7. CALCAREOUS NANNOFOSSILS RECOVERED BY LEG 114 IN THE SUBANTARCTIC SOUTH ATLANTIC OCEAN

Jason A. Crux

ABSTRACT

Ocean Drilling Program (ODP) Leg 114 recovered nannofossil-bearing sediments from seven sites in the high latitudes of the South Atlantic Ocean. Cretaceous sections were recovered from Sites 698 and 700, located on the Northeast Georgia Rise and its lower flanks, respectively. These contain distinctive high-latitude nannofossil floras similar to those from high-latitude areas of the Northern Hemisphere. Most of the biostratigraphic datums used to date the upper Campanian to Maestrichtian interval appear to lie at approximately the same level in both hemispheres. The FAD of Nephrolithus frequens is confirmed to be diachronous with an earlier occurrence in high latitudes. The LAD of Monomarginatus primus n. sp. also appears to be diachronous with a later LAD in the high latitudes of the Southern Hemisphere.

In the South Atlantic Ocean, which led to the thermal isolation of Antarctica and possibly the Pliocene, and possibly the Pliocene. Colder periods are inferred for the middle Eocene, most of the Oligocene, and the Miocene. Dramatic changes in the nannofossil floras of this section are of very low diversity, usually with fewer than eight species present.

Some stratigraphic ranges of importance are observed to be different in the high-latitude sections from those recorded from low-latitude areas. The LAD of Reticulofenestra bisecta, when calibrated by magnetostratigraphy, appears to occur earlier in Hole 699A (within Chron C6CR) than in Hole 703A and possibly Hole 704B and in other published accounts of lower latitude sites in the South Atlantic. The FAD of Nannopterinula julieni/Cristatula cristata appears to occur later in Hole 702B (chron C20R) than it does in other published accounts of lower latitude sites in the South Atlantic. Diachrony is also observed in the stratigraphic ranges of Chiasmolithus solitus and Chiasmolithus oamaruensis, although poor magnetostratigraphic results through the critical interval prevent confirmation of this.

Differences in the relative stratigraphic ranges of Isthmolithus recurvus and Cribrocentrum coenurum/C. reticulatum at Sites 699 and 703 are noted. These possibly suggest warmer surface waters on the eastern side (Site 703) of the middle to late Eocene South Atlantic than those on the western side (Site 699).

The diversities of the nannofossil floras and the presence of the warm-water genera Discoaster, Sphenolithus, Helicosphaera, and Amaurolithus reflect the changing surface water temperatures throughout the Cenozoic. Warmer periods are inferred for the late Paleocene to early middle Eocene, late middle Eocene to late Eocene, and possibly the Pliocene. Cold periods are inferred for the middle Eocene, most of the Oligocene, and the Miocene. Dramatic changes in the nannofossil floras of this section are thought to reflect a rapidly changing environment.

Monomarginatus primus, a new species from the Upper Cretaceous strata of Hole 700B, is described.

INTRODUCTION

Leg 114 of the Ocean Drilling Program (ODP) was the fourth cruise to the South Atlantic section of the Southern Ocean. This area is important to the understanding of the evolution of global climatic, glacial, and oceanographic systems since the late Mesozoic. Changes in these systems have probably been driven by tectonic events in the high southern latitudes, which led to the thermal isolation of Antarctica and the formation of the Antarctic ice sheets (LaBrecque et al., 1987). Twelve holes were drilled at seven sites during Leg 114, with the recovery of sediments from Santonian to Holocene age.

These sites are located on the Northeast Georgia Rise (Site 698), the lower flank of the Northeast Georgia Rise in the East Georgia Basin (Sites 699 and 700), Islas Orcadas Rise (Site 702), between the Islas Orcadas Rise and the Mid-Atlantic Ridge (Site 701), and the Meteor Rise (Sites 703 and 704) (Fig. 1).

Cretaceous sediments recovered from Sites 698 and 700 include an incomplete Santonian interval unconformably overlain by uppermost Campanian to Maestrichtian strata. The Maestrichtian section from Site 700 is the most complete recovered to date from the high-latitude Southern Ocean. An incomplete Cretaceous/Tertiary boundary section was recovered from Site 700 with the boundary lying between two cores. Incomplete sections of Paleocene strata were recovered from Sites 698, 699, 700, and 702. Almost complete Eocene sedimentary records were recovered from Sites 698, 699, 702, and 703, with partial records from Sites 698 and 701. Thick Oligocene to lowermost Miocene nannofossil-bearing sequences were recovered from Sites 699 and 703, and lower Oligocene to lowermost Miocene strata were recovered from Site 704. Site 701 penetrated a thick Oligocene section but this did not contain in-situ nannofossils.

Most of the Miocene to Quaternary sediments recovered are barren of nannofossils, as a result of their deposition below the carbonate compensation depth (CCD). The exception was the sediments from Site 704, which contain a distinctive high-latitude nannofossil record from the Oligocene to
Holocene, with only short hiatuses in the middle to upper Miocene and upper Pliocene. The recovered sedimentary records provide the greatest stratigraphic representation of the Upper Cretaceous to Cenozoic ever recovered from the Southern Ocean.

The aims of the present study are twofold. First, to construct a reliable biostratigraphic framework and to subdivide the fossiliferous sections penetrated using nannofossils. These subdivisions allow precise correlation of the cores recovered and help provide, together with other fossil groups and magnetostratigraphy, a chronostratigraphic framework within which sedimentologic, geochemical, and other studies can be set. The biostratigraphic zonation proposed in the present study is directly applicable to the sediments recovered. It is also possible to use the zonation to correlate with other sections from previous cruises to this area and lower latitudes.

The second aim of this study is to observe the biogeographic distribution of calcareous nannofossils in these high southern latitudes. Comparisons are made with the Cretaceous nannofossil floras of the Northern Hemisphere high-latitude areas. The presence and absence of Cenozoic low-latitude/warm-water nannofossil genera allow some observations about the changing of surface water temperatures through time. The study of high-latitude nannofossil floras is particularly interesting, because in these cold-water conditions the stratigraphic ranges of some biostratigraphic datum species are shortened or lengthened according to the temperature preferences of the living organisms. This provides us with valuable insights into the reliability of biostratigraphic data in lower latitudes.


Detailed nannofossil distribution tables are presented for Hole 699A and the Paleocene to Cretaceous of Hole 700B in Tables 1 through 4. These sections were chosen for more detailed study as they represent the best preserved and most complete sections recovered by ODP Leg 114. Together they form a nannofossil-bearing composite section from the lowermost Miocene to the Santonian.

METHODS

The nannofossil floras were examined with a transmitted-light microscope at magnifications between ×500 and ×1250. Crossed and partially crossed nicols were used routinely, and a gypsum first-order red plate was also used to identify certain species. All slides examined were smear slides, made using a toothpick and distilled water.

In constructing the range charts (Tables 1–4), the following codes were used to depict preservation of the nannofossils:

- **G = good**: well preserved, with only minor signs of dissolution and/or overgrowth.
- **M = moderate**: some overgrowth and/or dissolution of placoliths and discoasters.
- **P = poor**: strong overgrowth of discoasters and placoliths with secondary calcite. The assemblage is slightly biased toward solution-resistant forms.
- **VP = very poor**: severe overgrowth and/or dissolution. The assemblage is strongly biased toward solution-resistant forms.
- **E = etched nannofossils predominantly present.**
<table>
<thead>
<tr>
<th>Age</th>
<th>Eocene</th>
<th>Oligocene</th>
<th>Miocene-Quaternary</th>
<th>Hole 699A</th>
</tr>
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<tbody>
<tr>
<td></td>
<td>Quadricarinatus</td>
<td></td>
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<td></td>
<td>Chiasmolithus</td>
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<td></td>
<td>Callochammites</td>
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**Table 1. Distribution of Quaternary to Eocene calcareous nannofossils, Hole 699A.**

- **Abundance**
- **Preservation**

<table>
<thead>
<tr>
<th>Core, section, Zone</th>
<th>Sea level, internal term</th>
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**Legend:**
<table>
<thead>
<tr>
<th>Eocene</th>
<th>Palocene</th>
</tr>
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<tbody>
<tr>
<td>Discoaster</td>
<td>Multituberculata</td>
</tr>
<tr>
<td>47X-1, 124-127</td>
<td>5SX-CC, 1, 124-127</td>
</tr>
<tr>
<td>47X-1, 124-127</td>
<td>5SX-CC, 1, 124-127</td>
</tr>
<tr>
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<td>5SX-CC, 1, 124-127</td>
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<tr>
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<td>5SX-CC, 1, 124-127</td>
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<tr>
<td>52X-1, 124-127</td>
<td>5SX-CC, 1, 124-127</td>
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<td>5SX-CC, 1, 124-127</td>
</tr>
<tr>
<td>56X-1, 124-127</td>
<td>5SX-CC, 1, 124-127</td>
</tr>
</tbody>
</table>

**Note:** Abundance is characterized by VA = very abundant, A = abundant, C = common, F = few, R = rare, B = barren. For preservation, VP = very poor, P = poor, M = moderate, E = etched, O = overgrown.
Table 3. Distribution of Paleocene calcareous nannofossils, Hole 700B.

<table>
<thead>
<tr>
<th></th>
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</tr>
</thead>
<tbody>
<tr>
<td>Discocysta multiradiatus</td>
<td>26R-1, 40-61</td>
<td>A VP O</td>
<td>F F F</td>
<td>A R R R</td>
<td>R R R</td>
<td>R F C R</td>
<td>O = overgrown nannofossils predominantly present.</td>
<td>In the present study, similarities between the nannofossil floras of the high-latitude areas of both hemispheres are recognized. The experience gained in examining oil company sideway core samples from the North Sea is used in proposing a modified zonal scheme using species that occur in the two areas. Problems remain, however, with some species appearing to have different ranges in the Northern and Southern hemispheres. In addition, direct correlation with low-latitude areas is difficult, because no area is known that has both low- and high-latitude zonal indices present. The Cretaceous zonal scheme adopted here is graphically shown in Figure 2.</td>
<td>Lithastrinus floralis Zone</td>
</tr>
</tbody>
</table>
Table 4. Distribution of Cretaceous calcareous nannofossils, Hole 700B.

| Age     | Neophragmium | Eiffellithus | Monorhabdites | Monosaccus | Kamptnerius | Broinsonia | O. campanensis | Halobia | \( O. \) primus | \( O. \) flornlis |
|---------|--------------|-------------|----------------|------------|-------------|------------|----------------|--------|----------------|----------------|---|
| Zone    |              |             |                |            |             |            |                |        |                |                 |   |
| 37R-2, 31-32 |              |             |                |            |             |            |                |        |                |                 |   |
| 36R-CC  |              |             |                |            |             |            |                |        |                |                 |   |
| 35R-CC  |              |             |                |            |             |            |                |        |                |                 |   |
| 34R-CC  |              |             |                |            |             |            |                |        |                |                 |   |
| 33R-CC  |              |             |                |            |             |            |                |        |                |                 |   |
| 32R-CC  |              |             |                |            |             |            |                |        |                |                 |   |
| 31R-CC  |              |             |                |            |             |            |                |        |                |                 |   |
| 30R-CC  |              |             |                |            |             |            |                |        |                |                 |   |

Note: Abundance is characterized by VA = very abundant, A = abundant, C = common, F = few, R = rare, B = barren. For preservation VP = very poor, P = poor, M = moderate, E = etched, O = overgrown.

Helicothamnus anceps
Cretarhabdus angustiformis
Reinhardites antiquus
Reinhardites all. R. antipholosis/R. levii
Lucianorhabdus orcuttai
Watznaueria barnesae
Cylindricalithus bicornis
Watznaueria biporta
Orastrum campaniense
Lithographides camudensis
Lucianorhabdus cayaxii
Glaucolithus compactus
Cretarhabdus conicus
Biscutum coronum
Nephrithus corystis
Cretarhabdus creminatus
Pediscosphaera cretaea
Arkhangel'skella cymbiformis
Microhabdus decoratus
Biscutum dissipilis
Cribroosphera ehrenbergii
Biscutum elipsum
Brownsania enormis
Teichorhabdus ephesos
Eiffellithus eximius
Lithastrinus floridis
Nephrithus frequentus
Marathasterites furcatus
Tranosolithus gabolus
Quadrum gurtleri
Lithastrinus grailii
Lucianorhabdus spp.
Kamptnerius magnificus
Biscutum magnum
Micula sp.
Staurolithites michienensis
Micula murus
Gutierrezina obtliquum
Calciteles obscurus
Ahmuellerella octoradiata
Tranosolithus ornamentus
Watznaueria ovata
Brownsania parca
Monomarginatus pectinatus/M. quaternarius
Manivellina pennatoides
Serradiscusum primivium
Monomarginatus primus
Scarpanelia spp.
Lucianorhabdus scottus
Cylindricalithus sculptus
Brownsania signata
Zygophyllum bicuspidal
Zygophyllum spiralis
Micula staurophora
Pediscosphaera stoveri
Microhabdus stradneri
Thoracosphaera spp.
Helicothamnus trapezale
Eiffellithus turneifellii
Pervasithus varia
Remarks. Most authors consider the LAD of L. floralis to lie within the Santonian. Bukry (1969) and Crux (1980) found the range of L. floralis to extend into the Campanian, although the latter recorded a considerable reduction in its abundance within the Santonian. These younger records may be a result of reworking. For my concept of A. cymbiformis, see "Taxonomy" section.

Reference section. Samples 114-700B-54R-CC to 114-700B-52R-CC.

Seribiscutum primitivum Zone

Definition. From the LAD Lithastrinus floralis to the LAD of Seribiscutum primitivum.

Author. Base, Wise (1983); top, this study.

Age. Santonian–early Campanian.

Remarks. Wise (1983) recorded S. primitivum in sediments as young as early Campanian in the Falkland Plateau region. The last occurrence of this species at Site 700 is probably not its true extinction, as most of the Campanian is represented by a major hiatus. S. primitivum is reported to have preferred high-latitude/cooler surface-water areas (Wise and Wind, 1977). In the Northern Hemisphere, S. primitivum has not been reported above the Cenomanian (Forchheimer, 1968; Crux, 1982).

Reference section. Samples 114-700B-51R-CC to 114-700B-50R-CC.

Orastrum campanensis Zone

Definition. From the FAD to the LAD of Orastrum campanensis.

Author. Base, this study; top, Mortimer (1987).

Age. Campanian, possibly extending into the earliest Maestrichtian.

Remarks. Only Sample 114-700B-50R-2, 26–27 cm, was recovered from this zone in the present study. The lower part of the zone is not present as a result of the representation of most of the Campanian by a hiatus. Although poorly represented in the present study, this zone is easily defined in the Northern Hemisphere. The first occurrence of O. campanensis in the Northern Hemisphere is known to occur in the Campanian (Crux, 1982). It has also been reported from Santonian sediments by Mortimer (1987), but this was from oil company cuttings and sidewall core samples that were possibly contaminated. The last occurrence of this species was taken by Mortimer (1987) to mark the Campanian/Maestrichtian boundary, although this was based on North Sea well sections in which the boundary was not precisely known.

Reference section. Sample 114-700B-50R-2, 26–27 cm.

Eiffellithus eximius Zone

Definition. From the LAD of Orastrum campanensis to the LAD of Eiffellithus eximius.


Age. Early Maestrichtian, possibly extending into the late Campanian.

Remarks. Robaszynski et al. (1985) recorded the LAD of E. eximius in the lower Maestrichtian of the type Maestrichtian area. The LAD of E. eximius is found regularly in North Sea well sections to lie between the LADs of O. campanensis and Broinsonia parca. The age assigned to this zone in the present study contradicts evidence from the foraminiferal study of Huber (this volume). Huber recorded the LAD of Globigerinelloides impensus in Core 119-700B-43X, an event that he has previously correlated with magnetostratigraphic Chron C33N at Sites 689 and 690 on the Maud Rise. This would imply...
that the LAD of *E. eximius* must be older than Chron C33N at Site 700. Huber suggests that the LAD of *E. eximius* is as old as early Campanian, as it occurs in strata of reversed polarity. In this study I believe that a cosmopolitan datum such as the LAD of *E. eximius*, which has been widely recognized to lie near the Campanian/Maastrichtian boundary even in high-latitude areas, is more reliable than a local datum, which has yet not been extensively tested.

Further supporting evidence for a Maastrichtian for the *E. eximius* Zone is given by the presence of *Prediscosphaera stoveri*, which has an FAD in the upper Campanian (Perc-Nielsen, 1985). In addition, the occurrence of *Monomarginatus primus* in this zone also indicates a late Campanian or younger age, as this species has an FAD recorded in the upper Campanian in the Northern Hemisphere (Crux, 1982). This latter species did, however, prefer colder water environments and may have occurred earlier in the high-latitude South Atlantic.

**Reference section.** Samples 114-700B-49R-CC to 114-700B-47R-2, 93–94 cm.

**Broinsonia parca Zone**

**Definition.** From the LAD of *Eiffellithus eximius* to the LAD of *Broinsonia parca*.

**Author.** Base, Bukry and Bramlette (1970); top, Sissingh (1977).

**Age.** Early Maastrichtian.

**Remarks.** *B. parca* is only sporadic and rarely present toward the end of its stratigraphic range in the two holes that penetrated this zone.

**Reference section.** Samples 114-700B-46R-CC to 114-700B-46R-2, 69–70 cm.

**Monomarginatus primus Zone**

**Definition.** From the LAD of *Broinsonia parca* to the LAD of *Monomarginatus primus* n. sp.

**Author.** Base, Sissingh (1977); top, this study.

**Age.** Early Maastrichtian.

**Remarks.** *M. primus* n. sp. (*Staurolithites ellipticus* sensu Crux, 1982) has been recorded only from the upper Campanian in the Northern Hemisphere. In North Sea well sections, its last occurrence is located at approximately the same level as that of *O. campanensis*. It thus appears to have a longer range in the Southern Hemisphere.

**Reference section.** Samples 114-700B-45R-CC to 114-700B-43R-CC.

**Kampferiella magnificus Zone**

**Definition.** From the LAD of *Monomarginatus primus* n. sp. to the FAD of *Nephrolithus frequens*.

**Author.** Base, this study; top, Čepek and Hay (1969).

**Age.** Early? to late Maastrichtian.

**Reference section.** Samples 114-700B-43R-2, 32–33 cm, to 114-700B-41R-CC.

**Reinhardtites levis Zone**

**Definition.** From the FAD of *Nephrolithus frequens* to the LAD of *Reinhardtites levis*.

**Author.** Base, Čepek and Hay (1969); top, Sissingh (1977).

**Age.** Late Maastrichtian.

**Remarks.** Sissingh (1977) considered *N. frequens* and *R. levis* to be mutually exclusive and defined the *Arkhangelskia cymbiformis* Zone as an interval between the LAD of *R. levis* and the FAD of *N. frequens*. In both the Northeast Georgia Rise and the Falkland Plateau areas and in the North Sea area, the FAD of *Nephrolithus frequens* lies stratigraphically below the LAD of *Reinhardtites levis* (Wind, 1979; this study). This order of events was also noted by van Heck (1979b) in the type Maestrichtian, but she ascribed the co-occurrence of the two species to the reworking of *R. levis* into younger sediments. I consider it probable that *N. frequens*, a species known to prefer high-latitude colder water areas, has an earlier first occurrence in these latitudes, as reported by Wind (1979). The LAD of *R. levis* lies within Chron C31R.

**Reference section.** Samples 114-700B-39R-2, 32–33 cm.

**Biscutum magnum Zone**

**Definition.** From the LAD of *Reinhardtites levis* to the LAD of *Biscutum magnum*.

**Author.** Base, Sissingh (1977); top, Wind (1979).

**Age.** Late Maastrichtian.

**Remarks.** This zone can also be recognized in the North Sea area of the Northern Hemisphere. The LAD of *B. magnum* lies within Chron C31R.

**Reference section.** Sample 114-700B-39R-2, 32–33 cm.

**Nephrolithus frequens Zone**

**Definition.** From the LAD of *Biscutum magnum* to the LAD of *Nephrolithus frequens*.

**Author.** Base, Wind (1979); top, Čepek and Hay (1969).

**Age.** Late Maastrichtian.

**Remarks.** The LAD of *N. frequens* occurs in Chron C30N; its uppermost range is removed by a hiatus.

**Reference section.** Samples 114-700B-38R-CC to 114-700B-37R-2, 31–32 cm.

**Cenozoic**

The most widely used Cenozoic calcareous nannofossil zonal schemes (Martini, 1971; Bukry, 1973, 1975) are only partially applicable to the sedimentary sequences of the subantarctic South Atlantic. Progressive deterioration of the climate through the Cenozoic led to increased provincialism among the highly stenothermal calcareous nannoplankton. These algae respond quickly to changes in the thermal structure of the water masses. This is seen dramatically in the Miocene to Pleistocene of the subantarctic South Atlantic, where low diversity, endemic species, and poor preservation prevent the recognition of many biostratigraphic datums used to subdivide strata of low latitudes.

The zonation scheme employed for the Paleogene to earliest Miocene is largely based on that of Martini (1971), which in turn uses zones proposed by previous authors. Where datums used by Martini (1971) are not present, additional datums were used from the zonal schemes of Edwards (1971), Wise and Constans (1976), Romein (1979), Varol (1989), and Wise (1983). For the authorship of individual zones see Wise (1983), Bukry-Nielsen (1985), and Varol (1989). Datums not previously included in *Cenozoic* are those of *Nephrolithus frequens* Zone.

**Reference section.** Sample 704-41R-CC to 704-40R-CC.

**Prinsius dimorphosus Zone**

**Definition.** From the FAD to the last abundant occurrence of *Prinsius dimorphosus* (76.1 to 64.6–43.8 Ma, top in Chron C27N).
<table>
<thead>
<tr>
<th>Age</th>
<th>Zone</th>
<th>Subzone</th>
<th>Datum level</th>
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<td>C. abisectus</td>
<td>LAD C. abisectus</td>
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<td>late Oligocene</td>
<td>R. bisecta</td>
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<td>N. fulgens</td>
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<td>P. martinii</td>
<td>Last abundant P. martinii</td>
</tr>
<tr>
<td></td>
<td></td>
<td>P. dimorphosus</td>
<td>Last abundant P. dimorphosus</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>FAD P. dimorphosus</td>
</tr>
</tbody>
</table>

**Figure 3.** Paleocene to lower Miocene calcareous nannofossil zones used in this chapter.

**Age.** Early Paleocene.

**Remarks.** This zone is easily recognized in the North Sea region of the Northern Hemisphere.

**Reference section.** Samples 114-700B-33R-CC to 114-700B-32R-2, 83-85 cm.

**Prinsius maritini Zone**

**Definition.** From the last abundant occurrence of *Prinsius dimorphosus* to the last abundant occurrence of *Prinsius martini* (64.6–63.8 to 62.8 Ma, top in Chron C26R).

**Age.** Early Paleocene.

**Remarks.** This zone is easily recognized in the North Sea region.

**Reference section.** Samples 114-700B-33R-CC to 114-700B-32R-2, 83–85 cm.

**Prinsius bisulcus Zone**

**Definition.** From the last abundant occurrence of *Prinsius martini* to the FAD of the *Fasciculithus tympaniformis* group (62.7 Ma, within Chron C26R).

**Age.** Early to late Paleocene.

**Reference section.** Sample 114-700B-31R-CC.

**Fasciculithus tympaniformis group Zone**

**Definition.** From the FAD of the *Fasciculithus tympaniformis* group to the FAD of *Heliolithus kleinpellii*.
<table>
<thead>
<tr>
<th>Age</th>
<th>Marais (1971) zone</th>
<th>Hole 698A</th>
<th>Hole 699A</th>
<th>Hole 700B</th>
<th>Hole 701C</th>
<th>Hole 702B</th>
<th>Hole 703A</th>
<th>Hole 704B</th>
<th>Zone, this study</th>
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<tr>
<td>Oligocene</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>NP 23-24</td>
<td></td>
<td>27X-6, 60-61 cm</td>
<td>28X, CC</td>
<td>31X-1, 59-60 cm</td>
<td>43X-4, 34-35 cm</td>
<td>12H-6, 21-22 cm</td>
<td></td>
<td>R. daviesii</td>
<td></td>
</tr>
<tr>
<td>NP 19-21</td>
<td></td>
<td>31X-2, 135-136 cm</td>
<td>33X, CC</td>
<td>34X, CC</td>
<td>47X-2, 30-31 cm</td>
<td>13R-2, 74-75 cm</td>
<td>15R-2, 95-96 cm</td>
<td>C. subdistichus</td>
<td></td>
</tr>
<tr>
<td>NP 18</td>
<td></td>
<td>36X-2, 60-61 cm</td>
<td>47X, CC</td>
<td>3X, CC</td>
<td>15X-1, 20-21 cm</td>
<td>20X-2, 30-31 cm</td>
<td></td>
<td>I. recurvus</td>
<td></td>
</tr>
<tr>
<td>NP 14</td>
<td></td>
<td>39X-1, 66-61 cm</td>
<td>3R-1, 59-60 cm</td>
<td>4R, CC</td>
<td>6X-2, 69-70 cm</td>
<td>22X, CC</td>
<td></td>
<td>R. oamaruensis</td>
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</tr>
<tr>
<td>NP 13</td>
<td></td>
<td>42X-4, 4-5 cm</td>
<td>5R-2, 70-71 cm</td>
<td>6X-2, 59-60 cm</td>
<td>9X, CC</td>
<td>6X-2, 59-60 cm</td>
<td>22X, CC</td>
<td></td>
<td>J. recurvus</td>
</tr>
<tr>
<td>NP 10-12</td>
<td></td>
<td>42X-7, 4-5 cm</td>
<td>45X-1, 135-140 cm</td>
<td>49X-1, 27-28 cm</td>
<td>4X, CC</td>
<td>10X-5, 20-21 cm</td>
<td>23X-1, 20-21 cm</td>
<td>N. fulgens</td>
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</tr>
<tr>
<td>NP 9</td>
<td></td>
<td>1R, CC</td>
<td>3R-1, 26-21 cm</td>
<td>12R, CC</td>
<td>50X-1, 175-76 cm</td>
<td>15X-2, 60-61 cm</td>
<td>33X, CC</td>
<td></td>
<td></td>
</tr>
<tr>
<td>NP 8</td>
<td></td>
<td>49X-3, 55-56 cm</td>
<td>3R, CC</td>
<td>49X-3, 55-56 cm</td>
<td>3R, CC</td>
<td>50X-1, 175-76 cm</td>
<td>21X, CC</td>
<td></td>
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</tr>
<tr>
<td>NP 7</td>
<td></td>
<td>50X-4, 98-99 cm</td>
<td>3R, CC</td>
<td>50X-4, 98-99 cm</td>
<td>3R, CC</td>
<td>50X-1, 175-76 cm</td>
<td>21X, CC</td>
<td></td>
<td></td>
</tr>
<tr>
<td>NP 5</td>
<td></td>
<td>51X, CC</td>
<td>51X, CC</td>
<td>51X, CC</td>
<td>22X, CC</td>
<td>22X, CC</td>
<td>22X, CC</td>
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<tr>
<td>NP 4</td>
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<td>53X, CC</td>
<td>53X, CC</td>
<td>53X, CC</td>
<td>22X, CC</td>
<td>22X, CC</td>
<td>22X, CC</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Palaeocene</td>
<td></td>
<td>54X-2, 135-136 cm</td>
<td>10R, CC</td>
<td>20R-2, 106-107 cm</td>
<td>27X-1, 6-7 cm</td>
<td>27X, CC</td>
<td>27X, CC</td>
<td></td>
<td>D. multiradiatus</td>
</tr>
<tr>
<td>NP 9</td>
<td></td>
<td>54X-2, 135-136 cm</td>
<td>10R, CC</td>
<td>20R-2, 106-107 cm</td>
<td>27X-1, 6-7 cm</td>
<td>27X, CC</td>
<td>27X, CC</td>
<td></td>
<td>D. multiradiatus</td>
</tr>
<tr>
<td>NP 8</td>
<td></td>
<td>10R-2, 35-36 cm</td>
<td>56X-2, 40-41 cm</td>
<td>27X-1, 6-7 cm</td>
<td>27X, CC</td>
<td>27X, CC</td>
<td>27X, CC</td>
<td></td>
<td>D. multiradiatus</td>
</tr>
<tr>
<td>NP 5</td>
<td></td>
<td>12R, CC</td>
<td>32R-2, 83-85 cm</td>
<td>33R, CC</td>
<td>32X, CC</td>
<td>32X, CC</td>
<td>32X, CC</td>
<td></td>
<td>P. oamaruensis</td>
</tr>
<tr>
<td>NP 4</td>
<td></td>
<td>31R, CC</td>
<td>34R-2, 105-106 cm</td>
<td>35R, CC</td>
<td>32X, CC</td>
<td>32X, CC</td>
<td>32X, CC</td>
<td></td>
<td>D. saipaπensis</td>
</tr>
<tr>
<td>NP 2-3</td>
<td></td>
<td>32R-2, 83-85 cm</td>
<td>33R, CC</td>
<td>34R-2, 105-106 cm</td>
<td>35R, CC</td>
<td>32X, CC</td>
<td>32X, CC</td>
<td></td>
<td>D. saipaπensis</td>
</tr>
</tbody>
</table>

Figure 4. Paleocene to lower Miocene zonal assignments for Sites 698–704.
**Heliolithus kleinpellii Zone**

**Definition.** From the FAD of Heliolithus kleinpellii to the FAD of Heliolithus riedelli.  
**Age.** Late Paleocene.  
**Remarks.** This zone includes both the Heliolithus kleinpellii and Discoaster mohleri Zones of the Martini (1971) scheme. It is not possible to divide the zones because of the absence of D. mohleri in the sections studied. The reliability of the FAD of H. kleinpellii as a biostratigraphic datum is questioned by Wei and Wise (1989), who recognized some diachronieity and difficulties in differentiating it from other species of Heliolithus.

**Heliolithus riedelli Zone**

**Definition.** From the FAD of Heliolithus riedelli to the FAD of Discoaster multiradiatus.  
**Age.** Late Paleocene.  
**Remarks.** Both H. kleinpellii and H. riedelli are very rare in the sections studied, and thus these zonal assignments should only be regarded as approximate.

**Discoaster multiradiatus Zone**

**Definition.** From the FAD of Discoaster multiradiatus to the LAD of the Fasciculithus tympaniformis group.  
**Age.** Late Paleocene to earliest Eocene.  
**Remarks.** The absence of Tribrachiatas bramlettei in the sections studied does not allow recognition of the upper boundary of Martini’s (1971) Discoaster multiradiatus Zone. The substitution of the LAD of the F. tympaniformis group for the FAD of T. bramlettei extends this zone into the base of the Eocene.

**Tribrachiatas orthostylus Zone**

**Definition.** From the LAD of the Fasciculithus tympaniformis group to the LAD of Tribrachiatas orthostylus.  
**Age.** Early Eocene.  
**Remarks.** The absence of Tribrachiatas contortus and the rare occurrence of Discoaster lodoensis in the lower part of its range in this area prevents the recognition of Martini’s (1971) Tribrachiatas contortus and Discoaster binodosus Zones. These are grouped here with his overlying Tribrachiatas orthostylus Zone. Wei and Wise (1989) reported that the LAD of T. orthostylus was an unreliable biostratigraphic datum. It is used in the present study, however, as no other better biostratigraphic events occur to divide this interval.

**Discoaster lodoensis Zone**

**Definition.** From the LAD of Tribrachiatas orthostylus to the FAD of Discoaster sublodoensis.  
**Age.** Middle Eocene.  
**Remarks.** The absence of Rhabdosphaera gladius in the sections studied does not allow the recognition of the upper boundary of Martini’s (1971) Nannotetra fulgens (Chipsiragm-maltihus alatus) Zone. The FAD of R. bisecta used by Edwards (1971) to divide this interval appears to be a reliable datum in high-latitude sections. The FAD of R. bisecta/R. aff. R. bisecta lies within Chrons C19R-C18R in Hole 702B (Clement and Hailwood, this volume) with an age of 43.4 to 44.2 Ma.

**Chiasmolithus solituis Zone**

**Definition.** From the FAD of Reticulofenestra bisecta/R. aff. R. bisecta to the LAD of Chiasmolithus solituis.  
**Age.** Middle Eocene.  
**Remarks.** This zone is equivalent to the upper part of Martini’s (1971) Discoaster tanii nuddfer Zone.

**Discoaster saipanensis Zone**

**Definition.** From the LAD of Chiasmolithus solituis to the FAD of Chiasmolithus oamaruensis.  
**Age.** Middle Eocene.  
**Remarks.** This zone was not recognized in the present study. Its absence could be attributable to one of three causes: (1) too wide sample spacing, (2) a regional hiatus, or (3) the extension of the ranges of either or both of the two zonal indices. Perch-Nielsen (1985) and Wei and Wise (1989) reported the LAD of C. solituis to be diachronous, which may partially explain the absence of this zone.

**Chiasmolithus oamaruensis Zone**

**Definition.** From the FAD of Chiasmolithus oamaruensis to the FAD of Isthmolithus recurvus.  
**Age.** Late Eocene.  
**Remarks.** This zone is equivalent to Martini’s (1971) Isthmolithus oamaruensis Zone. The FAD of I. recurvus of this study is tentatively divided into the following three subzones.

**Isthmolithus recurvus Subzone**

**Definition.** From the FAD of Isthmolithus recurvus to the last common occurrence of Clausicoccus subdistichus.  
**Age.** Late Eocene to early Oligocene.  
**Remarks.** This zone is equivalent to Martini’s (1971) Isthmolithus recurvus, Sphenolithus pseudoradians, and most of the Clausicoccus (Ericsonia?) subdistichus Zones. These zones cannot be differentiated in the studied sections because of the absence and scarcity of the zonal indices. The I. recurvus Zone of this study is tentatively divided into the following three subzones.

**Isthmolithus recurvus Subzone**

**Definition.** From the FAD of Isthmolithus recurvus to the LAD of Discoaster saipanensis.  
**Age.** Late Eocene.  
**Remarks.** The rarity of D. saipanensis in the studied sections has possibly resulted in this species having an earlier LAD than in lower latitudes.

**Reticulofenestra oamaruensis Subzone**

**Definition.** From the LAD of Discoaster saipanensis to the LAD of Reticulofenestra oamaruensis.  
**Age.** Late Eocene to early Oligocene?  
**Remarks.** For the definition of the Chiasmolithus oamaruensis Zone.
Eocene

20-21 cm, indicates the presence of the Zone in sublodoensis in Samples 114-698A-1R-CC to 114-698A-3R-1, indicates the presence of the D. sublodoensis Zone in this interval. Sample 114-698A-3R-CC lies between the FAD of D. sublodoensis and the LAD of Tribrachiatus orthostylus; these events define the D. lodoensis Zone. The occurrence of T. orthostylus in Samples 114-698A-4R-CC to 114-698A-8R-CC indicates the presence of the T. orthostylus Zone.

Paleocene

The co-occurrence of the Fasciculithus tymaniformis group and Discoaster multidatus in Samples 114-698A-9R-1, 52–53 cm, to 114-698A-10R-1, 66–67 cm, indicates the presence of the D. multidatus Zone. The Eocene/Paleocene boundary lies very close to the top of this Zone. The absence of D. multidatus and the presence of Heliolithus riedelli in Samples 114-698A-10R-2, 35–36 cm, to 114-698A-10R-CC indicates the presence of the H. riedelli Zone. Samples between 114-698A-11R-CC and 114-698A-12R-CC continue to contain F. tymaniformis and are thus assigned to the Heliolithus kleinpellii to F. tymaniformis group Zones. The absence of H. kleinpellii from this interval is probably due to poor sample recovery and does not necessarily indicate a hiatus. A similar explanation for the absence of the Prinsius bisulcus and Prinsius martinii Zones is also probable. The abundant occurrence of Prinsius dimorphis in Sample 114-698A-13R-CC indicates the presence of the P. dimorphis Zone.

Cretaceous

A stratigraphic break probably occurs between Samples 114-698A-13R-CC and 114-698A-14R-1, 1–2 cm. This break separates lower Paleocene from upper Maestrichtian strata. The presence of Nephrolithus frequens between Samples 114-698A-14R-1, 1–2 cm, and 114-698A-15R-CC indicates the presence of the N. frequens Zone. The co-occurrence of N. frequens and Reinhardites lev is in Samples 114-698A-16R-CC and 114-698A-17R-CC indicates the presence of the Reinhardites lev Zone. The failure to recognize the Biscutum magnum Zone between the N. frequens and R. lev is Zones is probably due to insufficient sampling rather than the presence of a hiatus. A similar explanation is likely for the failure to recognize the Monomarginatus primus Zone below the Kumpinerus magnificus Zone at Sample 114-698A-20R-1, 48–49 cm. The presence of Bronsionia parca in Sample 114-698A-21R-CC indicates the presence of the B. parca Zone.

Site 699

Site 699 (51°32.537'S, 30°40.619'W; water depth 3705.5 m) is located on the northeastern slope of the Northeast Georgia Rise (Fig. 1). This site was drilled to obtain a continuously cored sequence of Late Cretaceous to Neogene age sediments, recording the history of deep-water communication between the Weddell and Georgia basins and the South Atlantic Basin. Other objectives were to document the development of the Antarctic Circumpolar Current and the southern high-latitude biosiliceous province.

The sediments recovered from Site 699 are predominantly pelagic in origin. The early Miocene to Quaternary interval (lithostratigraphic Unit I) consists of siliceous and clayey siliceous sediments. Nannofossils are recorded only from Sample 114-699A-2H-4, 5–6 cm, which contains a rare low-diversity assemblage. Nannofossils become common to abundant at Sample 114-699A-9H-CC and below (approximately lithostratigraphic Unit II), with nannofossil siliceous oozes down to Sample 114-699A-25X-CC; below this a short interval of graded gravel (lithostratigraphic Unit III) was recovered. This probably originated upheole and fell into the hole. Below the gravel, there is a further decrease in siliceous biogenic material downhill and an increase in nannofossil chucks, which become increasingly micritic (lithostratigraphic Units IV and V). At the bottom of Hole 699A, below Section
114-699A-54X-1, 0 cm, occurs a zeolite-bearing claystone with clay-bearing micritic nannofossil chalks (lithostratigraphic Unit VI). Nannofossils are poorly preserved throughout Hole 699A. Above Sample 114-699A-53X-CC, they are predominantly etched, and below this level they are predominantly overgrown with secondary calcite.

**Quaternary to Miocene**

Samples 114-699A-1H-2, 60–61 cm, to 114-699A-2H-2, 15–16 cm, are barren of nannofossils. Only Sample 114-699A-2H-4, 5–6 cm, yielded a sparse nannoflora, which includes Gephyrocapsa spp., indicating a Holocene to late Pliocene age. The interval between Samples 114-699A-3H-1, 60–61 cm, and 114-699A-9H-2, 5–6 cm, is assigned to the Cyclicargolithus abisectus Zone, whereas the interval between Samples 114-699A-9H-CC and 114-699A-11H-5, 8–9 cm, is assigned to the Cyclicargolithus abisectus Zone based on the presence of the nominate taxon.

**Oligocene**

The interval between Samples 114-699A-11H-CC and 114-699A-15H-2, 22–23 cm, is assigned to the Reticulofenestra bisecta Zone, whereas the interval between Samples 114-699A-15H-4, 23–24 cm, and 114-699A-27X-4, 115–116 cm, is assigned to the Chiasmolithus altus Zone. The Reticulofenestra daviesii Zone is present between Samples 114-699A-27X-6, 60–61 cm, and 114-699A-28X-CC; this is recognized by the first downhole occurrence of Reticulofenestra umbilicus. Also present in this interval is the stratigraphically significant species Isthmolithus recurvus. The presence of common Clausiococcus subdistichus in Sample 114-699A-29X-1, 59–60 cm, and below indicates the penetration of the C. subdistichus Subzone of the I. recurvus Zone. This is present down to Sample 114-699A-31X-2, 135–136 cm.

**Eocene**

The Oligocene/Eocene boundary probably lies somewhere in the Reticulofenestra oamaruensis Subzone of the I. recurvus Zone. The presence of this subzone is indicated between Samples 114-699A-31X-5, 138–139 cm, and 114-699A-33X-CC by the presence of the nominate taxon. The interval between Samples 114-699A-34X-CC and 114-699A-35X-CC is assigned to the I. recurvus Subzone of the I. recurvus Zone, based on the co-occurrence of I. recurvus and Discoaster saipanensis.

The interval between the FAD of I. recurvus and the FAD of Chiasmolithus oamaruensis defines the C. oamaruensis Zone. This is present between Samples 114-699A-36X-2, 60–61 cm, and 114-699A-38X-CC. A possible stratigraphic break occurs between the base of this zone and the top of the underlying Chiasmolithus solitus Zone, with the D. saipanensis Zone absent. This is discussed more fully under “Paleoenvironment and Provincialism.”

The C. solitus Zone is present between Samples 114-699A-39X-1, 60–61 cm, and 114-699A-42X-4, 4–5 cm, as indicated by the co-occurrence of C. solitus and Reticulofenestra bisecta. The Nannotetrina fulgens Zone is present between Samples 114-699A-42X-7, 4–5 cm, and 114-699A-45X-1, 139–140 cm. Samples 114-699A-45X-6, 139–140 cm, to 114-699A-49X-3, 55–56 cm, are assigned to the Discoaster sublodoensis Zone, which is the interval between the FAD of N. fulgens/Nannotetrina cristata and the FAD of D. sublodoensis. The Discoaster lodoensis Zone occurs between Samples 114-699A-49X-CC and 114-699A-50X-4, 98–99 cm. The presence of Tribrachiatus orthostylus between Samples 114-699A-51X-CC and 114-699A-53X-CC indicates the presence of the T. orthostylus Zone.

**Paleocene**

The co-occurrence of D. multiradiatus and the Fasciculithus tympaniformis group between Samples 114-699A-54X-2, 135–136 cm, and 114-699A-56X-2, 40–41 cm, indicates the presence of the Discoaster multiradiatus Zone. The Eocene/Paleocene boundary lies within this zone near the top.

**Site 700**

Site 700 (51°31.992'S, 30°16.697'W; water depth 3601 m) is in the western region of the East Georgia Basin on the northeastern slope of the Northeast Georgia Rise. The pur-

<table>
<thead>
<tr>
<th>Age</th>
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<th>Hole 700B</th>
</tr>
</thead>
<tbody>
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</tr>
<tr>
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<td>41R, CC</td>
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<td>20R-1-48.49 cm</td>
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Figure 5. Cretaceous zonal assignments for Sites 698–700.
pose of drilling Site 700 was to reach the deeper strata and basement not achieved by drilling Hole 699A. The objective was to obtain an Upper Cretaceous–Paleogene section recording the role of the Georgia Basin as an avenue for deep-water communication between the Weddell Sea and the South Atlantic. An additional objective was to further constrain the nature, age, and subsidence history of the Northeast Georgia Rise.

The sediments recovered from Hole 700B were divided into five lithostratigraphic units: Unit I, Quaternary–Upper Pliocene diatom ooze; Unit II, middle Eocene nannofossil ooze; Unit III, middle to lower Eocene nannofossil chalk that passes into micritic nannofossil chalk downhole; Unit IV, Paleocene micritic, indurated nannofossil chalk; and Unit V, lower Paleocene to Santonian micritic–nannofossil-bearing limestone that passes down into alternating micritic limestones and clay-bearing or clayey micritic limestones. These lithologic divisions are reflected in the preservation of the nannofossils. Unit I is barren of in-situ nannofossil flora, and Units II through V contain rare to abundant nannofossil floras, most of which are very poorly preserved with considerable secondary calcite overgrowth. There is some variation within the general category of very poor preservation, with some intervals in the Upper Cretaceous section containing the solution-resistant species *Micula staurophora* in abundant numbers. This probably indicates considerable selective dissolution of the original flora. In contrast, there is a marked improvement in preservation of the nannofossils from Cores 114-700B-50R to 114-700B-53R, where the increased clay content of the sediments has protected the nannofloras from the worst effects of dissolution.

### Quaternary to Miocene

Sample 114-700B-1W-CC is barren of in-situ nannofossils; isolated reworked Paleogene species were recorded.

### Eocene

The nannofossil floras from the interval between Samples 114-700B-3R-1, 59–60 cm, and 114-700B-4R-CC are assigned to the *Chiasmolithus solitus* Zone. The presence of this zone is indicated by the co-occurrence of *C. solitus* and *Reticulofenestra bisecta*. The interval between Samples 114-700B-5R-2, 70–71 cm, and 114-700B-11R-CC is assigned to the *Nannoteutinella fulgens* Zone, with the nominate zone boundary occurring between Samples 114-700B-7R-2, 33–34 cm, and 114-700B-11R-CC. The interval between the FAD of *N. fulgens/N. cristata* and the FAD of *Discosta beta subdolobensis* defines the *D. subdolobensis* Zone between Samples 114-700B-12R-CC and 114-700B-16R-CC. The interval between Samples 114-700B-18R-1, 80–81 cm, and 114-700B-18R-CC is assigned to the *Discosta beta lodoensis* Zone, as it lies between the FAD of *D. subdolobensis* and the LAD of *Trirachia multiradiata* orthostylus. The interval between Samples 114-700B-20R-2, 106–107 cm, and 114-700B-24R-CC is assigned to the *T. orthosystus* Zone. The nominate taxon is present in all samples examined except the bottom one.

### Paleocene

The co-occurrence of *Discosta multiradiatus* and the *Fasculicitus tympaniformis* group indicates the presence of the *D. multiradiatus* Zone between Samples 114-700B-26R-1, 40–41 cm, and 114-700B-26R-CC. The interval between the FAD of *D. multiradiatus* and the FAD of *Helolithus riedelli* defines the *H. riedelli* Zone, which lies between Samples 114-700B-27R-1, 30–31 cm, and 114-700B-29R-2, 122–124 cm. Only Sample 114-700B-29R-CC contains *Helolithus kleinpellii* and is assigned to the *H. kleinpellii* Zone. The interval between Samples 114-700B-30R-2, 120–121 cm, and 114-700B-31R-5, 115–116 cm, continues to contain the *F. tympaniformis* group, but without *H. kleinpellii*, and is thus assigned to the *F. tympaniformis* group Zone. Only Sample 114-700B-31R-CC lies in the interval between the FAD of the *F. tympaniformis* group and the last abundant occurrence of *Prinsius martini*. It is assigned to the *Prinsius bisculus* Zone. The interval between the last abundant occurrence of *P. martini* and *Prinsius dimorphosus* defines the *P. martini* Zone, which is located between Samples 114-700B-32R-2, 83–85 cm, and 114-700B-33R-CC. The abundant occurrence of *P. dimorphosus* between Samples 114-700B-34R-2, 105–106 cm, and 114-700B-36R-CC indicates the presence of the *P. dimorphosus* Zone.

### Cretaceous

A sample examined from the top of Core 114-700B-37R as part of the shipboard study contained an exclusively Cretaceous nannoflora; thus, the Cretaceous/Tertiary boundary is considered to lie within the break between Cores 114-700B-36R and 114-700B-37R. The occurrence of *Neophyliolithus frequens* and the absence of *Biscutum magnus* between Samples 114-700B-37R-2, 31–32 cm, and 114-700B-38R-CC indicates the presence of the *N. frequens* Zone. The occurrence of *B. magnus* together with *N. frequens* in Sample 114-700B-39R-2, 32–33 cm, indicates the presence of the *B. magnus* Zone. The *Reinhardtites levis* Zone is present between Samples 114-700B-39R-CC and 114-700B-41R-2, 23–24 cm, as indicated by the co-occurrence of *R. levis* and *N. frequens*. The interval between the FAD of *N. frequens* and the LAD of *Monomarginatus primus* defines the *Kamptnerius magnificus* Zone between Samples 114-700B-41R-CC and 114-700B-43R-2, 32–33 cm. The L. primus Zone is present between Samples 114-700B-44R-CC and 114-700B-45R-2, 61–62 cm; the *Borinsonia parca* Zone is present between Samples 114-700B-46R-2, 69–70 cm, and 114-700B-46R-CC; and the *Eifellithus eximius* Zone is present between Samples 114-700B-47R-2, 93–94 cm, and 114-700B-49R-CC. The tops of all three of these zones are defined by the LADs of their nominate taxa. Only Sample 114-700B-50R-2, 26–27 cm, contains *Orastra campanensis*, and it is assigned to the *O. campanensis* Zone.

A stratigraphic break possibly representing most of the Campanian and possibly some of the Santonian separates Samples 114-700B-50R-2, 26–27 cm, and 114-700B-50R-CC. The presence of *Seribiscutum primitivum* between Samples 114-700B-50R-CC and 114-700B-51R-CC indicates the presence of the *S. primitivum* Zone. The co-occurrence of *Lithastrinus floralis* and *Arkhangelskiella cymbiformis* between Samples 114-700B-52R-CC and 114-700B-54R-CC indicates the presence of the *L. floralis* Zone.

### Site 701

Site 701 (51°59.07’S, 23°12.73’W; water depth 4636.7 m) is on the western flank of the Mid-Atlantic Ridge about 160 km east of the Islas Orcadas Rise. The major objective of this site was to obtain a continuous sediment record of the development of an oceanic gateway for deep circulation between the South Atlantic and the Weddell Basin.

The sedimentary sequence penetrated at Site 701 was divided into three lithostratigraphic units. Unit I is predominantly a diatom ooze with varying additional clay, mud, and volcanic ash; Unit II is a diatom-bearing mud with less volcanic ash than above; and Unit III consists of indurated nannofossil chalk. Basalt, either from a sillon or oceanic basement, was recovered from the last core barrel. Only Samples 114-700A-4H-4, 12–13 cm, and 114-701C-25X-3,
**CALCAREOUS NANNOFOSSILS**

14–95 cm, were recorded with *in-situ* nannofossils in Unit I. Strongly etched nannofossils occur sporadically throughout Unit II, whereas in Unit III (below Sample 114-701C-49X-6, 29–30 cm), the nannofossils are overgrown with secondary calcite.

**Quaternary to Miocene**

Only Sample 114-701A-4H-4, 12–13 cm, from the Quaternary to Miocene strata in Hole 701A was found to contain nannofossils. The co-occurrence of *Pseudoemiliania lacunosa* and *Gephyrocapsa* spp. indicates a middle Pleistocene to Pliocene age. All other samples examined from Hole 701A were barren of nannofossils. Monospecific nannofossil floras of *Reticulofenestra perplexa* are present in Samples 114-701C-25X-3, 94–95 cm, to 114-701C-26X-CC. These indicate a late middle Miocene age.

**Oligocene to Eocene**

The interval from Samples 114-701C-27X-CC to 114-701C-43X-2, 34–35 cm, contains only reworked nannofossils. Strongly etched *in-situ* nannofossil floras occur sporadically in the interval between Samples 114-701C-43X-4, 34–35 cm, and 114-701C-47X-2, 30–31 cm. These nannofossil floras include *Isthmolithus recurvus*, *Reticulofenestra bisecta*, *Reticulofenestra umbilicus*, *Chiasmolithus altus*, *Clausocystis subdistichus* (only in Sample 114-701C-44X-2, 130–131 cm), *Cyclococcolithus floridanus*, *Blackites spinosus*, and *Chiasmolithus* sp. This interval is assigned to the undifferentiated *I. recurvus Zone* on the basis of the co-occurrence of *I. recurvus* and *C. subdistichus*.

Below this interval, *I. recurvus* is absent from the sparse nannofossil and *Chiasmolithus oamaruensis*, the FAD of which defines the base of the *C. oamaruensis Zone*, is present in samples from the interval between Samples 114-701C-47X-CC and 114-701C-48X-CC.

A stratigraphic break probably occurs between Samples 114-701C-48X-CC and 114-701C-49X-1, 27–28 cm, where there is also a change in the nannofossil floras from predominantly etched forms to strongly overgrown forms. The occurrence of *Nannotetrista fulgens*/Nannotetrista cristata in Samples 114-701C-49X-1, 27–28, and 114-701C-49X-3, 30–31 cm, indicates the presence of the *N. fulgens* Zone. Sample 114-701C-50X-1, 75–76 cm, is assigned to the *Discocastor sublodoensis Zone*. It has a nannofossil flora similar to that recorded in the overlying interval, but without *N. fulgens*/N. cristata. Sample 114-701C-51X-CC is barren of calcareous nannofossils and is believed to be of igneous origin.

**Site 702**

Site 702 (50°56.786'S, 26°22.117'W; water depth 3084 m) is located on the central part of the Islas Orcadas Rise. The major objectives of this site were (1) to determine the age, nature, and subsidence history of the Islas Orcadas Rise and (2) to investigate the influence of the shallow Islas Orcadas and Meteor rises on oceanic water mass communication between the southern high-latitude region and the South Atlantic.

The sedimentary sequence penetrated at Site 702 was divided into two lithostratigraphic units. Unit I (down to Section 114-702B-3H-4, 80 cm) consists of diatom mud to muddy diatom ooze overlying alternating nannofossil diatom ooze and diatom nannofossil ooze; Unit II is a sequence of nannofossil oozae, chalks, and indurated chalks with intercalated cherts, particularly in the lower part of the section. Samples above Sample 114-702B-6X-CC contain slightly etched nannofossil floras and those below that depth are overgrown with secondary calcite.

**Quaternary to Miocene**

The nannofossil floras of Sections 114-702B-1H-CC to 114-702B-2H-CC are dominated by *Reticulofenestra perplexa*, with lesser numbers of *Coccolithus pelagicus*. This assemblage indicates the presence of upper to middle Miocene strata.

**Eocene**

A stratigraphic break was recognized in both Holes 702A and 702B, with some Miocene, Oligocene, and upper Eocene strata absent. In Hole 702B, this break lies between Samples 114-702B-2H-CC and 114-702B-3H-CC. Samples 114-702B-3H-CC to 114-702B-5X-CC are assigned to the *Chiasmolithus oamaruensis* Zone, on the basis of the presence of *C. oamaruensis* and *Neococcolithes dubius* and the absence of *Isthmolithus recurvus*. No samples that could be assigned to the *Discocastor saipanensis* Zone were examined. The absence of this zone is discussed within "Paleoenvironment and Provinicialism."

The interval between Samples 114-702B-6X-2, 69–70 cm, and 114-702B-9X-CC is assigned to the *Chiasmolithus solitus* Zone. The interval between Samples 114-702B-10X-5, 20–21 cm, and 114-702B-14X-CC is assigned to the *Nannotetrista fulgens* Zone. This assignment is based on the presence of *N. fulgens*/Nannotetrista cristata in the lowermost two samples. Supporting evidence comes from the presence of *Sphenolithus furcatothioides* in Sample 114-702B-11X-CC and *Chiasmolithus gigas* in Samples 114-702B-12X-2, 20–21 cm, and 114-702B-12X-CC.

Samples 114-702B-15X-2, 60–61 cm, and 114-702B-21X-CC are assigned to the *Discocastor sublodoensis* Zone. This assignment is based on the FAD of *D. sublodoensis* in the lowest section. Sample 114-702B-22X-2, 20–21 cm, is assigned to the *Discocastor lodoensis* Zone. The occurrence of *Tridacnialithus orthostylus* between Samples 114-702B-22X-CC and 114-702B-26X-CC indicates the presence of the *T. orthostylus* Zone.

**Paleocene**

The co-occurrence of *Discocastor multiradiatus* and the *Fasciculithus typaniformis* group in Samples 114-702B-27X-1, 6–7 cm, to 114-702B-27X-CC indicates the presence of the *D. multiradiatus* Zone. The Eocene/Paleocene boundary lies near the top of this zone. The absence of *D. multiradiatus* and the presence of *Heliolithus riedelli* in Sample 114-702B-30X-CC indicates the presence of the *H. riedelli* Zone for the interval between Samples 114-702B-28X-CC and 114-702B-30X-CC.

Sample 114-702B-31X-CC contains *Heliolithus kleinpellii* and is thus assigned to the zone of that name. The continued presence of the *F. typaniformis* group below the FAD of *H. kleinpellii*, in Sample 114-702B-32X-CC, indicates the presence of the *F. typaniformis* group Zone. The abundance of *H. kleinpellii* is very low so its FAD in this hole may not be its true FAD.

**Site 703**

Site 703 (47°03.042'S, 07°53.679'E; water depth 1796 m) is located on the Meteor Rise. The major objectives of drilling at this site were to determine the nature, age, and subsidence history of the rise and to investigate the influence of the shallow Paleogene Meteor Rise, Islas Orcadas rise, and the adjacent fracture zones on oceanic communication between the high and temperate latitudes of the South Atlantic.
The sedimentary sequence consists of mainly nannofossil ooze with chalks in the lower part of the sequence (below Core 114-703A-26X). At the bottom of Hole 703A, basalt and tuff were found intermixed with sediments, although it was not possible to tell if this represented true basement. The Pleistocene to Pliocene nannofossil floras (above Sample 114-703A-2H-CC) comprise slightly etched fossils. The Miocene to Eocene nannofossil floras below are overgrown with secondary calcite deposition, which increases downward.

Quaternary to Pliocene

Samples 114-703A-1H-1, 1–2 cm, to 114-703A-2H-CC contain sparse nannofossil floras, including Pseudoemiliania lacunosa and Gephyrocapsa spp. The co-occurrence of these taxa indicates the presence of middle Pleistocene to upper Pliocene strata.

Lower Miocene to Oligocene

A stratigraphic break lies between Samples 114-703A-2H-CC and 114-703A-3H-2, 134–135 cm, with most of the Miocene and Pliocene absent. The nannofossil floras recovered from the interval between Samples 114-703A-3H-2, 134–135 cm, and 114-703A-4H-CC are dominated by Cyclamagolithus floridanus; also present are rare Cyclamicagolithus abisectus and Chiasmolithus altus. This may belong to the C. abisectus Zone or may represent reworking of C. abisectus and C. altus into younger Miocene strata. The presence of C. floridanus indicates a middle Miocene or older age.

The abundant occurrence of C. abisectus in Sample 114-703A-5H-2, 20–21 cm, indicates the presence of definite C. abisectus Zone strata. The interval between Samples 114-703A-5H-CC and 114-703A-7H-2, 10–11 cm, is assigned to the Reticulofenestra bisecta Zone, as the nominate taxon occurs in all samples from this interval. The first consistent downhole common occurrence of C. altus in Sample 114-703A-7H-5, 10–11 cm, indicates the presence of the C. altus Zone, which extends down to Sample 114-703A-11H-CC. The LAD of Reticulofenestra umbilicus defines the top of the underlying Reticulofenestra daviessii Zone, which lies between Samples 114-703A-12H-2, 22–23 cm, and 114-703A-12H-4, 24–25 cm. The abundant occurrence of Clasticoccus subdistichus in the interval between Samples 114-703A-12H-6, 21–22 cm, and 114-703A-12H-CC indicates the presence of the subdistichus Subzone of the Isthmolithus recurvus Zone.

Eocene

The Reticulofenestra oamaruensis Subzone is recognized below the LAD of the nominate taxon at Sample 114-703A-13H-2, 74–75 cm, down to Sample 114-703A-15H-2, 95–96 cm; the Oligocene/Eocene boundary probably lies within this subzone. Sample 114-703A-15H-CC, in which Discoaster saipanensis and Isthmolithus recurvus co-occur, is assigned to the J. recurvus Subzone. The interval between the FAD of J. recurvus and the FAD of Chiasmolithus oamaruensis defines the C. oamaruensis Zone between Samples 114-703A-16X-1, 20–21 cm, and 114-703A-20X-2, 30–31 cm. A possible hiatus separates the C. oamaruensis and Chiasmolithus solitus Zones, with the D. saipanensis Zone absent. This is discussed within “Paleoenvironment and Provincialism.” The co-occurrence of Reticulofenestra bisecta and C. solitus indicates the presence of the C. solitus Zone between Samples 114-703A-20X-CC and 114-703A-22X-CC. The interval between Samples 114-703A-23X-1, 20–21 cm, and 114-703A-30X-CC is assigned to the Nanotetra fulgens Zone, and the FAD of N. fulgens/N. cristata lies in the lowest sample in the interval. Supporting evidence for this zonal assignment is provided by the occurrence of Sphenolithus furcatus in Sample 114-703A-30X-2, 30–32 cm. The only sample available for study in the lower part of the hole is Sample 114-703A-33X-CC. This sample contains Reticulofenestra umbilicus, the FAD of which lies within the Discoaster sublodoensis to Discoaster lodoensis Zones.

Site 704

Site 704 (46°52.757′S, 07°25.250′E; water depth 2532.3 m) is located on the southern part of the Meteor Rise. Site 704 was located where a maximum thickness of Neogene and upper Paleocene sediment could be obtained in order to provide a high-resolution paleoceanographic-paleoclimatic record of an interval not well represented at the preceding Leg 114 sites.

The sedimentary sequence was divided into two lithostratigraphic units. Unit I consists of varying admixtures of two end-member components, namely calcareous oozes and chalks consisting of foraminifers and calcareous nannofossils and silicious oozes composed of diatoms with minor radiolarians and silicoflagellates. Unit II consists of micritic-bearing nannofossil chalk passing down into micritic indurated nannofossil chalk. The nannofossils from the upper part of Hole 704B (above Core 114-704B-10H) consist of slightly etched coccoliths. Below this depth, they become increasingly overgrown with secondary calcite.

Hole 704A

Quaternary

The upper part of Hole 704A is the subject of a separate study (Gard and Crux, this volume). This study has shown that the upper part of the hole is much younger than was reported by Ciesielcki, Kristoffersen, et al. (1988). Dominant Emiliania huxleyi in Sample 114-704A-1H-2, 24 cm, and above indicates the presence of sediments from oxygen isotope stage 5a or younger (Thierstein et al., 1977). The continued presence of E. huxleyi down to at least Sample 114-704A-1H-3, 104 cm, indicates an age no older than mid-oxygen isotope stage 8 (0.268 Ma; Thierstein et al., 1977). The LAD of Pseudoemiliania lacunosa has been recognized in Sample 114-704A-1H-5, 105 cm, and the interval from this sample down to Sample 114-704A-13H-CC is assigned to Zone NN19. Samples from the interval between Samples 114-704A-14H-5, 33–34 cm, and 114-704A-15H-CC additionally contain Helicosphaera sellii, a species with a LAD within NN19; thus, these samples can be assigned to the lower part of NN19 (lower Pliocene).
A stratigraphic break may occur between Samples 114-704A-22X-CC and 114-704A-23X-CC. This is reflected in the nannofossil flora by an abrupt change in the assemblages from those dominated by *Coccolithus pelagicus* to those dominated *R. pseudoumbilicus*. This change could alternatively be interpreted as a paleoceanographic change (Müller et al., this volume).

Samples 114-704A-23X-CC and 114-704A-24X-CC are assigned to Zones NN15-11 on the basis of the presence of *R. pseudoumbilicus* in both samples and *Amaurolithus delicatus* in the lower sample. Below Sample 114-704A-25X-CC, the strata are assigned to Zones NN15-77. The nannofossil floras are of low diversity and are dominated by *R. pseudoumbilicus*, *Reticulofenestra perplexa*, and *C. pelagicus*.

**Hole 704B**

The higher part of this hole has not been reexamined; the preliminary results are presented in Ciesielski, Kristoffersen, et al. (1988). However, the revisions to the younger part of Hole 704A probably apply to this hole as well.

The interval between Samples 114-704B-14X-CC and 114-704B-16X-CC contains *H. sellii*, a species with an LAD within Zone NN19.

The very rare occurrence of *D. broweri* and the presence of *C. macintyrei* in Sample 114-704B-17X-CC indicate the presence of NN18 or older strata. No older Pliocene zones were recognized in the interval between Samples 114-704B-17X-CC and 114-704B-21X-CC, although comparisons to Hole 704A would suggest that they are probably present.

A stratigraphic break or paleoceanographic change may occur between Samples 114-704B-21X-CC and 114-704B-22X-CC. This is reflected in the nannofossil floras by an abrupt change from assemblages dominated by *C. pelagicus* to those dominated by *R. pseudoumbilicus*.

Samples 114-704B-22X-CC to 114-704B-24X-CC are assigned to Zones NN14-9 (lower Pliocene–middle Miocene) on the basis of the presence of *Amaurolithus tricorniculatus* (NN14-11) in the highest sample and *Discoaster pentarradiatus* in the lowest sample. The underlying interval between Samples 114-704B-25X-CC and 114-704B-45X-CC is assigned to Zones NN14-77. Samples within this interval contain abundant nannofossil floras of low diversity that lack age-diagnostic species. The dominance of these floras by *C. pelagicus*, *R. perplexa*, and *R. pseudoumbilicus* together with their stratigraphic position confirms this age assignment.

An abrupt change in the nannofossil flora between Samples 114-704B-43X-CC and 114-704B-46X-CC indicates a stratigraphic break. The co-occurrence of *Sphenolithus heteromorphus* and *C. macintyrei* in the latter sample indicates the presence of Zones NN5-4. The nannofossil floras from this interval are dominated by *Cyclicargolithus floridanus* and *C. pelagicus*.

Samples below the FAD of *C. macintyrei* are assigned to between NN4 and the *Cyclicargolithus abisectus* Zone, between Samples 114-704B-50X-CC and 114-704B-63X-CC. No zonal index taxa are present, with the exception of *C. abisectus*. This species appears to range as high as Zone NN5 in this area, or it is reworked. Nevertheless, the top of the *C. abisectus* Zone cannot be accurately picked in Hole 704B. The *R. bisecta* Zone is recognized in Sample 114-704B-64X-CC by the occurrence of the nominate taxon. The top of the *Chiasmolithus altus* Zone is recognized in Sample 114-704B-65X-CC by the consistent and significant occurrence of the nominate taxon. This zone is present down to the lowest Sample 114-704B-72X-CC.

**PALEOENVIRONMENT AND PROVINCIALISM**

Cretaceous

The nannofossil floras recovered from the Upper Cretaceous sections penetrated by Holes 698A and 700B are poor to very poorly preserved. This should be taken into account in drawing any conclusions as to their paleoenvironmental and biogeographic significance. Another prerequisite for studies of the paleoenvironmental significance of these nannofossil floras is a reliable means of correlation between these sites and other areas both near and far. Previous studies (Ciesielski et al., 1977; Wind, 1979) on the paleoenvironmental significance of high-latitude nannofossil floras from the South Atlantic and Indian oceans have been hampered by the lack of a good zonal scheme for this area. These studies made comparisons of the nannofossil floras recovered in high-latitude areas using correlations derived from foraminiferal studies. They were also based on relatively incomplete sections.

Until Hole 700B was drilled, no sedimentary succession without major breaks had been recovered from the Maestrichtian of the high-latitude South Atlantic. Wind and Wise (1983) attempted to summarize the stratigraphic positions of the cores recovered by Deep Sea Drilling Project (DSDP) Legs 36 and 71 and the Islas Orcadas Core 10 07-75-44. This summary would appear to need modification in the light of the present findings.

In the present study, the following approach has been used to establish a nannofossil zonal framework with which to make direct comparisons of nannofossil floras of the same age from other areas. It is first assumed that cosmopolitan nannofossil datums, such as the LADs of *Effolithus eximius*, *Brionsonia parca*, and *Reinhardites levis*, are approximately synchronous in different latitudes. Second, nannofossil datums observed in high-latitude areas of the Northern Hemisphere, such as the LADs of *Orastrum campanensis* and *Biscutum magnum* and the FAD of *Nephrolithus frequens*, are considered to lie at approximately the same level in the Southern Hemisphere. The dating of cores and the positioning of the Maestrichtian/Campanian boundary relative to the nannofossil datums are critical for some of the previous paleoenvironmental interpretations. Ciesielski et al. (1977) compared cores recovered by DSDP Leg 36 at Site 327 with a series of cores from the Maurice Ewing Bank, in particular Islas Orcadas Core 10 07-75-44. They concluded that differences between the nannoflora of the two areas were accounted for by a major water-mass boundary lying between them. The findings of the present study would, however, suggest that the material recovered from DSDP Site 327 was younger than that recovered from the Islas Orcadas core. Thus, no paleoenvironmental explanations are necessary to account for the observed differences. The presence of *Effolithus eximius* and the absence of *Monomarginatia* spp. (not including *Monomarginatia primus*) and *Biscutum magnum* suggest an earliest Maestrichtian or older age for the Islas Orcadas core. The presence of *Monomarginatia* spp. and *B. magnum* and the absence of *E. eximius* suggest an early to late Maestrichtian age for the sediments recovered from DSDP Site 327.

Differences in taxonomic concepts of different authors prevent detailed comparisons of the nannofossils from material recovered by previous DSDP and Islas Orcadas cruises and those of the present study. In addition, the nannofossil floras recovered from Holes 698A and 700B are poorly preserved, as noted previously. The following observations can be made despite these limitations.
The nannofossil floras of the Santonian sediments recovered from Hole 700B are very similar to those of the same age in the Northern Hemisphere. The most notable difference is the presence of Seribiscutum primitivum in the Southern Hemisphere, whereas this species is not found above the Cenomanian in the Northern Hemisphere (Forchheimer, 1968; Crux, 1982). The LAD of Lithastrinus floralis, in contrast, may be later in the high latitudes of the Northern Hemisphere; Crux (1982) and Bukry (1969) recorded this species from the Campanian.

Only Sample 114-700B-50R-2, 26–27 cm, was recovered from strata that could be confidently assigned to the Campanian. The nannofossil flora of this sample was so poorly preserved that no useful comparisons can be made with other areas.

The nannofossil floras from the Maestrichtian sediments recovered from Holes 698A and 700B are very similar to those recorded by Wise and Wind (1977) and Wind and Wise (1983) (DSDP Legs 36 and 71), although less well preserved. Distinctive high-latitude species common to these studies and the present study include Biscutum coronum, Nephrolithus corsites, Biscutum dissimilis, Teichorhabdus eithmos, N. frequens, B. magnus, and Monomarginatus pectinatus/M. quaternarius. Species present only in the Falkland Plateau region include Boletuvelum candens, Centosphaera barbata, Lucianorhabdus arborius, Okkolithus australis, Orastrum asarotruncatum, Ottavianus giannus, ”Phanulithus” additus, and Pharus simulacrum. In Hole 703A, the abundant occurrence of Isthmolithus recurvus, Prinsius martius, and Reticulofenestra daviesii group, as in the chalks of southern England, where members of the genera Lucianorhabdus and Calcuttes are common (Crux, 1980).

Thus, it is possible the absence of some of the holococcoliths from the Falkland Plateau region could possibly be attributed to a deeper water environment than that of the Falkland Plateau region.

Holococcoliths are structurally weaker than heterococcoliths. This structural weakness commonly leads to their destruction before they are included in the sediment. In the sediments studied from the Northeast Georgia Rise, the poor preservation of the nannofossils is probably controlled by diagenetic processes, rather than destruction before burial, with progressive lithification downhole. Under such conditions, holococcoliths can, in fact, be preferentially preserved, as in the chalks of southern England, where members of the genera Lucianorhabdus and Calcuttes are common (Crux, 1980).

The abrupt LAD of C. coenurum and the closely related species C. reticulatum, which dominates the assemblages of the lowest upper Eocene in Hole 699A, is below the FAD of Isthmolithus recurvus. In Hole 703A, the abundant occurrence of C. coenurum continues to occur above the level of the FAD of I. recurvus. This latter sequence of events is the same as reported by Perch-Nielsen (1985) in low-latitude areas, whereas the former is typical of high latitudes.

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diachronous. *I. recurvus* is known to prefer high latitudes (Perch-Nielsen, 1985), whereas *C. reticulatum* (and presumably the closely related *C. coenurum*) was identified by Haq and Lohmann (1976) as typical of midlatitude assemblages. Thus we have two species with different environmental preferences. The FAD of *I. recurvus* is the harder event to define accurately, because of the very low numbers of specimens in the early part of its range. It is tempting to assume that this species occurred or appears to occur later at Hole 699A, because of the low number of specimens involved.

Depending on which event one considers to be isochronous, if either, the differences in the sequences of the events in the two areas may provide evidence for water circulation patterns for the late Eocene. It possibly suggests warmer surface waters on the eastern side of the South Atlantic to those of the western side, resulting in an earlier decline of *C. coenurum* in the west.

Another biostratigraphic problem caused by the environmental preferences of index taxa is the existence or absence of the *Discoaster saipanensis* Zone in the study area. This zone is defined as the interval between the LAD of *Chiasmolithus solitus* and *Chiasmolithus oamarunensis*. Both of these species belong to the genus *Chiasmolithus*, which has a preference for high-latitude/cold-water areas. In the present study the two events were found to occur very close to one another, but the two do not co-occur. It is assumed that this pattern is the result of the species both reaching their maximum stratigraphic ranges in this area. An alternative explanation could be the existence of a regional hiatus at this level causing the *D. saipanensis* Zone sediments to be absent. The *D. saipanensis* Zone is reported by Perch-Nielsen (1985) to have some very thick sections in low latitudes, where the stratigraphic ranges of the index taxa are shortened, thus expanding this zone. Wei and Wise (1989) have recently reported that *C. solitus* has a longer stratigraphic range in high latitudes, which explains at least some of the distribution pattern seen. The use of the LAD of *C. solitus* and FAD of *C. oamarunensis* within the study area as biostratigraphic datums is still considered valid, as there is little latitudinal variation. Comparisons with other areas should be made very carefully, as the *D. saipanensis* Zone increases in thickness in lower latitudes. It has not been possible to prove the total absence of the *D. saipanensis* Zone in the study area, although if present, it must be less than 2 m thick in Hole 699A.

The nannofossil floras recovered from the Eocene sections of the sites in the present study are, as previously noted, dominated by successive groups of palaeoliths, with the persistent occurrence of *Chiasmolithus* throughout. These are typical of high-latitude assemblages. Groups that are absent or rare in the material recovered include the warmer water/low-latitude *Discoaster, Helicosphaera*, and *Sphenolithus*. It is probable that the levels where these groups do occur were deposited during relatively warmer intervals, when low-latitude warmer waters were able to move farther south into the study area.

Discoasters are most common within the lower to lower middle Eocene (*Tribrachius orthostylus* to *Discoaster sublodoensis* Zone). They occur less commonly in the middle middle Eocene (*Nannoteirina fulgens* to *C. solitus* Zones), but become more common in the upper middle Eocene to upper Eocene (*C. solitus* to *I. recurvus* Zones). *Helicosphaera* is almost totally absent from the study area. It is present in a short interval in the lower Eocene (*T. orthostylus* to *D. lodoensis* Zones) and in a single sample in the middle Eocene (*C. solitus* Zone). *Sphenolithus* occurs rarely throughout the Eocene, but is more common in the lower to middle Eocene (*T. orthostylus* to *D. sublodoensis* Zones).

These patterns suggest that the early to early middle Eocene was a period with comparatively warm surface water. A cooler period occurred in the middle middle Eocene, and warmer conditions returned in the late middle to late Eocene. These findings are similar to the conclusions of the much more detailed study of Haq and Lohmann (1976), who recorded maximum poleward migrations of low-latitude nannofloras in the early middle Eocene (*D. sublodoensis* Zone) and the late Eocene (*I. recurvus* Zone, this study).

In addition to the temperature-sensitive groups described previously, the distribution of *Zygrhablithus bijugatus* is of interest. This species is a robust holococcolith with a total stratigraphic range from the upper Paleocene to the lowest Miocene. In the present study, it is found to occur in the upper Paleocene to lower Eocene and the upper middle Eocene. This might indicate that its distribution is also temperature controlled, as the intervals it occurs in are those influenced by warm surface water. The high abundance of this species in the upper Paleocene to lower Eocene may also be related to water depth, as this species is more commonly preserved in water depths of less than 1000 m (Edwards, 1973).

**Oligocene—Lowest Miocene**

Oligocene to lowest Miocene nannofossil floras were recovered from Sites 699, 701, 703, and 704. The most complete and thickest of these sections is from Hole 699A (Table 1). The preservation of the nannofloras in all the Oligocene sections is poor, with most of the assemblages etched by varying degrees. The Oligocene to lowest Miocene nannofossil floras are characterized by a lower diversity than those of the Eocene, with never more than 16 species present in one sample. The dominance of the assemblages alternates among the *Cyclicargolithus floridanus* group, the *Reticulofenestra daviessi* group, and *COCCOLITHUS PELAGICUS*. *Chiasmolithus* and *C. oamarunensis* have some common to abundant occurrences and are always a significant element of the assemblage.

The low diversity and near total absence of the genera *Discoaster, Helicosphaera,* and *Sphenolithus* suggest that very cold-water conditions prevailed during most of the Oligocene to earliest Miocene. The presence in both Holes 699A and 703A of rare *Discoaster deflandrei, Discoaster spp.*, and *Discoaster adamanteus* in the upper Oligocene to lowest Miocene (*R. daviessi* to *Cyclicargolithus abisectus* Zones) may indicate a period of warmer surface waters. Haq and Lohmann (1976) recorded a maximum poleward migration of warm-water/lower-latitude nannofloras in the upper Oligocene, corresponding to the base of the *R. daviessi* Zone.

**Miocene to Quaternary**

Miocene to Quaternary cores recovered during Leg 114, except for those from Site 704, contain only short intervals of nannofossil-bearing sediments. These contain nannofossil floras of only two or three species. Such low diversities probably indicate cold surface waters at the time of deposition and/or etching of the nannofossils.

**Site 704**

The lower Miocene section of Hole 704B contains abundant nannofloras of very low diversity (fewer than eight species/sample). These are dominated by *C. floridanus* and *C. pelagicus*. The low diversity is probably indicative of cold surface waters during deposition; however, warm-water elements do occur throughout the interval. These are *Sphenolithus heteromorphus*, *Sphenolithus moriformis*, and *Discoaster spp.*
A hiatus separates the lower Miocene from the middle to upper Miocene, where the nannofossil floras are again of very low diversity (fewer than eight species/sample). These are dominated by the cold-water species C. pelagicus and Reticulofenestra pseudoumbilicus, with lesser numbers of Reticulofenestra pseudoumbilicus. Sphenolithus abies is rarely present in the lower part of this interval between Cores 114-704B-45X and 114-704B-34X. Helicosphaera carteri is present in Sample 114-704B-32X-CC. Discoasters are totally absent. It would thus appear that the middle to upper Miocene interval was deposited during a period of cold surface waters that were coldest at the end of the Miocene.

The Pliocene interval between Core 114-704B-27X and Sample 114-704A-17X-CC also contains low-diversity nannofossil floras (fewer than eight species/sample). Discoasters are present but rare throughout the Pliocene, with rare Amauroolithus in the lower Pliocene and Helicosphaera in the upper Pliocene. The nannofossil floras of the lower Pliocene are dominated by R. pseudoumbilicus, and those in the upper Pliocene by C. pelagicus. The Pliocene would appear, from the sparse occurrence of the discoasters and other warm water genera, to be slightly warmer than the latest Miocene.

The Pleistocene interval above Sample 114-704A-17X-CC contains abundant low-diversity nannofossil floras (fewer than eight species/sample). These are dominated by C. pelagicus with common Gephyrocapsa spp., which increase uphole. Cacildiscus leptopus and Emiliania huxleyi are abundant in certain intervals in the top two cores. The paleoenvironmental significance of the nannofossils of this interval is the subject of a separate study (Gard and Crux, this volume).

**TAXONOMY**

Species discussed here are also listed in the Appendix. A number of species groupings were employed in the present study. These were used either because poor preservation did not allow the separation of the taxa concerned, or because the established taxonomic subdivisions of the groups are confused and based on dubious criteria. These groups are as follows.

**Mesozoic**

Arkhangelkiella cymbiformis—My concept of A. cymbiformis includes forms often assigned to Arkhangelkiella specillata. I believe these two species to be preservational conspecific morphotypes of one another.

Reinhardtites a.aff. R. anthophorus and R. levis—These two species are usually differentiated by the size of the two central openings either side of the central bridge. This division is difficult to apply in poorly preserved material where secondary calcite overgrowth can obscure the openings.

Monomarginatus pectinatus and Monomarginatus quaternarius—These two species are differentiated by the number of perforations bordering the outer rim. Secondary calcite overgrowth prevented the observation of these perforations.

**Cenozoic**

Fasciculithus tympaniformis group—Considerable overgrowth of secondary calcite on the members of the genus Fasciculithus prevented any reliable subdivision. The F. tympaniformis group thus includes all species of Fasciculithus present.

Chiasmolithus bidens and C. solitus—Even in well-preserved material, I have difficulty differentiating these two species with only the light microscope. In the Paleocene sections I have grouped them together.

Toweius pertusus and T. eminens—These two species are grouped together in the nannofossil distribution chart for the Paleocene of Site 700 (Table 3) because the very poor preservation prevents their reliable and consistent separation.

Nannofracta cristata and N. fulgens—The poor preservation of the sections studied does not allow the separation of these two species.

Reticulofenestra davisii group—This group includes all medium-sized Reticulofenestra with varying constructions of the central grid.

Cyclarolithus floridanus group—This group includes all round to subelliptical imperforate small- to medium-sized placoliths recorded in the Eocene to lower Miocene. One new species is proposed from the present study.

Genus MONOMARGINATUS Wind and Wise, 1977

Monomarginatus primus n. sp.

Staurolithes ellipticus (Gartner), Crux, 1982, p. 116, pl. 5.3, figs. 13–15; pl. 5.9, fig. 20.

**Description.** An elliptical cocolith with a loxolith rim and two cross bars parallel with the major and minor axes of the ellipse. These bars are composed of numerous crystal elements and can support a very robust spine composed of numerous elements. Each of the four quadrants formed by the cross is nearly filled by a diagonal bar that links the bars of the cross.

**Remarks.** This species differs from M. quaternarius by the more robust nature of the central structure, which in overgrown specimens appears to completely fill the central area. The thick multielement spine of this species is very distinctive and is often seen detached under the light microscope. This species is possibly ancestral to the other species of Monomarginatus. Monomarginatus primus differs from Staurolithes ellipticus by its much larger size and distinctive spine.

**Occurrence.** Maestrichtian of Holes 698A and 700B; upper Campanian of Norfolk, UK.

**Size.** Holotype 9.4 µm long, 5.9 µm wide. Size range of observed specimens between 9 and 12 µm long.

**Holotype.** From Crux (1982), pl. 5.3, fig. 20; negative number UCL-1028-22.

**Paratypes.** From Crux (1982), pl. 5.3, figs. 14 and 15; negative numbers UCL-1028-19 and UCL-1035-5.

**Type locality.** North Norfolk coast near West Runton, U.K.

**CONCLUSIONS**

Nannofossils are useful in dating and correlating the Cretaceous to Quaternary sediments deposited above the CCD in the high latitudes of the South Atlantic. The Cretaceous to the lowermost Miocene interval can be divided into 29 nannofossil zones and three subzones. This degree of biostratigraphic resolution is only slightly less than that achieved in low-latitude areas under more optimal conditions.

The LADs of Eifflolithus eximius, Broinsonia parca, Reinhardtites levis, and Biscutum mugan occur in the same order in both the Northern and Southern hemispheres and are considered to be reliable as biostratigraphic datums in high-latitude areas.

The diachronicity of the FAD of Nephrolithus frequens, first reported by Wind (1979), is confirmed. This datum occurs in older sediments than the LAD of Reinhardtites levis in high-latitude areas of both the Northern and Southern hemispheres, while in low-latitude areas this order is reversed (Sissingh, 1977).

Monomarginatus primus n. sp. appears to have a later LAD in the Southern Hemisphere than in the Northern Hemisphere. In the Southern Hemisphere this datum occurs above the LADs of Broinsonia parca and Eifflolithus eximius, while in the Northern Hemisphere it is found below them.

Some of the Tertiary biostratigraphic datums used show more restricted or expanded stratigraphic ranges in the high-latitude South Atlantic area. In particular, the ranges of Chiasmolithus solitus and Chiasmolithus oamaruensis appear...
to be longer in the high latitudes. The ranges of _Nannotetina fulgens_// _N. cristata_ and _Reticulofenestra bisecta_ appear to be shorter, when calibrated by magnetostratigraphy and compared to the findings of Wei and Wise (1989).

Differences in the relative positions of the LAD of _Cribrocentrum coenorum_ and the FAD of _Istiodolithus recurvus_ are observed between Holes 699A and 703A. This possibly suggests that warmer surface waters existed in the middle to late Eocene at Site 703 than at Site 699, as the warm-water species _Cribrocentrum coenorum_ and _C. reticulatum_ and _Cribrocentrum coenurum_ and _C. reticulatum_ appear to have later LADs there.

The presence or absence of temperature-sensitive calcareous nannofossil floras reflects the changing surface-water temperatures of the area throughout the Cenozoic. The highest diversities are recorded in the Paleocene to Eocene sections, with warmer-water genera present in the upper Paleocene to lower middle Eocene and in the upper middle to upper Eocene sediments. A marked drop in diversity occurs in the lower Oligocene to lowermost Miocene. Very low diversities are present in the middle Eocene and in the upper middle to upper Eocene sediments. Changes over short stratigraphic intervals in the nannofossil floras reflects the changing surface-water environment.

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**REFERENCES**


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Calcareous Nannofossils Considered in this Chapter

Mesozoic

Phanerolithus aditus Wind and Wise in Wise and Wind, 1977

Helicolithus ances (Görka, 1957) Noël, 1970

Cretarhabdus angustiforatus (Black, 1971) Bukry, 1973

Reinhardtites anthophorus (Deflandre, 1959) Perch-Nielsen, 1968

Reinhardtites aff. R. anthophorus sensu Prins and Sissingh in Sissingh, 1977

Lucianorhabdus arborius Wind and Wise in Wise and Wind, 1977

Lucianorhabdus arquatus Forchheimer, 1972

Orastrum asarotrum Wind and Wise in Wise and Wind, 1977

Centosphaera barbata Wind and Wise in Wise and Wind, 1977

Watznaueria barnesae (Black in Black and Barnes, 1959) Perch-Nielsen, 1968

Cylindricalithus bicus Bukry, 1969

Watznaueria biporta Bukry, 1969

Calcultites Prins and Sissingh in Sissingh, 1977


Boletovum candens Wind and Wise in Wise and Wind, 1977

Lithraphidites carioensis Deflandre, 1963

Lucianorhabdus cayetensis Deflandre, 1959

Glaukolithus compactus (Bukry, 1969) Perch-Nielsen, 1984

Cretarhabdus comicus Bramlette and Martin, 1964

Biscutum coronum Wind and Wise in Wise and Wind, 1977

Nephrolithus corytus Wind, 1983

Cretarhabdus crenulatus Bramlette and Martin, 1964

Prediscosphaera cratacea (Arkhangelsky, 1912) Gartner, 1968

Arkhangelskiella cymbiformis Vekshina, 1959

Microhabdus decoratus Deflandre, 1959

Biscutum dissimilis Wind and Wise in Wise and Wind, 1977

Cribrospira ephemerbergii Arkhangelsky, 1912

Biscutum ellipticum (Görka, 1957) Grun in Grun and Allemann, 1975

Broinsonia enormis (Shumenko, 1968) Manivit, 1971

Teichorhabdus ethmos Wind and Wise in Wise and Wind, 1977

Effellithus eurinius (Stover, 1966) Perch-Nielsen, 1968

Lithastrinus florialis Stradner, 1962

Nephrolithus frequens Görka, 1957

Marthasterites furcatus (Deflandre in Deflandre and Fort, 1954)

Deflandre, 1959

Tanolithus gabolus Stover, 1966

Quadrum gaudert Prins and Perch-Nielsen in Manivit, Perch-Nielsen, Prins, and Verbeek, 1977

Ottavianiana gunnisonis Risatti, 1973

Lithastrinus grilli Stradner, 1962

Reinhardtites levis Prins and Sissingh in Sissingh, 1977

Lucianorhabdus Deflandre, 1959

Kampineria magnificus Deflandre, 1959

Biscutum magnum Wind and Wise in Wise and Wind, 1977

Micula Vekshina, 1959

Staurolithus melinensis (Görka, 1957) Perch-Nielsen, 1968

Micula mutus (Martini, 1961) Bukry, 1973

Gartnerago obliquum (Stradner, 1963) Noël, 1970

Calculusites obscurus (Deflandre, 1959) Prins and Sissingh in Sissingh, 1977

Ahnmuellerella octoradiata (Görka, 1957) Reinhardt, 1964

Tanolithus orionatus (Reinhardt, 1966) Reinhardt, 1966

Watznaueria ovata Bukry, 1969

Broinsonia parca (Stradner, 1963) Bukry, 1969

Monomarginatus pectinatus Wind and Wise in Wise and Wind, 1977

Manivitella pemmatoides (Deflandre in Manivit, 1965) Thierstein, 1971


Monomarginatus primus n. sp.

Monomarginatus quaternarius Wind in Wise and Wind, 1977

Scampanella Forchheimer and Stradner, 1973


Cylindricalithus sculptus Bukry, 1969

Broinsonia signata (Noël, 1969) Noël, 1970

Pharus simulacrum Wind and Wise in Wise and Wind, 1977

Zeugrhabdotus sisyphus (Gartner, 1968) Crux, 1989

Arkhangelskiella specilata Vekshina, 1959

Zygodiscus adamas Bramlette and Sullivan, 1961

Forchheimer, 1972

Lucianorhabdus cayeuxii Vekshina, 1959

Micula murus (Martini, 1961) Bukry, 1973

Gartnerago obliquum (Stradner, 1963) Noël, 1970

Calculusites obscurus (Deflandre, 1959) Prins and Sissingh in Sissingh, 1977

Ahnmuellerella octoradiata (Görka, 1957) Reinhardt, 1964

Tanolithus orionatus (Reinhardt, 1966) Reinhardt, 1966

Watznaueria ovata Bukry, 1969

Broinsonia parca (Stradner, 1963) Bukry, 1969

Monomarginatus pectinatus Wind and Wise in Wise and Wind, 1977

Manivitella pemmatoides (Deflandre in Manivit, 1965) Thierstein, 1971


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Pharus simulacrum Wind and Wise in Wise and Wind, 1977

Zeugrhabdotus sisyphus (Gartner, 1968) Crux, 1989

Arkhangelskiella specilata Vekshina, 1959

Zygodiscus adamas Bramlette and Sullivan, 1961

Forchheimer, 1972
Calcareous Nannofossils

Chiasmolithus albus Bukry and Percival, 1971
Amaurolithus Gartner and Bukry, 1975
Markallus Markalius inversus (Lohmann, 1902) Hay and Mohler, 1967
Discosphaera braardiensis Tan, 1927
Chiasmolithus bidens (Bramlette and Sullivan, 1961) Hay and Mohler, 1967
Braarudosphaera bigelowii (Gran and Braarud, 1935) Deflandre, 1947
Zygochitina hujagatus (Deflandre in Deflandre and Fert, 1954) Deflandre, 1959
Discosphaera binodosus Martini, 1958
Reticulofenestra bisecta (Hay, Mohler, and Wade, 1966) Roth, 1970
Prinsus biscutus (Stradner, 1963) Hay and Mohler, 1967
Tribrachiatus tramletti (Bromann and Stradner, 1960) Proto Decima, Roth, and Todesco, 1975
Discosphaera brouweri Tan, 1927 endem. Bramlette and Riedel, 1954
Chiplagnamalithus calathus Bramlette and Sullivan, 1961
Chiasmolithus californicus (Sullivan, 1964) Hay and Mohler, 1967
Chiasmolithus Hay, Mohler, and Wade, 1966
Cribrocentrum coenatum (Reinhardt, 1966) Perch-Nielsen, 1971
Neochiastozygus concinnus (Martini, 1961) Perch-Nielsen, 1971
Chiasmolithus consuetus (Bramlette and Sullivan, 1961) Hay and Mohler, 1967
Tribrachiatus contortus (Stradner, 1958) Bukry, 1972
Toweius crassus (Bramlette and Sullivan, 1961) Perc-Nielsen, 1984

Neonannarina cristata (Martini, 1958) Perc-Nielsen, 1971
Chiasmolithus danicus (Brotzen, 1959) Hay and Mohler, 1967
Reticulofoenestra daviesii (Hay, Mohler, and Wade, 1966) Roth, 1970
Discosphaera deflandrei Bramlette and Riedel, 1954
Amaurolithus delicatus Gartner and Bukry, 1975
Reticulofoenestra dictydota (Deflandre in Deflandre and Fert, 1954) Stradner in Stradner and Edwards, 1973
Stradner in Stradner and Edwards, 1968
Neochiastozygus digitatus Perch-Nielsen, 1971
Helicosphaera dinesenii Perch-Nielsen, 1971
Discosphaera Tan, 1927
Neochiastozygus distentus (Bramlette and Sullivan, 1961) Perch-Nielsen, 1971
Ellipsolithus distichus (Bramlette and Sullivan, 1961) Perc-Nielsen, 1971
Helicosphaera euphratis Hay, 1966
Chiasmolithus expansus (Bramlette and Sullivan, 1961) Gartner, 1970
Cycloriococchi floraeanus (Roth and Hay in Hay, Mohler, Roth, Schmidt, and Boudreaux, 1967) Bukry, 1973
Coccolithus formosus (Kampfer, 1963) Wise, 1973
Sphenolithus furcatolithoides Locker, 1967
Toweius gommae (Bramlette and Sullivan, 1961) Romein, 1979
Chiasmolithus gigas (Bramlette and Sullivan, 1961) Radomski, 1968
Gephyrocapsa Kampfer, 1943
Rhabdosphaera gladius Locker, 1967
Chiasmolithus granulatus (Bramlette and Riedel, 1954) Radomski, 1968
Biscutum harrisonii Varol, 1989
Helicosphaera Kampfer, 1954
Pyrocycloïdes hermosus Roth and Hay in Hay, Mohler, Roth, Schmidt, and Boudreaux, 1967
Sphenolithus heteromorphus Deflandre, 1953
Reticulofoenestra hillae Bukry and Percival, 1971
Emilianna huxleyi (Loehmann, 1902) Hay and Mohler in Hay, Mohler, Roth, Schmidt, and Boudreaux, 1967
Markallus inversus (Deflandre in Deflandre and Fert, 1954) Bramlette and Martini, 1964
Pyrocycloïdes inversus Hay and Towe, 1962
Neochiastozygus jucnus (Bramlette and Sullivan, 1961) Perc-Nielsen, 1971
Heliolithus kleinielli Sullivan, 1964
Discosphaera kuepperi Stradner, 1959
Pseudonannolithus laconus (Kampfer, 1963) Gartner, 1969
Cricoplacolithus latipons Romein, 1979
Discosphaera lodoeensis Bramlette and Riedel, 1954
Cyclicargolithus luminis (Sullivan, 1965) Bukry, 1971
Calcisphaera macintyrei (Bukry and Bramlette, 1969) Loeblich and Tappan, 1978
Toweius magnificus (Bukry, 1971) Romein, 1979
Prinsus martini (Perch-Nielsen, 1969) Haq, 1971
Discosphaera moelri Yuky and Percival, 1971
Sphenolithus moriformis (Bronnimann and Stradner, 1960) Bramlette and Wilcoxen, 1967
Discosphaera multiarallata Bramlette and Riedel, 1954
Neochiastozygus Perc-Nielsen, 1971
Chiasmolithus oamaruensis (Deflandre in Deflandre and Fert, 1954) Hay, Mohler, and Wade, 1966
Reticulofenestra oamaruensis (Deflandre in Deflandre and Fert, 1954) Stradner in Haq, 1968
Thoracosphaera operculata Bramlette and Martini, 1964
Tribrachiatus orthostylus Shamrai, 1963
Coccolithus pelagicus (Wallich, 1877) Schiller, 1930
Discosphaera pantaradatus Tan, 1927 endem. Bramlette and Riedel, 1954
Reticulofenestra perplexa (Burns, 1975) Wise, 1983
Toweius pertusus (Sullivan, 1965) Romein, 1979
Cruciplacolithus primus Perch-Nielsen, 1977
Calcisphaera protoannulus (Gartner, 1971) Loeblich and Tappan, 1978
Neococcolithus proterus (Bramlette and Sullivan, 1961) Black, 1967
Sphenolithus pseudoradians Bramlette and Wilcoxen, 1967
Reticulofoenestra pseudoumbilicus (Gartner, 1967) Gartner, 1969
Sphenolithus radians Deflandre in Grasse, 1952
Isthmiolithus recurvus Deflandre in Deflandre and Fert, 1954
Cribrocentrum reticulatum (Gartner and Smith, 1967) Perl-Nielsen, 177
Heliolithus riedelii Bramlette and Sullivan, 1961
Neococcolithus rimosus (Bramlette and Sullivan, 1961) Varol, 1989
Discosphaera robustus Haq, 1969
Discosphaera saipanensis Bramlette and Riedel, 1954
Scampanella Forchheimer and Stradner 1973
Helicosphaera seminulum Bramlette and Sullivan, 1961
Euriolyssolithus selli Bukry and Bramlette, 1969
Placozoogyx sigmoideae (Bramlette and Sullivan, 1961) Romein, 1979
Chiasmolithus solitus (Bramlette and Sullivan, 1961) Loeblich, 1968
Sphenolithus Deflandre in Grasse, 1952
Blackites spinosus (Deflandre in Deflandre and Fert, 1954) Hay and Towe, 1962
Clasticococcus subdistichus (Roth and Hay, 1965) Prins, 1979
Discosphaera subiodoensis Bramlette and Sullivan, 1961
Coccolithus subpertusus (Hay and Mohler, 1967) Van Heck and Prins, 1987
Discosphaera surculus Martini and Bramlette, 1963
Discosphaera tani nodifer Bramlette and Riedel, 1954
Cricoplicolithus tenellus (Stradner, 1961) Hay and Mohler in Hay, Mohler, Roth, Schmidt, and Boudreaux, 1967
Hornbrookina teuriensis Edwards, 1973
Thoracosphaera Kampfer, 1927
Chiasmolithus titus Gartner, 1970
Toweius tovae Perch-Nielsen, 1971
Amaurolithus delicatus Gartner and Bukry, 1969
Cricoplacolithus tymaniformis (Sullivan, 1964) Hay and Mohler, 1967
Cricoplacolithus vanheckiae Perch-Nielsen, 1984
Discosphaera wemmelensis Achuthan and Stradner, 1969