A multi-proxy record of the Latest Danian Event at Gebel Qreiya, Eastern Desert, Egypt

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ABSTRACT – The Latest Danian Event (LDE) is a proposed early Palaeogene transient warming event similar to the Paleocene–Eocene Thermal Maximum, albeit of smaller magnitude. The LDE can be correlated with a carbon isotope excursion ('CIE-DS1') at Zumaia, Spain, and the 'top Chron C27n event' defined recently from ocean drilling sites in the Atlantic and Pacific, supporting a global extent. Yet, records of environmental change during the LDE (e.g. warming and sea-level fluctuations) are still rare. In this study, we focus on the micropalaeontology (calcareous nannofossils and benthic foraminifera), mineralogy and trace element geochemistry of the LDE in the Qreiya 3 section from the southern Tethyan margin in Egypt. In this section, the LDE is characterized by the occurrence of anomalous beds intercalated within upper Danian shales and marls. The event beds of the LDE are situated above an unconformity on top of a shallowing-upwards sequence deposited in a well-oxygenated outer neritic to upper bathyal marine palaeoenvironment. The lower LDE bed is barren of benthic foraminifera, but contains pyrite and fish remains, and is interpreted as an anoxic level formed during rapid relative sea-level rise. Incursion of a *Neoeponides duwi* (Nakkady, 1950) benthic assemblage in LDE bed II is interpreted as repopulation of the seafloor after anoxia. The sea-level cycle associated with the LDE is estimated at about 50 m maximum in the Qreiya 3 section. The environmental changes at Qreiya 3 are of supraregional extent, since a similar sequence of events has been observed at other southern Tethyan locations. *J. Micropalaeontol.* **30**(2): 167–182, September 2011.

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KEYWORDS: *Tethyan margin, Latest Danian Event, benthic foraminifera, calcareous nannofossils, trace element geochemistry*

INTRODUCTION

Recent studies provide strong evidence that several transient warming events have preceded and followed the Paleocene– Eocene Thermal Maximum (PETM; e.g. Speijer, 2000, 2003a; Thomas & Zachos, 2000; Lourens *et al*., 2005; Petrizzo, 2005; Bernaola *et al*., 2007; Nicolo *et al*., 2007; Quillévéré *et al*., 2008; Agnini *et al*., 2009; Stap *et al*., 2009; Coccioni *et al*., 2010). One of the proposed Paleocene hyperthermals is the Latest Danian Event (LDE), described from sections located at the southern Tethyan margin in Egypt (Sprong, pers.com.; Bornemann *et al*., 2009; Sprong *et al*., 2009; Youssef, 2009; 'el-Qreiya event' in Soliman & Obaidalla, 2010). Although it has been suspected that the LDE might reflect a hyperthermal (e.g. Speijer, 2000, 2003a; Bornemann *et al*., 2009; Youssef, 2009), a reliable high-resolution stable oxygen isotope record, necessary to prove warming, is difficult to obtain in the sediments from Egypt (Bornemann *et al*., 2009). In addition, the LDE has been associated with the Danian–Selandian transition because the stratigraphical position of the southern Tethyan LDE beds was not well enough constrained for correlation with other records (e.g. Speijer, 2003a; Guasti, unpublished thesis, University of Bremen, 2005; Soliman & Obaidalla, 2010). Since the Danian– Selandian boundary has been formally defined in the Zumaia section in Spain (Schmitz *et al*., 2008), and the biostratigraphical resolution of this interval has been refined (Agnini *et al*., 2007; Arenillas *et al*., 2008; Bernaola *et al*., 2009; Sprong *et al*., 2009), the LDE in Egypt and its correlative level in Tunisia (e.g. Steurbaut *et al*., 2000; Guasti *et al*., 2006; Van Itterbeeck *et al*., 2007) is found to be time-equivalent with the 'top Chron C27n event' in open ocean records (Westerhold *et al*., 2008, 2011) and with the 'CIE-DS1' event in Zumaia (Arenillas *et al*., 2008; Dinarès-Turell *et al*., 2010). Thus, the LDE as described from Egypt most probably represents one of the proposed Paleocene hyperthermals preceding the PETM.

The LDE is characterized by a 1–2‰ excursion towards lighter 13C values (Bornemann *et al*., 2009; Westerhold *et al*., 2011). The LDE beds at Gebel Qreiya correlate with other sections in the Nile Basin showing a set of event beds and the incursion of the benthic foraminifer *Neoeponides duwi* (Speijer & Schmitz, 1998; Speijer, 2000, 2003a, b). At the Qreiya 3 section, located near the southern peak of the Gebel Qreiya plateau (Fig. 1), the LDE is expressed as a 25 cm thick couplet consisting of a dark purplish-brown marl bed overlain by a dark grey shale bed, intercalated in the brown-grey upper Danian Dakhla shales. The basal LDE bed I is rich in organic carbon and fish remains, barren of benthic foraminifera and shows unusual, low-diversity planktic foraminiferal assemblages, strongly dominated by non-keeled, muricate and praemuricate taxa. The upper LDE bed II is dominated by *N. duwi* in association with shallow-water benthic foraminifera (Speijer, 2000, 2003a). The

Fig. 1. Location map: (**a**) overview; (**b**) detailed map. Gebel Qreiya is located north of the town of Qena, near the southern entrance of Wadi Qena. For coordinates of the Qreiya 3 section, see text. Also indicated are the Duwi and Aweina sections. Dotted line: approximate outline of the Nile Basin. Double arrows: Syrian Arc (unstable shelf).

LDE beds at Gebel Qreiya resemble the organic-rich dark brown layers at the PETM in the Esna Shale Formation (Speijer, 2000, 2003a; Knox *et al*., 2003), but no extinction of benthic foraminifera is observed during the LDE (e.g. Speijer, 2003a).

In this paper we present a high-resolution multi-proxy study of the LDE at the Qreiya 3 section. Benthic foraminifera, mineralogy and trace element geochemistry are used to reconstruct changes in the palaeoenvironment, and to constrain relative sea-level fluctuations across the LDE.

GEOLOGICAL BACKGROUND

Location and geological setting

Gebel Qreiya is situated east of the southern entrance of Wadi Qena in the Eastern Desert, Egypt (Fig. 1). Wadi Qena is a \sim 300 km long valley, extending from the slopes of the Galala mountains in the north to the town of Qena in the south, where it opens into the Nile Valley. Paleocene to lower Eocene marl–shale successions are well exposed at Gebel Qreiya. Section Qreiya 3 (26°N 27'42" and 33°E 1'54") is situated near the southern tip of Gebel Qreiya, about 50 km northeast of the town of Qena, north of the road from Qena to Safaga.

During Danian–Selandian times (~65–59 Ma; Luterbacher *et al*., 2004), the southern margin of the Nile Basin was situated close to the present southern border of Egypt (Luger, 1988). The basin was characterized by neritic to upper bathyal marl and shale deposition of the Dakhla Formation (Fm.) in a basin generally deepening to the northwest (Speijer, 2003a and references therein). Micropalaeontological data indicate Danian and Selandian palaeodepths ranging from 70–150 m at Gebel Duwi, near the Red Sea coast, to 400–600 m in Sinai. Deposition at

Gebel Qreiya took place at outer neritic to upper bathyal palaeodepths (150–250 m; Speijer, 2003a).

Lithology

The Paleocene succession of Gebel Qreiya consists of the upper part of the Dakhla, the Tarawan and the lower part of the Esna formations (Said, 1962; Hendriks *et al*., 1987). We studied 16.5 m of the upper part of the Dakhla Fm. (Fig. 2a, b). The Qreiya 3 section is a composite of two subsections (Fig. 2a). Upper subsection B, starting at the base of LDE bed I, is located 10 m to the northeast of lower subsection A. The interval between 8.05 m and 8.20 m consists of dark grey carbonate-poor shale of the Dakhla Fm., containing mm- to cm-sized burrows at the top. At 8.20 m, the LDE beds are intercalated within the Dakhla shales. The lithological transition is abrupt, is not bioturbated and contains (secondary) gypsum.

The LDE beds I and II are characterized as follows:

- \bullet Bed I (8.20–8.30 m) consists of dark purplish-brown laminated marl, rich in organic matter. It contains abundant fish remains (including fish imprints with partial scale preservation: Fig. 3), pyritized moulds of planktic foraminifera, and some coprolites. *In situ* benthic foraminifera are not present. The upper 7 cm of bed I contain dark grey clay lenses: these are burrows penetrating downwards from bed II containing benthic foraminifera. The sample at 8.29 m is very rich in planktic foraminifera.
- \bullet Bed II (8.30–8.45 m) consists of grey to dark grey shale, rich in organic matter. This bed contains planktic and benthic foraminifera, and pyritized moulds of bivalves and gastropods. Between 8.42 m and 8.45 m the sediment consists of

Fig. 2. Lithology and biostratigraphy of the Qreiya 3 section. (**A**) Photograph of the section showing the Dakhla, Tarawan, Esna and Thebes formations. The section was sampled in the Dakhla shales in two subsections A and B. The calcarenitic beds C1–C3, the Latest Danian Event beds (LDE) and the PETM are indicated. (**B**) Schematic lithology and biostratigraphical datums. Calcareous nannoplankton zonation follows Martini (1971) and Varol (1989); planktic foraminiferal biozonation follows Berggren & Pearson (2005). LDE, position of LDE beds I and II (grey shaded interval). Asterisk, LO of *M. velascoensis* was used to approximate the base of Zone P4, which is for this reason denoted as 'P4'; see text for explanation.

Fig. 3. Fish imprints and partially preserved fish scales, LDE bed I. Scale is a \in 2 coin.

purplish-brown marl and grey marly shale with laminae, flaser bedding and abundant pyritized moulds of planktic foraminifera.

Ten metres south of the main Qreiya subsection A, a several metres wide and up to 15 cm thick, lens-shaped calcarenitic bed cuts through the LDE beds (Fig. 2). This yellowish-grey upwards-fining bed (C1) rests on top of a 1 cm thick sand layer; it contains clay pebbles at the base, fish remains, and the upper surface is heavily bioturbated. A second lens-shaped calcarenitic bed (C2), about 30 cm thick and 10 m wide, is intercalated above the LDE beds at 11.4 m. Further up-section, at 13.8 m, a third calcarenitic bed (C3), about 50 m wide and up to 20 cm thick, is intercalated in the Dakhla shales. Both the bottom and top of this calcarenitic bed are heavily bioturbated. The calcarenite beds C1–C3 are interpreted as channel fills, incising the normal sedimentary succession.

METHODS

Samples were collected from 10–30 cm deep trenches or holes along the outcrop to obtain fresh sediment. Sample spacing was generally 25 cm; the interval comprising the LDE was sampled every 2–5 cm.

Foraminifera

For analysis of benthic foraminifera, 78 samples were selected: 59 from the main section and 19 from calcarenites C1–C3 (see Supplementary Material). The samples were dried at 60° C for 24 hours, weighed, disintegrated in a soda solution $(50g)$ ⁻¹ $Na₂CO₃$ and washed over a set of 2 mm and 63 µm sieves. The residues were dry-sieved over 630 µm and 125 µm sieves. Benthic foraminifer counts were obtained from the fraction 125–630 µm (see Supplementary Material). Representative aliquots containing 200–300 specimens were obtained with an ASC microsplitter. Foraminifera were picked from the washed residues, transferred to micropalaeontological slides and stored at the Department of Earth and Environmental Sciences, K.U. Leuven (Belgium) for future reference. Relative abundances were calculated, and absolute numbers of planktic and benthic foraminifera (PFN, BFN: planktic and benthic foraminiferal numbers per gram of dry sediment; Fig. 4 and Supplementary Material). Benthic diversity (Shannon index S[H]: Shannon & Weaver, 1949) and dominance (Simpson index: Hammer *et al*., 2001) were determined. The P/B ratio, derived from counting random squares in the picking tray, is expressed as $100 \times P/(P+B)$ and was used as a first estimate of relative palaeodepth (e.g. Grimsdale & Van Morkhoven, 1955; Berger & Diester-Haass, 1988; Van der Zwaan *et al*., 1990). In addition, water-depth preferences were assigned to species on the basis of literature data from Egypt and Tunisia (Fig. 5). The terms 'neritic' and 'bathyal' are commonly used to discriminate between palaeodepths less than, and in excess of 200 m (e.g. Van Morkhoven *et al*., 1986). Here, we use the terms inner neritic (IN), middle neritic (MN), outer neritic (ON) and bathyal (B) for palaeodepths of ≤ 50 m, $50-100$ m, $100-200$ m and ≥ 200 m, respectively.

For correspondence analysis (CA) the samples from the calcarenitic levels C1–C3, interpreted as channel fills, were excluded. PAST computer software (Hammer *et al*., 2001) was used to run CA on the relative abundance data. The barren sample at 8.2 m was omitted from the analyses; however, the sample at 8.25 m containing downwards bioturbated benthic foraminifera was included. Only taxa occurring in relative frequencies >2% in any sample were retained in the analysis. Correspondence analysis was run on 58 samples and 53 taxa.

Fig. 4. Foraminifera data. Calcarenitic levels C1–C3 not included. Left panel: P/B ratio expressed as %P; planktic foraminifera per gram of dry sediment (PFN); benthic foraminifera per gram of dry sediment (BFN), Shannon index H[S] for benthic foraminifer diversity, Simpson index for dominance in benthic foraminifer associations. Right panel: relative frequencies of selected benthic foraminifer taxa. Hatched bars indicate Latest Danian Event beds (LDE, bed I = 8.2–8.3 m and bed II = 8.3–8.45 m); grey bars indicate dissolution levels below LDE (7.85–8 and 8.05–8.2 m).

Calcareous nannofossils

Calcareous nannofossil biostratigraphy was determined from standard smear-slides following Martini (1971) and Varol (1989). Calcarenitic levels C1–C3 were excluded. Nannofossils were qualitatively analysed in 76 samples, using standard procedures as described in Perch-Nielsen (1985). Smear slides were examined with a light microscope at $1000 \times$ magnification. At least three traverses of each slide (~600 fields of view) were searched for zonal markers. Random settling slides, as described in Geisen *et al*. (1999), were used for a quantitative nannofossil investigation on the 76 samples (see Supplementary Material). In each slide, 300 individuals (or max. 200 fields of view) were identified and counted. Relative and absolute abundances (g^{-1}) were calculated.

Trace elements and mineralogy

The concentrations of selected trace elements (Ni, Cu, Zn, As, Rb, Sr, Zr, Pb, Th, U) were determined in 48 powdered samples at the University of Heidelberg (Germany) with the Energydispersive Miniprobe Multi-element Analyzer ('EMMA'; this Xray fluorescence analyser was developed by Andriy Cheburkin (see Cheburkin & Shotyk, 1996, for details)); it is not subject to matrix interferences, and the trace element analyses show very low detection limits (see Supplementary Material). The EMMA-XRF analyses were calibrated with several NIST reference materials and the relative analytical error is <10%.

The bulk rock mineralogy was determined at the University of Erlangen (Germany) on powdered samples (grain size $\leq 10 \mu m$) with a Siemens-Bruker D5000 X-ray diffractometer. This instrument is fitted with a copper tube (CuK α = 1.54178 Å), operating at 40 kV and 35 mA, and a post-diffraction graphite monochromator. Samples were side-loaded into a holder for random orientation and scanned from 5° to 65° 2 Θ in steps of 0.02° and 4 seconds scanning time. The X-ray diffractograms were evaluated by quantitative Rietveld analysis with BRUKER AXS TOPAS 4.0 software. The TOPAS Rietveld software provided very good fits of the diffractograms and the observed weighted residual errors R_{wp} are generally lower than 8%. The relative analytical error as determined by a comparison with several artificial mixtures and using ZnS as an internal standard is <3%.

BIOSTRATIGRAPHY

Planktic foraminifera

Planktic foraminifer biostratigraphy was studied by Sprong *et al*. (2009) by applying the biozonation criteria of Berggren & Pearson (2005). The studied interval covers planktic foraminiferal Zones P2 and P3 and the lower part of P4 (Fig. 2b). The base of the section is in Zone P2, identified by the occurrence of *Praemurica uncinata* (Bolli, 1957). Subzones P3a and P3b were identified by the lowest occurrences (LOs) of their respective markers, i.e. *Morozovella angulata* (White, 1928) (P3a; 3.5 m)

taxa	ΙN	MN ON	в		axis 1	axis 2
Lenticulina spp. costate				1,4	2.461	0.098
non-calcareous agglutinated taxa				1, 2, 4	2.317	-0.831
Ammodiscus cretaceus		---		1,2	1.429	-0.752
Neoeponides duwi				1,2	1.163	2.806
agglutinated sp. B					0.991	-0.694
Bulimina ovata-quadrata	----			2,4	0.830	-0.389
Siphogenerinoides esnehensis				1	0.434	1.113
Nodosaria spp. costate				1	0.406	0.303
Stilostomella plummerae				1.2	0.136	0.894
Marginulinopsis spp.				1,2	0.055	0.538
Lenticulina spp. smooth				1, 2, 3	0.000	0.208
Bulimina strobila/farafraensis				1,4	-0.035	0.725
Anomalinoides midwayensis				2,4	-0.181	-0.088
Alabamina midwayensis				$1 - 4$	-0.191	0.201
Bulimina midwayensis				$1 - 4$	-0.215	0.056
Tappanina selmensis				$\overline{2}$	-0.222	0.310
Hansenisca girardanus			۰.	1, 2, 4	-0.233	0.120
Oridorsalis plummerae				$1 - 4$	-0.245	0.028
Sporobulimina eocaena				1, 3, 4	-0.247	0.095
nodosariids				1,2	-0.252	-0.065
monothalamous taxa					-0.278	-0.075
Valvalabamina depressa				$1 - 4$	-0.294	-0.079
Anomalinoides praeacutus				1, 2, 4	-0.321	0.021
Cibicidoides cf. C. hyphalus				1, 3, 4	-0.333	-0.076
Angulogavelinella avnimelechi				1, 3, 4	-0.336	-0.156
Spiroplectinella dentata				1,2	-0.340	-0.131
Cibicidoides pseudoacutus				$1 - 4$	-0.347	-0.182
Valvalabamina planulata				1, 3, 4	-0.359	0.007
Alabamina cf. A. creta				1	-0.360	0.044
Gyroidinoides globosus				1	-0.374	-0.111
Cibicidoides rigidus				3,4	-0.388	-0.177
Gyroidinoides sp. A					-0.389	-0.079
Pullenia jarvisi group				1	-0.393	-0.195
Cibicidoides alleni				$\mathbf{1}$	-0.395	-0.294
Loxostomoides applinae			--	1.2.4	-0.432	-0.132
Gyroidinoides cf. G. tellburmaensis				$\overline{\mathbf{c}}$	-0.433	-0.109
Globocassidulina subglobosa				1, 3, 4	-0.450	-0.162
Tritaxia midwayensis				1,2	-0.460	-0.186
Quadrimorphina spp.				$\overline{2}$	-0.462	-0.109
Anomalinoides rubiginosus			----	1,2	-0.468	-0.190
Pulsiphonina prima				1,3	-0.469	-0.191
Eponides lunatus					-0.476	-0.058
Gaudryina pyramidata					-0.477	-0.287
Siphogenerinoides elegantus				1, 2, 4	-0.482	-0.317
Valvulineria? insueta					-0.484	-0.318
Osangularia plummerae				$1 - 4$	-0.485	-0.294
				1,2	-0.499	-0.167
Spiroplectinella knebeli Anomalinoides affinis					-0.504	-0.214
Gavelinella beccariiformis				1, 2, 4 1	-0.526	-0.326
Cibicidoides succedens						
				1, 2, 4	-0.528	-0.295
Coryphostoma cf. C. midwayensis Anomalinoides susanaensis			----	1 1,2	-0.532 -0.534	-0.296 -0.319
Nuttallides truempyi				1	-0.537	-0.268

Fig. 5. Correspondence analysis (CA). Columns 2–5, bathymetrical ranges in North Africa of most taxa used in CA, with literature references in column 6 (1, Speijer & Schmitz, 1998; 2, Schnack, 2000; 3, Ernst *et al*., 2006; 4, Stassen *et al*., 2009). IN, inner neritic – palaeodepths of <50 m; MN, middle neritic – palaeodepths of 50–100 m; ON, outer neritic – palaeodepths of 100–200 m; B, bathyal – palaeodepths of >200 m. The two final columns indicate the taxon loadings on CA axes, arranged according to the values on CA axis 1.

and *Igorina albeari* (Cushman & Bermudez, 1949) (P3b; 8.05 m, 15 cm below LDE bed I). *I. albeari* is the earliest igorinid developing a slight keel on the last chambers (cf. Sprong *et al*., 2009, Fig. 4, 10–12; Pl. 1, figs 4–6); in agreement with the concept of Quillévéré *et al*. (2002). The typical, large *Globanomalina pseudomenardii* Bolli, 1957, a marker of Zone P4 (Olsson *et al*., 1999), was not encountered, although small

Globanomalina specimens (<125 µm), referred to as *G*. cf. *pseudomenardii* (Pl. 1, figs. 1–3) occur from 11.9 m upward. Following Speijer (2003a), the LO of *Morozovella velascoensis* (Cushman, 1925) was used to approximate the base of Zone P4 (13.9 m; indicated as 'P4' in Fig. 2b) because *M. velascoensis* is traditionally thought to appear approximately simultaneously with *G. pseudomenardii* (e.g. Blow, 1979; Toumarkine & Luterbacher, 1985). However, *M. velascoensis* may appear somewhat below the base of Zone P4 (Premoli Silva *et al*., 2003; Steurbaut & Sztrákos, 2008) or even closer to the base of P3b (Olsson *et al*., 1999).

Calcareous nannofossils

According to the biozonation criteria of Martini (1971), the studied part of section Qreiya 3 spans Zone NP4 and the lower part of Zone NP5 (Fig. 2b). Following the zonation scheme of Varol (1989), the studied sequence covers Zones NTp6 to NTp9, some of which are subdivided into subzones. These zones and subzones have also been identified in correlative Danian sections in Egypt (Youssef, 2009) and in Tunisia (Steurbaut *et al*., 2000; Guasti *et al*., 2006; Van Itterbeeck *et al*., 2007; Sprong *et al*., 2009). The (sub-) zonal boundaries are marked by the highest occurrence (HO) of *Neochiastozygus eosaepes* Perch-Nielsen, 1981 (NTp6/NTp7), and the respective LOs of *Chiasmolithus edentulus* van Heck & Prins, 1987 (NTp7A/NTp7B), *Sphenolithus primus* Perch-Nielsen, 1971 (NTp7/NTp8), *Fasciculithus billii* Perch-Nielsen, 1971 or *F. ulii* Perch-Nielsen, 1971 (NTp8A/ NTp8B), and *F. janii* Perch-Nielsen, 1971 or *F. pileatus* Bukry, 1973 (NTp8B/NTp8C).

The NTp6/NTp7 Zonal boundary is situated 1 m above the base of the section and slightly postdates the HO of *Fasciculithus magnus* Bukry & Percival, 1971 (at ~0.7 m). Subzone NTp7A contains rich and highly diverse nannofossil assemblages with *Fasciculithus*. The base of Subzone NTp7B is positioned at 7 m. It is marked by a series of more-or-less coinciding nannofossil events, including the LOs of *C. edentulus*, small *Fasciculithus* taxa (*F. chowii* Varol, 1989 and others = first *Fasciculithus* radiation), medium-sized *Toweius* Hay & Mohler, 1967, and the start of the consistent occurrence of *Pontosphaera* Lohmann, 1902. The first specimens of *Sphenolithus primus*, defining the NTp7/NTp8 boundary, are recorded at 10.65 m. *Fasciculithus ulii* and *F. janii*, maring the boundaries of Subzones NTp8A/ NTp8B and NTp8B/NTp8C, first appear at 13.0 m and 13.45 m, respectively. The LOs of *F. ulii* and *F. janii* mark the second radiation in *Fasciculithus* and *F. tympaniformis* Hay & Mohler, 1967, the marker of the base of Zone NP5, appears in low numbers from 13.9 m onward, in association with relatively frequent *F. pileatus* and *F. involutus* Bramlette & Sullivan, 1961.

Stratigraphical position of the LDE beds

The contact between the carbonate-poor level underlying the LDE and lower LDE bed I is sharp in the Qreiya 3 section, and may represent an unconformity. Therefore, and contrary to Soliman & Obaidalla (2010) we do not include this carbonatepoor interval in the LDE.

The base of LDE bed I is 8.20 m above the base of the section (Fig. 2) in lower planktic foraminifer Zone P3b (LO of *I. albeari* at 8.05 m: Sprong *et al*., 2009), in calcareous nannofossil Zone NP4 (Martini, 1971) and in Subzone NTp7B (Varol, 1989; the LOs of *C. edentulus* and small *Fasciculithus* spp. coinciding in the section at 7 m). Planktic foraminifer biozonation in this interval is somewhat problematic due to differing taxonomical concepts (e.g. Arenillas *et al*., 2008; Sprong *et al*., 2009). The most reliable calcareous nannoplankton datums, according to Dinarès-Turell *et al*. (2010: 1st and 2nd radiation of *Fasciculithus*, LO of *C. edentulus* and LO of *F. tympaniformis*), are recognized in section Qreiya 3 and, on the basis of these datums, the section can be correlated to the Zumaia section in Spain (Bernaola *et al*., 2009). In 2008, the position of the Global Boundary Stratotype Section and Point (GSSP) of the basal Selandian at the base of the Itzurun Fm. in Zumaia was ratified (Schmitz *et al*., 2008; Bernaola *et al*., 2009; see https:// engineering.purdue.edu/Stratigraphy/gssp/); thereby replacing the conventional positioning of the Danian–Selandian (D–S) boundary at the P2/P3 boundary (e.g. Berggren *et al*., 1995; Steurbaut *et al*., 2000). Biostratigraphically, the newly defined Danian–Selandian boundary is situated between the second radiation of *Fasciculithus*, and the LO of *F. tympaniformis* marking the NP4/NP5 boundary (Schmitz *et al*., 2008; Bernaola *et al*., 2009). Consequently, the LDE beds are of late Danian age; they correlate with the lithological transition from the Crowded Member to the Stratified Member of the Aitzgorri Fm. at Zumaia, about 10 m below the D–S boundary (Steurbaut & Sztrákos, 2008; Bernaola *et al*., 2009; Sprong *et al*., 2009). This stratigraphical position is in agreement with chemostratigraphical correlations on the basis of carbon isotope records (Bornemann *et al*., 2009). Westerhold *et al*. (2011; using tuning option 1 in Westerhold *et al*., 2008) correlate the LDE with both the 'CIE-DS1' of Arenillas *et al*. (2008) and the 'Top Chron C27n event' of Westerhold *et al*. (2008), placing it at the very top of Chron C27n and the basis of Chron C26r (eccentricity cycle 37 in Zumaia: Dinarès-Turell *et al*., 2010) and estimate the duration of the LDE at \sim 200 ka, which is in agreement with the 191 ka estimated by Bornemann *et al*. (2009). The LDE postdates the Dan-C2 event (Quillévéré *et al*., 2008; Coccioni *et al*., 2010) and precedes the Mid Paleocene Biotic Event (MPBE) or Late Palaeocene Biotic Event (LPBE) (Westerhold *et al*., 2008, 2011; Marcantonio *et al*., 2010).

RESULTS

Foraminifera

Benthic foraminifera are moderately to well preserved below and above the LDE and the assemblages are rather diverse (Fig. 4). Some 25% of the species recorded have been described from bathyal open-marine environments (Fig. 5). The relative abundances of the bathyal species decrease towards the LDE, and *Gavelinella beccariiformis* (White, 1928) disappears above 7.5 m. Absolute abundances of planktic and benthic foraminifera (PFN, BFN) drop to extremely low values towards the LDE, the diversity decreases and the P/B ratio drops to 13% (Fig. 4). PFN, fluctuating between 200 and 4000 below and above the LDE drop immediately below, and in the lower part of LDE bed I, where PFN are ≤ 40 specimens g^{-1} ; whereas pyritized moulds of planktic foraminiifera are abundant. PFN increase in LDE bed II, and show peak abundance >10 000 at 8.29 m (top of LDE bed I). A maximum of ~9000 planktic specimens g^{-1} between

11 m and 14 m is followed by a decrease below 500 in the top 2.5 m of section. BFN generally fluctuate between 100 and 400 below and above the LDE, but only ~10 benthic foraminifera g^{-1} occur in the 15 cm just below the LDE beds, and LDE bed I is barren of *in situ* benthic foraminifera. In LDE bed II BFN increase to a temporary maximum of \sim 535 specimens g^{-1} at 8.4 m.

The relative abundances of non-calcareous agglutinated benthic foraminifera increase from <5% to 20–80% in the carbonate-poor interval below the LDE beds (7.8–8.2 m). Relatively common bathyal species, such as *Gavelinella beccariiformis, Osangularia plummerae* Brotzen, 1940 and *Anomalinoides affinis* (Hantken, 1875), disappear below the LDE, whereas the costate *Siphogenerinoides esnehensis* (Nakkady, 1950) has its lowest occurrence there. In LDE bed II *Neoeponides duwi* temporarily dominates the benthic assemblage (>50%) together with *Siphogenerinoides elegantus* (Plummer, 1927), *S*. *esnehensis* and stout lenticulinids. Above the LDE, at 9.5 m *G. beccariiformis* returns, but its abundance remains <10%. In contrast, *Nuttallides truempyi* (Nuttall, 1930) is more common above than below the LDE. No extinctions of benthic foraminifera are recorded (see Supplementary Material).

Correspondence analysis (CA) separates the LDE and the carbonate-poor interval below the LDE from the background samples (Fig. 6). The first two axes of the CA together explain \sim 49% of the variance in the data set: 29.6% and 19.1%, respectively. Running CA without the sample at 8.25 m (LDE bed I; with downwards bioturbated foraminifera) gives the same result. Relative abundances summed according to water-depth preferences (Fig. 5) suggest shallowing associated with the LDE interval (Fig. 7a, b). The high abundances of non-calcareous agglutinated taxa below and in the LDE are not observed in calcarenite C1 cutting through the LDE (Fig. 7c).

Calcareous nannofossils

The nannofossil assemblages are moderately to very well preserved up to 70 cm below the LDE (0.0–7.5 m) and moderately diverse (16 to 22 taxa). Small *Prinsius* spp. dominate these samples (40–75%) (Fig. 8; see Supplementary Material). Other commonly occurring taxa are *Coccolithus* spp. (17–45%), *Toweius* sp. (1–9%), *Ellipsolithus* spp. (1–8%), *Neochiastozygus* spp. (1–7%) and *Cruciplacolithus* spp. (1–4%). The preservation of the nannofossils rapidly deteriorates towards the LDE, and in this interval the assemblages are dominated by *Coccolithus pelagicus* (Wallich, 1877) (30–60%), occurring together with *C*. *subpertusus* (Hay & Mohler, 1967) (4–13%), small *Prinsius* spp. (4–25%), *Toweius* spp. (3–19%) and *Pontosphaera* sp. (4–10.5%). Rare, small *Fasciculithus* spp. appear in this interval. Absolute abundances drop from $>1 \times 10^9$ to $<1 \times 10^7$ specimens g⁻¹ between 8.05 m and 8.2 m, and remain low in LDE bed I and the lower part of LDE bed II.

In the LDE beds, the nannofossils are moderately preserved and the number of taxa ranges from 5 to 17. Overall the robust *C. pelagicus*–*C. subpertusus* group dominates the assemblage (40–85%). From 8.5 m upwards, the absence of signs of overgrowth and etching indicates improving nannofossil preservation. The assemblages are moderately preserved and diversified (10–27 taxa), and are again dominated by small *Prinsius* spp.

Fig. 6. Correspondence analysis (CA). Barren sample at 8.2 m in main section (LDE bed I) excluded. Sample at 8.25 m (LDE bed I, containing downwards bioturbated foraminifera) indicated by star. (**a**) Bi-plot of correspondence analysis (CA) axes 1 (explaining 29.6% of the variance) and 2 (19.1% of the variance). The dissolution interval, characterized by >20% non-calcareous agglutinated taxa (NC), the LDE and the 'background' outer neritic (ON) groups are separated by CA. (**b**) Trends in the first and second axes of CA, together explaining ~49% of the variance in the dataset.

Fig. 7. Benthic foraminifera abundances across the LDE. Relative abundances are summed per bathymetrical group (see Fig. 5). (**A**) Relative abundances of bathymetrical groups throughout the Qreiya 3 section. (**B**) As (A) but LDE interval enlarged. (**C**) Relative abundances of bathymetrical groups across calcarenite C1, for comparison with the LDE interval in the main section. Non-calcareous agglutinated taxa do not exceed 2.5% relative abundance in C1.

(40–80%). Other frequently occurring taxa are *C*. *pelagicus* (8–30%), *C*. *subpertusus* (1–9%) and *Toweius* sp. (9%).

Mineralogy and trace element geochemistry

The mineralogical composition of the shales is dominated by phyllosilicates (50–70 wt%; Fig. 9; see Supplementary Material). Calcite is the second dominant component $(15-50 \text{ wt}),$ whereas the quartz and feldspar content is low $\langle 4 \rangle$ and $\langle 6 \rangle$ wt $\%$, respectively). Substantial changes in the mineralogy occur ~35 cm below the LDE, where the drop in the calcite content to less than 3 wt% marks the base of the dissolution level. Within LDE bed I the calcite content rises abruptly to >50 wt%. In

Fig. 8. Relative abundances of calcareous nannofossil taxa, showing the distribution throughout the Qreiya 3 section, with an enlargement of the LDE interval on the right.

addition, the LDE beds show slightly higher amounts of quartz (up to 4 wt%), iron oxides, pyrite and anhydrite compared to the Dakhla shales. Above the LDE the calcite content rises to \sim 75 wt% in Zone 'P4'.

The Th/U ratio, a reliable proxy for palaeoredox conditions (e.g. Dypvik & Harris, 2001), is well above 1 below the LDE (Fig. 9; see Supplementary Material). It drops sharply in the 15 cm below the LDE and remains <0.1 in the LDE beds, rising again to values ~1 between 8.5 m and 10 m. In the upper part of the section Th/U remains <1. In concert with the Th/U ratio, the redox-sensitive trace elements U, As, Ni, Cu and Zn show a sharp ~10-fold increase in the LDE beds.

The Zr/Rb ratio, an indicator for relative grain-size changes (e.g. Dypvik & Harris, 2001) fluctuates between 2 and 2.5 in the lower 6 m of the section, and increases towards the LDE. A maximum ratio of 3.1 is found in bed I before the ratio drops to low values of ~2 in LDE bed II. Above the LDE beds the Zr/Rb ratio remains ~2.5 and, from 13 m upwards, oscillates around 2.

Calcarenite beds C1–C3

The lens-shaped calcarenitic beds C1–C3 are interpreted as channel fills, containing reworked material. In general these beds contain higher benthic and planktic foraminiferal numbers (see Supplementary Material). Whereas BFN drop towards the LDE in the main section, they rise in calcarenite C1 to a maximum of nearly 4000 specimens g^{-1} at 8.25 m. Calcarenites C2 and C3 both contain levels with relatively high BFN compared to the main section (C2: 1515 (11.4 m) vs. 246 (11.3 m); C3: 2026 vs. 1281, both at 13.7 m). The relative abundance of non-calcareous agglutinated taxa is $\leq 2.5\%$ in calcarenite C1, but an increased abundance of *N. duwi* and associated taxa is observed in calcarenite C1, indicating resedimentation of material from LDE bed II into the channel fill.

DISCUSSION

Onset of the LDE

Prior to the LDE our data point to an oligo-mesotrophic, well-oxygenated outer neritic to upper bathyal marine environment, sustaining rather diverse nannofloral and benthic foraminiferal assemblages. About 2 m below the LDE beds, coarsening of the sediments is indicated by the increasing Zr/Rb ratio, and the calcium carbonate content decreases. From about 70 cm below the LDE beds, the shift to lighter δ^{13} C coincides with changes in the abundance patterns of the nannoflora and benthic foraminifera (Figs 4, 7, 8). The interval below the LDE consists of two levels (7.85–8 m and 8.05–8.2 m) characterized by <3 wt% calcium carbonate, lowered P/B ratios, as well as

Fig. 9. Sediment geochemistry, showing (from left to right) the stable carbon isotopes, weight % CaCO₃, weight % clay minerals, major element ratios and redox-sensitive trace elements.

low diversity and absolute abundances of foraminifera and calcareous nannofossils (Figs 4, 8, 9). The nannofossil assemblages are dominated by robust *Coccolithus* species (Fig. 8). The benthic foraminiferal assemblages contain 20–80% noncalcareous agglutinated species (mainly *Haplophragmoides* spp. and trochamminids). Severe carbonate dissolution provides a parsimonious explanation for these observations.

In a thin level (8–8.05 m) dissolution of benthic foraminifera is less severe, indicated by higher benthic diversity, and lower dominance of the non-calcareous agglutinated taxa (Figs 4, 7). In this thin interval the relative abundance of outer neritic to bathyal species decreases to <5%, and *Gavelinella beccariiformis* and *Nuttallides truempyi* are not recorded, but there are no indications in the benthic fauna or the trace elements for dysoxia. The δ^{13} C shifts ~1‰ towards lighter values, but the changes in trace element concentrations below the LDE are relatively minor and probably largely associated with carbonate dissolution, and/or with leaching due to downward migration of a redox front; a diagenetic effect comparable to the 'burningdown' of sapropels (e.g. Higgs *et al*., 1994; Thomson *et al*., 1995). The Th/U-ratio, reflecting redox conditions at the seafloor (Dypvik & Harris, 2001), oscillates around 2 below 8.11 m, suggesting well-oxygenated bottom waters, and then rapidly declines towards the LDE.

The LDE

Considering the low PFN, BFN and nannofossil abundances in the LDE beds, dissolution probably affected sediments deposited during the LDE, but the primary palaeoenvironmental signal of the assemblages recorded during the event has been preserved.

Stable carbon isotopes. The negative carbon isotope excursion associated with the LDE in Egyptian sections (Bornemann *et al*., 2009; Fig. 9) can be correlated with Zumaia (Spain: Schmitz *et al*., 1996, 1997) and open-ocean stable carbon and oxygen isotope records (Arenillas *et al*., 2008; Bernaola *et al*., 2009; Bornemann *et al*., 2009; Westerhold *et al*., 2011). However, the magnitude of the excursion is larger in the Qreiya 3 section (>1‰ versus 0.5–1‰ at Zumaia and in the open-ocean records), and the double carbon isotope peak found at Shatsky Rise in the west-central Pacific (Westerhold *et al*., 2011) has so far not been resolved at the Qreiya 3 section. This may be due to the severe carbonate dissolution below the LDE and the absence of the species measured. From the oxygen isotope record at Shatsky Rise, Westerhold *et al*. (2011) estimate a bottom-water temperature rise of $\sim 2^{\circ}$ C. The authors mention the possibility of a causal relationship with the first phase of the North Atlantic Igneous Province.

Benthic anoxia: foraminifera and trace elements. Sedimentary lamination and the absence of benthic foraminifera in LDE bed I, in concert with organic enrichment, preservation of fish remains and presence of pyritized planktic foraminifera, point to temporary anoxia at the seafloor; an interpretation corroborated by the geochemical data. Very low Th/U-ratios in the LDE beds are due to significant uranium enrichment (Fig. 9). High uranium values are typical for anoxia (e.g. Dypvik & Harris, 2001; Tribovillard *et al*., 2006) and high concentrations of other redoxsensitive trace elements, including As, Ni, Cu and Zn further support this interpretation (Fig. 9; Calvert & Pedersen, 1993; Morford & Emerson, 1999; Nijenhuis *et al*., 1999; Tribovillard *et al*., 2006; see also Soliman & Obaidalla, 2010). In bed II the reappearance of benthic foraminifera and molluscs, trace element enrichment and the absence of lamination suggest at least intermittent availability of oxygen, enabling benthic repopulation. A low-diversity assemblage composed of rather robust, probably dissolution- and low oxygen-resistant foraminifera, including non-calcareous agglutinated taxa and *Neoeponides duwi* (Speijer, 2003b) dominates most of LDE bed II, but gives way to a more diverse assemblage upwards in bed II. The top of bed II (8.45 m) shows an increase in the redox-sensitive trace elements to levels approaching those in bed I, suggesting that a second brief anoxic interval occurred towards the end of the LDE. However, the aspect of the top centimetres of bed II and the correlation to channel C1 (Fig. 2) indicative of enhanced bottom currents, suggest some reworking and/or redeposition of sediments from the underlying LDE bed I. Comparing Bands 2 and 4 of the el-Qreiya bed in Soliman & Obaidalla (2010, fig. 5) raises the suspicion that Band 4 of these authors may also contain material resedimented from Band 2.

Nannoflora. *Coccolithus pelagicus* is relatively resistant to dissolution (Raffi & Rio, 1981), which explains its high relative abundance in the carbonate-poor interval preceding the LDE, and might in part explain its high relative abundance in the LDE beds. Eurytopic behaviour was suggested for *C. pelagicus* (Aubry, 1998). In addition, *C. pelagicus* has probably shifted its distribution since the Palaeogene (Haq & Lohmann, 1976), rendering interpretations difficult. Today *C. pelagicus* is abundant in the arctic/subarctic and in high-productivity regions off river mouths (Cachao & Moita, 2000) and in upwelling regions (Okada & McIntyre, 1979). In the Pliocene–Quaternary *C. pelagicus* was found in sapropels (e.g. Müller, 1985; Negri *et al*., 2003), commonly associated with water-mass stratification (e.g. Nijenhuis *et al*., 1999). In Paleocene–Eocene times, *C. pelagicus* was common at low latitudinal equatorial sites (Okada & Honjo, 1973). The abundance pattern of *C. pelagicus* across the LDE resembles the pattern across the PETM in Egypt (Monechi *et al*., 2000), which was interpreted as an increase in surface productivity (Speijer & Wagner, 2002). In addition, the abundance patterns of *C. pelagicus* and C. *subpertusus* during the LDE, of which the latter is generally associated with warm waters (Aubry, 1998), resemble those at the PETM at Demerara Rise in the western Atlantic (Jiang & Wise, 2006; Mutterlose *et al*., 2007). This suggests that increased productivity was associated with higher surface-water temperatures during the LDE.

In the upper part of LDE bed II the diversity and absolute abundance of nannoflora increase and the small *Prinsius* species dominate the nannofloral assemblages again, indicating both decreasing dissolution, and a return to more normal openmarine conditions.

Recovery after the LDE

The *N. duwi* assemblage already decreases in abundance in LDE bed II and the diversity and %P increase to pre-LDE values. The PFN and the benthic foraminiferal assemblages are restored more gradually (Figs 4, 7). Extinctions of benthic foraminifera are not recorded, suggesting that the impact of the LDE on bottom-water environments was less severe than the impact of the PETM. The Th/U-ratio is generally below 1 and the uranium content is slightly higher than pre-LDE levels, suggesting slightly lower oxygen levels at the seafloor. The PFN are high in this interval (Fig. 4), suggesting increased productivity. In Zone 'P4' the PFN drop, coinciding with lowered P/B ratios. The benthic foraminiferal assemblage changes significantly and *Bulimina strobila* Marie, 1941*/farafraensis* LeRoy, 1953 dominates in this interval. In recent environments, high abundances of buliminid taxa are commonly related to enhanced food levels or oxygen deprivation, or a combination of these factors (e.g. Corliss & Chen, 1988; Jorissen *et al*., 1995).

Sea-level across the LDE

The P/B ratio does not provide a reliable estimation of palaeodepth in the intervals affected by carbonate dissolution; as planktic and thin-walled, fragile benthic foraminifera will preferentially dissolve and more robust taxa will be enriched in the assemblage (e.g. Nguyen *et al*., 2009 and references therein). Alternatively, the relation of the benthic foraminifera assemblage composition with water depth may be used (Fig. 5). Well below and above the LDE, the abundances up to 25% of bathyal species such as *Gavelinella beccariiformis*, *Nuttalides truempyi* and *Gyroidinoides globosus* (von Hagenow, 1842) and the absence of (inner) neritic taxa (Figs 4, 7) indicate a palaeodepth of ~150–250 m (outer shelf to upper bathyal; see Speijer, 2003a). The lower abundances (<10%) of bathyal taxa directly below and above the LDE $(\sim 7.5-9.5 \text{ m})$ suggest shallowing preceding, and deepening following, the LDE. Within the carbonate-poor interval below the LDE a thin level around 8 m, where dissolution is not prominent, contains a moderately diverse assemblage with decreased (5%) abundances of typical bathyal to outer neritic taxa, such as *G. globosus* and *Angulogavelinella avnimelechi* (Reiss, 1952). Outer neritic species occur in this level, such as *Osangularia plummerae* and *Cibicidoides pseudoacutus* (Nakkady, 1950), together with increased abundances of shallower, neritic taxa, including 14–20% of the inner-middle neritic *Siphogenerinoides esnehensis*. LDE bed II (and the burrows penetrating from bed II down into bed I) contain high numbers of *Neoeponides duwi* and few other taxa, mainly restricted to neritic deposits. This benthic assemblage would indicate a water depth of ~30–70 m (e.g. Schnack 2000; Speijer, 2003a). If the succession of benthic assemblages would be attributed solely to variations in water depth, shallowing and subsequent deepening would amount to ~75–200 m. However, the evidence for increased supply of organic matter and anoxia outlined above may have affected the abundances of bathyal species including *G. beccariiformis* and *N. truempyi*, considered to prefer more oligotrophic conditions than inferred for the Egyptian Paleocene shelf during the LDE (e.g. Widmark & Speijer, 1997; Thomas, 1998). Therefore, extreme shallowing is a rather unlikely explanation for the temporary dominance of *N. duwi* within the LDE beds. The incursion of the *N. duwi* assemblage in Egyptian sections appears to occur in response to anoxia during the LDE (cf. Speijer, 2003a). Despite its size and stout test, *N. duwi* might be an opportunistic species, and its distribution may be driven

Fig. 10. Sketch summarizing the palaeoenvironmental reconstruction of the Late Danian Event beds at Qreiya-3. The transition from the underlying Dakhla Shales to the LDE beds is abrupt. The event beds are interpreted as representing a transgressive phase during a sea-level cycle. TOC, total organic carbon; C_{org}, organic carbon; NC, non-calcareous agglutinated taxa; ON, outer neritic; UB, upper bathyal. See text for further explanation.

by the 'delta effect' described by Pflum & Frerichs (1976). We suspect that *N. duwi* is able to withstand oxygen depletion, favouring its rapid entry as one of the first – calcareous – species in formerly vacated environments. Although documented present-day early recolonizers are often agglutinated taxa (e.g. Hess *et al*., 2001; Koho *et al*., 2007), these taxa are too fragile to yield fossil analogues. Alternatively, recolonization sequences of the Paleocene may have been genuinely different from today's.

Since *N. duwi* appears not to be confined to a particular depth range, water depth estimations based on this species in a context of severe environmental changes, such as anoxia, are not reliable. Consequently, we base the magnitude of the sea-level cycle across the LDE on relative abundances of bathyal and outer neritic taxa, estimating it to be around 50 m maximum, fluctuating between upper bathyal and outer neritic.

Sequence stratigraphy

The inferred palaeoenvironmental changes associated with the deposition of the LDE beds are summarized in Figure 10. The increase in the Zr/Rb ratio indicates that coarser sediments were deposited above 6.5 m, in agreement with shallowing towards the LDE and an interpretation as a late, prograding part of a highstand systems tract (HST) (Dypvik & Harris, 2001; Speijer, 2003a; Schulte & Speijer, 2009). Seafloor anoxia during deposition of LDE bed I suggests that the LDE beds were deposited during a sea-level rise and, therefore, could be assigned to a transgressive systems tract (TST) directly overlying the HST. This would imply that the base of the LDE constitutes a sequence boundary (SB) without evidence for deposition during the sea-level lowstand; an observation that is typical for sequences on the Egyptian shelf (e.g. Speijer & Wagner, 2002). This would also suggest that an unconformity is present at the base of the LDE. However, the duration of the possible hiatus associated with this unconformity cannot be constrained here. In the interval above the LDE, a maximum flooding surface (MFS) may be positioned around 9.5 m and the sediments further upsection may represent a HST. The calcarenitic channel fills indicate occasional variations of bottom-water currents and are not related to changes in sea-level.

A relative sea-level fall before deposition of the LDE beds and a subsequent rise in sea-level is also observed at Gebel Aweina (Speijer & Schmitz, 1998; Speijer, 2003a). In the area between Abu Tartur and Aswan a younger erosional surface (upper Zone P3a to lower P4 are missing), also observed at Gebel Aweina, seems to amalgamate with the older unconformity of the LDE, leading to a major hiatus, which was earlier related to a relative sea-level fall that started during early biochron P3a (Luger, 1985; Speijer 2003a).

CONCLUSIONS

The Qreiya 3 section comprises an almost complete upper Danian to lower Selandian succession in an outer neritic to upper bathyal depositional setting, and encompasses the Latest Danian Event (LDE: 8.20–8.45 m above base of section), characterized by anomalous lithology. Trace elements, nannofossils and benthic foraminifera indicate increased nutrient input and anoxic conditions in the lower part of the LDE (bed I). The upper part of the LDE (bed II) records intermittent oxygenation of the seafloor. Incursion of the *N. duwi* benthic foraminiferal assemblage in bed II is interpreted as repopulation during the recovery from anoxic conditions. Benthic foraminifera indicate shallowing preceding the LDE. LDE bed I was probably deposited during rapid relative sea-level rise.

Explanation of Plate 1.

Typical planktic and benthic foraminifera in the Qreiya 3 section. **figs 1–8**. Biostratigraphical marker species. **figs 1–3**. *Globanomalina* cf. *G. pseudomenardii*: **1,** spiral view, sample Q3–25 (12 m); **2,** umbilical view, sample Q3–25 (12 m); **3,** umbilical view, sample Q3–30 (14.5 m); **figs 4–6**. *Igorina albeari*: **4,** spiral view, Q3 sample C1–4-0; **5,** apertural view, Q3 sample C1–4-0; **6,** umbilical view, sample Q3–44 (15.85 m); **figs 7, 8**. *Morozovella angulata*: **7,** spiral view, sample Q3–16 (3.6 m); **8,** apertural view, sample Q3–17 (3.9 m). **figs 9–10**. Benthic foraminifera from the LDE interval: **9,** *Siphogenerinoides esnehensis*, sample Q3–35 (8.0 m); **10,** *Neoeponides duwi*, spiral view, sample Q3–44 (8.40 m); **figs 11–17**. Benthic foraminifera with an outer neritic and bathyal habitat: **11,** *Cibicidoides pseudoacutus*, spiral view, sample Q3–1 (0.0 m); **12,** *Angulogavelinella avnimelechi*, umbilical view, sample Q3–9 (1.95 m); **13,** *Anomalinoides affinis*, spiral view, sample Q3–20 (4.6 m); **14,** *Gavelinella beccariiformis*, umbilical view, sample Q3–1 (0.0 m); **15,** *Osangularia plummerae*, umbilical view, sample Q3–16 (3.6 m); **16,** *Nuttallides truempyi*, spiral view, sample Q3–51 (9.45 m); **17,** *Pulsiphonina prima*, spiral view, sample Q3–28 (6.5 m). Scale bars 100 µm; upper scale bar refers to figs 1–8 and lower scale bar to figs 9–17.

The magnitude of sea-level fluctuations cannot be constrained on the basis of the *N. duwi* assemblage, as the nominative species appears to be able to shift its depth distribution following a restoration of the benthic environment after widespread anoxia in the basin; however, based on the general benthic foraminiferal assemblage changes, the magnitude of shallowing and subsequent deepening across the LDE is estimated to be about 50 m maximum. A similar order of environmental changes is recorded in other Paleocene sections in the Nile Basin in Egypt. The event beds of the LDE are correlative with carbon isotope excursions in Zumaia, Spain, and in several open-ocean records from the Atlantic and Pacific and probably represent environmental changes associated with a Paleocene hyperthermal. Nannoflora and foraminifera record abundance shifts across the LDE, which are in certain aspects comparable to the PETM, but extinctions of benthic foraminifera do not occur; suggesting a lesser impact of the LDE compared to the PETM.

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APPENDIX A: SPECIES LIST

Several species are illustrated in Plate 1. For additional illustrations and references to taxonomical literature, see Speijer & Schmitz (1998) and Ernst *et al*. (2006).

Alabamina midwayensis Brotzen, 1948 *Alabamina* cf. *A. creta* Finlay, 1940 *Ammodiscus cretaceus* (Reuss, 1845) *Angulogavelinella avnimelechi* (Reiss, 1952) *Anomalinoides affinis* (Hantken, 1875) *Anomalinoides midwayensis* (Plummer, 1926) *Anomalinoides praeacutus* (Vasilenko, 1950) *Anomalinoides rubiginosus* (Cushman, 1926) *Anomalinoides susanaensis* (Browning, 1959) *Bulimina farafraensis* LeRoy, 1953 *Bulimina midwayensis* (Cushman & Parker, 1936) *Bulimina quadrata* Plummer, 1927 *Bulimina strobila* Marie, 1941 *Bulimina thanatensis* Cushman & Parker, 1947 *Cibicidoides alleni* (Plummer, 1927) *Cibicidoides* cf. *C. hyphalus* (Fisher, 1969) *Cibicidoides pseudoacutus* (Nakkady, 1950) *Cibicidoides rigidus* (Nakkady, 1959) *Cibicidoides succedens* (Brotzen, 1948) *Coryphostoma* cf. *C. midwayensis* (Cushman, 1936) *Eponides lunatus* Brotzen, 1948 *Gaudryina pyramidata* Cushman, 1946 *Gavelinella beccariiformis* (White, 1928) *Globocassidulina subglobosa* (Brady, 1881) *Gyroidinoides globosus* (von Hagenow, 1842) *Gyroidinoides* cf. *G. tellburmaensis* Futyan, 1976 *Hansenisca girardanus* (Reuss, 1851) *Loxostomoides applinae* (Plummer, 1927) *Neoeponides duwi* (Nakkady, 1950) *Nuttalides truempyi* (Nuttall, 1930) *Oridorsalis plummerae* (Cushman, 1948) *Osangularia plummerae* Brotzen, 1940

Pullenia jarvisi Cushman, 1936 *Pulsiphonina prima* (Plummer, 1927) *Siphogenerinoides elegantus* (Plummer, 1927) *Siphogenerinoides esnehensis* (Nakkady, 1950) *Spiroplectinella dentata* (Alth, 1850) *Spiroplectinella knebeli* (LeRoy, 1953) *Stilostomella plummerae* (Cushman, 1940) *Sporobulimina eocaena* Bykova, 1959 *Tappanina selmensis* (Cushman, 1933) *Tritaxia midwayensis* (Cushman, 1936) *Valvalabamina depressa* (Alth, 1850) *Valvalabamina planulata* (Cushman & Renz, 1941) *Valvulineria*? *insueta* (Cushman & Bermudez, 1951)

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