4. LOWER CRETACEOUS FORAMINIFERAL BIOSTRATIGRAPHY, PALEOECOLOGY, AND DEPOSITIONAL ENVIRONMENT AT DSDP SITE 397, LEG 47A

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ABSTRACT

Biostratigraphic results of foraminiferal studies for Site 397 indicate a probable Hauterivian-Barremian through Barremian-Aptian age for the Lower Cretaceous sedimentary section (Unit 5, Cores 52A-35A). Typical index species of the planktonic foraminifers include *Hedbergella hoterivica*, *Hedbergella sigali*, *Calvihedbergella aff. simplex*, and *Globigerinelloides blowi*. Important benthic species include *Epistomina caracolla*, *Epistomina hechti*, *Epistomina cretosa*, *Epistomina ornata*, *Lenticulina nodosa*, *Lenticulina saxonica*, *Lenticulina praegaultina*, *Lenticulina heiermanni*, *Lenticulina matilatus*, and *Conorboides* sp. Sedimentologic and ecologic data from the microfossils suggest prodelta shelf depth merging into upper bathyal environments. Benthic foraminifers (such as *Quinqueloculina* associated with ostracodes and small gastropods and pelecypods) may indicate a nearshore environment in front of a deltaic complex. Epistominids (such as *Epistomina caracolla* and the nodosariids such as *Lenticulina*, *Dentalina*, and *Planularia*) indicate an outer-shelf to upper bathyal environment. However, hedbergellids and a few radiolarian specimens suggest the epipelagic setting of an open-shelf water mass, and can be compared with Recent shelf pelagic biota. Paucity of fauna indicates neither stagnant bottom conditions (pelagic sapropel) nor oceanic depths below the calcite compensation depth (CCD). The barren character of the facies may indicate dilution by the sediments due to continuous terrigenous (cyclic sedimentation) input into the shelf basin. This in turn caused instability of the biophysiochemical conditions. Scarcity of the pelagic constituents in the sediment may also reflect low plankton productivity, possibly related to a restricted circulation due to the paleogeographic setting of the macroplates during the Early Cretaceous.

INTRODUCTION

During Leg 47A, Site 397 was drilled on the uppermost continental rise offshore from Cape Bojador. One of the main objectives at this site was to reconstruct the geologic history and evolution of the passive continental margin in the eastern North Atlantic during the Mesozoic. Through continuous drilling, about 1453 meters of sedimentary section were recovered at a water depth of 2900 meters nearly 100 km offshore from the Tarfaya-Aaiun coastal region. The lower part of the sedimentary section (153 m) is mainly of Barremian age and is separated from the Neogene at 1300 meters by a large hiatus (approximately Aptian-earliest Miocene). In this report, only the Cretaceous part of the section (Cores 52 to 34 of Hole 397A) is discussed. This section is defined lithologically as Unit 5 (see Site Report, this volume) and consists mainly of hemipelagic laminated strata consisting of fine-grained gray greenish to dark gray silty mudstones with occasional dolomitic and sideritic intercalation (Einsel and von Rad, this volume). Microfossils such as foraminifers are rare, and several samples are completely barren. The scarcity of fauna has been explained by stagnation of the bottom waters or by deposition below the CCD (Einsel and von Rad; Basov et al.; Roesler et al.; all this volume). The ecologic interpretations made in this report (which are based on a study of 150 samples from Cores 52 to 34 of Hole 397A) do not favor either of these views. Important disagreements also exist concerning the exact biostratigraphic position of the Lower Cretaceous sediments at Site 397 (see Site Report, this volume). On the basis of nannoplankton ranges, Čepík and Wind (this volume) propose a middle-late Hauterivian age, while Basov et al. (this volume) and Wiedmann (this volume) suggest Barremian-Aptian and Hauterivian-Barremian ages, respectively, on the basis of studying foraminifers and ammonites. However, the biostratigraphic results of this study support the views of Basov et al. and Wiedmann. However, three fundamental differences do exist: (1) disagreement of foraminiferal and nannoplankton results in general, (2) first appearance of the
genus *Favusella* and its association to pre-Albian planktonic foraminifers such as *Hedbergella hoterivica*, (3) the lack of an SEM study of the type specimens of *Hedbergella hoterivica*.

Because of the above conflicting results about Unit 5, the author has attempted to present a synthesis by comparing the sediments at Site 397 with the onshore Cretaceous sections in the Tarfaya-Aaui and Agadir-Essaouira regions of northwestern Africa.

**BIOSTRATIGRAPHY**

**Results**

Between 2 and 3 samples per section of Cores 52 to 34 of Hole 397A were analyzed to study their foraminiferal composition. Unfortunately, most of the samples were found to be barren, consisting mainly of terrigenous quartz and subordinate mica and pyrite grains. Plant fragments were common in the washed sample residue. In sections 47-1, 46-3, 46-2, and 42-2 (Figure 1) maximum numbers (30 to 50 specimens per sample), of microfossils including foraminifers, ostracodes, pelecypods, gastropods, radiolarians, and fish debris. The abundance of microfossils roughly correlates with the relative abundance of calcium carbonate in the cores (see Site Report, this volume). Foraminifers are generally small, but nodosariids such as *Lenticulina* display a normal size and moderately high diversity. Planktonic foraminifers include small fragile primitive “globigerine forms,” placed here under the genus *Hedbergella*, although some specimens of the genus *Globigerinelloides* could also be identified. The planktonic specimens are generally well preserved, but some show diagenetic corrosion of the shell wall. Furthermore, the faunas do not show any sign of reworking or hydrodynamic sorting. Stratigraphic ranges of the foraminifers and other microfossils in Cores 52 to 34 of Hole 397A are shown in Figure 1, while typical representative specimens are figured in Plates 1 to 4.

**Discussion of Results**

*Hedbergella hoterivica* (Ascoli, 1976, renamed it as *Caucasella hoterivica*; synonym *Globigerina kugleri* Bolli) in Hole 397A first appears in Sample 51A-1, 29-35 cm; while *Epistomina caracolla* first occurs in Sample 52A-2, 6-10 cm; *Epistomina cretosa* in Sample 50A-5, 21-25 cm; and *Epistomina hechti* and *Lenticulina saxonica* in Sample 50A-4, 95-100 cm. These suggest a Hauterivian-Barremian age. Supporting evidence for lowermost Barremian in Sample 50A-2, 0-6 cm comes with the occurrence of *Epistomina hechti* (aff. *Epistomina spinulifera*), *Lenticulina nodosa*, *Lenticulina praegigantula*, and *Conchorbodes* sp. This assemblage, along with *Hedbergella hoterivica* and *Epistomina caracolla*, may suggest the Barremian-Hauterivian boundary in Core 50A (Figure 1 and Plates 1 to 4).

The presence of *Lenticulina heiermanni* in Sample 47A-4, 51-55 cm; *Lenticulina matilatus* and *Lenticulina* sp. 1 in Sample 46-3, 145-149 cm; and *Hedbergella sigali*, *Clavihedbergella aff. simplex*, *Globigerinelloides blowi*, and *Hedbergella hoterivica* in Sample 46-2, 101-105 cm confirms the Barremian age of Cores 50 to 46 in Hole 397A. This conclusion agrees with van Hinte’s (1976) zonal scheme of the Cretaceous. Similar species from land deposits are described by several authors (Subbotina, 1953; Bolli, 1959; Moullade, 1966; Fuchs, 1971). *Hedbergella hoterivica*, *Hedbergella sigali*, *Clavihedbergella aff. simplex*, *Globigerinelloides blowi*, and a single specimen of *Frondicularia* sp. were identified again in Section 42-2 (Plates 1-4). In contrast to Cores 47, 46, and 42 of Hole 397A, Cores 41 to 35 contain very rare specimens of *Hedbergella hoterivica* and *Epistomina caracolla*, while Core 34 is contaminated throughout by Tertiary foraminifers. Sample 42-2, 5-10 cm consists of relatively abundant and diverse post-Hauterivian planktonic species (nearly 30 specimens). The relative abundance of *Hedbergella sigali* and *Globigerinelloides blowi* may suggest a Barremian-Aptian age for Cores 42 to 35. Due to the extreme contamination of Core 34, no age designation can be made.

Bolli (1959) described *Globigerinelloides blowi* as *Planomalina blowi* from ?Barremian-Aptian and *Globigerina kugleri* (synonym of *Hedbergella hoterivica*) from Barremian-Aptian sediments in Trinidad. *Globigerina hoterivica*, identified here as a species of the genus *Hedbergella*, was originally described by Subbotina (1953) from upper Hauterivian sediments in the northern Caucasus. Ascoli (1976) mentioned the same species as *Caucasella hoterivica* from the Valanginian-Barremian sediments in the Scotian Shelf of the western North Atlantic. Thus, the occurrence of *Hedbergella hoterivica* can be correlated on both sides of the North Atlantic. Thus the Lower Cretaceous benthic foraminifers, similar to the planktonics, indicate an identical ecologic niche in both eastern and western realms of the North Atlantic (Ascoli, 1976; Bartenstein, 1976; Butt in Wiedmann et al., 1978).

Daily (1973) described *Hedbergella hoterivica* as *Globigerina hoterivica* from Hauterivian-Barremian sediments in the Sacramento Valley of California, while Michael (1967) and Fuchs (1971) identified it in Barremian sediments in northwestern Germany and Vorarlberg (Austria), respectively. However, Dailey mentioned that some of these specimens of *Hedbergella hoterivica* show coarse perforations. It is still unknown whether these perforations in the shell wall of Dailey’s specimens correspond to the reticulated structure of the Albian genus *Favusella* of Michael (1972). Morphologically the specimens of *Hedbergella hoterivica* from Site 397 display a moderately low trochospiral test with two-and-a-half whorls of globular chambers. The spiral side is somewhat convex; the ventral convex side shows four visible chambers which rapidly increase in size and have a reduced umbilical area, as well as a large primary arch-shaped aperture (Plate 3, Figure 4). Most of the specimens on the umbilical side contain a small bulla-like fifth or last chamber, with an extra-umbilical aperture covering the entire umbilicus and primary aperture. Through the
### Figure 1. Distribution of Lower Cretaceous microfossils at Site 397.

<table>
<thead>
<tr>
<th>Upper Hauterivian-Lower Barremian</th>
<th>Barremian</th>
<th>Barremian-Aptian?</th>
<th>Stage</th>
</tr>
</thead>
<tbody>
<tr>
<td>Core</td>
<td>Section</td>
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- *H. hotavrica* 259
- *H. sigali* 259
- *C. simplex* 259
- *Globigerinelloides blowei* 259
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**Ostracodids**

**Pelecypods/Gastropods**

**Radiolarians**

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*Lower part of Upper Hauterivian*

- *Crucillites curtilli* (Barrian to Hauterivian)
- *Nannoconus kampteri* (Late Hauterivian) *Calcisphaera ablongata* (Early Barremian)

*Hauterivian-Barremian*

- Section 42-1: *N. mecomiensis* (Hauterivian)
- Section 39-2: *P. crebrilobicus* (Barremian-Aptian)

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- *Nannoplankton* (Copac and Wind, this volume)
- *Ammolites* (Viebemann, this volume)
light microscope, the calcareous wall of the specimens shows fine pores giving the impression of a smooth surface; but, through SEM study, the specimens reveal a coarse reticulate honeycomb sculpture with 3 to 4 (or more) pores in each polygonal structure (Plate 4, Figures 2 and 6), resembling the genus *Favusella* of Michael (1972). On the basis of these characters, Roesler et al. (this volume) called these specimens as new species: *Favusella stiftia*. The specimens also show weak to strong coarsening of the ornamentation (Plate 4), though some specimens display various destroyed stages of the reticulated shell wall (Plate 3, Figures 5 and 6). Still others consist only of calcareous casts (Plate 3, Figure 7). The test wall similarity of *Hedbergella hoterivica* with *Favusella* confirms the earlier opinion of Bolli (1959) and Bandy (1967) that *Globigerina kugleri* (here named *Hedbergella hoterivica*) gave rise to the Albian *Globigergerina washitensis* (renamed by Michael, 1972, *Favusella washitensis* type species of the genus *Favusella*) by developing a high spire, adding one more chamber in the last whorl, increasing in size, and developing a honeycomb sculpture. In view of the above morphologic affinity, it is not necessary to revise this species nor to erect a new one since the coarse sculpture may not be visible, under the light microscope, and the surface ornamentation often is not preserved leaving behind a calcareous mold or cast (compare Plate 3, Figure 7). Moreover, the global appearance of the genus *Favusella* also seems questionable. For instance, Ascoli (1976) reported *Favusella aff. washitensis* from Aptian sediments in the Scotian Shelf of northwestern Atlantic. It is interesting to note that Ascoli's figure of *Caucasella hoterivica* from the Valanginian (pl.1., fig. 3C of Ascoli, 1976) shows a coarse reticulated structure of the shell wall. This suggests that *Hedbergella hoterivica* is probably an ancestor of the so-called genus *Favusella*, and may be regarded as "prae Favusella" (Koch, personal communication).

**PALEOECOLOGY AND DEPOSITIONAL ENVIRONMENTS**

The general scarcity of microfossils in the Lower Cretaceous sediments at Site 397 caused confusion regarding their paleobathymetry. Two main views about the paleoenvironments have been presented: (1) paucity of the fossils indicates depths below the CCD, and (2) an expanded oxygen-minimum zone and stagnation of the bottom waters eliminated benthic life. Alternatively, based on the foraminiferal association, an outer shelf, upper bathyal depth range is suggested for the deposition of the Lower Cretaceous sediments. The sediments accumulated as a distal delta facies (Einsele and von Rad, this volume). This indicates a continuous high rate of terrigenous input (cyclic sedimentation), resulting in a scarcity of nutrients and biogenic calcite in the basin. Scruton (1955) noticed a scarcity of foraminifers (0.9%) in the modern prodelta sediments of the Mississippi Delta, where the dominant sediment components accompanying the fraction were mainly terrigenous sand and plant fragments.1

Miliolids such as *Quinqueloculina*, together with ostracodes, small gastropods, and peliscopods (Figure 1); Plate 1, Figures 24 to 27; Plate 2, Figures 13 to 17, 21, and 22), suggest proximity of the near-shelf shelf environment. Barren samples usually contain abundant terrigenous quartz and occasionally broken shell fragments. These constituents may suggest relatively higher rates of sedimentation, possibly related to climatic fluctuations on the nearby continent. The samples containing abundant foraminifers may represent periods of slow quiescent sedimentation, with an attendant environment of normal salinity, temperature, and an oxygenated bottom. The ostracode assemblage (Plate 2) exhibits a morphologic affinity with neritic assemblages reported from various land sections (e.g., see Brenner and Oertli, 1976). These ostracodes (Brenner and Oertli, 1976) are associated with foraminifers such as *Lenticulina heteromanni* and *Epistomina caracollia* as found in Cores 52 to 35.

Epistominids (such as *Epistomina caracollia*) and nodosariids (such as *Lenticulina, Astacolus, Planulalia*, etc.) may suggest an outer-shelf upper bathyal environment. *Epistomina caracollia* seems to be a homeomorph of a recent species: *Hoeglundina elegans*. According to Bandy (1967), this is an important isobathyal species which typically occurs at the shelf edge extending down to bathyal depths. Comparing the habitat of the Cretaceous foraminifers with recent homeomorphs, Sliter (1972) suggested that an outer-shelf assemblage is represented by nodosariids and *Hoeglundina*, though several genera such as *Gyroidina* (which Sliter included in the outer-shelf range) are not known globally in Lower Cretaceous sediments. The hedbergellids and globigerinellids at Site 397 may indicate epipelagic activity of a shelf water mass. In the study of the genus *Favusella*, Michael (1972) pointed out that this genus is usually reported in neritic deposits. Butt (in Wiedmann et al., 1978) reported *Favusella* (as *Hedbergella washitensis*) in Albian-Cenomanian shelf deposits of Agadir-Essaouira coastal sediments. It is concluded, therefore that *Hedbergella hoterivica* showing an affinity to *Favusella* may represent a pelagic shelf association.

**ONSHORE AND OFFSHORE CORRELATION**

*Tarfaya-Aaiun Region*

In contrast to the offshore prodelta facies at Site 397, the onshore Barremian-Aptian sediments from the Tarfaya-Aaiun region and a commercial well offshore from the Spanish Sahara represent a more than

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1In view of age determination of the Lower Cretaceous strata at Site 397, the author prefers to retain the name of the illustrated specimens (Plates 3,4) as *H. hoterivica*; however, it is likely that these specimens represent a subspecies of *H. hoterivica* bearing reticulated structures.
1200-meter-thick clastic sequence of a deltaic complex (CONOCO, 1969; Wiedmann et al., 1978; Einsele and von Rad, this volume). These sediments contain rare microfossils. The fauna consists mainly of ostracodes, pelecypods, gastropods, and echinoids. Lituolids (such as Cheffattella decipiens) are the only representative foraminifers mentioned in the unpublished data of CONOCO (1969). This onshore faunal association represents lagoonal to outer littoral environments relative to the offshore distal delta facies. Farther east, the landward outcrop data from the Tarfaya-Aaiun region indicates a lithofacies relation to sediments cored in the CONOCO well, representing fluviatile, lagoonal, and tidal deposits (Wiedmann et al., 1978). There, Aptian-lower Albian faunas consist mainly of abundant ostracodes and small mollusks. Foraminifers include agglutinated benthics such as Trochammina and Haplophragmoides. This faunal association provides evidence for a relative low salinity lagoonal environment in the Tarfaya-Aaiun onshore deltaic system. The above-mentioned Lower Cretaceous sediments, representing a shallow marine depositional environment, grade upward into a fine-grained sequence of hemipelagic sediments with interbedded shell beds in the Albian-Cenomanian. Overlying eupelagic sediments display laminated chert layers in the upper Cenomanian-Turonian (Wiedmann et al., 1978). Biogenic components of the sedimentary succession suggest a gradual paleodepth increase, indicating submergence of the continental margin under deeper water during the Middle Cretaceous.

**Agadir-Essaouira Region**

In contrast to the Tarfaya-Aaiun region, the Lower Cretaceous sediments of the Agadir-Essaouira region display an intertonguing of marine and non-marine sediments. Continental facies (such as clastic sediments and red beds) and marine shelf facies (such as silty marls and shell beds) reveal several progressive and transgressive episodes. Following Upper Jurassic shallow-water carbonate sedimentation, the Agadir and Essaouira basins subsided to a relative deep shelf environment. Faunal associations represent abundant, and large nodosarids, including Lenticulina nodosa, Lenticulina eichenbergi, Lenticulina guttata, Lenticulina ouachensis, Planularia crepidularis, Vaginulina schreiteri, etc. Epistominas (such as Epistoma coracolla), agglutinants (such as lituolids), and ostracodes frequently occur (Butt, in Wiedmann et al., 1978). This faunal association indicates an ecologic affinity with the foraminifers at Site 397. During the middle Hauterivian, the Agadir and Essaouira basins were marked by a local regression. In particular, the Essaouira Basin contains a red bed facies, comprising abundant brackish water foraminifers (Ammobaculites-Haplophragmoides).

During the late Hauterivian-Barremian, marine shelf conditions again prevailed, while the ecological spectrum is mainly represented by nodosarids as mentioned above. In particular, the Barremian sediments consist of relatively diverse planktonic foraminifers (such as Hedbergella infracretacea and Hedbergella signali), indicating biostratigraphic and paleoecologic affinities with the offshore fauna at DSDP Sites 397 and 370. But, during late Barremian-early Aptian, the coastal basins record another regression. Biocomponents such as charophytes, ostracodes, and agglutinated foraminifers there may indicate shallow marine to brackish environments (lagoonal-littoral zone) and a related lowering of sea-level. The upper Aptian-Albian sediments in the Agadir and Essaouira basins mark the beginning of a Middle Cretaceous transgression that later reached a maximum during the Turonian in both onshore basins. However, local shallowing trends (see bathymetric curves in Wiedmann et al., 1978) during the late Albian and late Cenomanian in the Agadir Basin and late Albian through Cenomanian shallowing (calcariaceous platform sedimentation) in the Essaouira Basin may suggest lateral bathymetric variations in the local coastal basins.

**Discussion of Results**

Onshore Lower Cretaceous sediments in both the Agadir-Essaouira and the Tarfaya-Aaiun regions reveal marine and non-marine sedimentary sequences. These facies display a deltaic character and can be correlated in terms of foraminiferal ecology and biostratigraphy with the offshore distal delta facies described as “turbidites” or “laminates” at DSDP Sites 370 and 397. In contrast to present depths at Sites 370 and 397, which are approximately 4000 and 3000 meters, respectively, the Lower Cretaceous foraminiferal association indicates a shallower paleodepth in a marginal setting of the eastern North Atlantic. These are possibly equivalent to outer-shelf and upper bathyal water depth levels (500 m). During the Middle Cretaceous, however, the offshore depositional sites probably subsided rapidly to bathyal-abyssal depth levels. Interestingly, this subsidence of the sea floor correlates with a synchronous Middle Cretaceous transgression and a related eustatic sea-level rise in the onshore basins (compare Sliter, 1976), eliminating the Lower Cretaceous shallow-water deltaic environments along the ancient coasts.

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PLATE 1

Figure 1  Spirillina sp.; Sample 42-2, 5-10 cm (150×).
Figure 2  Dentalina sp. 1; Sample 46-4, 80-84 cm (100×).
Figure 3  Dentalina sp. 2; Sample 46-3, 50-55 cm (50×).
Figure 4  Vaginulina sp. 2; Sample 42-2, 5-100 cm (150×).
Figure 5  Dentalina sp. 3; Sample 46-3, 50-55 cm (50×).
Figure 6  Astacolus sp. 1; Sample 60-4, 95-100 cm (100×).
Figure 7  Astacolus sp. 2; Sample 50-2, 15-19 cm (150×).
Figure 8  Vaginulina sp. 1; Sample 46-4, 50-55 cm (75×).
Figure 9  Astacolus sp. 3; Sample 50-1, 114-119 cm (100×).
Figure 10  Planularia sp. 2; Sample 39-2, 44-47 cm (75×).
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Figure 23  Pseudonodosaria sp.; Sample 46-4, 0-5 cm (150×).

Figures 24-27  Miliolids
24. Quinqueloculina sp.; Sample 50-3, 130-135 cm (150×).
25. Quinqueloculina sp.; Sample 48-2, 30-35 cm (150×).
26. Quinqueloculina sp.; Sample 50-3, 0-6 cm (150×).
27. Quinqueloculina sp.; Sample 50-4, 145-150 cm (150×).
PLATE 2

Figure 1  *Quinqueloculina* sp. showing enlargement of the apertural area; Sample 50-3, 130-135 cm (925×).

Figure 2  *Epistomina caracolla* (Roemer) ventral view; Sample 50-4, 145-150 cm (100×).

Figure 3  *Epistomina caracolla* (Roemer) dorsal view; Sample 46-3, 100-104 cm (100×).

Figure 4  *Epistomina hechti* Bartenstein, Bettenstaedt and Bolli; dorsal view; Sample 50-2, 0-6 cm (115×).

Figure 5  *Epistomina caracolla* (Roemer); ventral view (pyritized shell); Sample 46-3, 100-104 cm (100×).

Figure 6  *Epistomina* sp.; ventral view; Sample 50-4, 95-100 cm (150×).

Figure 7  *Epistomina hechti* Bartenstein, Bettenstaedt, and Bolli; dorsal view; Sample 50-5, 21-25 cm (100×).

Figure 8  *Epistomina cretosa* Ten Dam; dorsal view (compare Ascoli, 1976); Sample 50-5, 21-25 cm (100×).

Figure 9  *Conorboides* sp.; dorsal view; Sample 50-2, 0-6 cm (150×).

Figure 10  *Epistomina* sp.; ventral view; Sample 50-1, 80-85 cm (150×).

Figure 11  Ostracode; Sample 50-1, 80-85 cm (75×).

Figure 12  *Epistomina ornata* (Roemer) ?; Sample 50-1, 114-119 cm (150×).

13. *Orthonotacythere* ?; Sample 50-4, 95-100 cm (100×).
15. *Acrocythere* ?; Sample 41-2, 105-109 cm (100×).
17. *Schuleridea* sp. ?; Sample 50-2, 15-19 cm (50×).

Figure 18  Fish otolith; Sample 43-3, 25-30 cm (150×).

Figures 19, 20  Radiolarian specimens; Sample 46-3, 100-104 cm.
19. (150×).
20. (150×).

Figure 21  Micro-gastropod; Sample 47-3, 80-84 cm (150×).

Figure 22  Micro-pelecypod; Sample 50-2, 0-6 cm (100×).

Figure 23  Coccoliths attached to the shell wall of a planktonic foraminiferal chamber; Sample 46-3, 145-149 cm (3250×).
PLATE 3

Figures 1, 2  *Hedbergella hoterivica* (Subbotina); Sample 50-1, 80-85 cm.
1. Spiral view (250×).
2. Detail of the test wall (1500 ×).
   Note polygonal structures; pores are not visible.

Figure 3  *Hedbergella hoterivica* (Subbotina); umbilical view, showing small bulla-like chamber; Sample 46-3, 145-149 cm (250×).

Figure 4  *Hedbergella hoterivica* (Subbotina); umbilical view; showing primary umbilical aperture; Sample 47-3, 23-26 cm (250 ×) (compare Ascoli, 1976).

Figures 5, 6  *Hedbergella hoterivica* (Subbotina); Sample 47-4, 51-55 cm.
5. Spiral view, showing diagenetic destruction of the reticulated wall (250×).
6. Detailed view of the shell wall as in Figure 5 (2500×).

Figure 7  *Hedbergella hoterivica* (Subbotina); spiral view showing complete removal of the reticulated shell wall (Steinkern). This specimen resembles some land specimens of *H. hoterivica* showing smooth surface; Sample 46-2, 100-105 cm (250×).

Figure 8  *Hedbergella* sp.; Sample 42-2, 5-10 cm (250×).

Figure 9  *Hedbergella sigali* Moullade; spiral view; Sample 46-2, 101-105 cm (250×).

Figure 10  *Hedbergella sigali* Moullade; spiral view; Sample 42-2, 5-10 cm (200×).

Figure 11  *Clavihedbergella* aff. *simplex* (Morrow); spiral view; Sample 42-2, 5-10 cm (200×).

Figure 12  *Clavihedbergella* aff. *simplex* (Morrow); spiral view; Sample 46-2, 101-105 cm (250×).

Figure 13  *Clavihedbergella* sp.; spiral view; Sample 46-4, 80-84 cm (375×).

Figure 14  *Globigerinelloides blowi* (Bolli); side view; Sample 42-2, 5-10 cm (200×).

Figure 15  *Globigerinelloides blowi* (Bolli); peripheral view; Sample 42-2, 5-10 cm (200×).

Figure 16  *Globigerinelloides blowi* (Bolli); side view; Sample 42-2, 5-10 cm (200×).

Figure 17  *Globigerinelloides blowi* (Bolli); side view; Sample 42-2, 5-10 cm (200×).
Figures 1, 2  *Hedbergella hoterivica* (Subbotina), showing similarity of the ultrastructure with genus *Favusella* of Michael (1972); Sample 47-1, 10-14 cm.
1. Umbilical view; note bulla-like last chamber covering the umbilicus (350×).
2. Details of the shell wall; note polygonal structure with small pores (350×).

Figures 3, 4  *Hedbergella hoterivica* (Subbotina); Sample 47-3, 23-26 cm.
3. Umbilical view; note the last reduced chamber shows a lip-like extension, covering the primary umbilical aperture (350×).
4. Umbilical details as in Figure 3 (1400×).

Figures 5, 6  *Hedbergella* sp. aff. *H. Hoterivica* (Subbotina); Sample 46-2, 101-105 cm.
5. Spiral view (200×).
6. Micro-details of the shell wall as in Figure 5; note a lesser coarsening of the polygonal structures than in Figures 1 to 4, above (3500×).