Major shifts in calcareous phytoplankton assemblages through the Eocene-Oligocene transition of Tanzania and their implications for low-latitude primary production

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[1] A high-resolution record of exceptionally well preserved calcareous nannofossil assemblages from Tanzania is marked by two key transitions closely related to the climatic events of the Eocene-Oligocene transition (EOT). The first transition, at ~34.0 Ma, precedes the first positive shift in δ18O and coincides with a distinct interval of very low nannofossil abundance and a cooling in sea surface temperatures (SST). The second, at ~33.63 Ma, is immediately above the Eocene-Oligocene boundary (EOB) and is associated with a significant drop in nannofossil diversity. Both transitions involve significant reductions in the abundance of holococcoliths and other oligotrophic taxa. These changes in calcareous phytoplankton assemblages indicate (1) a widespread and significant perturbation to the low-latitude surface ocean closely tied to the EOB, (2) a potential role for reduced carbonate primary production at the onset of global cooling, and (3) a significant increase in nutrient availability in the low-latitude surface ocean through the EOT.


1. Introduction

[2] The transition between the Eocene and Oligocene epochs is the central point of the most significant climatic shift of the Cenozoic, as the Earth’s climate developed from the ice-free early Paleogene world of global warmth and high atmospheric pCO2 to the glaciated conditions of the early Oligocene with the rapid expansion of continental-scale ice sheets on Antarctica [Lear et al., 2000; Miller et al., 1991; Pagani et al., 2005; Zachos et al., 2001]. Eocene-Oligocene marine sedimentary records are characterized by large, ~1.0 to 1.5‰, positive excursions in benthic foraminiferal oxygen and carbon stable isotopes and a rapid deepening of the calcite compensation depth (CCD) [Coxall et al., 2005; Zachos et al., 1996]. Coincident with these changes in the deep sea, significant glaciomarine sedimentation commences around the margins of both East and West Antarctica indicating the first Cenozoic advance of continental-scale ice sheets in southern high latitudes [Brea and Wise, 1992; Ivany et al., 2006; Wise et al., 1991, 1992; Zachos et al., 1992]. Estimates of the increase in global ice volume through this transition, based on records of benthic foraminiferal δ18O, Mg/Ca ratios and sequence stratigraphy, are similar to or greater than the volume of the present-day Antarctic ice sheet [Billups and Schrag, 2003; Coxall et al., 2005; Katz et al., 2008; Lear et al., 2004, 2008; Miller et al., 2008; Pekar et al., 2002].

[3] Following Pearson et al. [2008] we use “Eocene–Oligocene transition” (EOT) to denote the extended interval of climatic and biotic change lasting about 500 ka between 33.5 and 34.0 Ma, which spans both the E/O boundary (EOB) at 33.7 Ma and the positive δ18O shift. Recent high-resolution records of the EOT have identified two distinct ~40 ka steps within the δ18O shift, termed here step 1 and step 2, which are separated by a ~200 ka plateau [Coxall et al., 2005]. Step 1 and the plateau occur in the uppermost part of magnetostratigraph C13r, while the maximum positive δ18O values at the end of step 2 directly correlate with the base of chron C13n and mark the end of the EOT and the start of the ~400 ka Early Oligocene Glacial Maximum (EOGM) [Coxall and Pearson, 2007; Coxall et al., 2005; Liu et al., 2004; Zachos et al., 1996]. The major ocean-wide increase in δ13C, which appears to lag the shift in δ18O by ~10 ka [Coxall et al., 2005], is compelling evidence for a significant perturbation in the global carbon cycle through the EOT [Merico et al., 2008; Salamy and Zachos, 1999; Zachos and Kump, 2005; Zachos et al., 1996]. This positive shift in δ13C is indicative of either an increase in the removal rate of light carbon from the ocean-atmosphere system, which has been attributed to the enhanced burial of organic carbon in marine sediments coupled to increased surface water productivity [Diester-Haass and Zachos, 2003; Salamy and Zachos, 1999; Zachos and Kump, 2005] or the input of isotopically heavy carbon via the increased weathering of neritic carbonates during a major
sea level fall [Merico et al., 2008]. Both of these hypotheses are associated with a drawdown of atmospheric carbon dioxide that may be the principal positive feedback mechanism that drove the climate system into a strongly glaciated state during the EOGM [Merico et al., 2008; Zachos and Kump, 2005].

[4] Evidence for significant increases in primary production through the EOT has been documented at a number of deep-sea sites in the Southern Ocean [Anderson and Delaney, 2005; Diester-Haass et al., 1996; Latimer and Filippelli, 2002; Salamy and Zachos, 1999], probably driven by cooling induced increases in surface-ocean divergence and the associated upwelling of nutrient-rich deep waters and/or enhanced fluxes of weathering derived nutrients during rapid sea level fall [Nilsen et al., 2003; Salamy and Zachos, 1999; Zachos and Kump, 2005]. There is however a lack of comparable paleoproductivity records from the low latitudes [but see Diester-Haass and Zachos, 2003; Nilsen et al., 2003; Ravizza and Paquay, 2008] where quantifying changes in primary carbonate and silica fluxes at many deep-sea sites is hampered by the dramatic deepening of the carbonate compensation depth (CCD) and the associated shift from siliceous to calcareous sediments through the EOT [Nilsen et al., 2003].

[5] Studies of modern coccolithophores (calcareous scale-producing haptophyte algae) have shown that the species makeup and community structure of assemblages are highly sensitive to surface water stratification, nutrient availability and nutricline depth. In the modern oceans there is a general correlation between high-productivity environments and low-diversity coccolithophore assemblages, dominated by bloom-forming species such as Emiliania huxleyi, while oligotrophic environments are associated with diverse assemblages, often with a clear depth zonation and a distinct lower photic zone (LPZ) flora [Broerse et al., 2000; Kinkel et al., 2000; McIntyre and Bé, 1967; Okada and Honjo, 1973]. The response of the calcareous phytoplankton to global change through the Eocene-Oligocene boundary is poorly known when compared to other critical intervals such as the Paleocene-Eocene boundary [e.g., Bradner, 2002; Gibbs et al., 2006a, 2006b; Raffi et al., 2005]. One set of high-resolution records from the Southern Ocean has shown a significant fall in nannofossil diversity and an increase in cold water taxa directly coincident with the onset of the EOGM [Persico and Villa, 2004] but there is no comparable data set from the low latitudes, with previous quantitative records being either at low resolution (~100 ka) [Wei et al., 1992; Wei and Wise, 1990] or severely compromised by poor preservation [Coccioni et al., 1988; Monchi, 1986].

[6] The recent recovery of a latest Eocene to earliest Oligocene sequence of hemipelagic clays from coastal Tanzania [Nicholas et al., 2006], which host exceptionally well preserved calcareous microfossils [Bown et al., 2008a; Pearson et al., 2008], provides a unique opportunity to examine the coccolithophore response through this critical climatic transition in a low-latitude setting. In this study we present a high-resolution record of coccolithophore assemblages through this succession and discuss their implications for changes in surface water productivity linked to major climate change. The deposition of these sediments in an outer shelf to upper slope environment, well above the paleolysocline, means they were unaffected by the dramatic variations in CCD depth through the EOT. Samples taken across the EOT consistently yield a highly diverse and exceptionally well preserved calcareous nannofossil assemblage (Figure 1) [Bown et al., 2008a; Dunkley Jones et al., 2008] and can be correlated directly to the Eocene-Oligocene boundary by planktonic foraminifera biostratigraphy [Wade and Pearson, 2008] and the onset of the EOGM by stable isotope stratigraphy [Pearson et al., 2008].

2. Materials and Methods

[7] This study is part of an ongoing paleoclimate research project, the Tanzanian Drilling Project (TDP), which is focused on recovering geochemical and paleobiological data using exceptionally well preserved calcareous microfossils from the Cretaceous and Paleogene sediments of the Kilwa Group, coastal Tanzania [Pearson et al., 2001, 2007, 2008]. The two cores examined in this study, TDP12 (9° 12' 54.74" S 39° 32' 53.60" E; total depth 147.44 m) and TDP17 (9° 11' 11.38" S 39° 33' 3.82" E; total depth 125.9 m) overlap to provide a continuous stratigraphic sequence across the EOT; both sites were located on the Pande Peninsula, approximately 3 km apart, in the Kilwa Administrative District, Tanzania (Figure 2). The cores are composed of a succession of dark greenish-gray clays with occasional limestone interbeds, typical of the middle Eocene to lower Oligocene Pande Formation [Nicholas et al., 2006], and span the EOB, nannofossil zones NP19/20 and 21 [Martini, 1971] and planktonic foraminifera zones E15/E16 and O1 [Berggren and Pearson, 2005].

[8] The in situ benthic foraminiferal assemblage of the Pande clays, recovered in TDP12 and TDP17, consists of “flysch”-style agglutinated forms, dominated by Bathysiphon, and a deep water calcareous fauna typical of outer shelf to upper slope environments (300–500 m water depth) [Nicholas et al., 2006]. Abundant allochthonous microscopic shell debris, transported from the innermost shelf, is present throughout the succession and often dominates benthic foraminiferal assemblages. The consistent presence of the specialist lower photic zone calcareous phytoplankton genera Gladiolithus and Algirosphaera (Figures 1i and 1j) within the studied sections, constrains the setting to a relatively deep, clear water environment [Jordan and Chamberlain, 1997; Takahashi and Okada, 2000].

[9] The TDP12 and TDP17 successions were correlated using planktonic (last occurrences of the Hantkeninidae and Turborotalia cerroazulensis group) and benthic foraminiferal (late Eocene last occurrence of a characteristic but undescribed Migros sp.) and calcareous nannofossil (last occurrence of Penna papillatum) bioevents. The age model for the two sites is based upon bioevents using the age assignments of Berggren et al. [1995] with additional control provided by tuning the composite TDP12 and TDP17 planktonic foraminifera (Turborotalia ampliapertura) oxygen isotope (δ18O) record to the deep sea benthic foraminiferal (Cibicidoides) δ18O record from
Figure 1. Scanning electron microscope images of selected calcareous nannofossils, all scale bars are 1 μm: (a) *Pontosphaera versa* TDP12/23–2, 79 cm; (b) *Orthozygus* sp. (holococcolith) TDP12/42–3, 86 cm; (c) *Varolha cistula* (holococcolith) TDP12/47–2, 64 cm; (d) *Cyclicargolithus floridanus* part-coccosphere TDP12/26–2, 62 cm; (e) *Calciosolenia* sp. TDP12/26–2, 62 cm; (f) *Rhabdosphaera* sp. TDP12/47–2, 64 cm; (g) small spinose murolith TDP12/26–2, 62 cm; (h) *Syracosphaera* sp. and *Blackites* sp. TDP12/26–2, 62 cm; (i) *Gladiolithus flabellatus* part coccosphere TDP12/26–2, 62 cm; and (j) *Algiriosphaera* sp. TDP12/26–2, 62 cm.
equatorial Pacific ODP Site 1218 [see Pearson et al., 2008]. All depths are in meters composite depth (mcd) unless otherwise stated. The last occurrence of Discoaster saipanensis (NP19-20/21 nannofossil zone boundary) at 130.35 mcd appears to be slightly high in the succession when compared to ages based on the correlation of stable isotope curves between these sites and ODP Site 1218. There are difficulties in placing this event as D. saipanensis is present at very low abundances in this material and may be subject to reworking, as a result this datum was not used within the age model. The base of planktonic foraminifera zones E15 and E16 are undifferentiated because of the absence of Globigerinatheca index in these sections as discussed by Wade and Pearson [2008].

Sediment samples, approximately 2 cm³ in volume, were extracted at ~1 m intervals and thoroughly scraped clean. Simple smear slides were made, taking care to maintain a consistent density of diluted sediment, using a small quantity of sediment and distilled/deionized water. This approach is the most suitable for the analysis of these diverse assemblages, which include a significant number of fragile taxa, as it minimizes the amount of sample processing and the associated risks of nannofossil fragmentation and dissolution. Duplicate smear slides from TDP12 indicate that transport and storage of samples did not have a deleterious effect on nannofossil preservation and that diversity measures are reproducible [Dunkley Jones and Bow, 2007]. All of the clay samples appeared homogeneous on the millimeter scale with no sign of laminations.

A total of 107 slides were examined for calcareous nannofossils from TDP12 (range in original depth scale 147.8 to 18.69 m below ground level (mbgl)), and 65 slides from TDP17 (range in original depth scale 101.11 to 31.15 mbgl), using standard light microscope techniques at a magnification of 1250x on an Olympus BX40 microscope. Up to 20 consecutive fields of view (FOV) were initially examined for each slide, with all whole identifiable nannofossil specimens counted. Specimens were identified to species level wherever possible, following the taxonomy of Dunkley Jones et al. [2008]. During this count, the number of whole reworked Cretaceous nannofossils and the number of cubic pyrite grains were counted per FOV. Counts were terminated when at least 400 complete in situ identifiable nannofossils had been counted; where necessary, counts were extended beyond 20 FOV, along continuous transects, until at least 400 specimens had been counted. Occasional samples from both boreholes contained very low nannofossil abundances (~4 or less per FOV); these were examined and counted as normal for 20 FOV to obtain nannofossil abundance, Cretaceous reworking and mineralogical counts but were then abandoned. Samples and slides are curated at the Department of Earth Sciences, University College London. The surfaces of broken rock chips were examined using a JEOL Digital JSM-6480LV scanning electron microscope, following the methodology of Lees et al. [2004]. A large number of new and novel taxa were encountered during this work (Figure 1) and these are described in a separate taxonomic study [Dunkley Jones et al., 2008].

Metric Multidimensional Scaling (MMDS) was performed on a reduced set of relative abundance data, for a total of 29 species, with the exclusion of species whose standard deviation of their relative abundances was less than 0.5%, followed by the rescaling of the data to sum to 100%. This eliminates rarely occurring species, with relative abundances typically less than 1%, and follows the standard procedure for the multivariate analysis of subcomposition data [Pawlowsky-Glahn and Egozcue, 2006]. Before analysis, zero data were replaced using the rounded zero replacement routine of the CoDaPack multivariate data analysis software [Thioulouse and Martin-Fernández, 2006] and then additive-log-ratio (ALR) transformed, with the most dominant species, the small form of Cyclicargolithus floridanus (<5 μm), used as the denominator. This allows the use of standard multivariate methods on percentage compositional data, although the denominator variable, Cyclicargolithus floridanus (<5 μm), is not present in the transformed data set [Pawlowsky-Glahn and Egozcue, 2006]. The transformed data were standardized before running the MMDS, which used the Euclidean distance measure to obtain a dissimilarity matrix between the multidimensional data and squared stress as the goodness-of-fit criterion between the multidimensional and two-dimensional distances. This method produces a two-dimensional representation of the samples that seeks to preserve and represent the distances between samples in the multidimensional space defined by the variables analyzed, in this...
The planktonic foraminiferal (Calcidiscus and the large form of Reticulofenestra does show a slight abundance decrease/C24 Varolia macleodii Reticulofenestra minuta, which all have low but consistent abundances in the late Eocene, whereas the relative abundance of Lanternithus minutus peaks to an abundance of 14.9% in the sample immediately below the EOB. Other species that show similar trends include Reticulofenestra dicytoda and two holococcolith taxa, Varolia macleodii and Zygrhablithus bijugatus, which all have low but consistent relative abundances in the Eocene, typically 2–5%, but decline to rare occurrences in the Oligocene, and Pemmatapillatum, which has a distinct last occurrence at 100 ka before the EOB at ~33.8 Ma. The mean abundances of total Discoaster spp. decrease across the EOB, relative to their general low abundance, with an Eocene value of 1.1% and an Oligocene value of 0.2% (Fig. 4b). Holococcolith taxa make up a considerable proportion of the late Eocene nannofossil assemblages but there is a marked decline in total holococcolith abundance at both ~34.2 Ma and then more significantly at the EOB. Mean holococcolith relative abundances for the Eocene and Oligocene are 12.9% and 2.8% respectively.

The three species Reticulofenestra minuta, Coccoclycitus pelagicus and Sphenolithus moriformis all have consistent moderate to low abundances (~2 to 10%) throughout TDP12 and TDP17 (Fig. 5b), although Reticulofenestra minuta does show a slight abundance decline into the Oligocene. The incertae sedis nannolith Pemmatapillatum, although rare, is abundant throughout the section but new detailed SEM imaging of this taxon strongly suggests that this is a disaggregated pentalith of the family Braarudosphaerae (Dunkley Jones et al., 2008), in which case its abundance is inflated fivefold and if rescaled would be more comparable to other minor assemblage components at ~2–3%.

On the basis of broadly increasing or declining relative abundances, two species groups can be defined. Group 1 consists of species with declining abundances across the EOB: Calcidiscus protoanulus, Reticulofenestra dicytoda, Pemmatapillatum, Lanternithus minutus, Varolia macleodii and Zygrhablithus bijugatus. Group 2 consists of Sphenolithus predistentus and the large form of Cyclicargolithus floridanus, which both increase in abundance across the EOB. These plots of grouped relative abundances (Fig. 6b) more clearly illustrate the dominant trends and major changes in assemblage composition by minimizing small-scale fluctuations in individual species data. The total relative abundance plot for group 1 species shows the marked transition across the EOB, from high abundances in the Eocene (mean 25.6%) to low values in the Oligocene (mean 1.2%). Conversely group 2 species have a low grouped relative abundance in the late Eocene.
Figure 3. Geochemical and smear slide count data across the Eocene-Oligocene transition. (a) Stable isotopes ($\delta^{18}$O and $\delta^{13}$C) [Pearson et al., 2008] and Mg/Ca ratios [Lear et al., 2008] from planktonic foraminifera Turborotalia ampliapertura and calculated near-surface $\delta^{18}$O$_{SW}$ [Lear et al., 2008]. (b) Nannofossil abundance, SHE diversity (Shannon’s $H$, left-hand curve, and $\ln(S)$, right-hand curve), and abundance of cubic pyrite and reworked Cretaceous nannofossils. Data are from TDP Site 12 (base of section, purple) and TDP Site 17 (top of section, light blue). The horizontal gray bars indicate the duration of the globally recognized first and second positive steps in $\delta^{18}$O, the pink shading indicates the duration of the EOT (34.0–33.5 Ma), and the position of the Early Oligocene Glacial Maximum (EOGM) is marked on the plot of $\delta^{18}$O. Planktonic foraminifera biozones of Berggren and Pearson [2005] and calcareous nannofossil biozones of Martini [1971] are marked on the right-hand side; there is some uncertainty in the placement of the NP19/20–21 nannofossil zone boundary.
(mean 1.4% below 33.8 Ma) but this increases dramatically at the onset of step 1 (~33.8 Ma) and into the Oligocene (mean 15.2% above 33.8 Ma).

The first two coordinates of the Metric Multidimensional Scaling (MMDS1 and MMDS2) analysis represent the major components of variation within the multivariate (species relative abundance) space, in a fashion similar to the first two components of a principal component analysis (Figure 6a). Two principal clusters of samples are clearly resolved: Eocene samples older than ~34.0 Ma ("Eocene") with generally negative values of MMDS1 and Oligocene samples with positive values. Samples between ~34.0 Ma and the EOB, labeled "Transitional", plot with intermediate MMDS1 values and high positive values of MMDS2. A plot of MMDS1 against age clearly resolves the pattern of nannofossil assemblage changes through time (Figure 6b), namely relative stability below ~34.0 Ma, a first transition (T1) to higher values between ~34.0 Ma and the EOB, and

![Figure 4](image-url)

**Figure 4.** Nannofossil species relative abundance data across the Eocene-Oligocene transition. (a) From left to right, *Cyclicargolithus floridanus* (<5 µm), *Cyclicargolithus floridanus*, *Sphenolithus predistentus*, and *Reticulofenestra* sp. A. (b) From left to right, *Calcidiscus protoannulus*, *Reticulofenestra dictyoda*, *Pemma papillatum*, and grouped *Discoaster* spp.; last occurrences of *P. papillatum* and *D. saipanensis* are marked. Data are from TDP Site 12 (base of section, purple) and TDP Site 17 (top of section, light blue). The horizontal gray bars indicate the duration of the globally recognized first and second positive steps in δ¹⁸O, and the pink shading indicates the duration of the EOT (34.0–33.5 Ma).
a second transition (T2) just above the EOB (~33.63 Ma) to the high positive values of the Oligocene.

4. Discussion

This discussion focuses on three key issues: the timing of major transitions in the nannofossil record, the nature of nannofossil assemblage changes and the potential paleoceanographic and climatic controls on these shifts.

4.1. Timing of Events Through the EOT

The variation in H diversity (Figure 3b), the first coordinate of the multidimensional scaling analysis (MMDS1) and the relative abundance plots of group 1 and 2 species (Figure 6b) summarize the major changes in

![Figure 5](image-url)
Figure 6. Results of the multivariate analysis of nannofossil assemblages. (a) First and second coordinates (MMDS1 and MMDS2) of the Metric Multidimensional Scaling, samples divided into “Eocene” (before 34.0 Ma), “Transitional” (34.0 Ma to 33.7 Ma) and “Oligocene” (after 33.7 Ma) groups. (b) From left to right, *Tuborotalia ampliapertura* δ18O [Pearson et al., 2008], first coordinate of the Metric Multidimensional Scaling analysis plotted against age (MMDS1), and grouped nannofossil abundance data (group 1, *Calcidiscus protoannulus*, *Reticulofenestra dictyoda*, *Pemma papillatum*, *Lanternithus minutus*, *Varolia macleodii*, and *Zygrhablithus bijugatus*; group 2, *Sphenolithus predistentus* and the large form of *Cyclicargolithus floridanus*). First (T1; ~34.0 Ma) and second (T2; ~33.63 Ma) major transitions in nannofossil assemblages are indicated on the plot of MMDS1. Data are from TDP Site 12 (base of section, purple) and TDP Site 17 (top of section, light blue). The horizontal gray bars indicate the duration of the globally recognized first and second positive steps in δ18O, and the pink shading indicates the duration of the EOT (34.0–33.5 Ma).
nannofossil assemblages through the EOT. The older, late Eocene part of this succession, below ~34.1 Ma, has stable nannofossil assemblages, as shown by the relatively constant H diversity, MMDS1 and group 1 and 2 abundances (Figures 3b and 6b). This is paralleled by relatively steady T. ampliapertura δ¹³C values (Figure 3a). The first disruption to this stability (T1) occurs across an interval of very low nannofossil abundances between ~34.0–33.95 Ma (four successive samples between 129.84 and 124.39 mcd in TDP12), which is coincident with a negative peak in δ¹³C, the onset of the first significant abundances of reworked Cretaceous nannofossils (~33.96 Ma; 125.56 mcd) and a ~2°C cooling of SSTs recorded by T. ampliapertura Mg/Ca ratios (~33.94 Ma; 122.4 mcd). This pronounced interval of low nannofossil abundances may be due to dissolution, either in the water column or during sediment diagenesis, dilution by other sedimentary constituents or low nannofossil productivity. Dissolution is unlikely, first because rare but well-preserved late Eocene nannofossils are still preserved within these sediments and second this is a relatively shallow (300–500 m paleowater depth) location significantly above the lysocline. Dilution of the nannofossil component is possible but would require a large change in sedimentation rates to account for the ~50 to 100 fold decrease in nannofossil abundance throughout this 5 m interval, in which there is no discernable change in the clay-rich lithology. The third possibility is an ecological event could account for both a decline in calcareous phytoplankton primary production and the observed transition (T1) in nannofossil assemblages. A fall in carbonate productivity, if widespread across the low-latitude oceans, could directly explain the negative trending δ¹³C values and the transient shoaling of the CCD observed immediately prior to the first δ¹⁸O step [Coxall et al., 2005] and may be coupled, via a weakened carbonate counter pump, to a transient decline in atmospheric pCO₂ and the SST cooling indicated by Mg/Ca data.

[22] The second, and more significant, assemblage transition (T2) occurs within the ~70 ka interval immediately above the EOB within the intermediate plate in δ¹⁸O. This involves an abrupt and permanent reduction in the H diversity of the nannofossil assemblages and a major shift in the taxonomic composition of assemblages, which is clearly seen in MMDS1 and group 1 and 2 abundances (Figure 6b). In contrast to T1 this transition is marked by high nannofossil abundances and does not correspond with any geochemical (stable isotopes or Mg/Ca) or sedimentological (Cretaceous reworking or pyrite abundance) signal. The T2 nannofossil event does, however, correspond with a wider biotic response across this relatively short ~70 ka interval, involving the extinction of five species of planktonic foraminifera and dwarfing of Pseudohastigerina [Wade and Pearson, 2008] and an overturn in larger benthic foraminifera [Pearson et al., 2008]. This largely biotic event precedes the δ¹⁸O maximum at the start of the OGGM by ~150–200 ka.

4.2. Nannofossil Assemblages and Paleoproducitivity

[23] A large component of the nannofossil assemblage shift across the EOT consists of a marked abundance decline across all the dominant late Eocene holococcolith species (Figure 5a). The production of fragile holococcoliths, formed of ~0.1 μm equidimensional calcite crystallites (Figures 1a and 1b), during the haploid phase of the coccolithophore life cycle, and more robust heterococcoliths (e.g., the placoliths of Figure 1d) during the diploid phase, has only recently been established [Billard, 1994; Young et al., 1999] and understanding the environmental trigger and adaptive function of the holococcolith-heterococcolith (haploid-diploid) transition in the coccolithophores is at an early stage [Cros et al., 2000; Noël et al., 2004]. Plankton sampling of the modern oceans has, however, established an association between high holococcolith abundances, near-surface habitats and generally oligotrophic conditions [Cros et al., 2000; Kleijne, 1991; Winter et al., 1994]. In spite of the potential environmental sensitivity of holococcoliths, their use in paleoceanographic reconstructions has been minimal because of their low preservation potential. The exceptional preservation of the Tanzanian sections through the EOT provides a unique deep-time record of the response of the holococcolith-bearing phase to a major environmental perturbation. The strong decline in the relative abundance of holococcoliths through the EOT can be interpreted as a significant shift away from oligotrophic surface water conditions. This assumes that there is no systematic decline in the preservational state, which would strongly affect the record of holococcoliths, but this is unlikely as very fragile holococcolith taxa, such as Orthozygus spp., are still well preserved in the basal Oligocene, whereas the abundances of more robust forms such as Lanterinitus and Zygryhablithus decline significantly.

[24] Further support for increasing productivity through the EOT is provided by the record of coccolithophore diversity. In modern coccolithophore assemblages, variations in the Shannon measure of ecological diversity (H) are correlated to changes in the nutrient content of surface waters, with r-selected, opportunist, usually placolith-bearing species, dominant in eutrophic conditions (low H diversity) while a more diverse flora of k-selected species develops under nutrient-limited conditions (high H diversity) [Broecker et al., 2000; Jordan, 2002]. The most rapid transition in the coccolithophore assemblages in this study is immediately after the EOB and is marked by a major and rapid decline in H diversity from late Eocene assemblages with a mean value of 2.67 to those of the earliest Oligocene with a mean of 2.41 (Figure 3b). This change in H is largely due to the increased dominance of *Cyclicargolithus floridanus* and the loss of important sub-dominant taxa, such as *Calcidiscus protoannulus* and *Lanterinitus minutus*. This diversity decline, coupled with the reduced abundance of oligotrophic taxa (holococcoliths and discoasters) and an increase in the abundance of the dominant, placolith-bearing species, *C. floridanus*, strongly indicates a marked increase in surface water nutrient concentrations across the EOB. This is consistent with a similar decline in the H diversity of planktonic foraminifera assemblages between the Eocene and Oligocene of this succession, which can also be related to more unstable and eutrophic conditions [Wade and Pearson, 2008].
The dominance of *C. floridanus* (small and large forms) and *Sphenolithus predistentus* in the early Oligocene of Tanzania, with combined abundances of ~60% or more, is similar to early Oligocene nannofossil assemblages of the equatorial Pacific Ocean, which are also dominated by *C. floridanus* and show increased sphenolith abundance during high-productivity intervals [Wade and Pálike, 2004]. *C. floridanus* has been associated with high-productivity environments [Aubry, 1992; Morechi et al., 2000], however sphenoliths have been shown to increase in abundance under both oligotrophic [Gibbs et al., 2004] and strongly eutrophic conditions [Wade and Bown, 2006]. In this study, although *S. predistentus* does show a strong increase in abundance across the EOT, the other moderately abundant sphenolith species, *S. moriformis*, shows no response through the same interval (Figure 5b), perhaps indicating distinct ecological adaptations within the genus.

### 4.3. Paleoceanography of the EOT

What are the possible causes of the surface ocean nutrient enrichment at this location and are they indicative of wider oceanographic-climate changes at this time? It is likely that the eustatic sea level fall, coincident with the onset of major continental-scale glaciation, released a significant pulse of nutrients around continental margins through the erosion of newly exposed organic-rich shallow shelf deposits. This is supported by the onset of significant reworking of Cretaceous nannofossils from the inland Kilwa Group at ~33.9 Ma. The input of freshly eroded nutrients may account for some of the enhanced productivity and nannofossil assemblage changes seen at this location, however the rapidity of the assemblage transition concentrated at the EOB and its independence from major phases of ice growth, as determined by the δ18O record, strongly indicate an independent biotic or physical transition in the surface water environment.

An alternative explanation would be a significant switch in regional ocean circulation patterns. In low southern latitudes the present-day western boundary of the Indian Ocean is dominated by the northward flowing East African Coastal Current (EACC), which is sourced from the westward flowing South Equatorial Current (SEC) [Swallow et al., 1991]. Toward the equator the EACC begins to be influenced by monsoonal variation as it interacts with the seasonally reversing Somali Current (SC), but it is relatively stable and not associated with significant upwelling south of the equator [Schott and McCreary, 2001]. Eocene general circulation models have a similar northward flowing EACC-type current along the western boundary of the Indian Ocean, which is fed by a strong westward flowing SEC-type current between 20 and 30°S [Huber et al., 2004; Thomas et al., 2006]. The consistent presence of a diverse lower photic zone coccolithophore assemblage in the Kilwa Group sediments, from the late Paleocene through to the EOB [Bown et al., 2008b], indicates a long-term stable oligotrophic paleoenvironment, which is consistent with the presence of a northward flowing western boundary current through much of the Paleogene. It is unlikely that the direction of this circulation pattern altered significantly across the EOT so other mechanisms for increasing the input of nutrients into the surface ocean at this location need to be considered.

Enhanced upwelling along the shelf break, driven by increased ocean-atmospheric circulation and associated turbulent mixing [Brunner and Biscaye, 2003], may have influenced this site located close to the shelf/slope transition (paleodepth ~300–500 m), but could not account for the more widespread disturbance of low-latitude planktonic ecosystems across the EOB. Evidence for this includes changes in equatorial Pacific radiolarian assemblages, which show a remarkably similar decline in H diversity and increase in cosmopolitan species abundance across the EOB [Funakawa et al., 2006] and the well-established intensive and globally synchronous extinctions within the planktonic foraminifera [Coxall and Pearson, 2000; Wade and Pearson, 2008]. One mechanism that could account for the geographical spread of this disturbance is the enhanced formation of Subantarctic Mode Water (SAMW), which in the modern ocean transports nutrients from the Subantarctic Zone to low-latitude regions of upwelling and may account for up to 75% of biological production north of 30°S [Sarmiento et al., 2004]. SAMW is formed by deep mixing in the Subantarctic Zone and is then transported at subthermocline depths throughout the subtropical gyres of the Southern Hemisphere [Ribbe, 2004; Sarmiento et al., 2004] and variations in the rate of formation of SAMW have been shown to connect high-latitude climate perturbations to the tropical Indian Ocean [Kiefer et al., 2006]. It is feasible that increased mixing in the Southern Ocean, coincident with Antarctic cooling and glaciation, enhanced the formation of SAMW which effectively connected southern high-latitude cooling with the tropical/subtropical oceans. New records are required to ascertain the extent of low-latitude organic carbon burial associated with these productivity increases [Ravizza and Paquay, 2008], which appears to be largely absent from the Southern Ocean [Rea and Lyle, 2005], and so test recent assertions that organic carbon burial did not play a significant role in the positive δ13C excursion and deepening of the CCD [Merico et al., 2008].

### 5. Conclusion

The new high-resolution record of calcareous nannofossils presented here clearly shows two marked shifts in coccolithophore assemblages closely related to the major climatic events of the Eocene-Oligocene transition. The first of these transitions, at ~34.0 Ma, coincides with a distinct, ~50 ka interval of very low nannofossil abundance, the first appearance of significant reworked Cretaceous nannofossils, indicative of a marked sea level fall and a distinct cooling in sea surface temperatures (SST). It precedes the first positive shift in δ18O values by ~150 ka. The second and larger transition, at ~33.63 Ma, includes a significant and long-term reduction in the diversity of nannofossil assemblages and is closely associated with the planktonic foraminiferal extinctions at the Eocene-Oligocene boundary [Wade and Pearson, 2008]. Both of these transitions are dominated by major reductions in the relative abundance of holococcolith and other oligotrophic taxa and an increase in the relative abundance of *Cyclicargolithus floridanus* and
Sphenolithus predistentus. These significant shifts contrast with the small number of existing EOT records from the low to midlatitudes, which document only a limited nanofossil response to this interval of pronounced global change [Monechi, 1986]. A large proportion of the nanofossil shift in the Tanzanian record is due to the decline of an extraordinarily abundant latest Eocene holococcolith assemblage. This, to our knowledge, the first documentation of long-term productivity changes between the haploid and diploid coccolithophore life cycle phases in response to pronounced paleoceanographic disturbance and we interpret this as representing increased nutrient availability in the low-latitude surface ocean. Many of these fragile and dissolution-susceptible holococcolith taxa are not usually preserved in typical carbonate-rich pelagic sediments, which is one of several preservational filters that can limit our understanding of past ecological change from an analysis of moderately to poorly preserved microfossil assemblages [Bown et al., 2008a; Young et al., 2005].

[30] Contrary to previous opinion [Diester-Haass and Zachos, 2003; Salamy and Zachos, 1999; Zachos and Kump, 2005], recent carbon cycle modeling has suggested that organic carbon burial associated with increased productivity across the EOT may have played little or no role in driving either the global positive shift in δ13C or the rapid deepening of the CCD, which are instead attributed to a reduction in shelf carbonate production and the weathering of exposed and relatively isotopically heavy neritic carbonates [Merico et al., 2008]. However, the calcareous nanofossil record presented here strongly indicates an increased availability of nutrients in the low-latitude surface ocean, adding to the growing number of studies that document an increase in surface ocean productivity directly coincident with the EOT [Anderson and Delaney, 2005; Diester-Haass et al., 1996; Diester-Haass and Zachos, 2003; Latimer and Filippelli, 2002; Ravizza and Paggiay, 2008; Salamy and Zachos, 1999]. This discrepancy between surface ocean productivity records and the extent of organic carbon burial across the EOT needs to be addressed and may indicate a widespread decoupling of productivity and carbon burial due to the increased ventilation of the deep ocean.

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