Original article

Bioturbational redistribution of Danian calcareous nannofossils in the uppermost Maastrichtian across the K-Pg boundary at Bidart, SW France

Remaniement par la bioturbation de nannofossiles calcaires daniens dans le Maestrichtien terminal autour de la limite C-Pg dans la coupe de Bidart (SO France)

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Abstract

In this paper we present the effects of different tracemakers on the redistribution of calcareous nannofossils throughout the K-Pg boundary at the Bidart section (SW France), along with their consequences for our knowledge of the K-Pg boundary event. Danian calcareous nannofossil assemblages are present in Maastrichtian samples due to infiltration into dark trace fossil infillings proceeding from the earliest Paleogene. This is evidenced by the appearance of abundant paleogene calcareous nannofossils just below the K-Pg boundary, showing the relevance of the trace fossil infillings in the context of the K-Pg boundary event.

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Keywords: Cretaceous-Paleogene boundary; Trace fossils; Calcareous nannofossils; Redistribution; Bidart section; SW France

Résumé

Dans cet article, nous présentons les effets des organismes bioturbateurs sur le remaniement de nannofossiles calcaires autour de la limite C-Pg dans la coupe de Bidart (SW France), ainsi que leurs conséquences pour la compréhension de l’événement de la limite C-Pg. Des assemblages de nannofossiles typiquement daniens sont présents dans des échantillons du Maestrichtien, remaniés par infiltration, dans des remplissages de couleur sombre de traces fossiles formées au début du Paléogène. Cela est monté par la présence d’abondants nannofossiles calcaires paléogènes juste sous la limite C-Pg, montrant tout l’intérêt de l’étude des remplissages de traces fossiles dans le contexte de l’événement de la limite C-Pg.

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Mots clés : Limite Crétagé-Paléogène ; Traces fossiles ; Nannofossiles calcaires ; Remaniement ; Coupe de Bidart ; SO France

1. Introduction

The Cretaceous-Paleogene (K-Pg) boundary has been profusely studied, mainly due to the major biological crisis associated with it (see Koutsoukos, 2005; Molina et al., 2006; Kring, 2007 for recent reviews). The extinction pattern is still controversial, mainly supported by gradual or catastrophic interpretations of the extinction of microorganisms, in which the existence of Danian taxa considered as survivors or reworked specimens must play a key role. This is especially true for the planktic calcareous organisms (foraminifers and calcareous nannofossils). Some authors interpret the presence of Maastrichtian taxa found in the Danian sediments as totally or partially reworked (e.g., Smit, 1990; Pospichal, 1994; Henriksson, 1996; Molina et al., 1998; Tantawy, 2003; Gallala et al., 2009), while others suggest that some Maastrichtian taxa survived the disaster but disappeared in the earliest Danian (e.g., Perch-Nielsen et al., 1982; Keller, 1988; Keller et al., 1995; Gardin and Monechi, 1998; Gardin, 2002; Bown, 2005).
Due to their very small size, calcareous nannofossils are comparatively easily resedimented and reworked. This fact has been brought forth in several papers on the K-Pg boundary, showing Cretaceous taxa in Danian samples and vice versa; but only occasionally has the incidence of the activity of trace makers on calcareous nannofossil redistribution been considered, and in almost all cases this aspect has been alluded to in general terms (e.g., Thierstein and Okada, 1979; Thierstein, 1981; Smit and Romein, 1985; Pospichal and Wise, 1990; Pospichal et al., 1990; Henriksson, 1996; Pospichal, 1996; Romein et al., 1996; Mai et al., 2003; Bown, 2005; Lamolda et al., 2005). Thus, the role of trace makers in the reworking of microfossils is usually underestimated. It has recently been demonstrated, however, that trace fossils throughout the K-Pg boundary interval are relatively abundant and diverse, colonizing lower Danian sediments and penetrating the light Maastrichtian marlstones (Rodríguez-Tovar and Uchman, 2004, 2008; Rodríguez-Tovar et al., 2004; Rodríguez-Tovar, 2005).

The K-Pg boundary section at Bidart is held to be a key section for interpreting the K-Pg boundary event (Galbrun and Gardin, 2004). Very recent planktic foraminifera analysis suggests a sudden catastrophic mass extinction at the K-Pg boundary affecting at least 53 species (Gallala et al., 2009).

Fig. 1. Location of the Bidart section between Hendaya and Biarritz (south-western France), and other important K-Pg boundary sections (Zumaya and Sopelana), and general stratigraphic column from the upper Maastrichtian to the lower Danian with location of the studied samples.

2. Geological setting

The Bidart section is located between Hendaya and Biarritz (south-western France), in the Basque-Cantabrian Basin (Fig. 1), cropping out along a beach a few kilometres long towards the north at Bidart (43° 27' N and 1° 35' E). The magnetostratigraphy and biostratigraphy of the Cretaceous-Tertiary boundary of Bidart was established by Delacotte et al. (1985). From a palaeogeographic point of view, the K-Pg boundary sections of the Basque Basin are located in a deep-basin environment, between 800 and 2.000 m depth (Orue-Itxebarría et al., 1996; Apellániz et al., 1997; Pujalte et al., 1998), with a dominant pelagic and/or hemipelagic sedimentation, showing local turbidites (Apellániz et al., 1997). The Bidart section is believed to be situated in close proximity to a palaeoslope to the northeast (Seyve, 1984), in a distal deep-sea fan environment with a 40 mm/ky sedimentation rate during the Maastrichtian (Nelson et al., 1991), showing increased activity on the slope – slumped blocks and turbidites are registered – during the Paleocene (Seyve, 1990). The study of benthic foraminifera points to the upper-middle part of the slope for sediment deposition at the Bidart section from the upper Maastrichtian to the lower Danian (Alegret et al., 2004). The palaeogeographic situation is particularly interesting, between the Tethys and the North Atlantic, showing a change in deposition influences from Tethyan during Maastrichtian, then to primarily North Atlantic during the Paleocene (Renard et al., 1982). Benthic foraminiferal assemblages reveal significant palaeoenvironmental changes throughout the K-Pg boundary interval, showing mesotrophic conditions during the late Maastrichtian and sufficient nutrient supply to the sea-bottom floor to sustain both infaunal and epifaunal morphogroups, with a strong decrease in food supply to...
the sea floor at the K-Pg boundary (Alegret et al., 2004). This food supply was not completely recovered for over 200 kyr, at least, after the K-Pg boundary (Alegret et al., 2004).

The planktic foraminiferal assemblage across the K-Pg boundary is highly diversified, containing up to 72 species, and with a catastrophic sudden mass extinction affecting at least 53 out of these species (Gallala et al., 2009). The K-Pg boundary extinct species include globotruncanids (e.g., *Contusotruncanana* spp., *Globotruncanana* spp., *Globotruncanita* spp.) and complex heterohelicids (e.g., *Racemiguembelina* spp., *Pseudotextularia* spp., *Gublerina* spp.). Specimens of 18 Maastrichtian species, small and even tiny in size, were found at the lowermost Danian. Few of these species belong to *Guembelitria* or *Hedbergella*, and it is likely that *Heterohelix* is the only real Cretaceous survivor species. Contrariwise, the specimens belonging to the rest of the 14 species (mainly *Heterohelix*, *Pseudoguembelina*, *Globotruncanella* and some species of *Rugoglobigerina* and *Globotruncanana*) seem to be reworked, according to their different taphonomic preservation (mainly different colour and fragmentation).

3. Material and method

The K-Pg boundary transition was studied at the Bidart section. The upper 4 m of the Maastrichtian contain grey and pinkish marls and marly limestones (Figs. 1 and 2). The K-Pg boundary is easily identifiable at the base of the boundary layer; a “rusty” layer, about 2 mm thick, is characterized by an Ir anomaly. Above the “rusty layer”, at the base of the Danian, a dark clay layer about 15 cm thick occurs, overlapped by about 23 cm of marls and marly limestones, which are reddish brown in the middle part of the layer and green and partly laminated towards the top. Above them, a 93 cm thick interval of white and reddish, partly parallel laminated limestones is present, followed by a few beds of white and reddish debris flow beds mostly made up of calcirudites. At the top of this interval the studied section consists of white and reddish bioturbated limestones. Continuous bed-by-bed detailed ichnological analysis was conducted 4 m below and above the K-Pg boundary, focusing on recognition of ichnotaxa and their vertical distribution, type of filling, cross-cutting relationships, ichnofabrics and other features. Moreover, isolated samples were collected for ichnological analysis in the laboratory.

Calcareaous nannofossils from sediments hosting trace fossils were analyzed in a span of about 5 m across the K-Pg boundary transition: 4 m in the uppermost Maastrichtian and 40 cm of the lowest part of the Paleogene (Figs. 1 and 2). Below the K-Pg boundary surface, samples were collected at −400 cm (Bd-1), −200 cm (Bd-6), −40 cm (Bd-11), −35 cm (Bd-9), and −3–0 cm (Bd-13). Above the K-Pg boundary surface, Bd-14 is located just above the boundary, and Bd-15, Bd-17, Bd-19 and Bd-22 are located at 6, 17, 29 and 40 cm, respectively. Analyses were focused on trace fossils filled with a dark-coloured sediment similar to that of the lowermost Danian marls and occasionally on light-coloured burrows occurring below the K-Pg boundary. Samples were selected on the basis of the type and size of trace fossils, and consequent possibilities for analyzing the filling material, location and depth of penetration; thus only *Thalassinooides* and *Zoophycos* structures were considered. Present in Bd-9 were dark *Thalassinooides* (Bd-9 df), in Bd-11 dark *Zoophycos* (Bd-11 dfz), and in Bd-13 dark (Bd-13 df) and light *Thalassinooides* (Bd-13 flf).

Samples for nannofossil investigation were prepared as follow: a small chip of sediment has been powdered and add distilled water. Few drops of the mixture were then spread on a coverslip and dried. The coverslip were glued on the glass slide
with a drop of Norland optical Adhesive (NOA-65). Unprocessed material was used in order to avoid mechanical and physical processes that could modify the original composition of the assemblages. The calcareous nannofossils were analyzed under a Zeiss Axioplan2 petrographic microscope at ×1560 magnification. Qualitative analyses have been performed on the studied samples in order to characterize the assemblage and at least three traverses of the slides were examined to find rare taxa.

Several detailed calcareous nannofossil studies describe the K-Pg transition of Bidart (Gorostidi and Lamolda, 1995 and reference therein; Galbrun and Gardin, 2004; Thibault et al., 2004; Minoletti et al., 2005). We refer to these papers for the complete biostratigraphic zonation, distribution and exhaustive palaeoecological calcareous nannofossil changes across the K-Pg boundary interval.

4. Results

4.1. Trace fossil assemblage

Relatively abundant and conspicuous dark-coloured trace fossils occur throughout the studied interval, including Chondrites, Planolites, Thalassinoides, Trichichnus and Zoophycos (Fig. 2), on totally bioturbated sediment background related to the activity of unknown organisms in the uppermost part of the substrate. Less abundant light-coloured ichnofossils include Planolites and Thalassinoides.

Chondrites isp. is observed mostly in cross section as patches of circular to elliptical spots and short bars, which some of them are branched, filled with a darker material than the surrounding rock. They are cross sections of a branched tubular burrow system. A larger form, 2–3 mm in diameter, and smaller form, up to 1 mm diameter, can be distinguished. Limited observations along the tunnels preclude more detailed determinations, however taking into account the size parameters (Uchman, 1999), the smaller form can belong to Chondrites intricatus (Brongniart, 1823) and the larger one to Chondrites targinii (Brongniart, 1828). The smaller form can be found in the fillings of some Thalassinoides or Planolites.

Chondrites von Sternberg, 1833 is a feeding system of unknown trace makers (Osgood, 1970) which, according to Kotake (1991a), are surface ingestors, packing their faecal pellets inside burrows. According to Seilacher (1990, 2007) and Fu (1991), the trace maker may be able to live at the aerobic-anoxic interface as a chemosymbiotic organism. Chondrites shows a wide stratigraphic and marine environmental range.

Planolites isp. occurs as horizontal to oblique, straight to slightly winding, simple flattened cylinders, 2.5–4 mm, without lining. They are filled with dark grey fine-grained material.

Planolites is a facies crossing form. It has been discussed by Pemberton and Frey (1982) and Keighley and Pickerill (1995) and interpreted as a pascichnion, probably produced by a number of different organisms.

Thalassinoides isp. is registered as straight or slightly winding, horizontal to oblique flattened, unlined cylinders, showing Y-shaped branches and shafts, filled with homogenous, or rarely meniscate, mostly darker sediment. The cylinders are swollen at the branching points. Smaller forms are 5–9 mm in diameter, with swellings up to 15 mm wide. Some forms, especially in the Danian, are larger, 15–25 mm in diameter, which swellings are up to 35 mm wide.

Thalassinoides Ehrenberg, 1944 is a domichnial and fodinichnial structure produced by crustaceans, mostly decapods (Frey et al., 1984). It occurs in a great variety of marine environments, yet is most typical for the shelf Cruziana ichnofacies. For further discussion of this ichnogenus and its ichnontaxonomic problems see Fürsich (1973), Ekdale (1992) and Schlirf (2000).

Trichichnus isp. is a winding or straight, differently oriented, rarely branched, 0.2–0.3 mm thick, thread-like cylinder filled with ferruginous substance, with a yellowish halo around the cylinder.

Trichichnus Frey, 1970 is a eurybathic marine trace fossil, which is common in fine-grained sediments. Its fillings display a strong tendency to pyritization (e.g., Werner and Wetzel, 1981; McBride and Picard, 1991). Trichichnus is regarded as the domicnial burrows of marine meiofaunal deposit-feeders (Frey, 1970). Possibly, the producer of Trichichnus was a chemosymbiont (Uchman, 1995), as in the case of the Chondrites producer (Seilacher, 1990; Fu, 1991). McBride and Picard (1991) suggest that Trichichnus had a more opportunistic character than Chondrites because it occurs more deeply in very poorly oxygenated sediments. For taxonomic discussion see Uchman (1999).

Zoophycos isp. occurs on parting surfaces as horizontal or oblique lobes and tongues, up to 40 cm wide, filled with spreite laminae encircled by a thin marginal tunnel, which are a part of a helical burrow system. In cross section it is seen as stacked stripes, 1.5–3 mm thick, which can be massive or filled with spreite manifested on the surface as menisci-like structures.

Zoophycos s.l. is generally considered as a structure produced by some yet undiscovered deposit-feeder, which is referred to as sipunculids (Wetzel and Werner, 1981; Olivero and Gaillard, 2007), polychaete annelids, arthropods (Ekdale and Lewis, 1991), or echinuran worms (Kotake, 1992). Kotake (1989, 1991b) showed that Zoophycos is produced by surface ingestors of organic detritus. The precise ethological interpretation of Zoophycos remains controversial. This is probably due to the assignment to Zoophycos s.l. of diverse structures generated by different tracemakers with different behaviour, or even a single tracemaker with diverse behaviours (Kotake, 1989, 1994; Bromley, 1991; Locklair and Savrda, 1998; MacEachern and Burton, 2000; Olivero and Gaillard, 2007). Bromley and Hanken (2003) suggested that the upper helical part of a large Pliocene Zoophycos from Rhodes, Greece, is a deposit-feeding structure, and lateral lobes developing from its lower part are sulphide wells for chemosymbiotic bacteria. The same interpretation refers to a similar but smaller Zoophycos from the Miocene of Austria, which displays very steep lobes in its lowermost part (Pervesler and Uchman, 2004).

The lower Danian dark-coloured trace fossil assemblage vertically penetrates the upper 40 cm of the light Maastrichtian
4.2. Calcareous nannofossil assemblages in the Maastrichtian and Danian sediment surrounding trace fossils

Samples from the Maastrichtian sediments, Bd-1 (−400 cm), Bd-6 (−200 cm), Bd-11 (−40 cm), Bd-9 (−35 cm), and Bd-13 (at −3–0 cm), show the classical latest Maastrichtian calcareous nannofossil assemblage of the *Micula prinskii* Zone (Gorostidi and Lamolda, 1995; Galbrun and Gardin, 2004; Minoletti et al., 2005). The assemblage is abundant, moderately preserved and highly diversified (Figs. 3–5). The most common
Fig. 5. Distribution of selected calcareous nannofossils in the studied samples. Nannofossil abundance (observation at \( \times 600 \)): A = abundant (>50 specimens per field of view); C = common (20-50 specimens per field of view); R = rare (<3 specimens per field of view). Species relative abundance observation: A = abundant (1 specimen per field of view); C = common (1 specimen per 2-10 fields of view); F = few (1 specimen per 11-20 fields of view); R = rare (1 specimen per 21-50 fields of view); RR = very rare (1 specimen per >50 fields of view). Preservation: P = poor; M = moderate.
taxa are *Watznaueria barnesiae*, *Prediscosphaera cretacea*, *Cribrophaerella ehrenbergii*, *Arkhangeliskiella cymbiformis*, *Micula* group (*M. decussata*, *M. concava*, *M. murus*), *Lithraphidites quadratus*, *Effelliellus turristiffeli*, *E. parallellus*, *Cribrocorona gallica* and *Micnorhabdulus decoratus*. In samples Bd-9 and Bd-11, very rare pieces of the calcareous dinocyst *Operculodinella operculata* (=*Thoracosphaera operculata* of previous authors) were recognized, while in Bd-13 we observed a few forms of *Operculodinella* spp., together with very rare *Cyclagelosphaera reinhardtii*, *Markalius inversus* and *M. apertus*. In sample Bd-14 (Fig. 5), just above the K-Pg boundary a change in the calcareous nannofossil assemblage is observed: *Operculodinella* increased in abundance, and *Biantholithus sparsus* and *Cyclagelosphaera reinhardtii* are present, together with the most common Maastrichtian taxa. A decrease in the abundance is observed together with carbonate fragments. Following the suggestion of Mai et al. (2003) and Bernaola and Monechi (2007), in order to unambiguously identify the basal Danian, the occurrence of the index species *B. sparsus* was used. It should be underlined that in these samples the preservation of the Cretaceous species is moderately good, especially for *A. cymbiformis* and *M. prinitsii*. This is also recognized in other K-Pg boundary sections, as in the Site 1262 (Bernaola and Monechi, 2007).

The following samples above the K-Pg boundary surface show a succession of abundant assemblages dominated by opportunistic forms (Fig. 3). The first sample, corresponding to Bd-15 (+6 cm), is characterized by abundant *Operculodinella* associated with *M. inversus*, *M. apertus*, *C. reinhardtii*, *C. alta* and few Maastrichtian taxa. *Operculodinella operculata* (calcareous dinoflagellate) represents a persistent taxon, surviving the K-Pg boundary, and the increase in abundance of this form reflects ecological stress after the rapid environmental changes at the boundary (Hildebrandt-Habel et al., 1999). The second appearance, in samples Bd-17 (+17 cm), Bd-19 (+29 cm) and Bd-22 (+40 cm), is characterized by an increased abundance of *Braarrudosphera bigelowii* and *B. alta*, displacing *Operculodinella*. The assemblage mainly consists of *Octolithus multilus*, *M. inversus*, *M. apertus*, *C. alta*, *C. reinhardtii* and a few Maastrichtian taxa.

4.3. Calcareous nannofossils from the trace fossil infillings

Dark-filled *Thalassinoideas* from Bd-9 shows an assemblage characterized by common Maastrichtian taxa and the presence of *Operculodinella* spp., *M. inversus*, and *B. sparsus* (?). In sample Bd-9, from the surrounding sediments, only very rare *Operculodinella* are recognized. Dark–filled *Zoophycos* from Bd-11 shows an assemblage of Maastrichtian taxa with very rare *M. inversus*. This assemblage is quite similar to that from the sediments surrounding trace fossils of samples Bd-9 and Bd-11 (Fig. 3). Dark–filled *Thalassinoideas* from Bd-13 shows an assemblage of abundant *Operculodinella* (=*Thoracosphaera*) associated with *M. inversus*, *M. apertus*, *C. reinhardtii* and a few Maastrichtian taxa (Fig. 5). This assemblage is very different from that of the surrounding Cretaceous sediments of sample Bd-13, in which Maastrichtian taxa are predominant (Fig. 5). The dark-filled *Thalassinoideas* shows the same assemblage as sample Bd-15.

Light-filled *Thalassinoideas* from Bd-13 shows an assemblage quite similar to the normal sedimentation of Bd-13, mainly characterized by Maastrichtian taxa with a few *Operculodinella* (=*Thoracosphaera*), *C. reinhardtii* and *M. inversus* (Fig. 3).

5. Discussion and conclusions

Only occasionally have trace fossils or bioturbation structures been mentioned in the K-Pg boundary section at Bidart (Smit and Ten Kate, 1982; Haslett, 1994; Gorostidi and Lamolda, 1995), without reference to a possible dislocation of microfossils by bioturbation (Nelson et al., 1991). Notwithstanding, analysis of calcareous nannofossil assemblages from the infillings of trace fossils and surrounding sediment would appear especially significant (Fig. 3). The calcareous nannofossil assemblages of the dark–filled *Thalassinoideas* from samples Bd-9 (−35 cm) and Bd-13 (−3–0 cm) are very different from those of the surrounding uppermost Maastrichtian sediment, while similar to ones from sample Bd-15 (6 cm above the K-Pg boundary; lowermost Danian). The calcareous nannofossil assemblages of the dark–filled *Zoophycos* and light-filled *Thalassinoideas* from samples Bd-11 (−40 cm) and Bd-13, respectively, show a similar composition to those from the surrounding uppermost Maastrichtian sediment. These data confirm the existence of two phases of macro-benthic colonization:

- the pre-K-Pg boundary event colonization phase, represented by the light-filled *Thalassinoideas*, showing an infilling nannofossil assemblage composition similar to that of the surrounding sediment;
- the post-K-Pg boundary colonization phase, represented by dark–filled *Thalassinoideas* and *Zoophycos*, where *Thalassinoideas* displays a nannofossil assemblage similar to the one from the lowermost Danian (characterized by abundant *Operculodinella* [=*Thoracosphaera*], *C. reinhardtii* and *M. inversus*).

Yet this is not the case for *Zoophycos*. The difference between the nannofossil assemblages from the dark-filled *Thalassinoideas* and dark–filled *Zoophycos* is related to the formation of these trace fossils; while *Thalassinoideas* is commonly interpreted as an open burrow passively filled by the overlying sediment (Bromley, 1996), the origin of *Zoophycos* remains controversial (Löwemark et al., 2004), the infilling being a large admixture of the *in situ* material.

Especially relevant is the good preservation of the nannofossil assemblages described here, with no significant differences found between those from the surrounding sediment and the *Thalassinoideas* infillings, either dark or light. This comes to support the general interpretation of *Thalassinoideas* as passively filled burrows, precluding any secondary modification of the infilling due to activity of the trace maker. Thus,
taphonomic features *per se* are not conclusive evidence of infiltration of the nannofossil specimens.

The story told by the Bidart section, as understood in the context of the present and other recent studies, comprises the following:

- a recognition of the reworking of Maastrichtian calcareous nannofossils within lowermost Danian sediments related to the drastic extinction event at the K-Pg boundary;
- patterns of abundance of the calcareous nannofossils species from passively filled burrows of the topmost Maastrichtian would lower the position of the earliest large occurrence of Danian calcareous nannofossils (such as *Opeculodinella*), leading one to misinterpret the K-Pg biotic turnover.

This evidence shows that the distribution of calcareous nannofossils in the Bidart section can be strongly dependent on the occurrence of particular ichnotaxa. Thus, we have to reconsider our knowledge on the K-Pg turnover event in a new light. The K-Pg reference sections should be re-analyzed for the presence of trace fossils with passive filling (such as *Thalassinoides*) with respect to micropalaeontological assemblages.

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