11. BENTHIC FORAMINIFERS AT SITE 397: FAUNAL FLUCTUATIONS AND RANGES IN THE QUATERNARY

Gerhard F. Lutze, Geol. Institute University Kiel, Olshausenstr. 40/60, 2300 Kiel, West Germany

ABSTRACT

Cyclic fluctuations of benthic foraminifers are not restricted to the youngest Pleistocene, but continue throughout the Pleistocene part of the section (137 m thickness) and into the late Pliocene. They are correlated with similar fluctuations in the planktonic foraminiferal assemblage, reflecting surface-water changes. Accordingly, changes in the Atlantic deep water circulation, believed to be responsible for the benthic fluctuations, must have begun in the late Pliocene/early Pleistocene. A zonation of the Pleistocene is tentatively proposed, enabling separation of the middle/late Pleistocene from the early Pleistocene by means of benthic foraminifers.

INTRODUCTION AND METHODS

Cyclic changes of benthic foraminifers in deep-sea cores have been reported by Streeter (1973), Lutze (1977), and Lutze, Sarnthein, et al. (this volume). Benthic foraminifers seemingly reflect drastic changes in Atlantic deep water circulation, which in turn are related to the well-known changes of the surface water during cool phases. This hypothesis needs much more refinement. Moreover, it is not known whether these cyclic changes are restricted to the glacial Pleistocene or even to the late Pleistocene (isotopic stage 1-5). DSDP Site 397 (off Cape Bojador, northwestern Africa; 2900 m water depth) offers a unique opportunity to extend the study of benthic fluctuation to the middle and early Pleistocene. Based on the extremely high rate of sedimentation, even the relatively low sampling density of one sample per section (i.e., per 15,000 years) would guarantee detection of the majority of cool and warm maxima, if they exist.

A further objective was to determine vertical ranges of important benthic species in order to identify potential guide fossils which might allow a local zonation of the Pleistocene.

The study is based on 111 samples, which were washed on a 63 µm sieve. Only the > 250 µm was statistically evaluated ("foram counts"), to avoid problems involved with changing amounts of allochthonous faunal components, which are more frequent in the finer fractions. Most samples were 15 cm³ in size. They usually contained between 80 and 150 tests, which were referred to some 60 species. Dominant species and species of stratigraphical or ecological importance were illustrated (Plates 1 to 3). A more conventional taxonomy was applied, based mainly on Phleger et al. (1953) and Parker (1964). Taxonomical remarks are given in the plate explanations.

To characterize faunal fluctuations related to cool or warm phases, the species were arranged in the same faunal groups used in the study of Meteor Core 12329 (Lutze, 1977). In this core, a very good correlation was found between a warm indicative group of benthic species and the climatic curve derived from planktonic foraminifers by Pflaumann (1975). To accentuate the fluctuations and to allow convenient comparison with other curves, the total proportion of this "warm" benthic species was expressed in percentages of both, the "warm" and the "cool" species group (whereas percentages of all other species were ignored). The resulting curve is called the warm benthic foraminifers (WBF) curve. "Cool" species are: Pyrgo murrhina, Cibicidoides wuellerstorfi, Uvigerina peregrina, and Globobulimina hoeglundi. WBF species are: Laticarinina pauperata, Cibicidoides kullenbergi, C. robertsonianus, Cibicidoides sp. 1, Hoeglundina elegans, and Bulimina aculeata.

FAUNAL FLUCTUATION

The general trend of faunal fluctuation is outlined in Figure 1 and is similar to that observed in the youngest Pleistocene Meteor Core 12309 of the same locality; see Lutze et al., this volume). Uvigerina/Globobulimina-maxima alternate with the WBF-assemblage peaks (mainly Cibicidoides sp. sp. and Hoeglundina). The fluctuations continue down to the late Pliocene. A total of about twenty warm maxima is marked by peaks of the WBF-curve (Figure 2).

Consequently, fluctuations in bottom water conditions, which are responsible for such drastic changes, must have existed throughout the entire time span under discussion here. This must be treated as a minimum estimate of the number of fluctuations, due to a sampling density of one sample per section (150 cm) which covers a time interval of roughly 10,000 to 18,000 years.

Comparison With δ¹⁸O Isotope Curve

The isotope curve given by Cita and Shackleton (this volume, δ¹⁸O based on benthic foraminifers) was compared with the WBF-curve as explained above.
Figure 1. Fluctuation of benthic foraminifera (< 250 µm.), cumulative percentage diagrams. Note the entirely different pattern observed in the top 10 meters, due to a higher sampling density in Meteor piston Core 12309. P = Pyrgo murrhina, W = Cibicidoides wuellerstorfi, M = Melonis pompilioides, B = Bulimina striata, R = Cibicidoides robertsonianus, C = Cibicidoides sp. 1.
BENTHIC FORAMINIFERS

Figure 2. Comparison of $\delta^{18}O$ isotope curve and the WBF curve ("warm benthic foraminifers"). The isotopic curve was taken from Cita and Shackleton (this volume). The WBF curve is based on benthic foraminifers, but different samples (see text).

(Figure 2). Both curves show approximately the same number of warm peaks and, in addition, similar positions of extended "cooler" periods. However, no strict peak for peak correlation is possible and several isotopic peaks correspond with depressions in the WBF curve. The correlation is slightly better in the early Pleistocene (Cores 10 to 16, Figure 3) but is weak in the late Pleistocene. This is partly due to differences in sampling positions. Shackleton's samples were usually taken 30 to 40 cm above the samples referred to in this study, which is equivalent to approximately 3000 to 4000 years. Distinctive shifts of the isotopic curve from cool towards warm may occur during such short intervals, as has been shown in Meteor Core 12309. To substantiate this explanation, an additional set of samples was studied from Cores 397-8 and 9 (new samples at 0.35 and 1.10 cm, original samples at 0.70 cm). The result is shown in Figure 4 with the two distinctive "warm" peaks (old position: broken line) separating into several peaks.

The discrepancy between warm isotopic values and "cool" foraminiferal evidence as found in Section 8-4 can now be explained by a "new" peak in lower Section 8-3 and upper 8-4. There is no doubt that an increased sampling density (20 cm distance) would give a resolution similar to that of Meteor Core 12309. It would result in the slopes of all peaks being much steeper and additional peaks being detected. Therefore, isotopic stage numbers (Shackleton and Opdyke, 1973) are applied only tentatively in Figure 2.

Correlation With Planktonic Foraminiferal Trend

The comparison with climatic evidence as indicated by planktonic foraminifer was restricted to the densely sampled interval (Cores 8 and 9) and only the $>250 \mu m$ fraction was used. A simple measure was found by calculating the percentage of typical tropical species from the sum of these and Globorotalia truncatuloides, which is a typical "transitional species" (Be and Tolderlund, 1971). Tropical species used in this
DIPLACED BENTHIC FORAMINIFERS

The amount of displaced benthic foraminifers decreases with an increase in grain size. Significant proportions of allochthonous species were found in the 63 to 125 µm fraction, fewer (up to 3%) in the 125 to 250 µm fraction. However, the true proportion of displaced specimens is difficult to estimate, since information on the taxonomy and the natural habitat of most species under discussion here is insufficient. The majority belongs to the genus Bolivina. Lutze shows that certain smaller Bolivinas B. pseudopectinata, B. difformis, and B. variabilis are true shelf dwellers. These species, at least in the smaller size fraction, must be considered allochthonous. They form considerable proportions in seven test samples selected to represent “cool” and “warm” maxima. The values for cool phases are about 14 per cent, those for warm intervals vary between 5 and 11 per cent. However, a more detailed study is required to find out whether these differences may be generalized. In the late Pleistocene (see Lutze et al., this volume) must higher proportions of allochthonous species were reported (up to 25%) but no significant difference between cool and warm intervals was noted.

Grain-by-grain transport, as discussed by Bein and Fütterer (1977), is suggested as being responsible for the displacement of these foraminifers. It is possible that sediment accumulation rates at continental margins are strongly influenced by this type of transport.

BIOSTRATIGRAPHY

The vertical ranges of the dominant species which are larger than 250 µm are given in Figure 7. Several typical Pliocene species disappear with the beginning of the Brunhes magnetic epoch: e.g., Stilostomella lepidula, Stilostomella sp. sp., Parafrondicularia advena, Orthomphila sp. 1, and Ellipsoglandulina laevigata. At this time, the first major cooling of Atlantic bottom water is also indicated by the isotopic record of Cita and Shackleton (this volume). Consequently this clear biostratigraphic differentiation is probably not locally restricted, which might therefore permit a subdivision of the Quaternary in this region. This would be exceptionally useful, since planktonic foraminiferal marker fossils in the Pleistocene have not proven to be reliable in all cases. It is also useful that the resulting boundary is equivalent to the early middle Pleistocene boundary as defined in the literature.

The subdivisions of the Pleistocene proposed here are named NB-5 and NB-6 according to the “faunal units” used in the Site 397 Report, this volume). A further subdivision is tentatively suggested: the first appearance of Bolivina quadrilaterta may enable the separation of NB-5b from NB-5a. A conspicuous feature within NB-6 (i.e., the middle and late Pleistocene) is the beginning of continuous occurrences of Hoeglundina elegans in the upper part of Core 4. This is used to differentiate between NB-6a and NB-6b. An additional subdivision might be possible with the last occurrence of B. quadrilaterta, separating NB-6c from NB-6b. This latter “boundary” would correspond with the beginning of the

study were Globorotalia cultrata and G. tumida, Pulteniatina obliquiloculata, and Globigerinoides sacculifer.

Figure 4 clearly demonstrates that both planktonic and benthic foraminifers follow the same trend of climatically induced cycling. The correlation between the benthic curve (WBF) and the planktonic curve is even better than in Meteor Core 12309 (Figure 5, r = 0.82).

RECENT DEPTH DISTRIBUTION, DISCUSSION

Benthic foraminiferal cycles and their possible origins have recently been discussed by Streeter (1973) and Schnitker (1974). New, detailed information has been reported recently by Streeter (1976; and May personal communication, 1977). He suggests that conditions typified by the present day shifted repeatedly to glacial situations such that the northern sources of oxygen-rich deep water were reduced or absent. During these periods oxygen-poor deep waters originated in the south. This “Pacific mode” of circulation is characterized by high frequencies of Uvigerina peregrina. The ecological importance of this species was emphasized by Lutze (1977, Leg 41) and is demonstrated again in the present study: U. peregrina and Globorotalia hof glohundi characterize cool periods. However, if Streeter’s explanation is correct, the present distribution of these low-oxygen tolerant species should not be strictly correlated with depth, since the oxygen content of North Atlantic deep water decreases with latitude (Dietrich et al., 1975, pl. 4; Wattenberg, 1939). The recent depth distribution of Uvigerina peregrina, as outlined on Figure 6 shows that the vertical boundaries of the distribution pattern and high abundances in low-oxygen areas support the hypothesis of Streeter.
Figure 4. Changes in the observed fluctuation pattern after additional sampling. Compare with the same interval on Figure 1. On the right: Correlation between the WBF-curve and tropical planktonic foraminifers (see text). Broken line: WBF curve after first sampling (1 sample per section).

Figure 5. Tropical planktonic foraminifers versus WBF percentages, Cores 8 and 9. Without Globigerinoides sacculifer.

Pleistocene. However, the somewhat scattered occurrence of B. quadrilatere does not suggest a more than local significance of this sub-unit.

ACKNOWLEDGMENTS
I would like to express my thanks to several friends and colleagues: S. Streeter, Lamont, provided access to unpublished manuscripts and valuable information. N. Shackleton made his oxygen-isotope curves available (see Cita and Shackleton, this volume). M. Sarnthein, Kiel, and U. Pflaumann, Kiel, discussed the results and made many suggestions. Financial support from the “Deutsche Forschungsgemeinschaft,” Bonn, is appreciated.

REFERENCES
Bein, A. and Fütterer, D., 1977. Texture and composition of continental shelf to rise sediments off the northwestern coast of Africa: An indication for downslope transporta-
leontol., v. 2, p. 55-84.
Parker, F. L., 1964. Foraminifera from the experimental Mo-
hole drilling near Guadalupe Island, Mexico, J. Paleon-\ttol., v. 38, p. 617-636.
Pflaumann, U., 1975. Late Quaternary stratigraphy based on planktonic foraminifera off Senegal. “Meteor” Forsch-
Pujos-Lamy, A., 1973. Repartition bathymétrique des foram-
Wattenberg, H., 1939. Atlas zu: Die Verteilung des Sauer-
stoffs im Atlantischen Ozean. Wissenschaftliche Ergebnisse der Deutschen Atlantischen Expedition auf dem Forschungs- und Vermessungs-schiff “Meteor,” 1925-
1927, Band IX, Atlas.
Figure 6 (a-d). Depth distribution of dominant species in modern surface sediments at the eastern North Atlantic slope and rise. Per cent of the death assemblage larger than 250 µm. The diagrams are part of a detailed study on depth ranges off West Africa (Lutze, Mascr.). Stations north of 45°N refer to literature (Pujos-Lamy, 1973; Caralp et al., 1970). Note the vertically shaped pattern of distribution in Figure 6a and 6c.
Figure 7. Ranges of dominant benthic foraminifers of the fraction 250 µm. The occurrence of several “warm” species (e.g., C. kullenbergi, C. robertsonianus, Laticarinina pauperata) is often restricted to warm phases (stippled intervals). The species numbers refer to the figures given on Plate 1 through 3. Nannofossil boundaries are given in the upper right, according to Čepel this volume. Magnetic stratigraphy according to Hamilton, this volume.
Figure 1  *Bulimina aculeata* Orbigny, 1926. SEM, Sample 397-4, 3-61 cm, size 0.45 mm.

Figure 2  *Bulimina striata* Orbigny, 1843. SEM, Sample 397-3, 5-75 cm, size 0.65 mm. Some authors use the name *B. costata*.

*Figure 3b*  *Uvigerina peregrina* Cushman, 1923. SEM, Sample 397-11, 5-69 cm, sizes 0.57 mm and 0.82 mm.

Figures 4a, 4b  *Pyro murrhina* (Schwager, 1886). Sample 397-4, 3-61 cm, sizes 1.35 mm.
4a. SEM, apertural detail.
4b. Light microscope.

Figures 5a, 5b  *Cibicidoides wuellerstorfi* (Schwager, 1866). Sample 397-6, 2-69 cm.
5a. SEM.
5b. Light microscope. Sizes 0.75 mm and 0.7 mm. placed in the genus *Planulina*; see Parker, 1964, p. 624.

Figure 6  *Globobulimina hoeglundi* Uchio, 1960. SEM, Sample 397-5, 3-69 cm, size 0.77 mm. Several authors referred to this species to *G. affinis* (Orbigny). I prefer Uchio’s name because it appears to be more precisely defined and topotypes are at hand.

Figure 7  *Melonis zandami* (Voorthuysen, 1952). SEM, Sample 397-10, 7-? cm, size 0.45 mm. Originally referred to *Anomalinoideae*. (For details, see Loebling and Tappan, 1953, Artic Foraminifera.) Most authors refer to this species to *M. baarleanum*. 
PLATE 2

Figures 1a, 1b *Melonis pompilioides* (Fichtel and Moll, 1798).
1a. SEM.
1b. Light microscope, apertural face. Sample 397-7, 4-70 cm, sizes 0.42 mm and 0.525 mm.

Figures 2a, 2b *Hoeglundina elegans* (Orbigny 1826).
2a. SEM, umbilical side.
2b. Spiral side, light microscope, Sample 397-2, 6-40 cm, sizes 1.17 mm and 0.87 mm.

Figures 3a, 3b *Cibicidoides* sp. 1, SEM.
3a. Spiral side, Sample 397-7, 4-70 cm, size 0.57 mm.
3b. Umbilical side, Sample 397-3-4-87 cm, size 0.5 mm.

Figures 4a, 4b *Cibicidoides robertsonianus* (Brady, 1881).
4a. Light microscope, spiral side.
4b. SEM, spiral side, Sample 397-4, 3-61 cm, sizes 0.82 mm and 0.57 mm.

Figures 5a, 5b *Cibicidoides kullenbergi* Parker, 1953.
5a. SEM, Sample 397-3, 4-87 cm, size 0.52 mm.
5b. Light microscope, Sample 397-9, 1-35 cm, size 0.6 mm. Both figures show spiral side.

Figure 6 *Laticarinina pauperata* (Parker and Jones, 1865).
Light microscope, Sample from Meteor Core 12309-2, 20 cm, size 1.75 mm.

Figure 7 *Oridorsalis umbonatus* (Reuss, 1851).
Sample 397-13, 2-70 cm, size 0.5 mm.
PLATE 2

BENTHIC FORAMINIFERS

1a  2a  2b
1b  3a  3b
4a  4b
5a  5b
6  7
PLATE 3

Figure 1  *Sigmoilopsis schlumbergeri* (Silvestri, 1904). SEM, Sample 397-7, 4-70 cm, size 0.72 mm.

Figure 2  *Sphaeroidina bulloides* Orbigny, 1826. Light microscope, Sample 397-6, 5-71 cm, size 0.87 mm.

Figure 3  *Bolivina quadrilatera* (Schwager, 1866). SEM, Sample 397-4, 1-69 cm, size 0.95 mm.

Figures 4a, 4b  *Pleurostomella brevis* (Schwager, 1866). SEM, Samples 397-12, 1-70 cm and 397-12, 5-70 cm, sizes 0.67 mm and 0.85 mm. The name was used in a rather wide sense. SEM studies of apertural details of various *Pleurostomella* species are required to enable a more exact identification and the separation of different taxons.

Figure 5  *Parafrondicularia advena* (Cushman, 1923). SEM, Sample 397-8, 3-67 cm, size 1.3 mm.

Figures 6a, 6b  *Orthomorphina? sp.* SEM. Sample 397-10, 1-70 cm; sizes 1.0 mm and 1.92 mm. Specimens with larger spines (Figure 20A) and those with small spines, nearly invisible under the light microscope (Figure 26B), were counted together. Close to *Orthomorphina? fistuca* (Schwager, 1866).

Figure 7  *Ellipsoglandulina laevigata* Schwager. 1900, light microscope, Sample 369-2, 5-60 cm, Pliocene, size 1.3 mm.