Pleistocene Calcareous Nannofossil Biostratigraphy and Gephyrocapsid Occurrence in Site U1431D, IODP 349, South China Sea

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Abstract: We reinvestigated the Pleistocene calcareous nannofossil biostratigraphy of Site U1431D (International Ocean Discovery Program (IODP) Expedition 349) in the South China Sea (SCS). Twelve calcareous nannofossil Pleistocene datums are identified in the site. The analysis confirms that the last occurrence (LO) of Calcidiscus macintyrei is below the first occurrence (FO) of large Gephyrocapsa spp. (>5.5 µm). The FO of medium Gephyrocapsa spp. (4–5.5 µm) is also identified in the samples through morphometric measurements, which was unreported in shipboard results. Magnetobiochronologic calibrations of the numerical ages of LO of Pseudoemiliania lacunosa and FO of Emiliania huxleyi are underestimated and need reassessment. Other potential markers such as a morphological turnover of circular to elliptical variants of Pseudoemiliania lacunosa and a small Gephyrocapsa acme almost synchronous with the FO of Emiliania huxleyi may offer biostratigraphic significance in the SCS. The morphologic changes in Gephyrocapsa coccoliths are also examined for the first time in Site U1431D. Placolith length and bridge angle changes are comparable with other ocean basins, suggesting that morphologic changes are most likely evolutionary novelties rather than being caused by local climate anomalies.

Keywords: calcareous nannofossils; South China Sea; Pleistocene; International Ocean Discovery Program; Site U1431; Gephyrocapsa

1. Introduction

The Pleistocene is characterized by worldwide sea level lows and highs due to increased amplitudes of glacial and interglacial episodes, respectively [1–3]. Ice volume fluctuations are well represented in deep sea sedimentary sequences by variations in proxies such as sediment properties, fossil assemblages, and/or isotope concentrations. The basis for proxy correlations is a precise and robust chronology constructed from natural cycles of sediment physical properties, radiometric ages, magnetic reversals, and/or biostratigraphic data. An accurate age model is important because major climatic shifts in the Pleistocene occurred within a few hundred thousand years, as exemplified by glacial–interglacial amplitude variations in δ18O records [2]. Furthermore, several periods in the Pleistocene are thought to be analogs to the modern-day climate, such as Marine Isotope Stage (MIS) 11 [4,5] and may contribute to a further understanding of past and future climate change dynamics. These climatic events can be put into geologic context through relative dating by using calcareous nannofossils.

Calcareous nannofossils are mostly made up of remains of marine, unicellular phytoplankton known as coccolithophores. Coccolithophores produce a robust, calcium carbonate exoskeleton called the coccosphere. The coccosphere disintegrates into its coccoliths (individual calcite plates)
upon death and is incorporated in fecal pellets and/or marine snow, thus dominating the carbonate fraction of deep-sea pelagic sediments. Calcareous nannofossils are utilized in sediment core studies because of their widespread geographic occurrence and stratigraphic significance. They underwent extinction and speciation events (datums) since the Triassic period, defining brief spans of geologic time [6]. The calibration of nannofossil datums through astronomical tuning and cyclostratigraphy has been substantial in creating a robust time scale for the Pleistocene [7–9]. Star-shaped discoasters are mainly used in subdividing the Early Pleistocene [7,10–14]. Gephyrocapsids are also major components of Pleistocene calcareous nannofossil assemblages [15–22]. Gephyrocapsids have been extensively studied because of complex morphological changes, which have been utilized to subdivide the Pleistocene [7,21–24]. However, semi-enclosed marginal seas may have different oceanographic parameters, resulting to a different calcareous nannofossil assemblage. It is therefore important to study the calcareous nannofossil biostratigraphy of marginal seas to verify if standard nannofossil zonations apply to these areas.

The South China Sea (SCS) is the largest marginal sea in the western Pacific area and has been subject to multiple paleoceanographic [25–30] and sedimentological studies [31–34]. The SCS is characterized by better carbonate preservation compared to other western Pacific marginal basins, providing a valuable sediment archive for East Asian Monsoon System (EAMS) variations since the Miocene [35]. Calcareous nannoplankton in surface sediments [36–39] have also been studied in the northern, western, and eastern SCS. Despite increasing scientific attention on the SCS, calcareous nannofossil biostratigraphic studies of Pleistocene deep sea sediments are limited. Most works are concentrated on the northern flanks of the SCS to acquire a record of the spreading history of the SCS [40–42].

International Ocean Discovery Program (IODP) Expedition 349 aimed to establish the ages of cessation of different spreading centers in the SCS Basin [43,44]. Site U1431D was the most successful in providing a continuous archive from the Miocene to Holocene, recovering around 400 m of sediment. Since Ocean Drilling Program (ODP) Leg 184 [42], IODP 349 is one of the very few research expeditions to obtain records since the start of ocean floor spreading of the SCS basin. The acquired cores, therefore, represent a valuable archive in which age constraints for different spreading events in the sub-basins can be studied. Furthermore, the thick sediment recovery provides a continuous sedimentary record of paleoceanographic changes in the SCS. Shipboard data have been able to estimate age dates of different sections of the whole core through biostratigraphic analysis using different microfossils [43]. Calcareous nannofossils have been useful in shipboard analysis to determine sedimentation rates and estimate the cessation of the sub-basin spreading. However, further refining of datum placements is needed to achieve a better time constraint for paleoceanographic changes.

This study reinvestigates the calcareous nannofossil biostratigraphy of the Pleistocene sediments acquired from IODP 349. Temporal size and bridge angle changes in the genus *Gephyrocapsa* are discussed in detail to assess the reliability of these bioevents in the SCS. The results of the research contribute to the assessment of nannofossil datums and their importance in Pleistocene correlation and chronology, especially in the SCS.

## 2. Materials and Methods

### 2.1. Present Day Circulation of the South China Sea

Surface water circulation in the SCS is currently influenced by the EAMS [30,45,46]. The EAMS is characterized by having a stronger winter and a weaker summer season. The winter season (December–February) is dominated by the northeast monsoon, causing a basin-wide cyclonic gyre (Figure 1). The winter monsoon is mainly driven by cold winds from Siberia, which runs southward along the coast of East Asia. Mountains near the East Asian coast force the winter monsoon to produce a southwest traveling jet over the SCS. The summer season (June–August) is characterized by a weaker southwest monsoon forming an anticyclonic gyre concentrated at the south and a cyclonic cell at the
north (Figure 1). The ocean jet separates near Vietnam: one branch travels northeast crossing the central basin while the other continues on the northern path. Both branches of the current exit the SCS through the Bashi Strait. The ocean dynamics of the SCS produce localized annual upwelling in Vietnam and in northwest Luzon, although upwelling during winter is generally stronger based on chlorophyll concentrations [30,45,46].

Eleven (11) lithostratigraphic units were defined by the shipboard team based on visual core descriptions, thin section analysis, biogenic materials, and physical properties. The first two units were investigated in this study, which spans the late Pliocene (section 15H) to the latest Pleistocene (section 1H; Figure 2). Unit I (0–101.16 m below sea floor (mbsf)) is characterized by a sequence of clay, silty clay, and clayey silt that has substantially more silt than Unit II. The silt layers are composed of quartz, feldspars (plagioclases and K-feldspars), and minor amounts of amphiboles, micas, and heavy minerals. The layers with more silt are greenish gray interbeds, usually with graded, fining upward sequences interpreted as turbidite deposits. Unit I contains multiple dark gray to brown volcanic ash layers, 0.5- to 5.0-cm thick.

The sediment cores were obtained from the central SCS by IODP Expedition 349 from 1 to 6 February 2014, at Site U1431 (Figure 1). Site U1431 is located near a relict spreading ridge where the youngest magnetic anomalies in the SCS are located [43]. The primary objective of the expedition was to determine the cessation of seafloor spreading at the East Sub-basin of the SCS. The site also provides an almost continuous sediment record, providing the opportunity to study a continuous, high-resolution paleoceanographic evolution of the SCS. Core U1431D was recovered from a mean seafloor depth of 4240.5 m below sea level (mbsl). About 402.11 m of core was recovered in Site U1431D in a 617-m interval (65.2% core recovery). The uppermost 135 m of core material (from section 1H to 15H) was sampled every 20 cm for the first two sections (1H and 2H), and every 150 cm for the rest of the sections. A total of 168 samples were analyzed for this study (Figure 2).

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sequences interpreted as turbidites [43]. Nannofossil oozes occur locally and often show a fining upward character, also identified as turbidite deposits. Unit I contains multiple dark gray to brown volcanic ash layers, 0.5- to 5.0-cm thick.

**Figure 2.** Schematic diagram of core sections (1H to 15H) examined in this study. It is generally composed of clays/claystone with interbeds of minor sand/sandstone, silt/siltstone, clayey silt, silty clay, and foraminiferal oozes. Age dates are determined from shipboard biostratigraphic data. White areas in the “Core Recovery” column approximate coring gaps. Horizontal lines in the “Samples” column approximate positions of the samples utilized for this study. Abbreviations listed: CSF = core depth below seafloor, H = advanced piston corer, Plio = Pliocene. Figure modified from [43].

Unit II (101.16–267.82 mbsf) is divided into two sub-units. Only samples from Sub-unit IIA (101.16–194.95 mbsf) are analyzed for calcareous nannofossils. This sub-unit is composed of clay, nannofossil-rich clay, and silty clay. The greenish gray clays are thickly bedded and are usually interbedded with greenish gray medium- to thickly-bedded nannofossil-rich clays and dark greenish gray thin- to medium-bedded silty clay layers. The clayey silt layers often show sharp, planar bases, which grade into clay and are interpreted as turbidite deposits [43]. Moreover, fine sand foraminiferal oozes with sharp, erosive bases, fining upward towards nannofossil-rich layers, are also interpreted as turbidites. Bioturbation from the *Nereites* ichnofacies are darker colored clay within lighter colored carbonate-rich sediments. Thin volcanic ash layers 0.5- to 2.0-cm thick are also identified but are much less compared to Unit I and are restricted at the upper portions of the sub-unit [43].

**2.3. Calcareous Nannofossil Analysis**

Smear slide replicates from unprocessed sediments were prepared following the standard procedure of [47]. At least 300 random fields of view (FOVs) were scanned using an Olympus BX51
were noted at 133.2–130.2 mbsf, 127.5 mbsf, 124.9 mbsf, 121.9–118.9 mbsf, and 99.9–98.3 mbsf. The study adopted the biometric subdivision of [21] which is mainly based on the maximum length of the coccolith. Forms <4 \mu m are small *Gephyrocapsa* spp.; medium *Gephyrocapsa* spp. are those with placolith lengths from \(\geq 4\) to \(<5.5\) \mu m, and large *Gephyrocapsa* spp. are \(\geq 5.5\) \mu m in length. All measurements are provided in the Supplementary Materials.

Morphological analysis of gephyrocapsids was done using smear slides prepared from unprocessed sediments. The *Gephyrocapsa* coccoliths were photographed and measured using Image-Pro Plus Version 7.0 mounted on an Olympus BX51 polarizing microscope under 1000× magnification. The *Gephyrocapsa* coccoliths were measured based on the parameters set by [52] (Figure 3), mainly the coccolith length and width, coccolith central area length and width, and bridge angle. At least 40 specimens per sample were analyzed. In samples with few nannofossils, 200 FOVs were scanned and all *Gephyrocapsa* placoliths seen were measured. The study adopted the biometric subdivision of [21] which is mainly based on the maximum length of the coccolith. Forms <4 \mu m are small *Gephyrocapsa* spp.; medium *Gephyrocapsa* spp. are those with placolith lengths from \(\geq 4\) to \(<5.5\) \mu m, and large *Gephyrocapsa* spp. are \(\geq 5.5\) \mu m in length. All measurements are provided in the Supplementary Materials.

![Figure 3](image-url) Measurements done on digitized photos of *Gephyrocapsa* placoliths under phase contrast research microscope at 1000× magnification.
3. Results

Twelve (12) calcareous nannofossil datums were detected for the Pleistocene section of Site U1431D. The preservation of calcareous nannofossils ranged from very poor to pristine specimens. Overgrowth and fragmentation of calcareous nannofossils (especially among discoasters) are evident. Foraminiferal tests, sponge spicules, radiolarian tests, and diatom frustules were observed in the first 3 m of the core. Samples barren of nannofossils were dominated by detrital materials such as clay minerals, biotite, plagioclase, quartz, and muscovite. Some barren intervals coincide with volcanic ash layers [43].

Calcareous nannofossil abundances were generally low throughout the core, but were high in portions of sections 1H and 2H (0 to 12.2 mbsf). Recrystallization of nannofossil specimens was uncommon throughout the core. Samples with partially to heavily dissolved placoliths coincide with decreases in calcareous nannofossil relative abundances. Figure 4 illustrates important calcareous nannofossil species used as biostratigraphic markers for Site U1431D. Table 1 summarizes all calcareous nannofossil biostratigraphic events documented in Site U1431D. The calcareous nannofossil distribution for all samples analyzed in Site U1431D is provided in the Supplementary Materials.

Figure 4. Photomicrographs of calcareous nannofossil marker taxa used in subdividing the uppermost Pliocene and Pleistocene. All specimens are photographed under 1000× magnification. Abbreviations listed: XPL = cross-polarized light image. PC = phase contrast image. (A) Emiliania huxleyi, XPL. Sample 2H-1W, 60–61 cm. (B) Pseudoemiliania lacunosa, XPL. Sample 2H-2W, 80–82 cm. (C) Gephyrocapsa sp. 3, XPL. Sample 4H-4W, 20–21 cm. (D) Gephyrocapsa spp. >4 µm, XPL. Sample 7H-4A, 20–21 cm. (E) Gephyrocapsa spp. >5.5 µm, XPL. Sample 8H-1W, 50 cm. (F) Calcidiscus macintyrei, XPL. Sample 10H-6W, 20–21 cm. (G) Discoaster brouweri, PC. Sample 12H-6W, 20–21 cm. (H) Discoaster triradiatus, PC. Sample 13H-7W, 20–21 cm. (I) Discoaster surculus, PC. Sample 15H-3W, 20–21 cm. White scale bar at the lower right of each photomicrograph is 5 µm in length.
Table 1. Summary of calcareous nannofossil biostratigraphic events of \cite{43} and this study for International Ocean Discovery Program (IODP) Expedition 349, Site U1431D. Age discrepancies from \cite{43} are due to the incorporation of Equatorial Pacific ages (compiled by \cite{9}) when available. Abbreviations listed: FO—first occurrence, LO—last occurrence, Ba—base acme, reemG—reentrance event of medium Gephyrocapsa, tlG—top (last occurrence) of large Gephyrocapsa, blG—base (first occurrence) of large Gephyrocapsa, bmG—base of medium Gephyrocapsa.

| IODP Expedition 349 Shipboard Results \cite{43} | This Study |
|-----------------------------------------------|------------|
| Age (Ma) Depth (mbsf) Nannofossil Event  | Age (Ma) Depth (mbsf) Nannofossil Event |
| 0.29 3.15  FO Emiliania huxleyi  | 0.29 3.905  FO Emiliania huxleyi |
| 0.44 12.15  LO Pseudoemiliania lacunosa  | 0.44 5.42  LO Pseudoemiliania lacunosa |
| 0.61 32.08  LO Gephyrocapsa sp. 3  | 0.61 25.925  LO Gephyrocapsa sp. 3 |
| 1.02 39.67  FO Gephyrocapsa sp. 3  | 1.026 57.605  FO Gephyrocapsa sp. 3 |
| 1.04 50.90  reemG  | 1.034 58.24  reemG |
| 1.24 60.70  tlG  | 1.062 60.60  tlG |
| 1.60 89.01  LO Calcidiscus macintyrei  | 1.541 78.985  blG |
| 1.62 96.10  blG  | 1.738 85.705  LO Calcidiscus macintyrei |
| 1.789 87.635 bmG  | 1.789 87.635 bmG |
| 1.905 94.15 LO Discoaster brouweri  | 1.905 94.15 LO Discoaster brouweri |
| 2.337 115.905 Ba Discoaster triradiatus  | 2.337 115.905 Ba Discoaster triradiatus |
| 2.491 127.855 LO Discoaster surculus  | 2.491 127.855 LO Discoaster surculus |

### 3.1. Calcareous Nannofossil Biostratigraphy

#### 3.1.1. LO of Discoaster surculus

The LO of *D. surculus* was previously assigned to Sample 15H-CC (135.20 mbsf) in the IODP 349 biostratigraphic results \cite{43}. Rare to few specimens are observed from Sample 15H-CC to Sample 15H-3W until they occur sporadically as reworked specimens throughout the younger portions of the sediment record (Figure 5A). There is a barren interval of nannofossils from 133.2 to 130.2 mbsf, but *D. surculus* reappeared as well-preserved specimens above this interval (Sample 15H-3W). It again reappears at 112.4 mbsf, but is regarded as reworked in this sample because not only is it higher than that of the bottom acme of *D. triradiatus*, a younger datum (Figure 5A), but its occurrence is also anomalously located above samples with consistent absences of *D. surculus*. Thus, the LO of *D. surculus* is placed between Samples 15H-3W, 20 to 21 cm and 15H-2W, 20 to 21 cm (127.85 mbsf).

![Figure 5](image_url)

Figure 5. Relative percentages of (A) *Discoaster surculus*, (B) *D. pentaradiatus*, (C) *D. triradiatus/D. brouweri + D. triradiatus*, and (D) *D. brouweri* with depth. Highlighted portions approximate the position of the datum from biostratigraphic data. Solid lines denote complete specimens while broken lines signify incomplete specimens. Abbreviations listed: LO—last occurrence, Ba—base acme.
3.1.2. LO of *Discoaster pentaradiatus*

The occurrence of *D. pentaradiatus* is very sporadic as they occur as complete and fragmented specimens throughout the deeper sections of the core (Sample 11H-CC to Sample 15H-CC; from 96 to 135 mbsf; Figure 5B). The LO of *D. pentaradiatus* may lie between 120 and 125 mbsf based on the sharp decrease in abundance between these depth intervals and the relative placement of the LO of *D. surculus* and Ba *D. triradiatus*. However, there is a pervasive occurrence of fragmented specimens above and below this depth. Similar problems regarding the LO of *D. pentaradiatus* is also reported in ODP 149 in the Iberia Margin (Holes 897C and 898A, [53]) and Deep Sea Drilling Project (DSDP) Site 662 of the eastern North Atlantic [10]. Based on discoaster counts for Site U1431D, the LO of this species is unclear and therefore not utilized in this study.

3.1.3. Base Acme (BA) of *Discoaster triradiatus*

Relative percentages based on counts of *D. triradiatus* against *D. brouweri* have been used in determining the base acme (>20%) of the former [51]. Counts at Sample 13H-7W (116.41 mbsf) show the first increased abundance of well-preserved specimens (Figure 5C). Because *D. triradiatus* was not observed below Sample 13H-CC (116.9 mbsf), the sudden increase of well-preserved specimens in Sample 13H-7W is considered the base acme event. Above 110 mbsf (Sample 13H-1W), *D. triradiatus* also increases in abundance greater than that of Sample 13H-7W. However, the paucity of well-preserved species and the abundance decrease of *D. triradiatus* from Sample 13H-6W to Sample 13H-2W (115–110 mbsf) make extending the acme event to younger sediments questionable (Figure 5C). Thus, this datum is placed at 115.905 mbsf, between Sample 13H-7W, 20–21 cm (116.41 mbsf) and Sample 13H-6W, 20–21 cm (114.91 mbsf).

3.1.4. LO of *Discoaster brouweri*

A steady decrease in complete *Discoaster brouweri* and increase in fragmented specimens are noted from the initial placement of the datum (Sample 12H-CC; 107.57 mbsf) until its complete absence in Sample 12H-2W, 20–21 cm (99.9 mbsf; Figure 5D). Only one complete specimen is found in 12H-3W, 20–21 cm (101.4 mbsf), probably due to decreased nannofossil abundance. Samples 12H-2W, 20–21 cm (99.9 mbsf) and 12H-1W, 19–20 cm (98.39 mbsf) are nearly barren of calcareous nannofossils. However, two consecutive samples above Sample 12H-1W (Sample 11H-5-CC, 140–145 cm; 96.10 mbsf and 11H-5W, 20–21 cm; 94.9 mbsf) still contain at least 2–3% *D. brouweri*. The species is absent in Sample 11H-4W, 20–21 cm (93.4 mbsf), and in four consecutive younger samples (6 m above Sample 11H-4W). Therefore, the LO of *D. brouweri* is reassigned between Sample 11H-5W, 20–21 cm (94.9 mbsf) and Sample 11H-4W, 20–21 cm (94.15 mbsf) from Sample 12H-CC (107.57 mbsf; [43]; Figure 5D).

3.1.5. First Occurrence (FO) of Medium *Gephyrocapsa* spp. (bmG)

The base of medium *Gephyrocapsa* (bmG) lies between Samples 10H-7W, 22 to 23 cm and 10H-8W, 20 to 21 cm (87.635 mbsf), which was previously unreported in shipboard results [43]. Placolith length measurements have been useful in determining the nannofossil datum. The gephyrocapsid specimens observed generally have a bridge oriented nearer the long axis of the coccolith (15°–40°) and a relatively open central area. These gephyrocapsids are most likely equivalent to *Gephyrocapsa oceanica*, *G. margereli*, and *G. muellerae*.

3.1.6. LO of *Calcidiscus macintyrei*

The species *C. macintyrei* generally occurs sporadically (only 1 or 2 specimens) in Sample U1431D-15H-CC, 10–16 cm to 7H-6W, 20 to 21 cm (135.33 to 57.9 mbsf), except in Samples 14H-CC (125.48 mbsf) and 10H-6W, 20 to 21 cm (85.8 mbsf) where they occur as few specimens. Recognition of *C. macintyrei* above Sample 10H-CC is difficult because siliciclastic materials dominate the samples and nannofossils are reworked to younger portions of the core. Coccolith preservation is moderate to good
in Samples 10H-CC to 10H-6W, while very few, poorly preserved and broken specimens of C. macintyrei are found above the interval and occur sporadically in younger sediment samples. Furthermore, coccoliths ~9 µm but similar to the morphology of C. macintyrei are noted in some samples. In this study, the last occurrence is placed between Samples 10H-6W, 20 to 21 cm and 10H-5W, 20 to 21 cm (85.705 mbsf).

3.1.7. FO of Large Gephyrocapsa spp. (blG)

The base (first occurrence) of large Gephyrocapsa (blG) had been documented at Sample U1431D 11H-CC (98.15 mbsf) based on IODP shipboard biostratigraphy (Li et al., 2014). However, this study did not observe large Gephyrocapsa at any sample within sections 11H and 10H (from 96.10 to 79.40 mbsf). It is detected much shallower (at Samples 9H-CC (78.91 mbsf) and above) and occurs above the LO of C. macintyrei rather than below it. Few to common large Gephyrocapsa specimens occur at Sample 9H-CC (78.91 mbsf) and they appeared abruptly in the sediment record. The first occurrence of large Gephyrocapsa morphotypes has also been documented above the LO of C. macintyrei in the Shatsky Rise in the northwest Pacific Ocean [54] and in ODP Leg 111 [55] and ODP Leg 138 [56] in the Equatorial Pacific. This pattern is also seen in the Caribbean Sea from ODP Leg 165 [57] and in the South Atlantic at ODP Sites 925 and 926 [58]. Thus, the blG is placed between Samples 9H-CC and 9H-7W, 36 cm (78.985 mbsf), above the LO of C. macintyrei. The distinct first appearance of large gephyrocapsids suggests that they are reliable markers in the SCS.

3.1.8. LO of Large Gephyrocapsa spp. (tlG)

The LO of large Gephyrocapsa (>5.5 µm) was reported in Sample U1431D 8H-1W, 50 cm (60.70 mbsf; [43]). Very good coccolith preservation is observed in this sample. Samples above and below this interval (8H-1W, 40 to 41 cm, 8H-1W, 20 to 21 cm and 8H-2W, 19 to 20 cm, respectively) are barren of nannofossils while Sample 7H-CC has no large Gephyrocapsa. Thus, the LO lies between Samples 8H-1W, 50 cm and 8H-1W, 40 to 41 cm (60.65 mbsf), and is consistent with shipboard results [43].

3.1.9. Reentrance Event of Medium Gephyrocapsa spp. (reemG)

Measurements of Gephyrocapsa coccoliths recognized the reentrance event of medium Gephyrocapsa (reemG) in Sample 7H-6W, 20 to 21 cm (57.90 mbsf). The placoliths are characterized by generally high bridge angle values (60°–70°) but are not exactly parallel to the short axis. The reemG is reassigned between Samples 7H-CC, 24 to 29 cm and 7H-6W, 20 to 21 cm (58.24 mbsf) from Sample 6H-CC reported in shipboard results (50.90 mbsf; [43]). In Site U1431D, the reemG event does not exactly correspond to the first occurrence (FO) of the omega-type Gephyrocapsa (Gephyrocapsa sp. 3). This trend is also observed in ODP 198, Site 1209B, in the Shatsky Rise [59] and in ODP 161 in the western Mediterranean [60].

3.1.10. FO and LO of Gephyrocapsa sp. 3

The FO of Gephyrocapsa sp. 3 was initially observed at Sample U1431D 5H-6W, 47 cm (39.67 mbsf; [43]). The occurrence of Gephyrocapsa sp. 3 in the sediment core shows a cyclical appearance and disappearance of nannofossils. Detrital material seen in smear slides probably diluted the nannofossil content. These barren layers are punctuated by samples with well-preserved, abundant calcareous nannofossils. The refinement of nannofossil biostratigraphy extended the FO further downcore to Sample 7H-5W, 60 to 61 cm (57.30 mbsf). The FO of Gephyrocapsa sp. 3 is then placed between Samples 7H-5W, 60 to 61 cm and 7H-6W, 20 to 21 cm (57.60 mbsf). This datum is 60 cm above the reemG event in Site U1431D. The LO of this nannofossil was initially placed at Sample U1431D 4H-CC, 29 to 34 cm (32.08 mbsf; [43]). The results of this study show that the LO is found between Samples 4H-3W, 20–21 cm, and 4H-3W, 120 to 121 cm (25.925 mbsf).
3.1.11. LO of *Pseudoemiliinia lacunosa*

Shipboard results from IODP Site U1431D placed the LO of *P. lacunosa* at Sample 2H-CC, 0 to 4 cm (12.15 mbsf; [43]). In this study, the datum is found at younger portions of the core—between Samples 2H-2W, 80 to 82 cm and 2H-2W, 60 to 62 cm (5.42 mbsf). Circular morphotypes are found more abundantly in the deeper sections of the core, while elliptical types are more common in younger sediments. Only the elliptical forms of *Pseudoemiliinia* are documented in the last two samples (Samples 2H-2W, 100 to 102 cm and 2H-2W, 80 to 82 cm; 5.7 and 5.5 mbsf, respectively).

3.1.12. FO of *Emiliania huxleyi*

The FO of *E. huxleyi* was previously reported at Sample 1H-CC, 7 to 14 cm (3.22 mbsf; [43]). The datum has been reassigned between Samples 2H-1W, 60 to 61 cm and 2H-1W, 80 to 81 cm (3.905 mbsf). Nannofossils above and below the event show a cyclical trend of barren and well-preserved assemblages, which made the recognition of this bioevent difficult. Dissolution in some of the nannofossils, especially among the placoliths, is also observed.

3.2. Morphologic Changes in Gephyrocapsids in Site U1431D

Small *Gephyrocapsa* is the only morphotype present from 135 to 87.94 mbsf (~2.58 to 1.79 Ma). It is characterized by a large central area and has varying bridge angles, although low average values are recorded in the interval. The overall trend shows an increasing small *Gephyrocapsa* coccolith length throughout the core. Medium *Gephyrocapsa* first appeared at 87.635 mbsf (~1.79 Ma) and does not show any long-term changes in length. The increase in the length of medium forms corresponds to an increase in the overall length of small morphotypes throughout most of the interval (Figure 6). Large *Gephyrocapsa* occurs consistently only from 78.91 to 60.70 mbsf. Exceptionally large forms (between 7.5 and 8.0 µm) are documented at around 0.8 Ma. No observable long-term trends in size changes are noted for the large morphotypes.

![Figure 6. Size measurements of *Gephyrocapsa* from Site U1431D correlated with the LR04 Stack of [2]. Dots in the leftmost line graph are sample points. Gray bars highlight onsets of relative abundance peaks in medium *Gephyrocapsa* forms. Blank intervals in the relative abundance graph indicate barren samples. The age of each sample was calculated through linear interpolation of data points following the results of nannofossil biostratigraphy. The dashed arrows in the relative abundance graph and LR04 Stack show trends in increasing amplitude fluctuations. Abbreviations listed: LO—last occurrence, FO—first occurrence, reemG—reentrance event of medium-sized *Gephyrocapsa*, tLG—top of large *Gephyrocapsa*, bLG—base of large *Gephyrocapsa*, bmG—base of medium *Gephyrocapsa.*](image-url)
Bridge angle changes across the *Gephyrocapsa* morphotypes range from 30° to 75° (Figure 6). A general decrease in bridge angle is documented from 2.58 to 1.79 Ma (small *Gephyrocapsa*) while an increase in bridge angle from 1.79 to 0.9 Ma is evident in the record (small and medium *Gephyrocapsa*). The highest values for bridge angles for small and medium forms are noted from 1.04 to 0.9 Ma and correlate with the first occurrence of *Gephyrocapsa* sp. 3. The bridge angle rapidly decreases in value from 0.9 to 0.8 Ma. An overall increase in bridge angle is seen from 0.8 Ma to present, with distinct lows between 0.5 and 0.45 Ma and at 0.2 Ma. In general, the bridge angle changes seem to respond similarly, regardless of the size fraction, and there is no apparent relation between the bridge angle with respect to average placolith length (Figure 6).

Relative abundances of small, medium, and large *Gephyrocapsa* morphotypes show oscillations in relative abundances of small and medium forms at around 1.79 and 0.65–0.6 Ma (Figure 6). Medium *Gephyrocapsa* comprise 20-30% of the total gephyrocapsid assemblage in this interval. From 0.65 to 0.6 Ma, oscillations of the shifting dominance of small and medium forms became more evident. When medium *Gephyrocapsa* is dominant, large morphotypes also appear in the record in few numbers, except for a spike in large *Gephyrocapsa* at around 0.2 Ma. The increasing amplitude fluctuation of medium *Gephyrocapsa* roughly coincides with the increasing glacial–interglacial variations trending towards younger sections (Figure 6). However, it is not clear if the driving mechanism of the changes in relative abundance fluctuations is correlated to glacial or interglacial stages, as both increasing and decreasing trends are evident in both conditions.

4. Discussion

4.1. Magnetobiochronologic Calibration of Nannofossil Datums in Site U1431D

Nannofossil events identified in Site U1431D are tied with magnetostratigraphic data from shipboard results to estimate a numerical age for the calcareous nannofossil datums. Magnetostratigraphic reversals are calibrated against [61], compiled in [9], while the calculation of the numerical ages using magnetostratigraphy is explained in the Supplementary Materials. Low- to mid-latitude Deep Sea Drilling Project (DSDP) and ODP sites are utilized to compare the nannofossil ages of datums observed in Site U1431D. The locations of these DSDP and ODP Sites are listed in Table 2 and are shown in Figure 7. The magnetobiochronologic calibrations are mostly in agreement with the compilation of [9], except for some datums (Table 1). These inconsistencies are focused on the discussion below.

| Table 2. Location of Deep Sea Drilling Project (DSDP), ODP Sites, and onshore references used for comparison. |
|---------------------------------------------------------------|
| **Hole/Name** | **Latitude** | **Longitude** | **Water Depth (m)** | **Reference(s)** |
| DSDP 552 | 56°02.56’ N | 23°13.88’ W | 2301 | [13,62] |
| DSDP 607 | 41°00.07’ N | 32°57.44’ W | 3427 | [21,22,63] |
| ODP 964 | 36°15.62’ N | 17°45.025’ E | 3650 | [64] |
| Boso Peninsula | 35° N | 140° E | - | [65] |
| ODP 1146 | 19°27.40’ N | 116°16.37’ E | 2092 | [42] |
| ODP 1148 | 18°50.17’ N | 116°33.93’ E | 3294 | [42,66] |
| ODP 769B | 8°47.12’ N | 121°17.68’ E | 3644 | [67] |
| ODP 768B | 8°00.05’ N | 121°13.19’ E | 4385 | [67] |
| ODP 925B | 4°12.25’ N | 43°29.35’ W | 3041 | [58] |
| ODP 677 | 1°12.03’ N | 83°44.16’ W | 3461 | [21,22] |
| ODP 848 | 2°59.6’ S | 110°28.8’ W | 3855 | [68] |
| ODP 710 | 4°18.7’ S | 60°58.8’ E | 3824 | [13,69] |
| DSDP 593 | 40°30.47’ S | 167°40.47’ E | 1068 | [70,71] |

The LO of *Calcidiscus macintyrei* in Pacific, Atlantic, Indian, and Mediterranean sections are correlated to Chron C1r.3r; however, the placement of the datum within the chron varies by location.
(Figure 8). Excluding Site 1146, the Indian, Atlantic, and Mediterranean records have younger age estimates than the Pacific and the SCS. In Site U1431D, correlation with shipboard magnetostratigraphy suggests that the event falls just above the top of the Olduvai Chron (C2n; 1.778–1.945 Ma, [9]), but still within the lowermost portions of Chron C1r.3r, while previous works document the LO of *C. macintyrei* at a slightly higher magnetostratigraphic position (lower C1r.3r but not directly on top of Olduvai—[7,9,22,58,64,70–72]). Discrepancies in the ages of the extinction event (0.2 Ma) may suggest diachronity.

Although the taxonomic definition of *C. macintyrei* is simple, several inconsistencies and differing species concepts exist in the literature. In Sites 768 and 769 in the Sulu Sea, [67] cited [73] for their adopted age for *C. tropicus* to subdivide NN19 of [74]. However, there is no written account of *C. tropicus* in [73]. The *C. tropicus* of [67] may be equivalent to the *C. macintyrei* of [73]. This equivalence may be unlikely because large age discrepancies exist in the literature between the two species (1.48 Ma—[67]; 1.57 Ma—[73]) and thus the differences may be an effect of varying species concepts. The stratigraphic inconsistency for the datum is also reported in the Japan Sea (1.33 Ma; [75]). The LO in the Japan Sea is younger by 0.22 Ma compared to the study of [51] and by 0.27 Ma compared to [7]. Size variations in *C. leptoporus* (5–8 μm, sometimes reaching up to 10 μm) may have caused confusion whether the larger end of the species belonged to *C. macintyrei* or a larger variant of *C. leptoporus* (e.g., [75]).

Studies concerning the LO of *Gephyrocapsa* sp. 3 are limited only to the Mediterranean. The limited occurrence and distribution of studies of the LO of *Gephyrocapsa* sp. 3 is the reason [7] did not include the event for global biostratigraphic correlation. The datum age (0.58 Ma) was first identified by [76] and calibrated through magnetostratigraphy. More recent calibrations in the Mediterranean correlated the event to MIS 15 (0.570–0.577 Ma; [64,77]). In the SCS, there is very little documentation on the distribution of this nannofossil downcore [43]. The LO of *Gephyrocapsa* sp. 3 is placed within the middle portions of Chron C1n (Brunhes Chron; 0–0.780 Ma) in Site U1431D, suggesting an age of 0.44 Ma for the datum (Figure 9). Although events using *Gephyrocapsa* are used in some studies in the SCS [41,42], this study presents one of the very few evidence for the occurrence and distribution of *Gephyrocapsa* sp. 3 in SCS Pleistocene sediments. Its clear appearance downcore implies its significance as a calcareous nannofossil marker in the SCS. It must be noted with caution that *Gephyrocapsa* sp. 3 is still present in the open ocean today [77]; thus, careful analysis of the bridge angles of *Gephyrocapsa* downcore may be necessary to determine the LO of this species.

**Figure 7.** Location of DSDP/ODP/IODP Sites used in this study for comparison of Pleistocene nannofossil datums. Site IODP 1431 is examined in the present study.
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The distribution of this nannofossil downcore [43]. The LO of the event to MIS 15 (0.570–0.577 Ma; [64,77]). In the SCS, there is very little documentation on the occurrence and distribution of studies of the LO of this species.

The open ocean today [77]; thus, careful analysis of the bridge angles of 3 in SCS Pleistocene sediments. Its clear appearance downcore implies its significance as a calcareous nannofossil datum (Figure 9). Although events using this study presents one of the very few evidence for the occurrence and distribution of studies of the LO of C. tropicus.

Studies concerning the LO of Gephyrocapsa sp. 3 are limited only to...the LO of P. lacunosa to be correlated to MIS 12 [17,21,22]. The upper portions of the core are defined mainly by C. leptoporus of [67] may be equivalent to the NN19 of [74]. However, there is no written account of or a larger variant of C. macintyrei or C. pacifica.

Comparison of stratigraphic ranges of the LO of Calcidiscus macintyrei from IODP/ODP/DSDP Sites. Magnetostratigraphic records from IODP Site U1431D, ODP Sites 1148, 848, 710 and DSDP Site 593 are calibrated to the astronomically tuned Geomagnetic Polarity Time Scale (GPTS) of [61] as compiled by [9]. Oxygen isotope data is used to determine the ages in ODP Sites 1146, 925, 964 and DSDP Site 607. These datums are calibrated to the benthic foraminiferal δ18O time scale of [2]. Horizontal bars represent the recalculated datum ages for each site. Med.—Mediterranean. Black—normal; white—reversed polarity.

Comparison of stratigraphic ranges of the FO and LO of Gephyrocapsa sp. 3 from IODP/ODP/ DSDP Sites. Magnetostratigraphic records from IODP Site U1431D and ODP Site 848 are calibrated to the astronomically tuned Geomagnetic Polarity Time Scale (GPTS) of [61] as compiled by [9]. Oxygen isotope data is used to determine the ages in ODP Site 964 and DSDP Site 607. These datums are calibrated to the benthic foraminiferal δ18O time scale of [2]. Horizontal bars represent the recalculated datum ages for each site. Upward facing horizontal bars represent FO datums, while downward facing bars signify LO datums. Med.—Mediterranean. Black—normal; white—reversed polarity.
Correlation with magnetostratigraphy places the LO of *P. lacunosa* within the upper portions of Chron C1n (Figure 10), which corresponds to an age of 0.092 Ma in Site U1431D. There is a considerable age discrepancy in the LO of *P. lacunosa* in Site U1431D compared to the literature. The difference ranges from 0.37 Ma between Sites U1431D and Sites 1146 (SCS; [42]), 925 (Atlantic; [58]), and 964 (Mediterranean; [77]) to as much as 0.55 Ma compared to the southwest Pacific [74,75]. The results of this study differ significantly compared from previous works which suggest the LO of *P. lacunosa* to be correlated to MIS 12 [17,21,22]. The upper portions of the core are defined mainly by interbedded to interlaminated clayey silt to silty clay which are interpreted as turbidite sequences [43]. Turbidity currents may have caused the resuspension of coccoliths of *P. lacunosa* to younger sediments of the core. Uncertainties in magnetostratigraphic calibration may also be a factor in the age discrepancy. Because the age calculated from magnetostratigraphy is too young compared to other sites, the standard age of 0.44 Ma [9] is adopted for the LO of *P. lacunosa* (Figure 10, gray datum).

The age of the FO of *E. huxleyi* in Site U1431D also has a large discrepancy. Correlation to Site U1431D magnetostratigraphy showed that the FO of *E. huxleyi* corresponds to the upper portions of Chron C1n at 0.066 Ma (Figure 11, black datum). The oldest age estimates are found in ODP Sites 1146 and 1148 followed by ODP Sites 768B and 769 in the Sulu Sea (Figure 11). Younger ages are noted from the open ocean (Pacific and Atlantic) and in the Mediterranean Sea; however, the age acquired from Site U1431D is much younger than all other sites. The uncertainty ranges from 0.305 Ma compared to Site 1148 [42] to 0.188 Ma compared to Site 925 [58]. Bioturbation and terrigenous input to the basin may have affected the signal of the FO. Bioturbation causes vertical sediment mixing which may reposition a datum to a slightly deeper depth. However, shipboard results reveal only slight bioturbation (10–30%) in the interval where the FO of *E. huxleyi* is documented [43] and may have had minor effect in the position of the datum. Terrigenous input may be a problem especially for less robust species like *E. huxleyi*. Previous reports suggest the sporadic occurrence of *E. huxleyi* near its first appearance [15,17,78]. Carbonate dilution through terrigenous input might further obscure the datum. Shipboard data suggest that carbonate percentages in Pleistocene sediments of Site U1431D are very low (1–10%; [43]). This interval is also defined by an abundance of plagioclase, mica, and amphibole.
seen in smear slides, suggesting increased detrital input. Magnetostratigraphic calibration may have also contributed to the age discrepancy recorded in IODP Site U1431D. Thus, the age of 0.29 Ma [9] is adopted for the datum (Figure 11, gray datum).

| Age (Ma) | Chron        | This Study | South China Sea | Sulu Sea | Pacific | Atlantic | Med. |
|----------|--------------|------------|-----------------|----------|---------|----------|------|
| 0.00     | C1n          | IODP       | ODP             | ODP      | DSDP    | ODP      | ODP  |
| 0.10     |              | U1431D     | U1148           | U1148    | 769B    | 768B     | 593  |
| 0.20     |              |            | 15°42′N         | 19°27′N  | 18°50′N | 8°47′N   | 8°00′N|
| 0.30     |              |            | 18°50′N         | 19°27′N  | 18°50′N | 18°50′N  | 8°00′N|
| 0.40     |              |            | 8°47′N          | 8°47′N   | 8°47′N  | 8°47′N   | 8°47′N|
| 0.50     |              |            | 8°00′N          | 8°00′N   | 8°00′N  | 8°00′N   | 8°00′N|
| 0.60     |              |            | 4°12′N          | 4°12′N   | 4°12′N  | 4°12′N   | 4°12′N|
| 0.70     |              |            | 36°16′N         | 36°16′N  | 36°16′N | 36°16′N  | 36°16′N|

Figure 11. Comparison of stratigraphic ranges of FO of Emiliania huxleyi from IODP/ODP/DSDP Sites. Magnetostratigraphic records from IODP Site U1431D, ODP Sites U1148, 769B, 768B, and DSDP Site 593 are calibrated to the astronomically tuned Geomagnetic Polarity Time Scale (GPTS) of [61] as compiled by [9]. Oxygen isotope data is used to determine the ages in ODP Sites 1146, 925 and 964. These datums are calibrated to the benthic foraminiferal δ18O time scale of [2]. Horizontal bars represent the recalculated datum ages for each site. Black datum in IODP Site U1431D represents age acquired from magnetostratigraphy. Gray datum in IODP Site U1431D represents placement of the datum from this study using the datum age reported in [9]. Med.—Mediterranean.

4.2. The Importance and Complexity of Gephyrocapsa Morphometry in Site U1431D

The overall time-transgressive size increase correlates well with those of previous works [19,21–24,59,79]. The work of [19] complements the data gathered; however, no large Gephyrocapsa were observed in the lower Pleistocene sections of the core studied. Only small Gephyrocapsa spp. placoliths are documented from Sample 15H-CC (~135.3 m; latest Pliocene) to 10H-8W, 20–21 cm (~87.94 m; early Pleistocene). The Pliocene–Pleistocene was defined at the base of the Calabrian (~1.8 Ma) by [19] and may account for the disparity. Nevertheless, the size increase is documented across ocean basins; thus, the morphologic change in Gephyrocapsa most probably points to an evolutionary trend. The time-transgressive size increase is applicable for biostratigraphic use and may explain the comparable record of Site U1431D sediments to other ocean basins. Global climate changes longer than glacial–interglacial cycles, such as ocean cooling brought on by a steady increase in ice cover, may have affected the long-term changes in the size of Gephyrocapsa coccoliths [79]. It is also noted that the changes in morphological parameters of the gephyrocapsids are not in one-to-one correspondence with glacial–interglacial variations. Higher resolution sampling, correlatable to suborbital variations in δ18O, is needed to further explain the response of Gephyrocapsa during these time scales.

Specimens of Gephyrocapsa coccoliths analyzed by [80] from tropical Atlantic waters generally have high bridge angles (parallel to the short axis of the coccolith). This observation is supported by [52], noting further that the bridge angle may be correlated with geographic distribution—the position of the bridge approaches the long axis towards high latitudes, in contrast to a mean bridge angle of 60° within 15 degrees north and south of the Equator. If temperature has a direct effect on bridge angle orientation, then there should be a glacial–interglacial trend with respect to bridge angle, with relatively colder periods shifting to lower bridge angle values. However, in Site U1431D there is
very little correlation between the average bridge angle values of all size fractions measured and the δ¹⁸O record (LR04 Stack of [2]; Figure 6). Most probably, environmental factors other than temperature may have had an influence on the development of the bridge.

The presence of medium-sized forms with high bridge angle values at around 1.0–0.6 Ma is documented in this study and is comparable to previous works [7,19,21,23,24,59,81–83]. This Gephyrocapsa morphotype is most probably equivalent to Gephyrocapsa sp. 3 of [19], Gephyrocapsa oceanica vertical of [23], and Gephyrocapsa sp. C of [24]. Small forms of Gephyrocapsa with a bridge angle parallel to the short axis are also documented mostly from 1.0 to 0.9 Ma near the “small Gephyrocapsa acme” [23,24]. These studies are correlatable and span different ocean basins, each with their own local climate. Thus, the changes in the morphology of Gephyrocapsa in the SCS may not be factors of regional climate perturbations. These results show the importance of both bridge angle and size for the determination of certain Gephyrocapsa morphotypes for Pleistocene biostratigraphy.

4.3. Age–Depth Model for Site U1431D

An age–depth model is constructed based on magnetostratigraphic and nannofossil datums identified in Site U1431D. The datum ages are calculated based on linear interpolation between magnetostratigraphic chronos (Supplementary Materials). The ages for the FO of E. huxleyi and LO of P. lacunosa are adopted from [9] because the calculated site-specific ages are too young compared to published figures. The LO of Gephyrocapsa sp. 3 is also taken from [9] because the only calibration done for this datum is from [76] and [64] in the Mediterranean. Thus, the age–depth model for Site U1431D is a combination of ages compiled by [9] and site-specific numerical ages.

The age–depth curve generated from the nannofossil datums using calcareous nannofossils shows consistency with other microfossil groups (Figure 12). From 135 mbsf to ~80 mbsf, planktonic foraminifera and calcareous nannofossil data generally agree with each other (Figure 12), with slight divergence at the 100–135 mbsf interval. Above 90 mbsf, planktonic foraminiferal datums imply younger ages than the calcareous nannofossils. Further analysis of planktonic foraminiferal biostratigraphy is needed to verify the age discrepancies between the two microfossil groups. The age model from radiolarians for the uppermost layers (0–10 mbsf) agrees with the nannofossils, reflecting a slow sedimentation rate within the interval.

![Age–Depth Models](image)

**Figure 12.** Comparison of age–depth models using different microfossil groups for Site U1431D. Symbols represent specific microfossil groups found in the site. The nannofossil age model incorporates published ages for FO of E. huxleyi, LO of P. lacunosa, and LO of Gephyrocapsa sp. 3 found in [9] and magnetostratigraphic ages acquired from this study for the older datums. The planktonic foraminifera and radiolarian curves use the ages from [9].
4.4. Other Possible Nannofossil Events

There are other possible nannofossil events found in the core which may need further reexamination. These events are recorded as shifts in the abundance values of nannofossil species, or a cross-over of the abundance of variants within a species. These shifts may offer biostratigraphic significance if assessed further and documented across other basins. The genus *Pseudoemiliania* had been differentiated based on the number of distal slits [23] and the degree of ellipticity [84]; these morphotypes are not routinely used in biostratigraphy. However, large, circular forms only appeared in the late Pleistocene and the changes in their morphology may be related to temperature fluctuations [48,85]. Absolute abundance counts from Site U1431D show a shift in abundance between *Pseudoemiliania* circular and elliptical variants between 120 to 130 mbsf (Figure 13). Circular variants of *P. lacunosa* are dominant from 135 to 130 mbsf and become replaced by elliptical types. This observation is different from the work of [84] wherein the circular forms of *Pseudoemiliania* dominate the younger portions of DSDP Leg 4 cores. It is possible that ecological preferences and the marginal basin setting of the SCS may have affected the distribution of *Pseudoemiliania* compared to the open-ocean setting of DSDP Leg 4 [84].

![Figure 13](https://example.com/figure13.png)

**Figure 13.** Absolute abundance of the calcareous nannofossil *Pseudoemiliania lacunosa* from 0 to 135 mbsf. Broken line shows abundance of circular variants. Solid line shows abundance of elliptical morphotypes. The gray highlighted area estimates the transition of circular-dominated to elliptical-dominated *Pseudoemiliania* in the core.

The first and last occurrences of the small *Gephyrocapsa* acme (1.24 to 1.02 Ma, respectively [9]), was not detected in Site U1431D. Possible carbonate dilution through terrigenous input may have been the cause of the absence of the event. However, Zone NN21 [74] consists of several other *Gephyrocapsa* acme intervals, one of which may be recorded in the SCS. Nannofossil biostratigraphy from the Kings Trough at the northeastern Atlantic Ocean show several acme intervals corresponding to different species of small *Gephyrocapsa* [85]. They have documented the dominance of *G. aperta* near the FO of *E. huxleyi*. In Sample 2H-1W, 60–61 cm (3.80 mbsf), an increase in small *Gephyrocapsa* placoliths is observed (Figure 14). These small placoliths have a large central area and a bridge near the long
axis, in agreement with the taxonomic description of *G. aperta*. Interestingly, the first occurrence of *E. huxleyi* is assigned to this sample and correlates well with observations in Kings Trough. A minor small *Gephyrocapsa* acme event occurs between Zones NN20 and NN21 [86] and was also documented by [24], most likely corresponding to the earlier portions of their Assemblage “B”.

The reliability of these secondary events can be assessed through correlation with oxygen isotope stratigraphy or cyclostratigraphy using the physical properties of sediments in the SCS. These minor events have been noted in some studies [24,84–86]. Details of these events in the SCS can be studied further through higher resolution sampling in the intervals where they are noted. Additional data from other ocean basins are needed to verify whether these nannofossil events are recorded on a regional or a global scale.

**Figure 14.** Absolute abundance of small *Gephyrocapsa* morphotypes from 0 to 8 mbsf. Gray highlighted area estimates an acme interval of the nannofossil. The secondary event coincides with the first occurrence of *Emiliania huxleyi*, which also lies within the gray area.

### 5. Conclusions

Twelve (12) calcareous nannofossil datums are documented in Pleistocene sediments of Site U1431D. This study is one of the first attempts to describe the distribution of *Gephyrocapsa* sp. 3 in SCS sediments and is suggested to be a well-defined marker for the subdivision of the Pleistocene in the marginal sea. The FO of *Gephyrocapsa* sp. 3 is near the reentrance event of medium *Gephyrocapsa* (reemG), which is also observed in several studies [7,21–23,60,81,83]. Nannofossil analysis suggests that the FO of large *Gephyrocapsa* (blG) is found above the LO of *Calcidiscus macintyrei*, which differs from shipboard results [43]. The LO of *Discoaster* species are determined using a semi-quantitative method to reduce the effect of reworking; however, the LO of *D. pentaradiatus* is very sporadic and therefore is not used as a marker for the Pleistocene in Site U1431D. The numerical ages of nannofossil datums are generated through correlation with magnetostratigraphy. The ages of the FO of *E. huxleyi* and LO of *P. lacunosa* are underestimated by 0.22 and 0.35 Ma, respectively, while bmG and LO of *C. macintyrei* are slightly overestimated by 0.12 and 0.15 Ma, respectively. The results of the magnetobiochronological calibration suggests further analysis using other stratigraphies (e.g., oxygen isotope stratigraphy, cyclostratigraphy).

Morphometric analysis of *Gephyrocapsa* coccoliths from the late Pliocene to Pleistocene sediments of Site U1431D allowed for the recognition and refinement of bioevents of the gephyrocapsids. These events are useful in updating the nannofossil biostratigraphy of the site. The events documented are the FO of medium *Gephyrocapsa* morphotypes (87.635 mbsf), the FO and LO of large *Gephyrocapsa* (78.985 and 60.65 mbsf, respectively), the reemG event (58.24 mbsf), and the FO and LO of *Gephyrocapsa* sp. 3 (57.605 and 25.925 mbsf, respectively). The trends of placolith length and bridge angles of *Gephyrocapsa* in Site U1431D are correlatable with previous studies on other ocean basins. The data
suggest that the change in the size of the gephyrocapsids points to an evolutionary trend rather than being caused by local or regional climate anomalies.

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Appendix A. Taxonomic Remarks

Calcidiscus leptoporus (Murray and Blackman, 1989) Loeblich and Tappan, 1978
Circular to subcircular proximal shield, birefringent. Distal shield non-birefringent with curved sutures. Shields easily separated. Closed central area.
REMARKS: Species is of little biostratigraphic value for the Pleistocene sediments of Site U1431D.

Calcidiscus macintyrei (Bukry and Bramlette, 1969a) Loeblich and Tappan, 1978
Circular proximal shield, birefringent. Distal shield non-birefringent with curved sutures, >10 µm with central opening.

Ceratolithus cristatus Kamptner, 1950
Strongly birefringent, c-axis lies in plane of horseshoe, perpendicular to length. Variable in size and degree of ornamentation.
REMARKS: Species is of little biostratigraphic value for the Pleistocene sediments of Site U1431D.

Ceratolithus telesmus Norris, 1965
Strongly birefringent, c-axis lies in plane of horseshoe, perpendicular to length. Long, sinuous arms.
REMARKS: Species is of little biostratigraphic value for the Pleistocene sediments of Site U1431D.

Coccolithus pelagicus (Wallich, 1877) Schiller, 1930
Elliptical placolith; central area open or with simple bar on the proximal surface.
REMARKS: Specimens of Coccolithus pelagicus with simple bar are only noted from section 10H and below (79.2 mbsf and below). The disappearance of C. pelagicus with bar may be a result of the migration of the species to higher latitudes within the Pleistocene. However, this event was not detected in the samples.

Cyclicargolithus floridanus (Roth and Hay in Hay et al., 1967) Bukry, 1971a [Coccolithus]
Coccoliths circular to sub-circular with small central area, <11 µm in width.
REMARKS: Occurs as reworked specimens.

Discoaster asymmetricus Gartner, 1969c
Asymmetric five-rayed variety of D. brouweri.
REMARKS: Occurs as reworked specimens.
Discoaster bellus Bukry and Percival, 1971
Five-rayed discoasters without any central knob or central area development. Straight rays that taper slightly and terminate in points. All rays lie on the same plane.
REMARKS: Occurs as reworked specimens.

Discoaster berggrenii Bukry, 1971b
Symmetric; five-rayed; rays curved; without bifurcations; well-developed central boss; central area 1-2x free ray length; adjacent rays separated at central-area.
REMARKS: Occurs as reworked specimens.

Discoaster blackstockae Bukry, 1973
four-rayed form with inter-ray angles of 60° and 120°. Considered a variant of D. brouweri.
REMARKS: Distribution within the core is very limited and does not parallel that of D. brouweri.

Discoaster brouweri Tan, 1927b emend. Bramlette and Riedel, 1954
Ray tips non-bifurcate but with marked proximal extensions giving it a distinctive concavo-convex form; six-rayed asterolith species.

Discoaster calcaris Gartner, 1967
Six-rayed asterolith, bifurcating asymmetrically at tip. Longer branch of bifurcation curved sharply proximally but extending only slightly laterally beyond tip of ray.
REMARKS: Occurs as reworked specimens.

Discoaster kugleri Martini and Bramlette, 1963
Central-area wide and flat; free rays short with notched ends rather than true bifurcations.
REMARKS: Occurs as reworked specimens.

Discoaster pentaradiatus Tan, 1927b
Five-rayed acute bifurcations; strongly concavo-convex and birefringent (each ray is a separate unit, with c-axes inclined slightly from vertical toward radial).
REMARKS: The distribution of D. pentaradiatus is sporadic in Site U1431D sediments, thus the LO of the species was not used for biostratigraphy.

Discoaster quinqueramus Gartner, 1969c
Five-rayed with concavo-convex rays. Central area with prominent distal sutural ridges and large proximal boss.
REMARKS: Occurs as reworked specimens.

Discoaster surculus Martini and Bramlette, 1963
Six-rayed and rarely with fewer rays with a stellate knob in the central area. From the knob, small ridges extend to the margin between the rays on one side of the asterolith and along the arms on the side. The slender rays are slightly enlarged at the end, and show a trifurcation, with the central spine extending beyond the arms.

Discoaster tamalis Kamptner, 1967
Concavo-convex rays. Calcite body consists of four arms arranged to form an orthogonal cross. There is no boss in the central area.
REMARKS: Occurs as reworked specimens.

Discoaster triradiatus Tan, 1927b
Symmetric three-rayed variety of D. brouweri.
Discoaster variabilis Martini and Bramlette, 1963
Central-area with proximal and distal ridges but weak bosses. Bifurcations well developed.
REMARKS: Occurs as reworked specimens.

Emiliania huxleyi (Lohmann, 1902) Hay and Mohler in Hay et al., 1967
No bridge, slits between distal shield elements.

Florisphaera profunda Okada and Honjo, 1973
Coccoliths formed of single angular 5-sided calcite units.
REMARKS: Species is of little biostratigraphic value for the Pleistocene sediments of Site U1431D.

Gephyrocapsa Kamptner, 1943
Like Reticulofenestra but with bridge formed from inner tube.
REMARKS: Size differences of coccoliths are used to separate the genus into different morphotypes for biostratigraphic purposes. The size criteria assigned by [21] is adopted in this study.

Gephyrocapsa sp. 3 Rio, 1982 (synonym: G. parallela Hay and Beaudry, 1973)
Large (4.0–6.4 µm) circular to sub-circular placolith, open central-area spanned by a bridge aligned 60°–90° to the long axis.
REMARKS: In this study, only those with a bridge aligned at 85°–90° to the long axis are considered as Gephyrocapsa sp. 3.

Helicosphaera carteri (Wallich, 1877) Kamptner, 1954
Medium to large size; flange ends in wing; two pores in central-area.
REMARKS: Species is of little biostratigraphic value for the Pleistocene sediments of Site U1431D.

Helicosphaera carteri var. wallichi (Lohmann, 1902) Theodoridis, 1984 [H. wallichi]
Like H. carteri but central-area with inclined pores.
REMARKS: Species is of little biostratigraphic value for the Pleistocene sediments of Site U1431D.

Helicosphaera inversa (Gartner, 1977 ex Gartner, 1980) Theodoridis, 1984
Like H. carteri but with large pores separated by prominent conjunct sub-horizontal bar usually with inverse direction.
REMARKS: Only one specimen of H. inversa is seen in Site U1431D samples; thus, it is not utilized for biostratigraphy.

Helicosphaera sellii (Bukry and Bramlette, 1969a) Jafar and Martini, 1975 [Helicopontosphaera]
Like H. carteri but pores large.
REMARKS: Distribution is sporadic in Site U1431D samples; thus, it is not utilized for biostratigraphy.

Neosphaera coccolithomorpha Lecal-Schaluder, 1950
Circular, ring-shaped coccoliths with open central-area. Single shield and tube, no distal shield. Superficially similar in light microscope to Umbilicosphaera rotula but shield paler in phase contrast and distinctive pseudoextinction cross seen in cross-polars at high focus.
REMARKS: Species is of little biostratigraphic value for the Pleistocene sediments of Site U1431D.

Pseudoemiliania lacunosa (Kamptner 1963) Gartner, 1969 [Ellipsoplacolithus]
Circular to broadly elliptical with variable number of slits in distal shield. Abundant in late Pliocene but well-developed, large circular forms occur only in the Pleistocene.
REMARKS: [84] observed that circular forms became much abundant during the late Pleistocene. However, circular forms are more abundant in the early Pleistocene in samples from Site U1431D.
Reticulofenestra minuta Roth, 1970
<3 µm which probably includes several different species.
REMARKS: Species is of little biostratigraphic value for the Pleistocene sediments of Site U1431D.

Reticulofenestra haqii Backman, 1978
3–5 µm which intergrades with R. pseudoumbilicus.
REMARKS: Species is of little biostratigraphic value for the Pleistocene sediments of Site U1431D.

Reticulofenestra pseudoumbilicus (Gartner, 1967b) Gartner, 1969c [Coccolithus]
>5 µm but for biostratigraphy it is better to separate the >7 µm specimens.
REMARKS: Occurs as reworked specimens.

Rhabdosphaera clavigera Murray and Blackman, 1898
Robust, club-shaped, 7–10 µm long.
REMARKS: Species is of little biostratigraphic value for the Pleistocene sediments of Site U1431D.

Scyphosphaera Lohmann, 1902
Vase-like coccoliths, generally large.
REMARKS: Species is of little biostratigraphic value.

Sphenolithus abies Deflandre in Deflandre and Fert, 1954
Similar to S. moriformis but more elevated and with cuspate outline.
REMARKS: Occurs as reworked specimens.

Sphenolithus moriformis (Brönnimann and Stradner, 1960) Bramlette and Wilcoxon, 1967a (Nannoturbella)
Generalized form, no spine, upper and lower parts of similar size.
REMARKS: Occurs as reworked specimens.

Syracosphaera histrica Kamptner, 1941
Oblong or irregularly elliptical with bright rim and central-area.
REMARKS: Species is of little biostratigraphic value for the Pleistocene sediments of Site U1431D.

Umbellosphaera tenuis (Kamptner, 1937) Paasche in Markali and Paasche, 1955 [Coccolithus]
Trumpet-shaped coccoliths, with ridges on the distal surface
REMARKS: Species is of little biostratigraphic value for the Pleistocene sediments of Site U1431D.

Umbilicosphaera sibogae var. sibogae (Weber-van Bosse, 1901) Gaarder, 1970 [Coccosphaera]
Central opening wide, proximal shield monocyclic, wider than the distal shield.
REMARKS: Species is of little biostratigraphic value for the Pleistocene sediments of Site U1431D.

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