CAMPANIAN TO PALEOCENE AGGLUTINATED FORAMINIFERA FROM FRESHWATER INFLUENCED MARGINAL MARINE (DELTAIC) SEDIMENTS OF SOUTHERN EGYPT

by

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With 3 figures and 2 plates

ZUSAMMENFASSUNG

Einige artenarme Assoziationen agglutinierender Foraminiferen aus obercampanen bis paleozänen Sedimenten Südägyptens werden vorgestellt und diskutiert. Die Gattung *Ammoastuta* wird erstmalig aus Nordostafrika beschrieben. Der Vergleich mit anderen rezenten und fossilen Foraminiferenassoziationen, in denen *Ammoastuta* vorkommt, zeigt die Verwendbarkeit dieser Gattung als Faziesleitfossil für küstennahe, brackische Flachwasserbereiche warme Paläoklimate mit hohen Niederschlägen, auch wenn es sich bei ausschließlich aus agglutinierenden Foraminiferen bestehenden Assoziationen größtenteils um Oryktozönosen (Rückstandsgemeinschaften) mit diagenetisch reduzierten Faunenspektren handelt.

ABSTRACT

Low diversity agglutinated foraminiferal assemblages from the Campanian to Early Paleocene of southern Egypt are described and discussed. The occurrence of the genus *Ammoastuta* is reported for the first time from northeast Africa. *Ammoastuta* is regarded as a facies-index fossil for brackish littoral conditions in warm climates with high rainfall and high runoff.

INTRODUCTION

From Aptian/Albian to Eocene time, marine transgressions from the northern Tethyan Sea invaded the large intracratonal basins of southern Egypt. The major transgression started during the Campanian and, interrupted by minor regressive phases, culminated during the Late Paleocene. Due to a constantly high input of terrestrial material from the south, the Campanian to Middle Paleocene sediments can be subdivided into a northern pelitic basinal and a southern psammitic marginal lithofacies with very gradual interfingering of sedimentary environments. The psammitic marginal lithofacies shifted southwards with increasing transgression. This facies differentiation was studied in a number of sections of Maastrichtian to Middle Paleocene sediments along the 350 km long, NNE/SSW trending escarpment of the great limestone plateau west of the River Nile (Luger 1985). The Campanian data are derived from a different area further to the north: Gebel Qreiya, at the southern end of Wadi Qena, Eastern Desert (figure 1).

THE FORAMINIFERAL ASSOCIATIONS OF SOUTHERN EGYPT

The sediments of the upper Hawashya Formation (mainly pelitic facies, Campanian), Dakhla Formation (mainly pelitic basinal facies,

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Fig. 1. Occurrence of Ammoastuta assemblages in Egypt.

Maastrichtian to Middle Paleocene), Shab Member of Kiseiba Formation (mainly psammitic marginal/deltaic facies, Maastrichtian) and Kurkur Formation (mainly psammitic marginal/deltaic facies, Paleocene) yielded foraminiferal assemblages which may roughly be grouped into the following three categories (for stratigraphic correlation see figure 2):

1. Midway fauna (Paleocene) and Midway-like (Maastrichtian) assemblages:

Composition: Calcareous and agglutinated benthics, abundant to common planktonics. Agglutinated benthics use calcareous and siliciclastic material for building their tests; forms using calcareous material dominate the agglutinated association. Benthic foraminiferal species diversity is high.

Origin: Middle to outer shelf.

Occurrence: Dakhla Formation (Maastrichtian to Middle Paleocene).

2. Shallow shelf assemblages:

Composition: Calcareous and agglutinated benthics, planktonics rare to absent. Agglutinated benthics use calcareous and siliciclastic material for building their tests; forms using siliciclastic material dominate in the agglutinated association. Benthic species diversity is low.

Origin: Shallow euryhaline shelf or lagoons.

Occurrence: Dakhla Formation (Maastrichtian part only).

3. "Littoral" associations:

Composition: Almost exclusively agglutinated foraminifera, using only siliciclastic material for building their tests. Foraminiferal species diversity is very low.

Origin: ?euryhaline lagoons to mixohaline lagoons, marshes and coastal swamps.

Occurrence: Dakhla Formation (Maastrichtian part only), Kurkur Formation, upper Hawashya Formation, Shab Member of Kiseiba Formation.

DESCRIPTION OF THE "LITTORAL" ASSOCIATIONS

The "littoral" associations are found mainly in noncalcareous pelitic sediments which may have been decalcified in early diagenetic stages. Therefore they cannot be interpreted as taphocoenosis but instead as oryktocoenosis (=relic assemblages). Among them the following types occur:

A)Composition: Ammobaculites div. sp., Haplophragmoides div. sp., Reophax div. sp., Saccammina sp., Textularia? sp., Trochammina sp.

Species diversity: 1-6

Occurrence: Basinal and marginal facies, Campanian to Middle Paleocene.

B)Composition: Miliammina sp., div. Ammobaculites div. sp., Ammodiscus sp., Haplophragmoides sp., Trochammina sp. Species diversity: 1-8.

Occurrence: Transition between basinal and marginal facies as well as marginal facies, Campanian to Maastrichtian.

C)Composition: Ammoastuta div. sp., Ammobaculites div. sp., Ammodiscus sp., Haplophragmoides div. sp., Miliammina div. sp., Reophax sp., Trochammina sp. Species diversity: 1-8

Occurrence: Interfingering of basinal and marginal facies as well as marginal facies, Campanian to Early Paleocene.

The most important constituents of the associations A-C are figured on plates 1 and 2. Most of the components of the associations of type A may cooccur in the shelf assemblages of categories 1 and 2. As decalcification may have occurred, some of the associations of type A could represent oryktocoenosis of normal marine shelf assemblages.

The important constituents of types B and C, Ammoastuta div. sp. and Miliammina div. sp., (plates 1 and 2), occur exclusively in associations of category 3. The associations type B and C are usually found in pelitic sediments which contain high amounts of terrestrial plant debris (mainly Gymnosperms). Occasionally they occur in the pelitic parts of fining-upwards sequences of shallowing cut-and-fill channel systems. Sandstones with root relics sometimes were observed in the vicinity of the sediments bearing the low-diversity agglutinated associations. Very rarely calcareous rotaliid foraminifera (Anomalinoides sp. only, plate 2), occur in associations of type C. This indicates that a possible decalcification did not drastically change the generic composition of the associations of this type.

NE	Kharga - Paric	Bir Murr - Toshka
Esn	a Formation	BII Mull - IOSIKa
	. <u> </u>	Garra Formation
Tarawan Form		
Dakhla Form	ation	Kurkur Formation* 🗙 ★ 🛧 🛧
		Hiatus
Dakhla Formation \star		
		Kiseiba
Rakhiyat Formation	Duwi Formation	Formation
Hawashya	Mut Formation	Lower Member
	NE Qena Esn Tarawan Form Dakhla Form Dakhla Form Dakhla Form Rakhiyat Formation Hawashya	NE Qena Kharga - Baris Esna Formation Tarawan Form. Dakhla Formation Dakhla Formation Rakhiyat Duwi Formation Hawashya * * * Mut Formation

★ = Occurence of Ammoastuta assemblages

Fig. 2.

Simplified stratigraphic table of the Campanian to Paleocene lithological units in southern Egypt. Not scaled.

In associations of type C, the genus Ammoastuta in Egypt is represented by different species. In the Campanian occurs Ammoastuta sp., in the Maastrichtian (Ammoastuta megacribrostomoides Luger, 1985) and in the Early Paleocene Ammoastuta aegyptiaca Luger, 1985:see plate 2). These species differ from each other in the shape of the test and in the form of the secondary cribrate aperture. None of the different species occur together in the few localities where the genus is observed. It is not clear whether this phenomenon is controlled by biostratigraphical or paleocological reasons, for example, possible restriction of species to certain sub-environments controlled by salinity, substrate oxygenation, pH etc.

Palynological investigations carried out by Schrank (*in*: Schrank and Perch-Nielsen 1985) proved the existence of different species of *Spinizonocolpites* in Maastrichtian deposits of several localities in the middle latitudes of Egypt. *Spinizonocolpites* is the pollen of the *Nypa*-palm, a mangrove-like plant, known from the tropical regions of southeast Asia in Recent time. The occurrence of *Spinizonocolpites* in open marine sediments of the Maastrichtian clearly indicates the existence of extended mangrove-type coastal swamps along the southern sea shore during that time. This facies may be represented by some sediments of the Shab Member of the Kiseiba Formation.

COMPARISON WITH RECENT ASSEMBLAGES

Associations very similar to those of types B and C were recorded by Phleger (1954, 1955) from Recent sediments of the Gulf of Mexico and the Mississippi Delta. There, Ammoastuta, Ammobaculites, Arenoparella, Miliammina, Haplophragmoides, Leptodermella, Proteonina (=Reophax), Trochammina and Urnulina live in mixohaline marshes and bays. Rare calcareous foraminifera such as Elphidium, Rotalia and miliolids also occur. Hiltermann and Tüxen (1978) carried out a statistical analysis based on Phleger's results and named assemblages containing living Ammoastuta "Ammoastutetum ineptae". They demonstrated the restriction of living Ammoastuta to the mixohaline sedimentary environments mentioned above. Dislocated dead specimens of this genus were rarely found in the Mississippi Sound. Recent Ammoastuta are known from mixohaline marshes and bays of the Mississippi Delta area and Trinidad as well as from brackish mangrove swamps of Brazil (Hiltermann et al. 1981; Brönnimann 1986). Unlike Miliammina, the genus does not occur in hypersaline environments. If the genera not known before the Miocene (Arenoparella, Leptodermella, Urnulina)

and the calcareous forms which might not be preserved in Recent brackish assemblages given by Phleger (1954, 1955) and Hiltermann and Tüxen (1978) are excluded, the remaining generic composition of the Recent Ammoastuta assemblages ("Ammoastutetum ineptae") appears to be almost identical to the fossil record.

FOSSIL AMMOASTUTA ASSEMBLAGES

Fossil Ammoastuta assemblages, consisting of Ammoastuta, Ammobaculites, Haplophragmoides, Trochammina and others have been described from the Eocene of Rio Guamal, Columbia (Petters, V. 1954) and the Turonian-Santonian Pindinga Formation in northwestern Nigeria (Petters, S.W. 1979). They were interpreted to be of brackish lagoonal to marsh origin by the authors mentioned above. The Nigerian, Columbian and the Egyptian Ammoastuta assemblages are found in the vicinity of the Late Cretaceous to Early Paleocene paleoequator, *i.e.* in areas in which warm climates with high rainfalls and high runoff have to be assumed (figure 3).

Some fossil foraminifera, such as "Ammoastuta" curfsi Hofker (1966, from marine strata of the Maastrichtian-Danian from Denmark) and "Ammoastuta" sakhalinica Voloshinova (1961, Miocene) have erroneously been ascribed to the genus Ammoastuta. These forms do not fit the generic definition of Ammoastuta as they lack the typical secondary aperture (for emended generic definition see: Brönnimann 1986).

CONCLUSIONS

Recent living Ammoastuta is known only from brackish littoral environments. The fossil and Recent Ammoastuta assemblages and the (paleo-)climatic conditions of the areas they are found in are very similar. This clearly indicates the value of the genus Ammoastuta as an ecological index fossil for brackish littoral environments. Therefore, associations consisting mainly or entirely of agglutinated foraminifera (type C of this paper) in which Ammoastuta occurs, may be interpreted to be of mangrove-swamp, marsh or lagoonal origin with considerably reduced salinity in warm climates with high runoff. The possibility of dead Ammoastuta specimens having been transported into normal marine environments can be disregarded as far as the Egyptian material is concerned. The studied specimens have a very delicate test which would not have withstood long-distance transport. Also, no specimens have been found in associations, which indicate euryhaline conditions during sedimentation.



Known fossil occurrences of Ammoastuta assemblages (\blacktriangle). Maps after SMITH et al. (1982). Left = Late Cretaceous; right = Paleogene.

The Ammoastuta associations are known from the Late Cretaceous to Recent times. Future investigations may show whether or not the different species of this genus can be used as stratigraphical markers in extreme environments which are usually devoid of any index fossils.

Low-diversity agglutinated for a miniferal associations with *Miliammina* as the main constituent, in which *Ammoastuta* is not present, may be interpreted as being of marsh to lagoonal origin. However, if no further information is available, it cannot be decided whether they were formed under mixohaline or hypersaline conditions (type B). Low-diversity agglutinated associations without any of the two forms mentioned above may indicate extreme environmental conditions during sedimentation, but may just as well represent oryktocoenosis of euryhaline shallow shelf assemblages (type A).

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PLATE 1

Figure 1	Saccammina sp., distorted specimen, Maastrichtian, Dakla Formation, about x135.
Figure 2	Ammodiscus cretaceus (Reuss), Maastrichtian, Shab Member of Kiseiba Formation, about x135.
Figure 3	Ammodiscus sp., Campanian, upper Hawashya Formation, about x340.
Figure 4	Miliammina telemaquensis Saunders, Maastrichtian, Shab Member of Kiseiba Formation, about x340.
Figure 5	Miliammina onyeamensis Petters, Maastrichtian, Shab Member of Kiseiba Formation, about x340.
Figure 6	Reophax texanus Cushman and Waters, Maastrichtian, Shab Member of Kiseiba Formation, about x135.
Figure 7	Reophax texanus Cushman and Waters, Maastrichtian, Shab Member of Kiseiba Formation, about x135.
Figure 8	Haplophragmoides calculus Cushman and Waters, Maastrichtian Shab Member of Kiseiba Formation, about x135.
Figure 9	Ammobaculites sp., Campanian, upper Hawashya Formation, about x135.
Figure 10	Ammobaculites fragmentarius Cushman, Shab Member of Kiseiba Formation, about x135.
Figure 11	Ammobaculites subcretaceus Cushman and Alexander, Maastrichtian, Shab Member of Kiseiba Formation, about x85.
Figure 12	Ammobaculites coprolithiformis (Schwager), Paleocene, Kurkur Formation, about x125.

2



PL.	ATE	2
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Figures 1, 2	Ammobaculites texanus Cushman, Maastrichtian, Dakhla Formation, about x50.
Figure 3.	Ammoastuta sp., Campanian, upper Hawashya Formation. a) lateral view, about x135.
	b) same specimen, front view of last chamber, note position of primary oval aperture at about one-third of chamber length from the secondary cribrate aperture, about x145.
	c) same specimen, view of secondary cribrate aperture, about x340.
Figures 4,5	Ammoastuta aegyptiaca Luger, Paleocene, Kurkur Formation, about x135. 4a) lateral view.
	4b) same specimen as 4a, semi-lateral view, note small secondary cribrate aperture restricted to the last chamber.
	b) lateral view.
Figures 6,7	Ammoastuta megacribrostomoides Luger, Maastrichtian, Shab Member of Kiseiba Formation, about x135.
Figure 8	Trochammina afikpensis Petters, umbilical view, Paleocene, Kurkur
	Formation, about x135.
Figure 9	Trochammina sp., Campanian, upper Hawashya Formation, about x135.
Figure 10	Anomalinoides sp., Maastrichtian, Dakhla Formation, about x135.

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