unreliable biohorizon (Agnini et al., 2014).

**Base of *Isthmolithus recurvus***

The base of NP19 and of CP15b is defined by this event, though at low latitudes it is quite rare, having a preference for cool waters (Bukry, 1973; Wei and Wise, 1990). Moreover an early occurrence of *I. recurvus* in Chron C17n.1n has been highlighted by Backman (1987), Villa et al. (2008), and Fornaciari et al. (2010), with a re-entry in upper Chron C17n.1n (Agnini et al., 2014). At Site 711 Okada (1990) indicates the Base of *I. recurvus* at 163.87 mbsf, at the top of Chron C16n. Due to the extreme scarcity and the poor preservation of this species in the studied material, we do not use this event, otherwise useful in other areas.

**Top common of *Cribozentrum reticulatum***

This event is considered diachronous between mid-low and high latitudes, at high latitudes it occurs in Chron C16n (Wei, 1991; Wei and Thierstein, 1991; Marino and Flores, 2002; Channell et al., 2003; Persico et al., 2012). In Hole 711A, it is recorded in Chron C15n (at 159.71 mbsf, Fig. 3b) with an age of 34.647 Ma, in agreement with previous low-mid latitude studies (Backman, 1987; Wei and Wise, 1989; Fornaciari et al., 2010; Agnini et al., 2014). Our data show that the Tc of *C. reticulatum* is an excellent event in low and mid latitudes, although diachronous with respect to high latitudes, being environmentally controlled (Wei et al., 1992; Villa et al., 2008).

**Top common of *Discoaster barbadiensis***

Here, this event occurs in the lower part of Chron C13r (34.575 Ma) and slightly precedes the Top of *Discoaster saipanensis*, in agreement with data from Site 1218 (Blaj et al., 2009), Site 522 (Poore et al., 1984) and Site 516 (Wei and Wise, 1989). Both *D. barbadiensis* (Plate 1, Fig. 18) and *D. saipanensis* Tops are useful events, being easily identified even when the specimens are overgrown.

### **Base of *Sphenolithus akropodus***

DeKaenel and Villa (1996) first observed this taxon and defined its stratigraphic range as restricted to lower Oligocene Zones NP22 to NP23. In the equatorial Pacific Bown and Dunkley Jones (2012) record the stratigraphic range of this species in the lower Oligocene, NP21- NP22 Zones. *Sphenolithus akropodus* (Plate 2, Figs 23-24) likely corresponds to *Sphenolithus* aff. *S. distentus* in Okada (1990) and to *Sphenolithus* sp.1 in Fornaciari et al. (1990), who report this form restricted to the lower Oligocene and define it as a transitional form to *S. distentus*. We detect *S. akropodus* from 157.12 mbsf (34.476 Ma), therefore our data extend the range of this taxon down to the late Eocene, in Chron C13r (Fig. 3c).

### **Base common of *Reticulofenestra daviesii***

Studies on nannofossil paleoecology indicate that *Reticulofenestra daviesii* is a cold (e.g., Wei and Thierstein, 1991) or cool-water taxon (e.g., Persico and Villa, 2004) that shows an abrupt increase at the Oi-1 glacial event (Wei et al., 1992; Monechi et al., 2000; Persico and Villa, 2004; Villa and Persico, 2006; Villa et al., 2008). In the studied section we observe an increase in abundance at 157.12 mbsf (Fig. 3c),

corresponding to the lower part of Chron C13r (34.476 Ma), preceding then the EOT.

### **Top of *Discoaster saipanensis***

This event marks the top of the NP20 and CP16 Zones and is considered the closest nannofossil event to approximate the Eocene-Oligocene boundary, defined by the HO of the hantkeninids (Nocchi et al., 1988). There is a general consensus about the Top of this taxon at low-middle latitudes occurring in Chron C13r (Poore et al., 1984; Shackleton et al., 1984; Wei and Wise, 1989; Coxall et al., 2005; Toffanin et al., 2013; Agnini et al., 2014). Our data substantially agree with previous results and give an age of 34.187 Ma (at 155.26 mbsf) to this event. A few specimens in the lower Oligocene sediments are considered to be reworked, like some of the planktonic foraminifera reported at this site by Premoli Silva and Spezzaferri (1990).

### **Base common of *Clausicoccus* spp.**

The Base common of *Clausicoccus subdistichus* defines the top of CNE21 Zone (Agnini et al., 2014) and approximates the base of the Oligocene as it is correlated with the top of Chron C13r (Toffanin et al., 2013). In our samples, calcite etching/overgrowth make it difficult to classify *Clausicoccus* at the species level, however the genus increases by 2% of the assemblage (Fig. 3c), just after the Top of *D. saipanensis*, at the base of C13n (33.509 Ma).

**Top of *Ericsonia formosa***

This event marks the top of the NP 21 Zone; in Hole 711A the Top of *E. formosa* (Plate 1, Fig. 9) is recorded at 141.26 mbsf (Fig. 3c) in the lower middle part of Chron C12r (32.24 Ma), in agreement with data from the equatorial Pacific (Blaj et al., 2009; Bown and Dunkley Jones, 2012; Toffanin et al., 2013) and South Atlantic (Poore et al., 1984; Backman, 1987; Wei and Wise, 1989).

**Top of *Reticulofenestra umbilicus***

Both Martini (1971) and Okada and Bukry (1980) used this bioevent in their zonal schemes, defining the base of the NP23/CP17 Zones. In Hole 711A, towards the end of its range the species is rare and scattered. Despite this abundance pattern, we have identified its Top at 130.86 mbsf (Fig. 3a) with an age of 31.059 Ma, in upper Chron C12r, and consider the event consistent and reproducible over wide distances, in general agreement with previous age assignments (e.g., Marino and Flores, 2002; Persico and Villa, 2004; Wei, 2004; Blaj et al., 2009).

**4. Paleocceanographic implications****4.1 Middle Eocene CCD fluctuation in the equatorial Indian Ocean**

The Eocene climate cooling, that witnessed the transition from greenhouse to icehouse conditions, is punctuated by climate reversals, the most striking of which is the MECO (Bohaty and Zachos, 2003) at ~ 40 Ma (Villa et al., 2008; Bohaty et al., 2009). This warm event is characterized by worldwide CCD shoaling (Bohaty et al., 2009; Pälike et al., 2010; Pälike et al., 2012). At Site 711 the carbonate

content in sediments dated from 42 to 31 Ma averages ~72%, ranging between 0 and over 90%, and shows the same trend as the nannofossil total abundance. It has been hypothesized that the total loss of carbonate content, recorded also in other equatorial deep water sites, is mostly related to carbonate dissolution rather than to changes in paleoproductivity (Bohaty et al., 2009), revealing a strong relationship of the nannofossil records with CCD paleodepth history (Pälike et al., 2012). Our data show that the total abundance, expressed as number of nannofossils N/mm<sup>2</sup>, considerably decreases in an interval about 3m thick in Core 711A-20 (from 184.27 mbsf to 187.51 mbsf) nearly corresponding to an interval of low CaCO<sub>3</sub> content, indicating a shoaling of the CCD (Peterson and Backman, 1990). The onset of the dissolution event coincides with the Tops of *S. spiniger* and *S. furcatolithoides* as detected in the Pacific Ocean (Toffanin et al., 2013). This interval is almost barren in nannofossils and the only taxa present are rare specimens of dissolution-resistant *Discoaster* spp. and *C. floridanus*. There is a general agreement about the *Discoaster* dissolution resistance, while our data disagree with Blaj et al. (2009), who consider *C. floridanus* a dissolution-prone taxon.

The absence of calcareous microfossils spans the interval from the top of Chron C18r to the mid part of Chron C18n (Fig. 4) and matches the minimum of CaCO<sub>3</sub> content indicated by MAR data (Peterson and Backman, 1990), the slight difference between the two curves is ascribable to different sample sets.

The onset of the dissolution event at the top of Chron C18r has been recorded at other deep sites, e.g., the Pacific equatorial Sites 1218 and 1219 (Bohaty et al., 2009), and correlated with the MECO, however at Site 711, the C17/C18 interval is

not well defined (Fig. 2) and together with the low sedimentation rate of this interval, a precise correlation of the peak of the dissolution event, with the peak of the MECO, is hampered.

#### **4.2 The Eocene / Oligocene Transition and Nannofossil assemblage variations**

The Eocene-Oligocene Transition (EOT) encompasses the interval between ~34 and 33.5 Ma, including the E/O boundary (33.7 Ma) and the positive  $\delta^{18}\text{O}$  shift which defines the transition to a fully glaciated state (Miller et al., 1987; Zachos et al., 2001; Miller et al., 2005; Cramer et al., 2009; Coxall et al., 2005; Coxall and Wilson, 2011), with sea surface temperatures decreasing both at high and low latitudes (e.g., Zanazzi et al., 2007; Lear et al., 2008; Liu et al., 2009; Wade et al., 2012) and a global sharp deepening in the CCD, with a shift from siliceous to calcareous sediments (Pälike et al., 2010).

Long-term Eocene cooling was not entirely monotonic, rather, intervals of rapid climate variations have been documented (Vonhof et al., 2000; Bohaty and Zachos, 2003; Villa et al., 2008; Coxall and Wilson, 2011; Villa et al., 2014).

It is well known that some nannoplankton species reflect a strong sensitivity to environmental conditions (Okada and Honjo, 1973; Winter et al., 1994; Gibbs et al., 2006; Kalb and Bralower, 2012; Villa et al., 2014). In order to evaluate the response of the nannofossil assemblage around the EOT at the studied equatorial latitudes, the abundance patterns of selected taxa have been plotted as  $\text{N}/\text{mm}^2$  vs. age (Fig. 4), and their variations are assumed to represent a paleoecological

response to paleoclimatic and/or trophic changes. The environmental preference of selected taxa show clear variation in this interval:

- *Discoaster* has long been assumed to be a warm-water taxon, declining with high nutrient availability (Gibbs et al., 2004), the same preference for oligotrophic conditions are suggested for *Cribocentrum reticulatum* (Villa et al., 2008 and reference herein). At Site 711, both *Discoaster* and *C. reticulatum* show a clear decrease just before the EOT at 34.5 Ma.

- Sphenoliths are widespread in low latitudes and are considered to belong to warm-water taxa, more debated is their behaviour with regard to trophic conditions. They have been thought to be adapted to oligotrophic stable environments (Aubry, 1998; Bralower, 2002; Gibbs et al., 2004; Kalb and Bralower, 2012). Nevertheless they are also considered adaptable to mesotrophic/eutrophic conditions (Wade and Pälike, 2004; Wade and Bown, 2006; Dunkley Jones et al., 2008; Pälike et al., 2010). This genus, and in particular *Sphenolithus predistentus*, shows an increase in abundance just before the EOT, at the same level as the *Discoaster* decrease, with a noticeable similarity with the trend identified in Tanzania by Dunkley Jones et al. (2008).

- *Cyclicargolithus floridanus* is considered indicative of high productivity environments (Aubry, 1992; Monechi et al., 2000; Dunkley Jones et al., 2008), and at Site 711 it shows an increase before the beginning of the EOT.

- *Reticulofenestra daviesii* is considered a cool-water taxon (Wei et al., 1992; Persico and Villa, 2004; Villa and Persico, 2006) with preference for eutrophic waters (Villa et al., 2014). It increases significantly in the SO at the Oi-1. This



species is present in the studied section with low percentages, and increases considerably in abundance before the beginning of the EOT.

- The genus *Helicosphaera* has been previously related to nutrient-rich waters (DeKaenel and Villa, 1996). Studies on living coccolithophorids confirm the relationship of helicosphaerids with high primary productivity rates (Haidar and Thierstein, 2001; Toledo et al., 2007), preferring near-shore/eutrophic or mesotrophic environments (Ziveri et al., 2004; Guerreiro et al., 2005). At Site 711, although this genus is present with very low abundances (in terms of percentages, it represents less than one percent of the total assemblage), we note an increase starting from ~34.5 Ma. Nevertheless, helicosphaerids are preservation sensitive taxa and their increase could be related to the CCD deepening, which is a feature of many EOT sections (e.g., Pälike et al., 2010). At Site 711 an increase in the absolute abundance of calcareous nannofossils, coupled with an increase in MAR  $\text{CaCO}_3$ , suggests a deepening of the CCD also in the equatorial Indian Ocean. The general trend of the described taxa shows a decrease in oligotrophic taxa (*Discoaster* and *C. reticulatum*), and a concurrent increase in eutrophic taxa (*R. daviesii*, *C. floridanus*, *S. predistentus* and *Helicosphaera* spp.) at about 34.5- 34.1, confirming an early response of the nanoplankton assemblage to paleoceanographic changes predating the EOT (~34-33.5 Ma) (Dunkley Jones et al., 2008). This assemblage variation could be linked to the formation at high latitudes of nutrient-rich waters: in particular the Subantarctic Mode Water (SAMW) is a water mass created in the SO by deep mixing, that sinks below the ocean surface and moves northward, reaching most of the upper ocean. SAMW is responsible for

nutrient transport, from the SO to the low-latitude regions (Sarmiento et al., 2004). The enhanced mixing in the SO connected to the late Eocene Antarctic cooling could have strengthened the SAMW that in turn could have influenced the productivity of surface waters in the western tropical Indian Ocean (Kiefer et al., 2006) as proposed for the increased primary productivity recorded in Tanzania through the EOT (Dunkley Jones et al., 2008). However, we cannot exclude a link between the recorded nannofossil assemblage change and the hypothesized “Late Eocene event” at about 34.1 Ma, representing an Antarctic ice-volume increase that predates the EOT (Katz et al., 2008; Coxall and Wilson, 2011).

## 5. Conclusions

Quantitative analyses on 109 samples from Hole 711A, partly correlated with magnetostratigraphy, encompassing a time interval from middle Eocene to early Oligocene, enabled us to discuss 31 biohorizons across a time interval of about 11 Myr, with an average biostratigraphic resolution of 0.42 kyr, and to verify their reliability and traceability. A summary on the reliability of the discussed biohorizons is reported in Tab. 1. Our biostratigraphic results have shown that some of the traditional markers proposed by Okada and Bukry (1980) and Martini (1971) are absent or very rare in equatorial environments (e.g., *I. recurvus*, *C. solitus*), and in contrast, additional bioevents may increase the biostratigraphic resolution in this time interval. For example the numerous *Sphenolithus* species provide potential biohorizons for the middle to late Eocene time interval. The discussed bioevents have been compared with previous studies from low and high

latitudes in different oceanic basins, in order to define their reliability on a regional scale or over a wide area. One of the most relevant results concerns the Base common of *Cribozentrum erbae*. This species, recently defined in the Tethyan realm, has been proposed as the biohorizon that best defines the base of the Priabonian (Agnini et al., 2011). Our results confirm the reproducibility of this event also in equatorial Indian Ocean paleolatitudes. The strict link between calcareous nannofossil preservation and carbonate content in the deep-water mass (e.g., Toffanin et al., 2013) permits an interpretation of the CCD fluctuation in the Equatorial Indian Ocean at this time. A middle Eocene interval, about 3 m thick, with no carbonate content, almost barren in nannofossils, may be interpreted as a CCD shoaling. This interval partially overlaps the MECO event, the peak of which is globally recognized as coincident with a CCD shoaling of 500-1500 meters (Bohaty et al., 2009).

Also, we document a shift in the nannofossil assemblage, starting before the inception of the EOT, at about 34.5-34.1 Ma, evidenced by an increase in eutrophic species and a decrease in oligotrophic/warm taxa. This change highlights a response of nannofossils to the paleoceanographic and climatic change, interpreted as a shift toward increased nutrient availability at the surface waters of the equatorial Indian Ocean, likely driven by the equatorial upwelling of the SAMW. Future works on low latitude sediments could elucidate the response of calcareous nannofossils to the climate changes that predate the EOT in order to investigate the different response of low latitude and high latitude calcareous nannofossil

assemblages, the latter having been shown to respond abruptly to the EOT (Villa et al., 2014).

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## Figure captions

FIG. 1 – Location map showing the position of the studied site.

FIG. 2 - Calcareous nannofossil biostratigraphic summary of the middle Eocene-lower Oligocene study section, plotted against depth, variations in the declination and inclination and magnetostratigraphic interpretation (Savian et al. 2013). Simplified lithology is from Backman and Duncan (1988). Biostratigraphic events are indicated with arrows as B (Base), Bc (Base common), T (Top), Tc (Top common). \*According to Perch-Nielsen (1985).

FIG. 3 - Calcareous nannofossil total abundance and abundance patterns of selected nannofossil species in Hole 711A, expressed as Number of specimens per  $\text{mm}^2$ , are plotted against depth (mbsf),  $\text{CaCO}_3$  MAR, magnetostratigraphy (Savian et al. 2013), and chronostratigraphy. *Cribrrocentrum erbae* and *Clausicoccus* spp. are expressed also as percentage. The light yellow bar indicates the interval with strong nannofossil dissolution.

FIG. 4 -Abundance patterns of selected calcareous nannofossil taxa plotted against age. Data of carbonate MAR and carbonate percentage are from Peterson and Backman (1980). The blue line indicates the Eocene/Oligocene boundary. The light yellow bar indicates the interval of calcareous nannofossil dissolution of the middle Eocene. Light blue bar indicates the assemblage variation interval between ~ 34.5 and 34.1 Ma. Arrows indicate the abundance trend from this interval.

Magnetostratigraphy from Savian et al. (2013) correlated with the GPTS of Gee and Kent (2007).

TAB. 1 - Calcareous nannofossil biohorizons identified in Hole 711A. The age of the bioevents is defined using available magnetostratigraphic records (Savian et al., 2013) by applying a linear interpolation between nearest reversal boundaries.



Calculated ages are also compared with recent published data. B=Base; T= Top; Bc= Base common; Tc = Top common. \*datum from Okada, 1990. A summary of the reliability of the considered biohorizons is reported in the “reliability and biostratigraphic remarks” column.

Plate 1: Photomicrographs of selected calcareous nannofossils from Site 711. Figs 1-17, 19-21: Light microscope, cross-polarized light, 100X magnification; Fig. 18: Light microscope, plane light, 100X magnification; Figs 22-24: Scanning Electron Microscope, scale bar on Figs. 1 – *C. reticulatum*, Sample 160.21 mbsf, 2 – *C. reticulatum*, Sample 179.42 mbsf; 3 – *C. erbae*, Sample 176.89 mbsf; 4 – *C. erbae*; 5 - *C. reticulatum*, Sample 180.42 mbsf; 6-8 – *C. grandis*, Sample 198,29 mbsf; 9 – *E. formosa*, Sample 191,71; 10 – *C. eopelagicus*, Sample 145.2 mbsf; 11 – *D. stavensis*, Sample 153.39 mbsf; 12 – *D. bisectus*, Sample 191.71 mbsf; 13,14 – *R. samodurovii*, Sample 198.29 mbsf; 15 – *R. umbilicus*, Sample 198.29 mbsf; 16 – *R. umbilicus*, Sample 158.21 mbsf; 17 – *H. compacta*, Sample 151.39 mbsf; 18 – *D. barbadiensis*, Sample 196.17 mbsf; 19,20 – *S. kempii*, Sample 233.74 mbsf; 21-24 – *S. cf. S. kempii*, Sample 233.29 mbsf.

Plate 2: photomicrographs of selected calcareous nannofossils from Site 711. All samples taken in cross-polarized light, 100X magnification 1, 2 – *S. strigosus*, Sample 202.98 mbsf; 3, 4 – *S. obtusus* Sample 182.11 mbsf; 5, 6 - *S. obtusus* Sample 183.91 mbsf; 7 – *S. furcatolithoides* Sample 198.29 mbsf; 8-10 - *S. furcatolithoides* Sample 196.17 mbsf; 11,12 – *S. radians* Sample 202.89 mbsf; 13,14 – *S. spiniger*, Sample 202.89 mbsf; 15 – *S. intercalaris*, Sample 179.48 mbsf; 16 – *S. intercalaris*, Sample 191.71 mbsf; 17,18,21 – *S. predistentus*, Sample 128.83 mbsf; 19 – *S. predistentus*, Sample 145.2 mbsf; 20 – *S. predistentus*, Sample 151.39 mbsf; 22 – *S. pseudoradians*, Sample 128.83 mbsf; 23,24 – *S. akropodus*, Sample 145.2 mbsf.

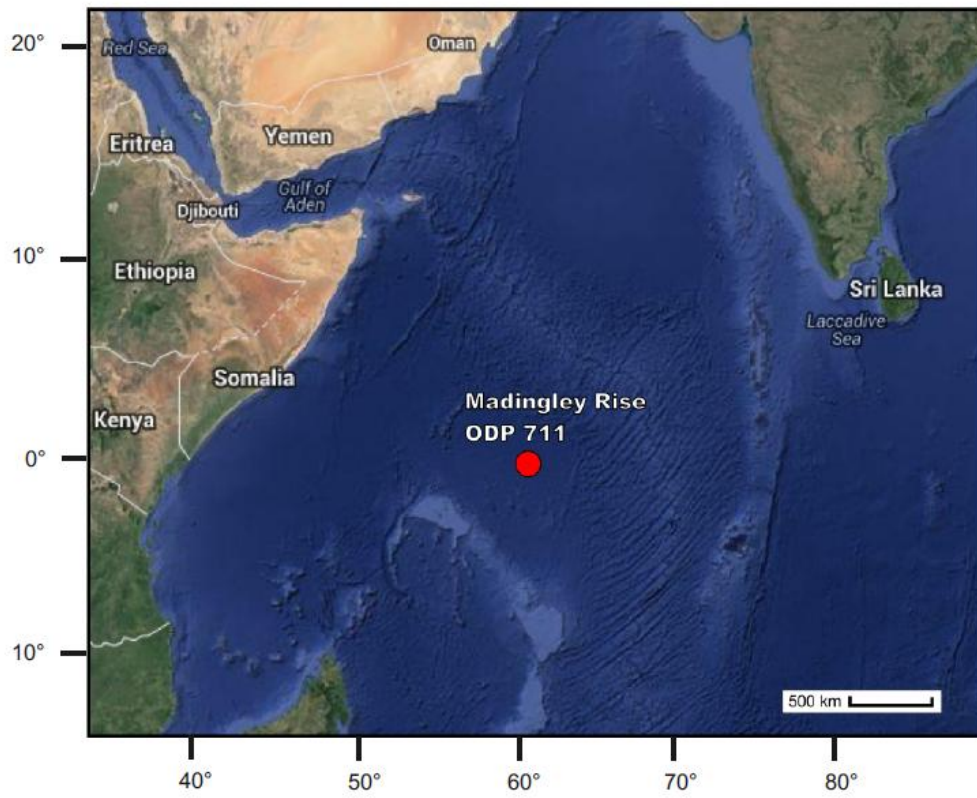


FIG. 1

Fig. 1

ACCEPTED

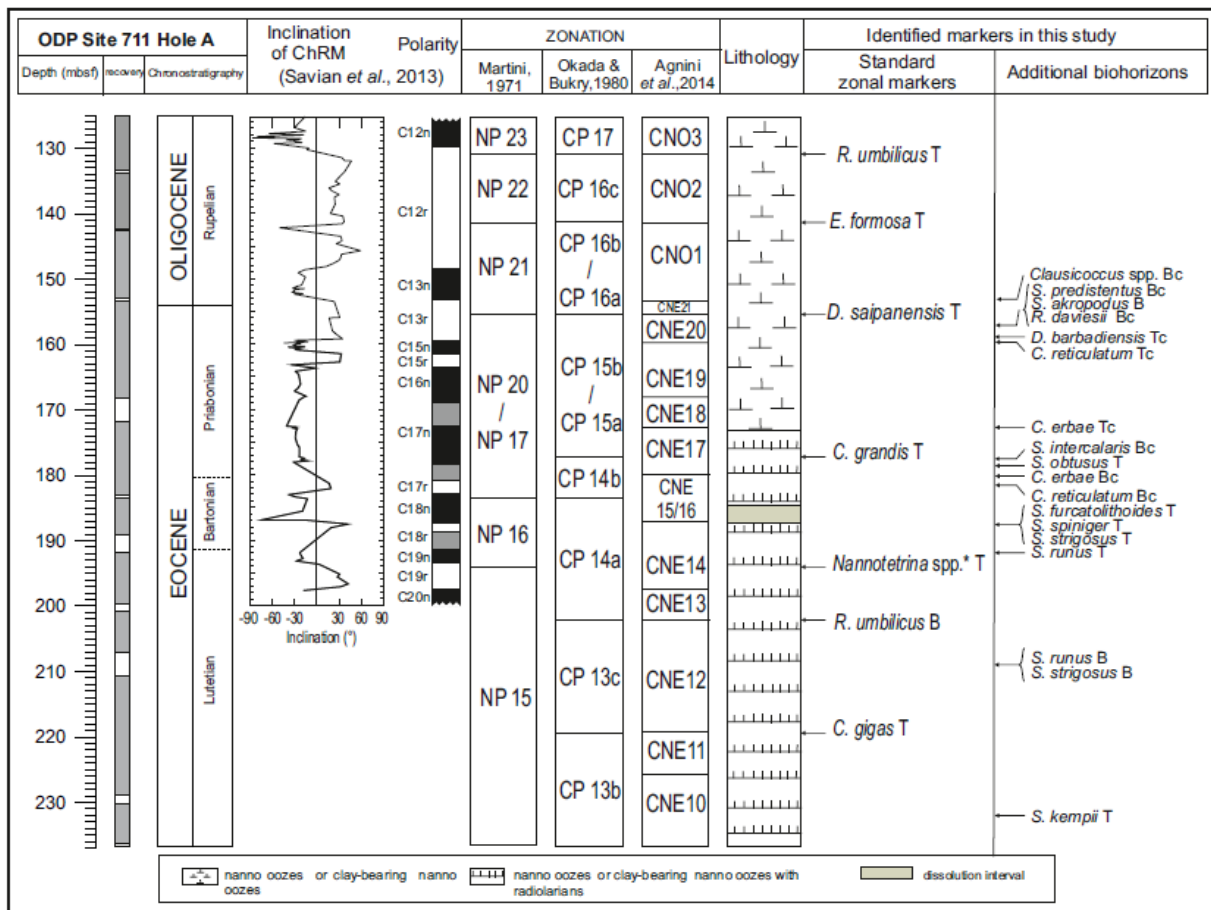


Fig. 2

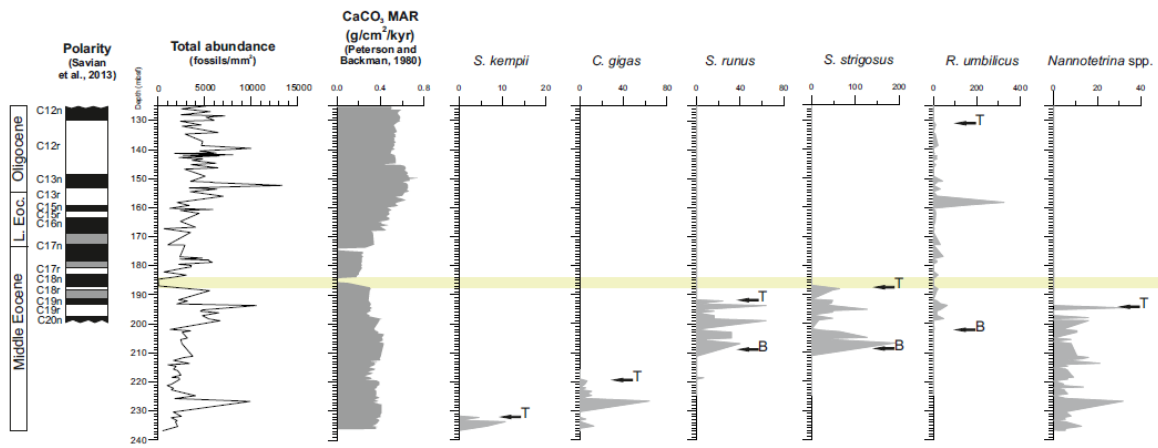


FIG. 3a

Fig. 3

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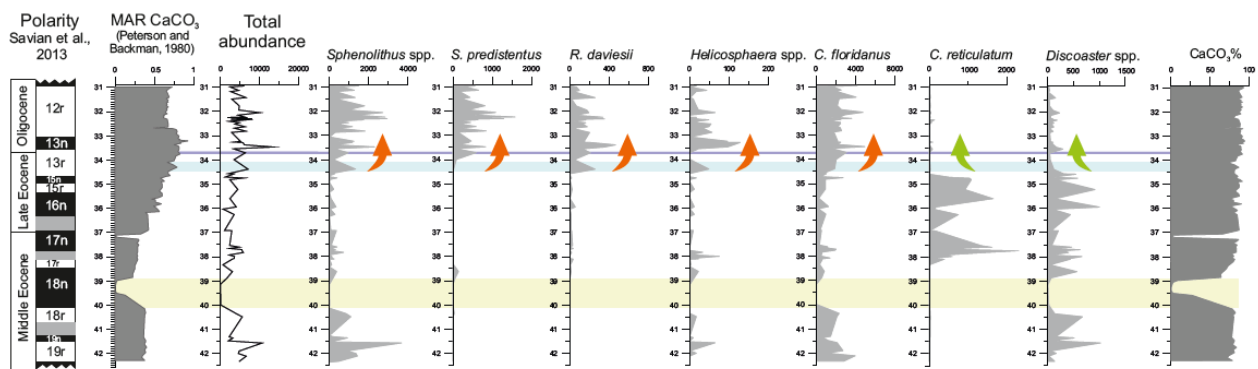


FIG. 4

Fig. 4

ACCEPTED MANUSCRIPT

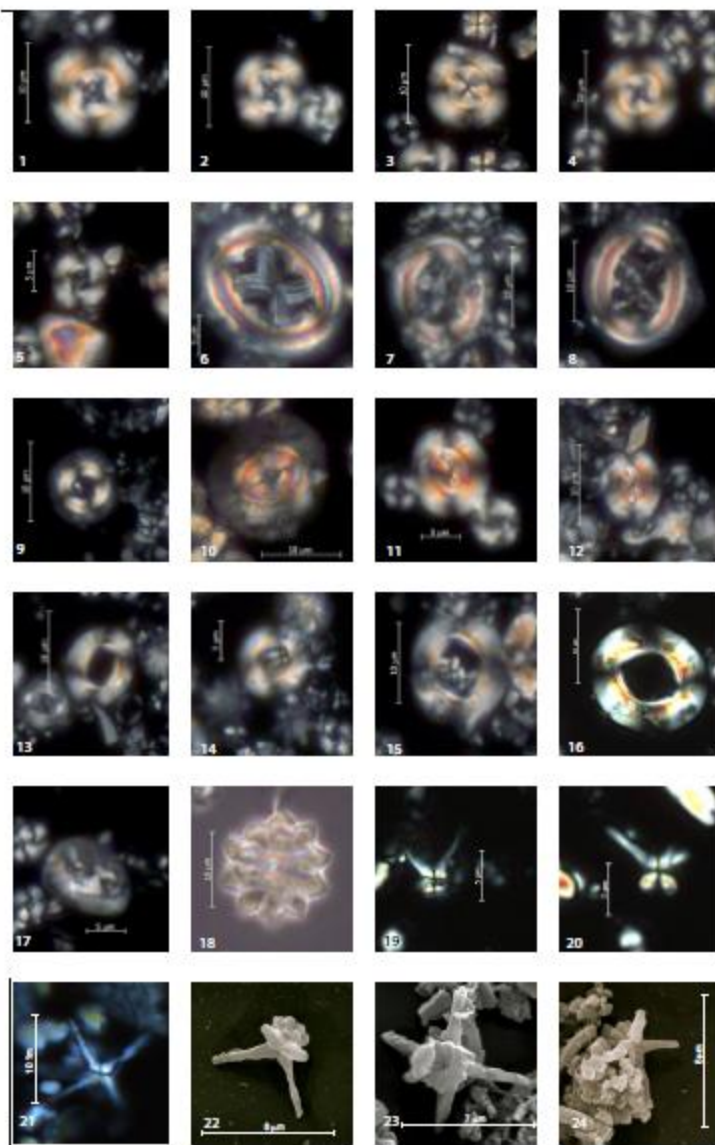


Plate 1

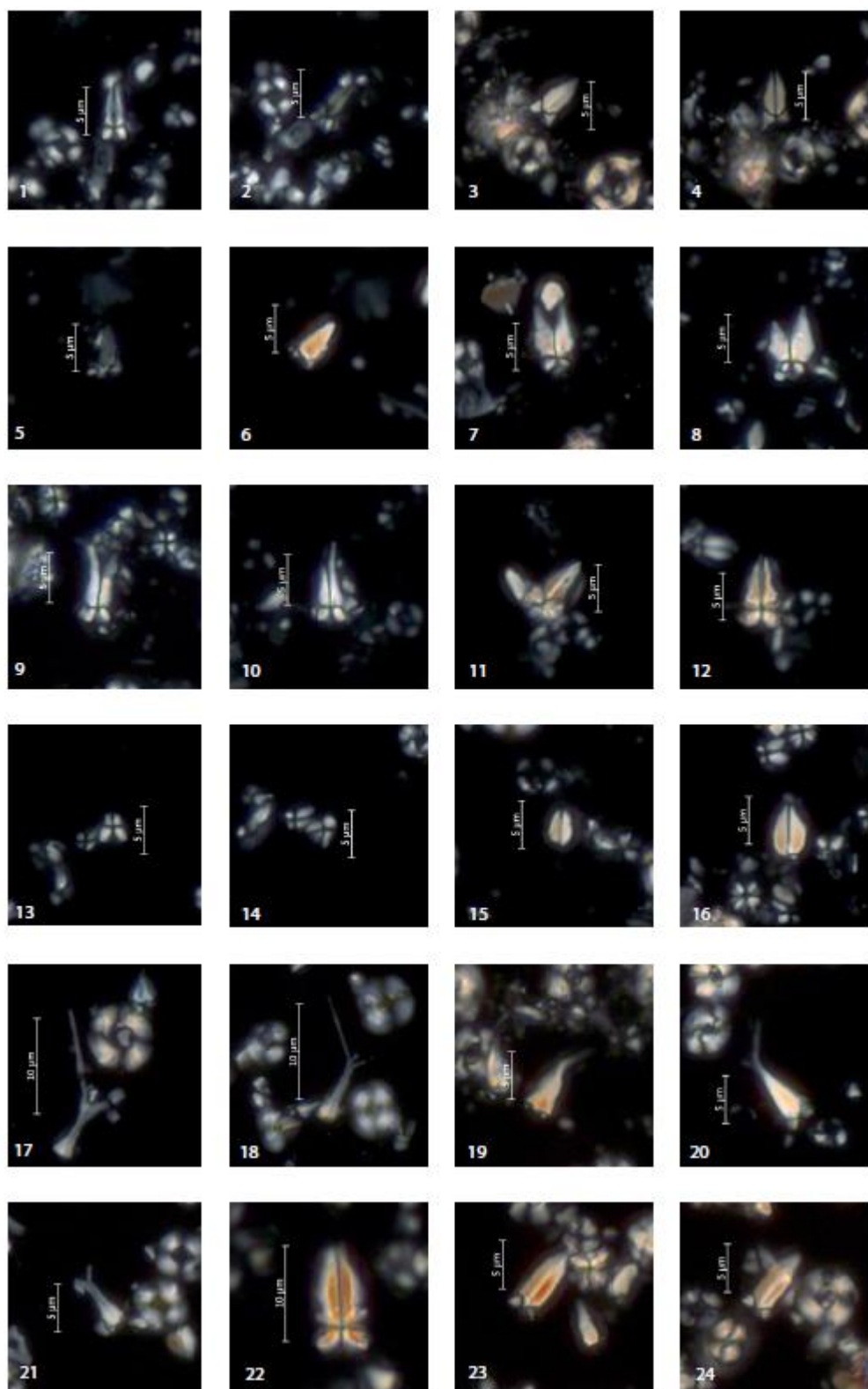


Plate 2

Table 1

Bioevents	Depth				Age (Ma)						Reliability and biostratigraphic remarks	
	Top depth (mbsf)	Bottom depth (mbsf)	Depth error (m)	mean depth (mbsf)	This study	Sarvan et al. (2013)	Jovane et al. (2010)	Toffin et al. (2013)	Formicari et al. (2010) Site 1052	Alano		Agimi et al. (2014)
T <i>Reticulofenestra umbilicus</i>	130.22	131.51	1.29	130.86	31.059	32.3					32.02	Good
T <i>Elkonia formosa</i>	141.21	141.31	0.10	141.26	32.24	32.8			33.13		32.92	Good
B <i>Clauacoccus</i> spp.	152.31	153.20	0.89	152.75	33.509				33.91		33.88 (C. subobtusus)	Enough good, slightly diachronous
T <i>Discosaster saepepinnatis</i>	154.49	155.04	1.55	155.26	34.187						34.44	Good
B <i>Reticulofenestra daviesii</i>	156.04	158.21	2.17	157.12	34.476							Good
B <i>Sphenolithus elongatus</i>	156.04	158.21	2.17	157.12	34.476							Potentially good, needs further investigation
B <i>Sphenolithus predilectus</i>	156.04	158.21	2.17	157.12	34.476							Good
T <i>Discosaster barbadensis</i>	158.21	159.21	1.0	158.71	34.678						34.77 (T)	Good
To <i>Cribrocentrum reticulatum</i>	159.21	159.21	1.0	159.21	34.647						35.24 (T)	Good at mid-low latitudes
B <i>Isthmolithus recurvus</i> *	163.12	164.62	1.5	163.87	35.454	36		36.08			36.84	Too rare
To <i>Cribrocentrum erbae</i>	172.79	172.86	0.07	172.82	36.929						37.46	Good
T <i>Chiasmolithus granulosus</i>	176.69	177.34	0.65	177.11	37.57	37.1		36.61	37.71	37.72	37.77	Unreliable
B <i>Sphenolithus intercalaris</i>	177.34	177.71	0.37	177.52	37.632			36.08	37.42	37.45	37.77	Potentially good, needs further investigation
T <i>Sphenolithus obtusus</i>	178.31	178.81	0.5	178.56	37.788			36.96	36.45	36.25	36.47	Slightly diachronous
B <i>Cribrocentrum erbae</i>	179.89	180.42	0.53	180.15	38.03				37.82	37.83	37.88	Good
B <i>Cribrocentrum reticulatum</i>	180.61	182.11	1.5	181.36	38.215						42.37	Good at mid-low latitudes
T <i>Chiasmolithus solitus</i>	183.21	183.91	0.70	183.56	38.744	40.4		38.79	38.40	38.49		Too rare, diachronous
B <i>Dicryococcolites obliquus</i>	183.91	184.63	0.72	184.27	39.007				40.42			Time transgressive
B <i>Sphenolithus obtusus</i>	183.91	184.63	0.72	184.27	39.007				40.13	39.63		Probably good but here masked by a dissolution event
T <i>Sphenolithus spiniger</i>	186.92	188.11	1.19	187.51	40.174				40.10	39.63		Probably good but here masked by a dissolution event
T <i>Sphenolithus fuscescens</i>	186.92	188.11	1.19	187.51	40.174		40.51	40.61	40.38	40.76	40.51	Probably good but here masked by a dissolution event
T <i>Sphenolithus striatus</i>	186.92	188.11	1.19	187.51	40.174						40.78	Too rare, diachronous
B <i>Sphenolithus predilectus</i>	191.71	192.13	0.4	191.92	41.335				40.35			Too scattered and rare
B <i>Cribrocentrum reticulatum</i>	191.71	192.13	0.4	191.92	41.335					41.119 (Bc)		Potentially good, needs further investigations
T <i>Sphenolithus rufus</i>	191.71	192.13	0.4	191.92	41.335							Good but rare
T <i>Nannoflorina</i> sp.	193.70	194.50	0.8	194.10	41.739							Good
B <i>Reticulofenestra umbilicus</i>	201.80	202.40	0.6	202.10	/	42.67		42.35			43.06 (Bc)	Potentially good, needs further investigations
B <i>Sphenolithus striatus</i>	206.82	211.15	4.33	208.98	/			41.83				Potentially good, needs further investigations
B <i>Sphenolithus rufus</i>	206.82	211.15	4.33	208.98	/							Potentially good, needs further investigations
T <i>Chiasmolithus gladius</i>	219.10	219.72	0.62	219.41	/		44.79					Good but rare
T <i>Sphenolithus kempfi</i>	231.81	232.30	0.49	232.05	/							Potentially good, needs further investigations

## Highlights

Revision of the Nannofossil biostratigraphy of the middle Eocene- lower Oligocene at Site 711.

Discussion on the reliability of bioevents for the late Paleogene at low latitudes.

Magnetobiochronology of Site 711.

Evaluation of the Nannofossils response during the MECO and the EOT.