5. RADIOLARIAN FAUNAL CHARACTERISTICS IN OLIGOCENE SEDIMENTS OF THE KERGUELEN PLATEAU, LEG 183, SITE 1138¹

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ABSTRACT

Three sites from Ocean Drilling Program (ODP) Leg 183 (Kerguelen Plateau) have been analyzed to document faunal change in highlatitude radiolarians and to compare the faunal change to Eocene-Oligocene climatic deterioration. Radiolarians are not preserved in Eocene sediments. In Oligocene sediments, radiolarian preservation improves in a stepwise manner toward the Miocene. A total of 115 species were found in lower Oligocene samples from Site 1138; all are documented herein. Radiolarian preservation is presumably linked to productivity triggered by climatic cooling during the early Oligocene. Similar patterns of improving preservation through the Eocene/Oligocene boundary are documented from several Deep Sea Drilling Project and ODP sites in the Southern Ocean, indicating a general pattern. In contrast to the Southern Kerguelen Plateau, however, proxies for productivity are more divergent at Site 1138 (Central Kerguelen Plateau). Whereas carbonate dissolution, as indicated by poor preservation of foraminifers and common hiatuses, is very pronounced in the upper Eocene-lowermost Oligocene, the quality of radiolarian and diatom preservation does not significantly increase until the uppermost lower Oligocene. Multiple measures of radiolarian diversity in the Oligocene from Site 1138 closely parallel radiolarian preservation, indicating that preserved radiolarian diversity is controlled by productivity.

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INTRODUCTION

The role of the Eocene–Oligocene transition in Cenozoic radiolarian evolution is still insufficiently known, especially when compared to that for other microplankton groups such as planktonic foraminifers and calcareous nannoplankton. Some faunal change has been observed in low latitudes (Riedel and Sanfilippo, 1986; Sanfilippo et al., 1985), but it appears minor in comparison to the severe taxonomic turnover reported in planktonic foraminifers (Keller et al., 1992) and calcareous nannoplankton (Aubry, 1992). The climatic deteriorations around the Eocene/Oligocene boundary usually play a major role in the scenarios explaining plankton changes in the Southern Ocean. Most commonly, changes are attributed to an increase in productivity (Baldauf, 1992; Baldauf and Barron, 1990; Diester-Haass, 1995), eventually favoring opal preservation and promoting carbonate dissolution.

As part of a larger project concerning radiolarian faunal change, the original aim of this study was to document the radiolarian faunal change during the late Eocene to the early Oligocene in the Southern Ocean in relation to the outstanding climate change during this time period. Previous studies were devoted mostly to developing a stratigraphic zonation (Caulet, 1991; Takemura, 1992; Takemura and Ling, 1997) but did not characterize overall faunal change in relation to Southern Ocean cooling (Zachos et al., 1999). So far, this has only been done by Lazarus and Caulet (1993), who reported a considerable faunal turnover.

Unfortunately, no radiolarians of Eocene age are preserved in any of the studied sites. Thus, our study is limited to the Oligocene suite and can only document faunal changes within this interval. We document qualitative and quantitative radiolarian faunal characteristics through a large part of the Oligocene and relate this to published paleoclimatic and productivity data. Taxonomy, stratigraphy, diversity, abundance, and preservation of radiolarians are described. A significant accomplishment of this study is the documentation, albeit preliminary, of the complete recovered Oligocene radiolarian assemblage, including (in open nomenclature) several previously undescribed species.

SAMPLES AND METHODS

During Ocean Drilling Program (ODP) Leg 183, the Eocene/Oligocene boundary interval was drilled four times (Table T1). Samples from three sites (1138, 1139, and 1140) (Fig. F1) were studied and, according to shipboard stratigraphic results, contain a sedimentary record of the Eocene/Oligocene boundary. A total of 64 samples were studied for this paper.

Only Hole 1138A (53°33.105′S, 75°58.493′E; Central Kerguelen Plateau) yielded a good radiolarian record within the time interval of interest, and, even at this site, only Oligocene radiolarians could be identified. The interval is situated within lithofacies Unit III as identified by the Shipboard Scientific Party (2000). This unit ranges in age from about mid-Campanian to late Oligocene and consists of white to light gray and light greenish gray foraminifer-bearing nannofossil chalk. According to the current age model (Fig. F2), the Eocene– Oligocene time period is represented in Cores 183-1138A-29R through 48R. The base of the cored interval for Core 183-1138A-36R (top of Chron C13r) is interpreted as the Eocene/Oligocene boundary, **T1.** General data of Leg 183 sites, p. 31.

F1. Kerguelen Plateau and Leg 183 sites, p. 22.







but there is no core recovery. As indicated by calcareous nannoplankton and paleomagnetic studies, a hiatus or condensed section is present close to the Eocene/Oligocene boundary (Shipboard Scientific Party, 2000). Glauconite-filled foraminifers in Core 183-1138A-36R are further evidence for stratigraphic condensation. No radiolarians were found below Core 183-1138A-36R (Fig. F3). The oldest poorly preserved radiolarians are observed in Sample 183-1138A-36R-1, 41– 43 cm (Chron C11r; early Oligocene; ~30.2 Ma). Preservation gradually improves upward through Core 183-1138A-35R and is generally good in Core 183-1138A-34R.

At Site 1139, 15 samples from the upper Eocene–lower Oligocene (Sections 183-1139A-38R-2 to 41R-1) do not contain identifiable radiolarians (Fig. F4). The radiolarian record starts in the uppermost Oligocene (Core 183-1139A-19R), and well-preserved faunas are identified (Cores 183-1139A-17R and 18R) only in the lower Miocene.

Eight samples of early Oligocene to early Miocene age were studied from Site 1140. Oligocene samples are barren of radiolarians. Samples from Site 1140 of early Miocene age had well-preserved faunas in Core 183-1140A-14R. The faunas are dominated by spumellarians with abundant Actinommidae, Litheliidae, and Pyloniidae. Common nassellarians are Artostrobiidae and Plagiacanthidae (Fig. F5).

All material was cleaned with hot hydrochloric acid and H_2O_2 and then was wet sieved (38-µm mesh). After short ultrasonic treatment, the samples were neutralized with distilled water. The radiolarian slides were prepared by following standard procedures for random grain distribution (Moore, 1973), including an improved coverslip holder for preparing microslides (Lazarus, 1994). The dry SiO₂ material was then embedded in Canada balsam and dried at 60°C. The radiolarian slides were examined with a transmitted light microscope. Pictures were captured directly from this microscope using a high-resolution black-andwhite video camera connected to a computer. All images were transferred to a species-oriented image database (Cumulus, Canto Software) for further analysis.

Species counts were based on >400 specimens whenever possible. If <400 specimens were found on the slide, the total slide was counted. Radiolarians were counted along random transects of the slides. Presence/absence data for the biostratigraphic analysis are based on examination of larger portions of the slides than used for the species counts. Total radiolarian abundance was determined on the counts of radiolarians, including fragments. Abundance data are reported as numbers of individuals per gram of dry bulk sediment (except for a rough evaluation of family data) (Figs. F3, F4, F5) rather than our usual interval (Abundant [A], Common [C], Rare [R]) classification.

Preservation of radiolarians was evaluated as follows:

Poor = <10%, Poor to moderate = 10%–30%, Moderate = 30%–50%, Moderate to good = 50%–70%, and Good = >70% of the radiolarians can be identified to the genus level.

In addition, we suggest a new approach to evaluating radiolarian preservation. We tentatively use the relative abundance of robust nassellarians (*Siphocampe* and *Artostrobus*) as an indicator of preservation.

F3. Family level faunal characteristics, Site 1138, p. 25.



F4. Family level faunal characteristics, Site 1139, p. 26.



F5. Family level faunal characteristics, Site 1140, p. 27.



The percentage of diatoms was determined on the counts of all radiolarian fragments and all diatom fragments. A minimum of 10% of the siliceous tests of single specimens had to be preserved to be included in the counts.

Radiolarian diversity was assessed in several ways based on species counts from random views of the radiolarian slides. Other than species richness, diversity has been evaluated by the following criteria:

- 1. The Shannon index: $H' = -\Sigma (P_i \times \ln P_i)$, where P_i is the fraction of the i_{th} species of the total fauna. This index provides a rough measure of diversity, which is much less biased by sample size than species richness.
- 2. Evenness: $J' = H'/H'_{max}$, where H' is the Shannon index as defined above, $H'_{max} = \ln S$, and S is the number of species observed. This index determines how evenly the proportions of taxa are distributed in a sample.
- 3. Margalef's index: $SR = (S 1)/\ln N$, where N is the number of individuals. This index provides a measure of species richness that is roughly normalized for sample size without using more complex rarefaction techniques.

Additional faunal indices with a potential paleoceanographic signal are nassellarian/spumellarian ratios, measured as the percentage of spumellarians in the total radiolarian fauna, and the radiolarian/diatom ratio, measured as the percentage of diatoms in the whole siliceous microfauna. Statistical tests were carried out using the statistical software package, SPSS 9.0. Radiolarian slides are curated at the Museum für Naturkunde in Berlin (Germany).

RADIOLARIANS AND DIATOMS AT SITE 1138

A total of 120 radiolarian species were recorded in the samples from Hole 1138A, as summarized in Table T2. A full documentation of the taxonomy of recovered radiolarians is given in "Appendix," p. 16, and in Plates P1, P2, P3, P4, P5, P6, P7, P8, P9, P10.

Core 183-1138A-34R

Radiolarians are abundant and usually well preserved in Sections 183-1138A-34R-1 through 34R-3. Sporadically, an abundant diatom flora is present. Diatoms dominate the siliceous microfossil assemblage in Sample 183-1138A-34R-3, 20–22 cm. Stratigraphic markers are present (e.g., *Axoprunum irregularis, Lychnocanoma conica, Amphistylus*? sp. sensu Takemura, 1992) but are only moderately preserved and rare. Even the relatively well preserved samples of Sections 183-1138A-34R-1 through 34R-3 are dominated by fragmented radiolarians and diatoms rather than by complete specimens.

The high abundance of Litheliidae in all samples of Sections 183-1138A-34R-3 through 34R-1 is remarkable. *Lithelius* sp. A gr. (Pl. **P3**, figs. 1–4), *Lithelius* sp. C (Pl. **P3**, fig. 6), *Lithelius* sp. D (Pl. **P3**, fig. 7), and Pylonid sp. 1 (Pl. **P2**, fig. 8) are common throughout. Spumellaria generally dominate the radiolarian assemblage. Among the nassellarians, Artostrobiidae are the most abundant family, and within the Artostrobiidae, *Siphocampe* and *Artostrobus* dominate. In all samples of Sections 183-1138A-34R-3 through 34R-1, *Siphocampe* and *Artostrobus* are the T2. Abundance data for all early Oligocene-aged radiolarians, Hole 1138A, p. 32.

P1. Actinommidae, p. 39.



P2. Actinommidae and Pyloniidae, p. 40.



P3. Litheliidae, p. 41.



P4. Spongodiscidae and Sponguridae, p. 42.



dominant nassellarian genus, and *Artostrobus pusillum* gr. is the most frequent species (Pl. **P9**, figs. 6–10).

Core 183-1138A-35R

Radiolarians are generally rare to common and poorly to moderately preserved in Core 183-1138A-35R. Stratigraphic markers are present (e.g., A. irregularis, L. conica, Amphistylus? sp. sensu Takemura, 1992, and *Eucyrtidium spinosum*). Spumellarians in general and Litheliidae in particular dominate the assemblages in this interval. *Lithelius* sp. A (Pl. P3, figs. 1–4), Lithelius sp. C (Pl. P3, fig. 6), Lithelius sp. D (Pl. P3, fig. 7), and Pylonid sp. 1 (Pl. P2, fig. 8) are especially conspicuous. Additional common spumellarians are Actinommidae, especially Cenosphaera sp. A gr. (Pl. P2, figs. 6-7), Amphistylus? sp. A (Pl. P1, fig. 2), Actinomma henningsmoeni (Pl. P2, fig. 1), and Stylosphaera radiosa gr. (Pl. P1, figs. 6-8). Among the nassellarians, Siphocampe and Artostrobus are most common, particularly A. pusillum (Pl. P9, figs. 6-10), Siphocampe nodosaria (Pl. P9, fig. 11), Siphocampe acephala gr. (Pl. P9, figs. 14-17), and Siphocampe arachnea gr. (Pl. P9, figs. 12, 13). Peridium? sp. A and Peridium? sp. B (Pl. P5, figs. 18, 19) are generally abundant in these samples except for the two levels at which diversity declines (see below).

The overall radiolarian diversity is generally lower than that found in Sections 183-1138A-34R-1 through 34R-3. The highest diversity in Core 183-1138A-35R is found in Samples 183-1138A-35R-2, 105–107 cm, and 35R-2, 24–26 cm. There is a conspicuous diversity decline in Sample 183-1138A-35R-1, 140–142 cm, where the assemblage is dominated by Litheliidae. Another diversity drop is observed in Sample 183-1138A-35R-3, 101–103 cm, where Artostrobiidae and Litheliidae dominate the poorly to moderately preserved fauna. Diatoms are common to abundant in this core (Table T3), but diatom diversity is remarkably lower than in Sections 183-1138A-34R-1 through 34R-3.

Stratigraphic markers are present but rare. *A. irregularis* was observed in Samples 183-1138A-35R-5, 51–53 cm; 35R-4 (all); 35R-3, 101–103 cm; 35R-2 (all) to 34R-3 (all); 34R-2, 23–25 cm; and Section 34R-1 (Table **T2**). *E. spinosum* is present only in Sample 183-1138A-35R-1, 103–105 cm (Table **T2**). *L. conica* is present in all samples of Section 183-1138A-35R-4; in Samples 183-1138A-35R-2, 105–107 cm, and 35R-1, 140–142 cm; all samples of Section 183-1138A-34R-3; and in Sample 183-1138A-34R-1, 103–105 cm (Table **T2**).

Core 183-1138A-36R

Radiolarians are rare and poorly preserved in Samples 183-1138A-36R-1, 0–2 cm, through 36R-1, 41–43 cm. Almost exclusively, radiolarian fragments rather than complete specimens were observed. Artostrobiidae are dominant, and some additional Actinommidae are present, which contribute most to the observed diversity. Although there is a continuous trend toward declining diversities downcore in Section 183-1138A-36R-1, the very top of the section is characterized by a profound diversity decline. Below interval 183-1138A-36R-1, 40–42 cm, samples are barren of radiolarians. Diatoms are generally rare in all samples of Core 183-1138A-36R. P5. Plagiacanthidae, p. 43.



P6. Plagiacanthidae, Pterocorythidae, Plagonidae, Cannobotridae, and Theoperidae, p. 44.



P7. Theoperidae, p. 45.



P8. Acanthodesmiidae, p. 46.



STRATIGRAPHY

The current age model based on paleomagnetic reversals, planktonic foraminifers, and nannofossil biostratigraphy (Fig. F2), suggests that the radiolarian-bearing Paleogene samples of Site 1138 are of middle early Oligocene to late early Oligocene age (Fig. F6). Owing to a great degree of endemism (Lazarus and Caulet, 1993), direct correlation of recently established Paleogene radiolarian zones of the Southern Ocean (Takemura and Ling, 1997) with lower-latitude chronozones is not possible at a fine scale. Furthermore, the heterogeneous preservation does not allow a straightforward application of Takemura and Ling's (1997) zonation.

The age-depth model (Fig. F2) interprets the magnetic reversal in Section 183-1138A-36R-1 as the C11r/C11n boundary. The upper 50 cm of Core 183-1138A-36R is thus assigned to Chron C11n. According to the number of magnetic reversals and nannoplankton data, the top of our section in Core 183-1138A-34R should be assigned to Chron C10n.

Only a few radiolarian marker taxa help to constrain the age of our samples, and they give equivocal results, especially compared to the paleomagnetic data. *A. irregularis*, the zonal marker for the *A.? irregularis* Zone of Takemura and Ling (1997), is found more or less regularly from Samples 183-1138A-35R-5, 20–22 cm, to 34R-1, 103–105 cm. The vertical distribution of this species agrees well with the age-depth model, indicating a late early Oligocene age. However, *Amphistylus*? sp. sensu Takemura (1992) (= our *Amphistylus*? sp. A) is found in most samples of Cores 183-1138A-34R to 35R, which are placed in Chron C10n to C10r (Fig. F2). This species was previously thought to disappear shortly after the first appearance datum (FAD) of *A. irregularis*, presumably within Chron C12r or earlier, according to Takemura and Ling (1997).

E. spinosum, the zonal marker of the *E. spinosum* Zone, is recorded in Sample 183-1138A-35R-1, 103–105 cm, assigned to Chron C10n (Fig. **F2**). Based on preliminary paleomagnetic calibrations, Takemura and Ling (1997) gave a range for the last appearance of *E. spinosum* between Chrons C11r and C13n. Further complicating the issue is the first appearance of *L. conica* in Sample 183-1138A-35R-4, 103–105 cm. This species is supposed to appear first in Chrons C11 or C12 (Takemura and Ling, 1997), but its first appearance is assigned to C10r in Hole 1138A (Fig. **F2**). Other stratigraphic markers of Takemura and Ling (1997) could not be found.

The combination of these somewhat equivocal radiolarian data, the shipboard calcareous nannofossil and foraminifer dates (Shipboard Scientific Party, 2000), and the paleomagnetic results render it likely that some identified radiolarian ranges need to be revised. The revision affects both the absolute ranges and the relative ranges. First, the stratigraphic overlap between Amphistylus? sp. and A. irregularis is likely to be greater than previously believed. The most parsimonious explanation is that Amphistylus? sp. has a later last appearance datum (LAD) than previously believed. Second, E. spinosum ranges significantly longer into the Oligocene than implied by Takemura and Ling (1997). Although Takemura (1992) reported finding E. spinosum in samples as young as ours, he interpreted those occurrences as reworked. However, we found complete specimens of *E. spinosum* in a well-preserved assemblage; hence, we interpret the occurrence as in situ. Even with the adjustments, the interpretation of the paleomagnetic data, based on radiolarians, is hardly compatible with the interpretations based on calcareous

P9. Artostrobiidae, p. 47.



P10. Artostrobiidae from the Kerguelen Plateau, p. 48.



T3. Summary of radiolarian faunal indices, Hole 1138A, p. 37.

F6. Oligocene radiolarian marker taxa, Hole 1138A, p. 28.



nannoplankton and planktonic foraminifers (according to the current age model) (Fig. F2). The reasons for these discrepancies cannot at present be determined.

PALEOECOLOGICAL PARAMETERS

Preservation and Abundance

Although not a paleoecological parameter per se, careful examination of preservation is important to evaluate the bias in any paleoecological signal, and given that preservation is correlated to productivity, changes in preservation may provide important paleoceanographic information. Preservation can be analyzed based on fragmentation, dissolution, and recrystallization. Whereas recrystallization is not of major concern in the material (no chert was observed above Core 183-1138A-41R; 383 mbsf), fragmentation and dissolution of tests seriously affect the overall preservation of the faunas. As tectonic strain is virtually absent, the observed fragmentation can be used as a proxy of opal dissolution. The traditional classification of preservation into poor, moderate, and good applied to ODP micropaleontological samples thus may be translated into strongly dissolved/fragmented, moderately dissolved/ fragmented, and weakly dissolved/fragmented faunas.

We tentatively classified preservation by visual estimation (Table T3) as done in most other ODP reports, but additionally we suggest that a more rigorous classification of dissolution can be applied. Only with a more quantitative measure of preservation can we hope to statistically test correlations between shell dissolution and productivity. As a first approximation, we calculated the percentage of Siphocampe and Artostrobus in our samples (Table T3). These genera are among the most robust taxa in our material and were observed in a fairly high absolute abundance throughout (Pl. P9, figs. 3-18; Pl. P10, figs. 1-8). Although the percentage of Siphocampe and Artostrobus may be controlled by additional factors and there are occasional other robust taxa in the samples, it is thought to be an independent quantitative proxy of fragmentation/dissolution. The higher the proportion of Siphocampe and Artostrobus, the higher dissolution is thought to be. Because Siphocampe is by far the more abundant of the two genera in our samples, we define the name "Siphocampe index" for the cumulative percentage of Siphocampe and Artostrobus in a sample. The significant correlation between the qualitative preservation evaluation and our *Siphocampe* index (Fig. F7) supports the suggestion that the latter may represent a proxy for radiolarian faunal preservation in the Oligocene. Both the qualitative preservation index and the Siphocampe proxy indicate a significant upward increase of preservation in the studied interval. The percentage of Siphocampe and Artostrobus declines from >80% in the lowest three samples to <10% in the upper part. The first well-preserved faunas are recorded in interval 183-1138A-35R-2, 105-107 cm, according to qualitative studies, whereas the Siphocampe index suggests that dissolution/fragmentation is reduced already in interval 183-1138A-35R-5, 20-22 cm.

Radiolarian abundance ranges from ~2,500 individuals per gram to >150,000 individuals per gram of dry sediment. Abundance increases upcore as significantly as preservation. Core 183-1138A-36R consistently yields abundances of <10,000 individuals per gram; abundance in Core 35R varies between ~12,000 and ~80,000 individuals per gram; and Core 34R always shows values >50,000 individuals per gram. Both

F7. Radiolarian faunal indices, Hole 1138A, p. 29.



the *Siphocampe* proxy and the qualitative estimate of preservation are significantly correlated with radiolarian abundance (Fig. **F7**). Following common arguments (Baldauf and Barron, 1990), both opal preservation and abundance of siliceous plankton groups are thought to co-vary with productivity, although the relationship is not simple (Nelson et al., 1995). Lazarus and Pallant (1989) have shown that Oligocene radiolarian abundance in the Labrador Sea is very well correlated to total organic carbon content and other independent proxy indicators of productivity. This, however, is not true for diatoms that are generally thought to be even better productivity proxies (Ragueneau et al., 2000).

Faunal Indices

All diversity indices are strongly correlated with each other and with our preservational proxies (Figs. F7, F8). This observation poses problems for the paleoecological interpretation of our data. All diversity indices and the abundance data are largely explained by fluctuations in preservation. Based on R^2 values, up to 90% of the variation in diversity can be explained by variations in preservation. Important exceptions are evenness and the percentage of diatoms, which are only weakly correlated with our qualitative measure of preservation but still exhibit very high correlations with our *Siphocampe* proxy. The weakest correlation in the whole data set is between the percentage of diatoms and radiolarian abundance. If so, simple measurements of opal flux are probably not sufficient to characterize the productivity and interpret the signal, a point already emphasized by Diester-Haass (1995) and Ragueneau et al. (2000).

Even a principal component approach (Varimax rotation) does not help greatly to constrain primary patterns. Only one factor has an eigenvalue of >1, explaining 77% of the total variance in the data set. Three factors explain 91% of the total variance. The first two factors are best interpreted as preservation. It is only the third factor that has the highest loadings on the percentage of diatoms in the assemblage, again indicating a somewhat decoupled pattern of diatom and radiolarian abundances.

When comparing only diversity patterns of equally well-preserved faunas in the section, no significant trend through the lower Oligocene is evident. The maximum diversity (Shannon and Margalef indices and species richness) is reached in Sample 183-1138A-34R-3, 105–107 cm, well below the top of the investigated time interval.

DISCUSSION

We test the hypothesis that changes in radiolarian faunal indices correspond to climatic change across the Eocene/Oligocene boundary interval and within the lower Oligocene, the null hypothesis being that radiolarian faunal indices develop independently from reconstructed changes in Earth-system parameters. In contrast to low latitudes, the sedimentary record of Eocene siliceous microplankton is generally patchy in southern high latitudes. This applies to diatoms as well as radiolarians (Baldauf, 1992; Baldauf and Barron, 1990). The observation that radiolarian preservation and accumulation rates increase gradually from the Eocene to the Oligocene has been made at many Southern Ocean Deep Sea Drilling Project and ODP sites (Table T4) including **F8**. Oligocene-aged radiolarian faunas, Hole 1138A, p. 30.



T4. Mean preservation of radiolarian faunas, p. 38.

Antarctica. In nearly all regions where both Eocene and Oligocene sediments were studied, the Oligocene yields better-preserved faunas than the Eocene. This pattern is quite opposite to that in the tropics where the average preservation is much better in the late Eocene (Nigrini and Sanfilippo, 2000), but it is coherent with findings from northern high latitudes (Lazarus and Pallant, 1989). Although well-preserved faunas were recovered from the Eocene at several localities including the Kerguelen Plateau (Caulet, 1991), the mean preservation of radiolarians is significantly (p < 0.001, based on t-test) different between Eocene and Oligocene samples. One of the few exceptions is reported from the Falkland Plateau (Weaver, 1983), where well-preserved faunas are apparently present throughout the Eocene and Oligocene, although preservation varies between holes.

The general paucity of silica accumulation/preservation in many Eocene age southern high-latitude sites has been attributed to low productivity. Enhanced Oligocene silica accumulation and productivity are usually explained by high-latitude cooling, which increased latitudinal temperature gradients and led to stronger oceanic turnover and thus higher nutrient supply (Kennett, 1977). Tectonic uplift and enhanced weathering may also have contributed to increasing nutrient concentrations (Zachos et al., 1999). The development of the Antarctic Bottom Water formation may additionally have aided radiolarian preservation and declination of planktonic foraminiferal preservation (Diester-Haass, 1995). Although authors have often argued for a continuous and gradual climatic deterioration (Keller et al., 1992), it is now clear that cooling was punctuated in the earliest Oligocene, at least in the Southern Ocean (Wei, 1991; Zachos et al., 1999; Zachos et al., 2001).

Considering the arguments above, we are tempted to invoke climatic cooling and associated productivity fluctuations as the prime control of radiolarian preservation, abundance, and diversity. Although the relationship may not be as straightforward (Diester-Haass, 1996), the lower Oligocene opal maximum is usually associated with high values of other productivity proxies such as benthic foraminiferal accumulation rates and carbonate dissolution (Diester-Haass, 1996; Diester-Haass and Zahn, 1996). Carbonate dissolution is invoked as a paleoproductivity proxy owing to the increase of calcite dissolution with increasing organic carbon supply in a well-oxygenated environment (Diester-Haass, 1995). In our material, a temporal decoupling of the productivity proxies is evident. Carbonate dissolution appears to be most substantial in the upper Eocene and earliest Oligocene, judging from the incomplete stratigraphic record in Core 183-1138A-36R and the pronounced hiatus at the top of Core 183-1138A-37R (Fig. F2). Radiolarian preservation, in contrast, is not significantly enhanced before the uppermost lower Oligocene (Section 183-1138A-35R-2; Chron C10r or possibly Subchron C11n.1), immediately after the end of the condensed sequence of Core 183-1138A-36R. This pattern suggests regional differences on the Kerguelen Plateau. The sediments from Southern Kerguelen Plateau show a profound synchronous increase in productivity in the earliest Oligocene, Chron C13n (Sites 738 and 744; Diester-Haass, 1995, 1996; Zachos et al., 1999; Site 748; Wise et al., 1992), whereas the central and Northern Kerguelen Plateau (Sites 1138, 1139, and 1140) sediments record an increase in radiolarian and diatom preservation 3 to 4 m.y. later. This interpretation is consistent with the observation of Lazarus and Caulet (1993) that endemic radiolarian faunas, indicative of a distinct surface water mass, developed first during the late Eocene only close to the Antarctic continent on the Southern Kerguelen Plateau and

spread throughout the Antarctic region only later, during the Oligocene.

CONCLUSIONS

Radiolarians of late early Oligocene age from the Central Kerguelen Plateau exhibit an upsection tendency toward better preservation, higher abundance, and higher diversity. All these faunal characteristics are likely to be linked to cooling during the early Oligocene and a higher productivity of siliceous microplankton. In contrast to the Southern Kerguelen Plateau, the central and Northern Kerguelen Plateau exhibit a temporally decoupled pattern of different productivity proxies. Carbonate dissolution as identified by reduced carbonate accumulation rates and a pronounced hiatus occurred significantly earlier (late Eocene to early Oligocene; Chrons C17n to C11r) (Fig. F2) than the first record of well-preserved and abundant radiolarians (late early Oligocene; Chron C10r). We speculate that the productivity rise on the Central Kerguelen Plateau was first governed by nonsiliceous phytoplankton and was only later dominated by enhanced radiolarian and diatom abundance and preservation.

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APPENDIX

Taxonomic List

- Actinomma holtedahli Bjørklund gr. (Pl. P2, figs. 2–3); Bjørklund, 1976, p. 1121, pl. 20, figs. 8–9.
- Actinomma henningsmoeni Goll and Bjørklund (Pl. P2, fig. 1); Goll and Bjørklund, 1989, p. 734, pl. 2, figs. 10–15.
- Amphistylus angelinus (Clark and Campbell) (Pl. P1, fig. 3); Chen, 1975, p. 453, pl. 21, figs. 3–40.
- Amphistylus? sp. A (Pl. P1, fig. 2).
- *Amphistylus*? sp. Takemura, 1992, p. 741, pl. 5, fig. 9–10; Takemura and Ling, 1997, p. 108, pl. 1, fig. 3.
- *Amphycraspedum proxilum?* Sanfilippo and Riedel (Pl. **P4**, fig. 9); Sanfilippo and Riedel, 1973, p. 608, pl. 10, figs. 7–11; pl. 11, figs. 1–5, pl. 28, figs. 3–5
- Amphymenium splendiarmatum Clark and Campbell (Pl. P4, fig. 8); Clark and Campbell, 1942, p. 46, pl. 1, fig. 12; Sanfilippo and Riedel, 1973, p. 524, pl. 11, figs. 6–8; Petrushevskaya, 1975, p. 577, pl. 7, fig. 1; pl. 37, figs. 1–3; Caulet, 1991, p. 537; O'Connor, 1993, p. 40, pl. 2, figs. 16–17.
- Antarctissa longa Petrushevskaya (Pl. P5, fig. 7); Petrushevskaya, 1975, p. 618, pl. 11, figs. 8–10.
- Antarctissa robusta Petrushevskaya (Pl. P5, fig. 8); Petrushevskaya, 1975, p. 591, pl. 11, figs. 21–22.
- *Antarctissa* sp. cf. *A. conradae* Chen (Pl. **P5**, figs. 9–10); cf. Chen, 1975, p. 457. pl. 17, figs. 1–5.

Remarks: Both specimens look like the species described by Chen, but poor preservation prevents a definite assignment.

- Artostrobus annulatus (Bailey) gr. (Pl. P9, figs. 3-4).
- *Cornutella annulatus* Bailey, 1856, p. 3, fig. 5; Petrushevskaya, 1975, p. 579, pl. 10, figs. 4–5.

Artostrobus stathmeporoides (Petrushevskaya) nov. comb. (Pl. P10, figs. 1–2); Lithomitrella stathmeporoides Petrushevskaya, 1979, p. 151, fig. 411, not fig. 410.

Remarks: Comparison with the type material of Ehrenberg showed that the species has no resemblance with *Eucyrtidium acephala*, the type species of *Lithomitrella*, but has to be assigned to *Artostrobus*. Petrushevskaya described this species as lacking a cephalic horn. Its depicted holotype, however, bears a horn, just like specimens.

Artostrobus pusillum (Ehrenberg) gr. (Pl. P9, figs. 6–10); Eucyrtidium pusillum Ehrenberg, 1873, p. 232, pl. 11, fig. 6; not Petrushevskaya, 1971, pl. 92, fig. 5, not A. pusillum (Ehrenberg), Petrushevskaya, 1975, p. 578, pl. 26, figs. 1–2.

Remarks: The holotype of *A. pusillum* in the Ehrenberg collection (*E. pusillum*) shows no similarity with *A. pusillum* described by Petrushevskaya, 1975.

Artostrobus pretabulatus Petrushevskaya (Pl. P9, fig. 5); Takemura, 1992, p. 745, pl. 5, fig. 12; Crouch and Hollis, 1996, p. 26.

Axoprunum bispiculum (Popofsky) (Pl. P1, fig. 13)

Stylocontarium bispiculum Popofsky, 1912, p. 91, pl. 2, fig. 2; Chen, 1975, p. 454, pl. 21, figs. 1–2.

Axoprunum bispiculum (Popofsky) Takemura, 1992, p. 741, pl. 1, figs. 1–2; Hollis et al., 1997, p. 43, pl. 1, fig. 14.

Axoprunum irregularis Takemura (Pl. **P1**, figs. 15–16); Takemura, 1992, p. 752, pl. 3, figs. 8–11.

Axoprunum pierinae (Clark and Campbell) (Pl. P1, fig. 14)

Lithatractus pierinae Clark and Campbell, 1942, p. 34, pl. 5, fig. 25.

Axoprunum pieridae (Clark and Campbell), gr., Sanfilippo and Riedel, 1973, p. 488, pl. 1, figs. 6–11; pl. 23, fig. 3; Petrushevskaya, 1975, p. 571; Caulet, 1991, p. 537; Takemura, 1992, p. 742, pl. 6, figs. 3–6; Strong et al., 1995, p. 208, fig. 10c; Crouch and Hollis, 1996, p. 26; Takemura, 1997, p. 112, pl. 1, fig. 1; O'Connor, 1999, pl. 10, fig. 12.

- Botryocella pauciperforata O'Connor (Pl. P6, fig. 11); O'Connor, 1999, p. 42, pl. 5, figs. 20a-24.
- Botryostrobus kerguelensis Caulet (Pl. P9, fig. 2); Caulet, 1991, p. 535, pl. 3, figs. 6–8.
- Ceratocyrtis mashae Bjørklund (Pl. P6, fig. 5); Bjørklund, 1976, p. 1125, pl. 17, figs. 1–8.
- *Ceratocyrtis robustus*? Bjørklund (Pl. P6, figs. 3–4); Bjørklund, 1976, p. 1125, pl. 17, figs. 9–10.

Remarks: Our specimens are very similar to Bjørklund's species, but poor preservation prevents a definite statement.

Ceratocyrtis sp. aff. C. stigi (Bjørklund), nov. comb (Pl. P6, fig. 2); aff. Lithomelissa stigi Bjørklund, 1976, p. 1125, pl. 15, figs. 12–17.

Ceratocyrtis stigi (Bjørklund), nov. comb (Pl. P6, fig. 1)

Lithomelissa stigi Bjørklund, 1976, p. 1125, pl. 15, figs. 12-17.

Ceratocyrtis panicula Petrushevskaya, Petrushevskaya and Kozlova, 1972, p. 115.

Lithomelissa sp. C Chen, 1975, p. 458, pl. 11, figs. 4-5.

Lithomelissa sp. Bjørklund, 1976, pl. 15, figs. 9-11.

Ceratocyrtis stigi (Bjørklund), Nigrini and Lombari, 1984, p. N13, pl. 15, fig. 7.

Cornutella profunda Ehrenberg gr. (Pl. P6, figs. 13–15)

Cornutella clathrata s.s. profunda Ehrenberg, 1854b, pl. 35BIV, fig. 24.

Cornutella verrucosa Ehrenberg, 1872a, p. 287; Ehrenberg, 1872b, pl. 9, fig. 16.

Cornutella profunda Ehrenberg, 1858, p. 31; Riedel, 1958, p. 232, pl. 3, figs. 1–2; Petrushevskaya, 1975, p. 587, pl. 13, figs. 32–33.

- *Corythospyris fiscella* Goll gr. (Pl. **P8**, figs. 1–4); Goll, 1978, p. 178, pl. 5, figs. 1–21; Abelmann, 1990, p. 695, pl. 4, figs. 4a, 4b, 7.
- *Corythospyris jubata* Goll (Pl. **P8**, fig. 5); Goll, 1978, p. 177, pl. 4, figs. 1, 2, 4, 5, 7–17.
- *Cyrtolagena laguncula* Haeckel (Pl. **P7**, fig. 14); Haeckel, 1878, p. 1451, pl. 75, fig. 10; Petrushevskaya, 1975, p. 583, pl. 14, figs. 3–4.

- Dendrospyris stabilis Goll (Pl. P8, figs. 7-8); Goll, 1968, p. 1422-1423, pl. 173, figs. 16-18.
- *Dictyophimus archipilium* Petrushevskaya (Pl. **P7**, fig. 10); Petrushevskaya, 1975, p. 583, pl. 25, figs. 1–2.
- *Dictyophimus craticula* Ehrenberg (Pl. **P7**, fig. 9); Sanfilippo and Riedel, 1973, p. 609, pl. 19, fig. 1; pl. 33, fig. 11.
- Dictyophimus sp. aff. D. archipilium Petrushevskaya (Pl. P7, fig. 11); aff. Petrushevskaya, 1975, p. 583, pl. 25, figs. 1–2.
- *Dictyoprora physothorax* Caulet (Pl. P10, figs. 9–10); Caulet, 1991, p. 535, pl. 3, fig. 11.
- *Drymyomma? elegans* Jørgensen gr. (Pl. P2, figs. 4–5); Bjørklund, 1976, p. 1132, pl. 3, figs. 1–4.
- *Eucyrtidium spinosum* Takemura (Pl. P7, fig. 5); Takemura, 1992, p. 746, pl. 5, figs. 5–8.

Eucyrtidium punctatum (Ehrenberg) (Pl. **P7**, fig. 6)

Lithocampe punctata Ehrenberg, 1844, p. 84.

Eucyrtidium punctatum (Ehrenberg), Ehrenberg, 1847, p. 43; Ehrenberg, 1854c, pl. 22, fig. 24.

Artostrobus zitteli Vinassa de Regny, 1900, p. 586, pl. 3, fig. 19.

Eucyrtidium punctatum (Ehrenberg) group, Chen, 1975, p. 495, pl. 15, fig. 8; Sanfilippo et al., 1973, p. 221, pl. 5, figs. 15–16.

Eucyrtidium punctatum (Ehrenberg) group, Weaver et al., 1976, p. 581, pl. 4, figs. 1–2; pl. 8, figs. 4–6.

Eucyrtidium punctatum Caulet, 1986, p. 852, pl. 5, fig. 9.

Eurystomoskevos petrushevskaae Caulet (Pl. P7, fig. 13)

Diplocyclas sp. A Petrushevskaya and Kozlova, 1972, p. 541, pl. 33, figs. 14–16; Petrushevskaya, 1975, p. 587, pl. 24, fig. 4.

Eurystomoskevos petrushevskaae Caulet, 1991, p. 536, pl. 3, figs. 14–15.

- *Lithomelissa gelasinus* O'Connor (Pl. **P5**, fig. 4); O'Connor, 1997, p. 69, pl. 2, figs. 3–6; pl. 6, figs. 6–9, text-fig. 4; Hollis et al., 1997, p. 52, pl. 3, figs. 15–16; O'Connor, 2000, p. 206, pl. 1, figs. 7a, 7b, 8a, 8b, 9a, 9b.
- Lithomelissa sp. aff. Lithomelissa ehrenbergi Bütschli (Pl. P5, fig. 5); aff. Bütschli, 1882, p. 519, fig. 21.

Lithomelissa dupliphysa Caulet (Pl. P5, fig. 1); Caulet, 1991, p. 534, pl. 2, fig. 4.

- *Lithomelissa tricornis* Chen (Pl. **P5**, fig. 3); Chen, 1975, p. 458, pl. 8, figs. 6–7; Abelmann, 1990, p. 695, pl. 5, fig. 3; Takemura, 1992, p. 744, pl. 2, figs. 11–12; Hollis et al., 1997, p. 53.
- *Lophocyrtis (Apoplanius) klydus*? Sanfilippo and Caulet (Pl. **P7**, fig. 8); Sanfilippo and Caulet, 1998, p. 12, pl. 3a, figs. 11–12; pl. 3b, figs. 10–11; pl. 5, figs. 4a, 4b, 5a, 5b, 8, 10, 11.
- *Lophophaena capito* Ehrenberg (Pl. **P6**, fig. 6); Ehrenberg, 1873, p. 242; 1875, pl. 8, fig. 6; Crouch and Hollis, 1996, p. 26.

Lychnocanoma conica (Clark and Campbell) (Pl. **P7**, fig. 1)

Lychnocanoma conicum Clark and Campbell, 1942, p. 71, pl. 9, fig. 38.

Lychnocanella conica Petrushevskaya, 1975, p. 583, pl. 12, figs. 2, 11-15.

Lychnocanoma conica Abelmann, 1990, p. 697, pl. 6, fig. 8; pl. 7, fig. 1a, 1b.

Lychnocanoma sp. cf. *Lychnocanoma babylonis* (Clark and Campbell) (Pl. **P7**, fig. 2); cf. *Dictyophimus babylonis* Clark and Campbell, 1942, p. 67, pl. 9, figs. 32, 36.

Lychnocanoma babylonis (Clark and Campbell).

Lychnocanoma tripodium (Ehrenberg) (Pl. P7, fig. 3)

Lychnocanium tripodium Ehrenberg, 1875, pl. 7, fig. 2.

Lychnocanoma tripodium (Ehrenberg), in Haeckel, 1887, p. 1229.

- *Perichlamydium limbatum* Ehrenberg (Pl. **P4**, fig. 3); Ehrenberg, 1854b, pl. 22, fig. 20; Haeckel, 1887, p. 514; Petrushevskaya, 1975, p. 575, pl. 6, fig. 11; pl. 39, figs. 1–4.
- *Plannapus hornibrooki* O'Connor (Pl. **P10**, fig. 12); O'Connor, 1999, p. 7, pl. 1, figs. 7a–10; pl. 5, figs. 8a–11, text-fig. 2; O'Connor, 2000, p. 207, pl. 1, figs. 10a, 10b–11a, 11b.
- *Plannapus mauricei* O'Connor (Pl. **P10**, fig. 13); O'Connor, 1999, p. 8, pl. 1, figs. 11–14; pl. 5, figs. 12a–15; O'Connor, 2000, p. 208, pl. 1, figs. 12a, 12b–13a, 13b.

Genus PRUNOPYLE Stöhr

Remarks: Spumellarians with pylomes, a well-developed cortical shell, and internal spiral or spongy structures ("prunoids") were first clearly described by Stöhr (1880) from Sicily. His drawings, however, do not clearly match any known material from more recent studies either of the Mediterranean region or the Antarctic. Stöhr also did not leave behind any type material for reanalysis. Thus, his genus concepts cannot be used with any degree of confidence. Dreyer (1889) described several prunoid species and genera, working with the same set of Challenger expedition materials used by Haeckel. As these materials are still available for restudy, it seems best to use this publication for determining the priority of generic concepts. Dreyer's family level concepts were the artificial ones introduced by Haeckel, and partly in consequence, he included many distantly related forms in his analysis. He also considered that any larger than average cortical shell pore was the same homologous structure—a pylome. Thus, his new genera often would now be assigned to widely different families. Dreyer's type species for the genus Prunopyle appears, on examination of the illustration and text description, to actually be a stylosphaerid, thus not suitable for prunoid taxa, as defined here. Although the distinction between Dreyer's genera that do refer to prunoid morphologies is not yet fully clear, the closest match to our material is his genus *Larcopyle*, and we tentatively assign our material to it.

Prunopyle? fragilis (Stöhr) (Pl. P3, fig. 13)

Ommatodiscus fragilis Stöhr, 1880, p. 116, pl. 6, fig. 10.

Lithocarpium fragilis (Stöhr), Petrushevskaya, 1975, p. 572, pl. 4, figs. 2-4.

Prunopyle frakesi Chen, 1975, p. 454, pl. 10, figs. 1–3.

Prunopyle sp. B gr. Abelmann, 1990, p. 693, pl. 4, fig. 3a, 3b.

Prunopyle fragilis (Stöhr), Crouch and Hollis, 1996, p. 26; Hollis et al., 1997, p. 47, pl. 2, figs. 28–29; O'Connor, 1999, pl. 8, fig. 8.

Prunopyle? hayesi Chen (Pl. P3, fig. 9); Chen, 1975, p. 454, pl. 9, figs. 4-5.

Prunopyle? polyacantha Clark and Campbell gr. (Pl. P3, figs. 11, 12)

Larnacantha polyacantha Campbell and Clark, 1944 p. 30, pl. 5, fig. 4; Caulet, 1991, p. 539; Crouch and Hollis, 1996, p. 26.

Lithocarpium polyacantha (Campbell and Clark), Petrushevskaya, 1975 (in part), p. 572, pl. 3, figs. 6–8, fig. 9 (not pl. 29, fig. 6); Abelmann, 1990, p. 694, pl. 4, fig. 2; O'Connor, 1993, p. 37, pl. 2, figs. 12–13.

- *Prunopyle? titan* Clark and Campbell (Pl. P3, fig. 14); *Prunopyle titan* Campbell and Clark, 1944a, p. 20, pl. 3, figs. 1–3; Caulet, 1986, p. 853; Abelmann, 1990, p. 693, fig. 16.
- *Prunopyle* cf. *titan* Clark and Campbell; O'Connor, 1993, p. 33, pl. 1, figs. 16–17, pl. 10, fig. 1; Crouch and Hollis, 1996, p. 26; Hollis et al., 1997, p. 49, pl. 2, figs. 31–32; O'Connor, 1999, pl. 1, fig. 9.

Prunopyle? trypopyrena Caulet (Pl. **P3**, fig. 10); Caulet, 1991, p. 533, pl. 1, figs. 5–7.

Pseudodictyophimus gracilipes (Bailey) gr. (Pl. **P7**, fig. 12)

Dictyophimus gracilipes Bailey, 1856, p. 4, pl. 1, fig. 8.

Pseudodictyophimus gracilipes (Bailey), Petrushevskaya, 1971, p. 93, figs. 47–49; 1975, p. 592, pl. 11, fig. 17; Caulet, 1986, p. 853.

Pterosyringium hamata O'Connor (Pl. P5, fig. 2); O'Connor, 1999, p. 27, pl. 4, figs. 16–21b; pl. 7, figs. 20a–23.

Siphocampe acephala (Ehrenberg) gr. (Pl. P9, figs. 14–17)

Eucyrtidium elegans Ehrenberg, 1854b, pl. 36, fig. 17; 1875, pl. 11, fig. 12.

Siphocampe acephala (Ehrenberg) gr., Hollis et al., 1997, p. 54, pl. 4, figs. 8–20.

Siphocampe cf. acephala (Ehrenberg), Nigrini, 1977, p. 254, pl. 3, fig. 5.

Siphocampe missilis O'Connor, 1994, p. 340, pl. 1, figs. 7, 9, 12; pl. 3, figs. 8–12; O'Connor, 1999, p. 36, pl. 9, fig. 41.

Siphocampe arachnea (Ehrenberg) gr. (Pl. P9, figs. 12–13)

Eucyrtidium lineatum arachneum Ehrenberg, 1861, p. 229.

Lithomitra arachnea (Ehrenberg), Riedel, 1958, p. 242, pl. 4, figs. 7–8.

Siphocampe arachnea (Ehrenberg) gr., Nigrini, 1977, p. 255; Caulet, 1991, p. 539.

Siphocampe nodosaria (Haeckel) (Pl. P9, fig. 11)

Lithomitra nodosaria Haeckel, 1887, p. 1484, pl. 9, fig. 1.

Siphocampe arachnea Abelmann, 1990, p. 698, pl. 8, fig. 4a, 4b.

Siphocampe imbricata Caulet, 1991, p. 539, pl. 3, fig. 13.

Siphocampe nodosaria Hollis et al., 1997, p. 55, pl. 4, figs. 28-32.

- *Siphocampe? elizabethae* sensu Hollis (Pl. **P9**, fig. 18); Hollis et al., 1997, p. 55, pl. 4, fig. 27.
- Spirocyrtis greeni O'Connor (Pl. P9, fig. 1); O'Connor, 1999, p. 8, pl. 1, figs. 15–20b; pl. 5, figs. 16a–19.
- Spongodiscus sp. aff. Spongodiscus maculatus Clark and Campbell (Pl. P4, fig. 1); aff. Clark and Campbell, 1942, emend. Blueford, 1988, p. 254, pl. 7, figs. 6–7.
- *Spongopyle osculosa* Dreyer (Pl. **P4**, fig. 2); Dreyer, 1889, p. 42, pl. 11, figs. 99–100; Abelmann, 1990, p. 693, pl. 3, fig. 11.
- *Stylodictya aculeata* Jørgensen (Pl. **P4**, fig. 6); Jørgensen, 1905, p. 119, pl. 10, fig. 41; Petrushevskaya, 1967, p. 35, pl. 17, figs. 1–3; Abelmann, 1990, p. 693, pl. 3, fig. 9.
- *Stylodictya ocellata*? Ehrenberg (Pl. **P4**, fig. 5); Ehrenberg, 1873, p. 258; Ehrenberg, 1875, pl. 23, fig. 7.
- *Stylodictya validispina* Jørgensen (Pl. **P4**, fig. 7); Jørgensen, 1905, p. 119, pl. 10, fig. 40; Petrushevskaya, 1967, p. 33, fig. 17, IV–V; Abelmann, 1990, p. 693, pl., 3, fig. 10.

Stylosphaera radiosa Ehrenberg gr. (Pl. **P1**, figs. 6–8)

Stylosphaera radiosa Ehrenberg, 1854a, p. 256; Ehrenberg, 1875, pl. 24, fig. 5.

Druppatractus? agostinelli Carnevale, 1908, p. 20, pl. 3, fig. 10.

Amphisphaera radiosa (Ehrenberg) group Petrushevskaya, 1975, p. 570, pl. 2, figs. 18–20.

Stylosphaera coronata coronata Ehrenberg, Chen, 1975, p. 455, pl. 5, figs. 1–2.

Tripodiscinus clavipes (Clark and Campbell) (Pl. P6, fig. 9)

Tripilidium clavipes Clark and Campbell, 1942, p. 64, pl. 9, fig. 29.

Tripodiscinus clavipes Hollis et al., 1997, p. 53, pl. 3, figs. 28–29.

Figure F1. Location of the Kerguelen Plateau with Leg 183 sites and other sites discussed in the text. Underlined numbers indicate Leg 183 sites that have been examined for radiolarians. This study describes Oligocene-aged radiolarians from Site 1138.



Figure F2. Age-depth plot for the Eocene–Oligocene interval of Hole 1138A. The curve is based on biostratigraphic and magnetostratigraphic data (M. Antretter et al., unpubl. data). The magnetostratigraphic timescale is taken from Cande and Kent (1995). Normal polarity is shown in black; reversed polarity is shown in gray. The black line is the estimated age-depth curve; the uncertainty interval is shaded yellow. All magnetostratigraphic chrons that can be assigned to a specific polarity interval of a core are plotted as boxes at the same depth as the core interval. The age curve must run through one of these boxes. Biostratigraphic events and zones used to constrain the magnetostratigraphic assignments are as follows: Nannofossils: N20: FO Discoaster multiradiatus (457.30–458.45 mbsf; 56.2 Ma); N21: LO Fasciculithus tympaniformis (449.60–450.35 mbsf; 5.3 Ma); N22: FO Tribrachiatus orthostylus (449.60–450.35 mbsf; 53.6 Ma); N23: FO Discoaster lodoensis (441.92–448.85 mbsf; 52.8 Ma); N24: FO Discoaster sublodoensis (441.92–448.85 mbsf; 49.7 Ma); N25: FO Nannotetrina spp. (429.55–430.38 mbsf; 47.3 Ma); N26: FO Chiasmolithus gigas (422.08– 429.55 mbsf; 46.1 Ma); N27: LO C. gigas (405.90–406.65 mbsf; 44.5 Ma); N28: FO Reticulofenestra bisecta (356.20-357.19 mbsf; 38.0 Ma); N29: LO Chiasmolithus solitus (351.10-351.94 mbsf; 37.9 Ma); N30: FO Chiasmolithus (346.60–347.35 mbsf; 37.1 Ma); N31: FO Chiasmolithus oamaruensis (345.85–346.60 mbsf; 37.0 Ma); N32: FO Isthmolithus recurvus (344.35–345.10 mbsf; 35.7–36.3 Ma); N33: FO Reticulofenestra oamaruensis (342.85-343.60 mbsf; 35.4 Ma); N34-N39: Chiasmolithus altus zone (278.21-330.78 mbsf; 26.0-31.3 Ma); N40: R. bisecta zone (269.41 mbsf; 23.8–26.1 Ma); and N41: Zone CN2–CN1 (261.81 mbsf; 18.3–23.9 Ma). Foraminifers: F17-F19, F21: Zone AP11-AP10 (357.2-383.3 mbsf; 37.7-42.9 Ma); F20: LO Acarinina bullbrooki (354.5-365.6 mbsf; 40.5 Ma); F22: LO Acarinina primitiva (349.35-350.85 mbsf; 39.0 Ma); F23: LO Acarinina collactea (339.8–343.4 mbsf; 37.7 Ma); F24: LO Subbotina linaperta (339.8–343.4 mbsf; 37.7 Ma); F25: LO *Globigerinatheka index* (339.8–343.4 mbsf; 34.3 Ma); F26: Zone AP13 (340.60 mbsf; 30.0–34.3 Ma); F27: LO Subbotina angiporoides (324.45–335.25 mbsf; 30.0–30.5 Ma); F28–F29: Zone AP16–AP15 (278.21– 285.85 mbsf; 23.8–28.5 Ma); F30: LO Globigerina euapertura (261.81–269.41 mbsf; 23.8 Ma); and F31: Zone NK1 (261.81 mbsf; 21.6–23.8 Ma). (Figure shown on next page.)



Figure F2 (continued). (Caption shown on previous page.)

Figure F3. Family level faunal characteristics of studied samples (late Eocene to Oligocene age) at Site 1138. Samples of early Eocene to Maastrichtian age were also studied but contained no radiolarians. Preservation: P = poor, M = moderate, G = good. Abundance: R = rare, C = common, A = abundant.



Figure F4. Family level faunal characteristics of studied samples (late Oligocene to early Miocene age) at Site 1139. Early Miocene to late Oligocene aged samples; late Eocene to early Oligocene aged samples were also studied but contained no radiolarians. Preservation: P = poor, M = moderate, G = good. Abundance: R = rare, C = common, A = abundant. See Figure F2, p. 23, for legend.



Figure F5. Family level faunal characteristics of studied samples at Site 1140. Preservation: P = poor, M = moderate, G = good. Abundance: R = rare, C = common, A = abundant. See Figure F2, p. 23, for legend.



Figure F6. Oligocene radiolarian stratigraphic marker taxa in Hole 1138A. The ranges of the marker taxa of Takemura and Ling (1997) are indicated. There is a discrepancy between the published radiolarian zonation and our radiolarian ages. See Figure F2, p. 23, for magnetostratigraphic interpretation. Dashed line indicates species would be expected but was not found.



| | | Depth | Number of species | Number of specimens | Margalef index | Shannon index | Evenness | % Spumellaria | % Diatoms | Preservation | % Siphocampe + Artostrobus | Abundance |
|-------------------------|------------------------------|-----------|----------------------|------------------------|-------------------|---------------|----------|---------------|-----------|--------------|-------------------------------|-----------|
| Depth | Pearson Correlation | 1 | | | | | | | | | | |
| | Sig. (2-tailed) | | | | | | | | | | | |
| Number of species | Pearson Correlation | 796(**) | 1 | | | | | | | | | |
| | Sig. (2-tailed) | < 0.001 | | | | | | | | | | |
| Number of | Pearson Correlation | 668(**) | .941(**) | 1 | | | | | | | | |
| specimens | Sig. (2-tailed) | 0.001 | < 0.001 | | | | | | | | | |
| Margalef index | Pearson Correlation | 804(**) | .997(**) | .920(**) | 1 | | | | | | | |
| • | Sig. (2-tailed) | < 0.001 | < 0.001 | < 0.001 | | | | | | | | |
| Shannon index | Pearson Correlation | 726(**) | .955(**) | .908(**) | .962(**) | 1 | | | | | | |
| | Sig. (2-tailed) | < 0.001 | < 0.001 | < 0.001 | < 0.001 | | | | | | | |
| Evenness | Pearson Correlation | 671(**) | .893(**) | .853(**) | .904(**) | .983(**) | 1 | | | | | |
| | Sig. (2-tailed) | 0.001 | < 0.001 | < 0.001 | < 0.001 | < 0.001 | | | | | | |
| % Spumellaria | Pearson Correlation | 764(**) | .824(**) | .735(**) | .841(**) | .842(**) | .833(**) | 1 | | | | |
| | Sig. (2-tailed) | < 0.001 | < 0.001 | < 0.001 | < 0.001 | < 0.001 | < 0.001 | | | | | |
| % Diatoms | Pearson Correlation | 469(*) | .511(*) | 0.429 | .536(*) | .565(**) | .578(**) | .647(**) | 1 | | | |
| | Sig. (2-tailed) | 0.032 | 0.018 | 0.052 | 0.012 | 0.008 | 0.006 | 0.002 | | | | |
| Preservation | Pearson Correlation | 823(**) | .765(**) | .708(**) | .750(**) | .628(**) | .542(*) | .629(**) | 0.25 | 1 | | |
| | Sig. (2-tailed) | < 0.001 | < 0.001 | < 0.001 | < 0.001 | 0.002 | 0.011 | 0.002 | 0.274 | | | |
| % Siphocampe | Pearson Correlation | .723(**) | 908(**) | 860(**) | 917(**) | 952(**) | 946(**) | 903(**) | 570(**) | 597(**) | 1 | |
| + Artostrobus | Sig. (2-tailed) | < 0.001 | < 0.001 | < 0.001 | < 0.001 | < 0.001 | < 0.001 | < 0.001 | 0.007 | 0.004 | | |
| Abundance | Pearson Correlation | 822(**) | .796(**) | .765(**) | .781(**) | .661(**) | .562(**) | .699(**) | 0.234 | .800(**) | 679(**) | 1 |
| | Sig. (2-tailed) | < 0.001 | < 0.001 | < 0.001 | < 0.001 | 0.001 | 0.008 | < 0.001 | 0.306 | < 0.001 | 0.001 | |
| ** Correlation is sign | ificant at the 0.01 level (2 | -tailed). | | | | | | | | | | |
| * Correlation is signif | icant at the 0.05 level (2- | (tailed | | | | | | | | | | |

Figure F7. Correlations between radiolarian faunal indices in Hole 1138A, Cores 183-1138A-36R through 34R (early Oligocene).

Figure F8. Quantitative characterization of Oligocene-aged radiolarian faunas in Hole 1138A. The Shannon diversity index, evenness, number of species, percentage of spumellarians, percentage of diatoms, preservation, and radiolarian abundance per gram of dry sediment are indicated. In addition, a newly proposed proxy for radiolarian preservation (% *Siphocampe + Artostrobus = Siphocampe* proxy) is indicated. Note the distinct trends of all parameters and strongly fluctuating diversity, abundance, and preservation in the lower part of section.



| Site: | 1138 | 1139 | 1140 |
|-------------------------------------|---|---|---|
| Geography (Fig. F1, p. 22): | Central Kerguelen Plateau, 180 km east-southeast of Heard Island | 350 km west-southwest of the Kerguelen archipelago, Skiff Bank (Leclaire Rise) | Northernmost Kerguelen Plateau, 270 km north of the Kerguelen archipelago |
| Core depth (mbsf): | 842.7 | 694.2 | 321.9 |
| Water depth (m): | 1141 | 1415 | 2394 |
| Basement: | Basalts overlain by volcaniclastic rock layers were recovered below 689 mbsf from the lower 144 m of the hole; age: Late Cretaceous | Altered volcaniclastic rocks, basalts, and one minor sedimentary bed from the lower 233 m of the hole; age: late Eocene? | Pillow basalts on three minor sediment beds from the lower 87.38 m of the hole; age: late Eocene–early Oligocene |
| Sediments recovered down to (mbsf): | 689 | 461 | 234 |
| Sedimentology: | 655-m pelagic ooze, chalk, and calcareous claystone overlying 43 m of glauconitic calcareous sandstone and silty clay interbedded with sandstone and conglomerate | 383-m calcareous claystone and chalk with thin intervals of calcareous ooze and chalk at the top and base overlying a <10-m interval of sandy packstone and 77-m grainstone | Pelagic ooze and chalk, whole sedimentary section |
| Eocene/Oligocene (Fig. F2, p. 23): | Base of Core 183-1138A-36R | Sections 183-1139A-38R-2 to 41R-1 | Cores 183-1140A-37R to 26R |

Note: All data are from Shipboard Scientific Party (2000) except Eocene/Oligocene data for Holes 1138A and 1139A.

| | | | | | | | | | | | А | ctinor | nmida | ae | | | | | | | | Pyloniidae | | | | | | |
|-----------------|--|--|---|---|--------------------|---|--------------------------------------|-------------------------|-------------------------|---|------------------------|-------------------------|-------------------|----------------------|--------------------|---|---|----------------------|---|---|--|---------------|---------------|---------------------------|---------------|---------------|-------------------------------------|--|
| Subepoch | Core, section, interval (cm) | Depth (mbsf) | Abundance | Preservation | Amphisphaera sp. A | Amphistylus? sp. A | Amphistylus angelinus | Stylatractus sp. A | Stylatractus sp. B | Stylosphaera radiosa gr. | Stylosphaera sp. A gr. | Hexacontium sp. A | Hexacontium sp. B | Axoprunum bispiculum | Axoprunum pierinae | Axoprunum irregularis | Actinomma henningsmoeni | Actinomma holtedahli | Drymyomma? elegans gr. | Cenosphaera sp. A gr. | Pylonid sp. 1 | Pylonid sp. 2 | Pylonid sp. 3 | Pylonid sp. 4 | Pylonid sp. 5 | Pylonid sp. 6 | Pylonid sp. 7 | |
| early Oligocene | 183-1138A- 34R-1, 103-105 34R-2, 23-25 34R-2, 101-103 34R-3, 20-22 34R-3, 105-107 35R-1, 24-026 35R-1, 103-105 35R-1, 140-142 35R-2, 24-26 35R-2, 105-107 35R-3, 23-025 35R-3, 101-103 35R-4, 20-022 35R-4, 104-106 35R-5, 20-022 35R-5, 51-53 36R-1, 0-2 36R-1, 10-12 36R-1, 20-22 36R-1, 31-33 36R-1, 41-43 | 315.14 315.84 316.62 317.31 318.16 323.95 324.74 325.11 325.45 326.26 326.94 327.72 328.41 329.25 329.91 330.22 333.01 333.11 333.21 333.21 333.22 | A A A A R R C-R C C R R R R R R R R R R R R R R | G G G P P P G G P P P P P P P P P P P P | 6 11 8 3 | 5 6 15 2 4 17 19 45 26 3 | 2 1 4 5 1 1 3 2 | 29 17 4 3 3 | 8 3 12 2 17 | 3 22 13 19 13 22 13 6 15 14 26 28 7 6 1 | 1 | 1 6 25 34 1 | 9 | 8 7 | 1 7 6 | 2 1 2 21 2 3 28 2 1 1 3 3 3 | 1 9 22 20 9 3 2 20 9 3 2 20 9 3 2 22 26 | 3 2 7 2 | 3 1 10 7 14 3 3 7 4 | 40 11 42 30 8 19 21 7 27 9 22 54 2 2 | 63 23 23 44 19 24 22 8 16 20 22 2 16 32 24 | 8 | 27 22 2 | 3 12 12 12 27 | 9 12 | 74 | 12 14 7 12 9 4 27 | |

Table T2. Abundance data for all early Oligocene-aged radiolarians, Hole 1138A. (Continued on next four pages.)

Notes: Abundance: R = Rare, C = Common, A = Abundant. Preservation: P = Poor, M = Moderate, G = Good. * = indeterminate species of nassellaria.

Litheliidae Spongodiscidae Sponguridae Spongodiscus maculatus Amphymenium splendiarmatum Prunopyle? polyacantha gr. Amphycraspedum proxilum Perichlamydium limbatum Prunopyle? trypopyrena Stylodictya validispina Spongodiscus sp. aff. C gr. Spongopyle osculosa Stylodictya ocellata? Stylodictya aculeata Prunopyle? cf. titan Prunopyle? hayesi Prunopyle? fragilis Spongopyle sp. A \checkmark В Lithelius sp. A gr Prunopyle? sp. Prunopyle? sp. Prunopyle? sp. Lithelius sp. B υ Lithelius sp. D Lithelius sp. E Preservation Abundance Lithelius sp. Core, section, Depth Subepoch interval (cm) (mbsf) 183-1138Aearly Oligocene 34R-1, 103–105 315.14 А G 50 32 32 19 4 9 4 4 3 1 G 27 21 23 18 34R-2, 23-25 315.84 А 16 4 2 1 1 16 1 3 4 4 4 34R-2, 101-103 316.62 А G 17 12 14 29 2 18 6 2 19 34R-3, 20-22 Μ 9 23 27 317.31 А 2 4 1 4 1 3 4 34R-3, 105-107 318.16 А G 32 12 22 3 9 7 4 2 1 3 7 15 35R-1, 24-026 323.95 R Р 27 22 14 26 3 8 2 1 1 10 4 35R-1, 103-105 R Р 9 12 2 9 324.74 27 9 1 20 1 1 1 C–R Р 4 35R-1, 140-142 325.11 14 23 8 2 3 18 1 1 35R-2, 24-26 325.45 A–C G 42 8 12 7 2 2 3 4 С 35R-2, 105-107 G 18 34 47 1 9 7 4 1 326.26 26 C–R Р 2 35R-3, 23-025 326.94 17 9 4 35R-3, 101-103 327.72 R Μ 42 42 4 8 С 35R-4, 20-022 Р 35 9 22 4 328.41 4 6 1 1 35R-4, 104-106 329.25 С Μ 57 39 32 2 7 2 15 11 34 4 4 35R-5, 20-022 329.91 С Р 52 7 47 2 5 7 2 8 1 7 1 35R-5, 51-53 330.22 R Р 3 4 7 4 1 1 36R-1, 0-2 333.01 R Р 36R-1, 10-12 333.11 R Р 8 1 1 36R-1, 20-22 333.21 R Р 36R-1, 31-33 333.32 R Р 1 36R-1, 41-43 333.42 R Р 1

Table T2 (continued).

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Radiolarian Faunal Characteristics

Plagiacanthidae Pterocorythidae Lithomelissa sp. aff. Lithomelissa ehrenbergi Ceratocyrtis sp. aff. Ceratocyrtis stigi Nassellaria fam. gen. sp. indet l conradae Ceratocyrtis robustus gr. Lithomelissa dupliphysa Pterosyringium hamata Lithomelissa gelasinus Carpocanarium sp. A Antarctissa sp. cf. A. Ceratocyrtis mashae Lophophaena capito Antarctissa sp. C gr. Antarctissa spp. gr. Lithomelissa? sp. A Antarctissa robusta Lamprocyclas sp. A Antarctissa? sp. A Antarctissa? sp. B Antarctissa longa Antarctissa sp. D Ceratocyrtis stigi Peridium? sp. A Peridium? sp. B Preservation Abundance Core, section, Depth Subepoch interval (cm) (mbsf) 183-1138Aearly Oligocene 34R-1, 103–105 G 10 4 2 6 57 29 16 2 315.14 А 1 1 5 1 G 34R-2, 23-25 315.84 А 4 23 18 37 26 10 2 34R-2, 101–103 G 13 19 316.62 А 6 8 3 4 2 37 2 34R-3, 20-22 317.31 А Μ 4 4 13 4 11 4 19 6 2 4 2 19 34R-3, 105-107 318.16 А G 1 12 6 13 7 14 4 2 2 35R-1, 24-026 323.95 R Р 1 22 17 27 R 24 35R-1, 103-105 Р 12 324.74 1 35R-1, 140-142 325.11 C-R Р 10 35 14 1 1 35R-2, 24-26 325.45 A–C G 3 1 13 26 7 4 1 6 С 35R-2, 105-107 326.26 G 3 7 39 19 4 4 4 4 35R-3, 23-025 326.94 C-R Р 2 15 19 1 35R-3, 101-103 327.72 R М 1 35R-4, 20-022 328.41 С Р 1 8 6 32 13 35R-4, 104-106 329.25 С Μ 2 4 3 27 1 С 35R-5, 20-022 329.91 Р 1 5 27 2 13 2 35R-5, 51-53 330.22 R Р 2 16 24 36R-1, 0-2 333.01 R Р 1 1 36R-1, 10-12 333.11 R Р R Р 36R-1, 20-22 333.21 36R-1, 31-33 333.32 R Ρ 36R-1, 41-43 333.42 R Р

Table T2 (continued).

M. APEL ET AL. Radiolarian Faunal Characteristics

| | | | | | Plago | nidae | Cannol | ootridae | | | | | | | | | Theop | erida | 5 | | | | | | | | Aca | anthoo | lesmii | dae |
|-----------------|--|--|--|---|------------------------|---------------------|----------------------------|--------------------|---|----------------------------|-----------------------------------|------------------------|-------------------|----------------------|-----------------------|-------------------|----------------------------------|------------------------|--------------------------|--------------------------------------|-------------------------------|--------------------------------|-----------------------|-------------------|--------------------|-------------------------------------|--|----------------------|--------------------|--|
| Subepoch | Core, section, interval (cm) | Depth (mbsf) | Abundance | Preservation | Tripodiscinus clavipes | Tripodiscinus sp. A | Botryocella pauciperforata | Botryocella? sp. A | Cornutella profunda gr. | Lychnocanoma conica | Lychnocanoma sp. cf. L. babylonis | Lychnocanoma tripodium | Thyrsocyrtis? sp. | Eucyrtidium spinosum | Eucyrtidium punctatum | Eucyrtidium sp. A | Lophocyrtis (Apoplanius) klydus? | Dictyophimus craticula | Dictyophimus archipilium | Dictyophimus sp. aff. D. archipilium | Pseudodictyophimus gracilipes | Eurystomoskevos petrushevskaae | Cyrtolagena laguncula | Cyrtolagena sp. A | Cycladophora sp. A | Nassellaria fam. gen. sp. indet II* | Corythospyris fiscella gr. | Corythospyris jubata | Dendrospyris sp. A | Dendrospyris stabilis |
| early Oligocene | 183-1138A- 34R-1, 103-105 34R-2, 23-25 34R-2, 101-103 34R-3, 20-22 34R-3, 105-107 35R-1, 24-026 35R-1, 103-105 35R-1, 140-142 35R-2, 24-26 35R-2, 24-26 35R-2, 105-107 35R-3, 23-025 35R-3, 101-103 35R-4, 20-022 35R-4, 104-106 35R-5, 20-022 35R-5, 51-53 36R-1, 0-2 36R-1, 10-12 36R-1, 20-22 36R-1, 31-33 36R-1, 41-43 | 315.14 315.84 316.62 317.31 318.16 323.95 324.74 325.11 325.45 326.26 326.94 327.72 328.41 329.25 329.91 330.22 333.01 333.11 333.21 333.22 333.42 | A A A A C-R C C C R R R R R R R R | G G G M G P P G G P M P P P P P P P P P | 2 | 1 | 17 8 1 1 | 1 | 9 3 11 6 12 23 18 16 13 14 12 12 17 | 2 1 3 1 1 1 | 1 | 2 1 2 | 3 | 2 | 1 | 2 2 | 1 8 3 3 | 3 | 10 3 1 | 1 | 1 | 1 | 1 | 1 | 2 | 3 | 8 4 22 2 7 5 6 3 5 4 1 | 2 1 1 | 6 | 3 2 21 20 15 28 8 4 6 6 13 |

Table T2 (continued).

Table T2 (continued).

| | | | | | | | | | | | | | | Arto | strob | iidae | | | | | | | | | |
|-----------------------|---|--|---|-----------------------------------|--------------------|----------------------------|--|--|---|--|---|-------------------------|-----------------------------|------------------|------------------|------------------|----------------------------|------------------|------------------------------|---------------------------|-------------------|---|--|-----------------|-------------------|
| Subepoch | Core, section, interval (cm) | Depth (mbsf) | Abundance | Preservation | Spirocyrtis greeni | Botryostrobus kerguelensis | Artostrobus pretabulatus | Artostrobus pusillum gr. | Siphocampe nodosaria | Siphocampe arachnea gr. | Siphocampe acephala gr. | Siphocampe? elizabethae | Artostrobus stathmeporoides | Siphocampe sp. A | Siphocampe sp. B | Siphocampe sp. C | Siphocampe sp. D | Siphocampe sp. E | Siphocampe sp. F | Dictyoprora physothorax | Dictyoprora sp. A | Plannapus hornibrooki | Plannapus mauricei | Plannapus sp. A | Dicolocapsa sp. A |
| 1: early Oligocene | 83-1138A- 34R-1, 103-105 34R-2, 23-25 34R-2, 101-103 34R-3, 20-22 34R-3, 105-107 35R-1, 24-026 35R-1, 103-105 35R-1, 140-142 35R-2, 24-26 35R-2, 24-26 35R-2, 24-26 35R-2, 24-26 35R-3, 23-025 35R-3, 101-103 35R-4, 20-022 35R-4, 104-106 35R-5, 20-022 35R-5, 51-53 36R-1, 0-2 36R-1, 10-12 36R-1, 20-22 | 315.14 315.84 316.62 317.31 318.16 323.95 324.74 325.11 325.45 326.26 326.94 327.72 328.41 329.25 329.91 330.22 333.01 333.11 333.21 | A A A A A R R C_R C_R C C R R R R | G G G M G P P G G P M P P P P P P | 1 | 3 | 10 3 9 13 32 22 17 28 | 17 8 17 43 22 17 18 9 19 37 55 82 38 32 27 96 19 78 69 | 1 17 4 12 22 1 7 22 3 | 17 18 14 24 2 13 18 8 1 18 8 1 8 | 5 10 22 3 8 4 16 10 25 4 22 12 14 3 2 | 2 | 2 1 1 2 4 | 4 | 1 | 1 | 9 4 8 4 1 2 | 28 | 2 4 18 1 12 6 | 14 12 14 17 4 | 1 3 | 7 6 7 12 17 25 19 22 7 1 14 12 | 3 2 3 6 12 9 7 12 1 2 7 1 | 6 | 2 |

| Core, section, interval (cm) | Depth (mbsf) | Species (N) | Individuals (N) | Margalef index | Shannon index | Evenness | Spumellarians* (%) | Diatoms† (%) | Preservation | Siphocampe (%) | Radiolarians per gram |
|---------------------------------|-----------------|----------------|--------------------|-------------------|------------------|----------|-----------------------|-----------------|--------------|-------------------|--------------------------|
| 183-1138A- | | | | | | | | | | | |
| 34R-1, 103–105 | 315.14 | 42 | 490 | 6.62 | 2.92 | 0.78 | 59.02 | 13.58 | G | 3.61 | 153,891 |
| 34R-2, 23–25 | 315.84 | 50 | 430 | 8.08 | 3.34 | 0.85 | 61.41 | 30.99 | G | 6.07 | 127,942 |
| 34R-2, 101–103 | 316.62 | 37 | 363 | 6.11 | 3.17 | 0.88 | 44.25 | 21.62 | G | 13.22 | 64,859 |
| 34R-3, 20–22 | 317.31 | 42 | 516 | 6.56 | 3.13 | 0.84 | 39.63 | 56.38 | M–G | 12.40 | 81,898 |
| 34R-3, 105–107 | 318.16 | 53 | 488 | 8.40 | 3.59 | 0.90 | 47.42 | 30.22 | G | 11.13 | 58,786 |
| 35R-1, 24–26 | 323.95 | 36 | 402 | 5.84 | 3.13 | 0.87 | 48.26 | 17.11 | PM | 5.97 | 57,118 |
| 35R-1, 103–105 | 324.74 | 36 | 353 | 5.97 | 3.38 | 0.94 | 36.54 | 18.29 | P-M | 8.78 | 46,246 |
| 35R-1, 140–142 | 325.11 | 28 | 198 | 5.11 | 2.82 | 0.85 | 63.78 | 59.50 | Р | 8.67 | 15,875 |
| 35R-2, 24–26 | 325.45 | 36 | 372 | 5.91 | 3.08 | 0.86 | 37.33 | 23.29 | M–G | 10.31 | 39,039 |
| 35R-2, 105–107 | 326.26 | 42 | 535 | 6.53 | 3.23 | 0.87 | 46.62 | 19.10 | G | 14.90 | 79,855 |
| 35R-3, 23–25 | 326.94 | 28 | 345 | 4.62 | 2.86 | 0.86 | 22.26 | 15.33 | Р | 32.34 | 18,894 |
| 35R-3, 101–103 | 327.72 | 16 | 238 | 2.74 | 2.02 | 0.73 | 41.77 | 20.47 | P-M | 36.29 | 11,708 |
| 35R-4, 20–22 | 328.41 | 37 | 421 | 5.96 | 3.08 | 0.85 | 33.75 | 18.26 | Р | 15.14 | 62,517 |
| 35R-4, 104–106 | 329.25 | 44 | 610 | 6.70 | 3.25 | 0.86 | 40.16 | 24.41 | М | 9.67 | 62,604 |
| 35R-5, 20–22 | 329.91 | 33 | 385 | 5.38 | 2.95 | 0.84 | 47.01 | 30.13 | P-M | 7.01 | 37,388 |
| 35R-5, 51–53 | 330.22 | 22 | 254 | 3.79 | 2.24 | 0.72 | 30.24 | 35.25 | Р | 58.06 | 16,554 |
| 36R-1, 0–2 | 333.01 | 3 | 22 | 0.65 | 0.49 | 0.44 | 4.55 | 8.33 | Р | 86.36 | 1,371 |
| 36R-1, 10–12 | 333.111 | 12 | 128 | 2.27 | 1.45 | 0.58 | 8.59 | 14.67 | Р | 63.28 | 7,131 |
| 36R-1, 20–22 | 333.21 | 7 | 89 | 1.34 | 0.92 | 0.47 | 0.00 | 6.32 | Р | 92.13 | 7,528 |
| 36R-1, 31–33 | 333.32 | 8 | 66 | 1.67 | 0.86 | 0.42 | 1.52 | 1.50 | Р | 80.30 | 4,123 |
| 36R-1, 41–43 | 333.42 | 3 | 15 | 0.74 | 0.49 | 0.44 | 6.67 | 0.00 | Р | 86.67 | 2,447 |

 Table T3. Summary of radiolarian faunal indices, Hole 1138A.

Notes: Preservation: P = poor, M = Moderate, G = Good. * = percentage relative to nassellarians. † = Percentage relative to total radiolarians. N = Number. Diversity and evenness measures are defined in text.

Table T4. Mean preservation of Eocene–Oligocene radiolarian faunas from Southern Ocean sites.

| | Leg Kerguele | 120 n Plateau | Leg Kerguele | 119 n Plateau | Leg Falkland | 71 Plateau | Ross | Leg 183 Kerguelen Plateau | | |
|-----------------|-----------------|------------------|-----------------|-------------------|-----------------|------------------|-------|---------------------------------|-------|-------|
| Time slice | 748B* | 749B* | 738B† | 744A [†] | 511‡ | 512 [‡] | 274** | 267** | 264** | 1138A |
| late Oligocene | 2.6 | 2 | | 3 | | | | 2.2 | | |
| early Oligocene | 2.5 | 2 | 1.8 | 3 | 2.6 | | 2.3 | | | 1.8 |
| late Eocene | 1.5 | 1.5 | 1.8 | 1 | 2.5 | | | | 1.4 | 1 |
| middle Eocene | 1.6 | 1.5 | 1.6 | 1 | 1.0 | 2.9 | | | 1.4 | 1 |

Notes: Evaluations of preservation were taken from all samples of the reported data, transformed into numerical values, and averaged for the time intervals indicated. 1 = preservation poor or no radiolarians preserved, 2 = preservation moderate, 3 = preservation good. Sources: * = Takemura (1992), † = Caulet (1991), ‡ = Weaver (1983), ** = Chen (1975).

Plate P1. Oligocene Actinommidae from the Kerguelen Plateau, Site 1138. **1**. *Amphisphaera* sp. A; Sample 183-1138A-36R-1, 10–12 cm. **2**. *Amphistylus*? sp. A; Sample 183-1138A-34R-2, 101–103 cm. **3**. *Amphistylus angelinus* (Clark and Campbell); Sample 183-1138A-34R-3, 20–22 cm. **4**. *Stylatractus* sp. A; Sample 183-1138A-36R-1, 10–12 cm. **5**. *Stylatractus* sp. B; Sample 183-1138A-36R-1, 10–12 cm. **6–8**. *Stylosphaera radiosa* Ehrenberg gr.; Sample 183-1138A-36R-1, 41–43 cm. **9–10**. *Stylosphaera* sp. A gr.; Sample 183-1138A-34R-1, 103–105 cm. **11**. *Hexacontium* sp. A; Sample 183-1138A-34R-1, 103–105 cm. **12**. *Hexacontium* sp. B; Sample 183-1138A-34R-1, 103–105 cm. **13**. *Axoprunum bispiculum* (Popofsky); Sample 183-1138A-34R-1, 103–105 cm. **14**. *Axoprunum pierinae* (Clark and Campbell); Sample 183-1138A-34R-1, 103–105 cm. **15–16**. *Axoprunum irregularis* Takemura; Samples 183-1138A-35R-4, 20–22 cm, and 34R-2, 23–25 cm.

Family: Actinommidae



Plate P2. Oligocene Actinommidae and Pyloniidae from the Kerguelen Plateau, Site 1138. **1**. *Actinomma henningsmoeni* Goll and Bjørklund; Sample 183-1138A-34R-3, 20–22 cm. **2–3**. *Actinomma holtedahli* Bjørklund gr.; Sample 183-1138A-34R-1, 103–105 cm. **4–5**. *Drymyomma? elegans* Jørgensen gr.; Sample 183-1138A-35R-1, 140–142 cm. **6–7**. *Cenosphaera* sp. A gr.; Sample 183-1138A-35R-4, 104–106 cm. **8**. Pylonid sp. 1; Sample 183-1138A-36R-1, 31–33 cm. **9**. Pylonid sp. 2; Sample 183-1138A-34R-3, 20–22 cm. **10**. Pylonid sp. 3; Sample 183-1138A-35R-5, 20–22 cm. **11**. Pylonid sp. 4; Sample 183-1138A-35R-4, 104–106 cm. **12**. Pylonid sp. 5; Sample 183-1138A-35R-1, 24–26 cm. **13**. Pylonid sp. 6; Sample 183-1138A-35R-2, 24–26 cm.

Family: Actinommidae

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Plate P3. Oligocene Litheliidae from the Kerguelen Plateau, Site 1138. **1–4.** *Lithelius* sp. A gr.; Sample 183-1138A-35R-2, 105–107 cm. **5.** *Lithelius* sp. B.; Sample 183-1138A-34R-3, 20–22 cm. **6.** *Lithelius* sp. C; Sample 183-1138A-35R-2, 105–107 cm. **7.** *Lithelius* sp. D; Sample 183-1138A-35R-2, 24–26 cm. **8.** *Lithelius* sp. E; Sample 183-1138A-35R-2, 105–107 cm. **9.** *Prunopyle? hayesi* Chen; Sample 183-1138A-35R-1, 103–105 cm. **10.** *Prunopyle? trypopyrena* Caulet; Sample 183-1138A-34R-2, 23–25 cm. **11–12.** *Prunopyle? polyacantha* Clark and Campbell gr.; Sample 183-1138A-34R-3, 105–107 cm. **13.** *Prunopyle? fragilis* (Stöhr); Sample 183-1138A-35R-2, 24–26 cm. **14.** *Prunopyle? titan* Clark and Campbell; Sample 183-1138A-34R-1, 103–105 cm. **15.** *Prunopyle*? sp. A; Sample 183-1138A-35R-1, 140–142 cm.

Family: Litheliidae



Plate P4. Oligocene Spongodiscidae and Sponguridae from the Kerguelen Plateau, Site 1138. 1. *Spongodiscus* sp. aff. *Spongodiscus maculatus* Clark and Campbell; Sample 183-1138A-35R-3, 23–25 cm. 2. *Spongopyle osculosa* Dreyer; Sample 183-1138A-34R-2, 101–103 cm. 3. *Perichlamydium limbatum* Ehrenberg; Sample 183-1138A-34R-2, 23–25 cm. 4. *Spongopyle* sp. A; Sample 183-1138A-35R-1, 140–142 cm. 5. *Stylodictya ocellata*? Ehrenberg; Sample 183-1138A-35R-4, 104–106 cm. 6. *Stylodictya aculeata* Jørgensen; Sample 183-1138A-35R-4, 104–106 cm. 7. *Stylodictya validispina* Jørgensen; Sample 183-1138A-34R-2, 23–25 cm. 8. *Amphymenium splendiarmatum* Clark and Campbell; Sample 183-1138A-35R-1, 103–105 cm. 9. *Amphycraspedum proxilum*? Sanfilippo and Riedel; Sample 183-1138A-35R-2, 105–107 cm.

Family: Spongodiscidae



Family: Spongodiscidae



Family: Sponguridae





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Plate P5. Oligocene Plagiacanthidae from the Kerguelen Plateau, Site 1138. **1**. *Lithomelissa dupliphysa* Caulet; Sample 183-1138A-34R-3, 105–107 cm. **2**. *Pterosyringium hamata* O'Connor; Sample 183-1138A-34R-2, 23–25 cm. **3**. *Lithomelissa tricornis* Chen; Sample 183-1138A-34R-1, 103–105 cm. **4**. *Lithomelissa gelasinus* O'Connor; Sample 183-1138A-34R-3, 20–22 cm. **5**. *Lithomelissa* sp. aff. *L. ehrenbergi* Bütschli; Sample 183-1138A-35R-5, 20–22 cm. **6**. *Lithomelissa*? sp. A; Sample 183-1138A-35R-1, 24–26 cm. **7**. *Antarctissa longa* Petrushevskaya; Sample 183-1138A-34R-3, 105–107 cm. **8**. *Antarctissa robusta* Petrushevskaya; Sample 183-1138A-34R-3, 105–107 cm. **8**. *Antarctissa robusta* Petrushevskaya; Sample 183-1138A-34R-3, 20–22 cm. **12**. *Antarctissa*? sp. B; Sample 183-1138A-35R-4, 20–22 cm. **11**. *Antarctissa*? sp. A; Samples 183-1138A-35R-2, 24–26 cm. **15–17**. *Antarctissa*? sp. gr.; Sample 183-1138A-35R-5, 51–53 cm. **18**. *Peridium*? sp. A; Sample 183-1138A-35R-3, 23–25 cm. **19**. *Peridium*? sp. B; Sample 183-1138A-35R-3, 23–25 cm. **19**. *Peridium*? sp. B; Sample 183-1138A-35R-2, 24–26 cm. **15**–17. *Antarctissa*? sp. gr.; Sample 183-1138A-35R-2, 24–26 cm. **19**. *Peridium*? sp. B; Sample 183-1138A-35R-3, 23–25 cm. **19**. *Peridium*? sp. B; Sample 183-1138A-35R-2, 24–26 cm. **15**–17. *Antarctissa*? sp. gr.; Sample 183-1138A-35R-2, 24–26 cm. **19**. *Peridium*? sp. B; Sample 183-1138A-35R-3, 23–25 cm. **19**. *Peridium*? sp. B; Sample 183-1138A-35R-1, 140–142 cm.

Family: Plagiacanthidae



 $100 \ \mu m$

Plate P6. Oligocene Plagiacanthidae, Pterocorythidae, Plagonidae, Cannobotridae, and Theoperidae from the Kerguelen Plateau, Site 1138. 1. *Ceratocyrtis stigi* (Bjørklund); Sample 183-1138A-35R-5, 20–22 cm. 2. *Ceratocyrtis* sp. aff. *C. stigi* (Bjørklund); Sample 183-1138A-35R-2, 105–107 cm. **3–4**. *Ceratocyrtis robustus*? Bjørklund; Sample 183-1138A-35R-5, 20–22 cm. **5**. *Ceratocyrtis mashae* Bjørklund; Sample 183-1138A-34R-1, 103–105 cm. **6**. *Lophophaena capito* Ehrenberg; Sample 183-1138A-35R-2, 105–107 cm. **7**. *Carpocanariu* sp. A; Sample 183-1138A-35R-1, 103–105 cm. **8**. *Lamprocyclas* sp. A; Sample 183-1138A-35R-2, 24–26 cm. **9**. *Tripodiscinus clavipes* Clark and Campbell; Sample 183-1138A-34R-2, 101–103 cm. **10**. *Tripodiscinus* sp. A; Sample 183-1138A-35R-2, 105–107 cm. **11**. *Botryocella pauciperforata* O'Connor; Sample 183-1138A-35R-4, 104–106 cm. **12**. *Botryocella*? sp. A; Sample 183-1138A-34R-1, 103–105 cm. **13–15**. *Cornutella profunda* Ehrenberg gr.; Sample 183-1138A-35R-5, 51–53 cm.

Family: Plagiacanthidae & Pterocorythidae (8)



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Plate P7. Oligocene Theoperidae from the Kerguelen Plateau, Site 1138. **1**. *Lychnocanoma conica* (Clark and Campbell); Sample 183-1138A-35R-4, 20–22 cm. **2**. *Lychnocanoma* sp. cf. *Lychnocanoma babylonis* (Clark and Campbell); Sample 183-1138A-34R-3, 20–22 cm. **3**. *Lychnocanoma tripodium* (Ehrenberg); Sample 183-1138A-34R-3, 105–107 cm. **4**. *Thyrsocyrtis*? sp.; Sample 183-1138A-34R-2, 23–25 cm. **5**. *Eucyrtidium spinosum* Takemura; Sample 183-1138A-35R-1, 103–105 cm. **6**. *Eucyrtidium punctatum* (Ehrenberg); Sample 183-1138A-35R-2, 105–107 cm. **7**. *Eucyrtidium* sp. A; Sample 183-1138A-34R-3, 105–107 cm. **8**. *Lophocyrtis* (*Apoplanius*) *klydus*? Sanfilippo and Caulet; Sample 183-1138A-34R-1, 103–105 cm. **9**. *Dictyophimus craticula* Ehrenberg; Sample 183-1138A-35R-2, 105–107 cm. **11**. *Dictyophimus* sp. aff. *D. archipilium* Petrushevskaya; Sample 183-1138A-35R-2, 105–107 cm. **12**. *Pseudodictyophimus gracilipes* Bailey gr.; Sample 183-1138A-34R-3, 105–107 cm. **13**. *Eurystomoskevos petrushevskaae* Caulet; Sample 183-1138A-35R-5, 20–22 cm. **14**. *Cyrtolagena laguncula* Haeckel; Sample 183-1138A-35R-5, 51–53 cm. **15**. *Cyrtolagena* sp. A; Sample 183-1138A-34R-2, 101–103 cm. **16**. Nassellaria Gen. et sp. indet. II; Sample 183-1138A-34R-2, 23–25 cm.

Family: Theoperidae



Plate P8. Oligocene Acanthodesmiidae from the Kerguelen Plateau, Site 1138. 1-4. Corythospyris fiscella Goll gr.; Sample 183-1138A-36R-1, 20–22 cm. 5. Corythospyris jubata Goll; Sample 183-1138A-35R-5, 20–22 cm. 6. Dendrospyris sp. A; Sample 183-1138A-35R-4, 20-22 cm. 7. Dendrospyris stabilis Goll; Sample 183-1138A-35R-2, 24–26 cm. 8. D. stabilis Goll (bottom and top view); Sample 183-1138A-36R-1, 31–33 cm.

Family: Acanthodesmiidae





3a











1b



3b



5b



2a



4a



6a



8a





4b



6b



8b

Plate P9. Oligocene Artostrobiidae from the Kerguelen Plateau, Site 1138. 1. *Spirocyrtis greeni* O'Connor; Sample 183-1138A-35R-2, 105–107 cm. 2. *Botryostrobus kerguelensis;* Sample 183-1138A-35R-4, 104–106 cm. 3–4. *Artostrobus annulatus* (Bailey) gr.; Sample 183-1138A-34R-2, 101–103 cm. 5. *Artostrobus pretabulatus* Petrushevskaya; Sample 183-1138A-35R-4, 104–106 cm. 6–10. *Artostrobus pusillum* (Ehrenberg) gr.; Sample 183-1138A-35R-4, 104–106 cm. 12–13. *Siphocampe arachnea* (Ehrenberg) gr.; Sample 183-1138A-34R-2, 23–25 cm. 14–17. *Siphocampe acephala* (Ehrenberg) gr.; Sample 183-1138A-35R-2, 24–26 cm. 18. *Siphocampe? elizabethae* sensu Hollis; Sample 183-1138A-36R-1, 20–22 cm.

Family: Artostrobiidae

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Plate P10. Oligocene Artostrobiidae from the Kerguelen Plateau, Site 1138. **1–2**. *Artostrobus stathmeporoides;* Sample 183-1138A-35R-3, 23–25 cm. **3**. *Siphocampe* sp. A; Sample 183-1138A-35R-3, 23–25 cm. **4**. *Siphocampe* sp. B; Sample 183-1138A-35R-2, 24–26 cm. **5**. *Siphocampe* sp. C; Sample 183-1138A-35R-2, 24–26 cm. **6**. *Siphocampe* sp. D; Sample 183-1138A-35R-1, 140–142 cm. **7**. *Siphocampe* sp. E; Sample 183-1138A-35R-1, 140–142 cm. **7**. *Siphocampe* sp. E; Sample 183-1138A-35R-1, 140–142 cm. **8**. *Siphocampe* sp. F; Sample 183-1138A-35R-4, 104–106 cm. **9–10**. *Dictyoprora physothorax* Caulet; Sample 183-1138A-35R-2, 105–107 cm. **11**. *Dictyoprora* sp. A; Sample 183-1138A-34R-3, 105–107 cm. **12**. *Plannapus hornibrooki* O'Connor; Sample 183-1138A-34R-2, 23–25 cm. **13**. *Plannapus mauricei* O'Connor; Sample 183-1138A-35R-4, 104–106 cm. **14**. *Plannapus* sp. A; Sample 183-1138A-35R-3, 23–25 cm. **15**. *Dicolocapsa* sp. A (Haeckel); Sample 183-1138A-34R-3, 20–22 cm.

Family: Artostrobiidae



 $100 \ \mu m$