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1	Multiproxy analysis of paleoenvironmental, paleoclimatic and paleoceanographic
2	changes during the early Danian in the Caravaca section (Spain)
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14	ABSTRACT
15	
16	After the Chicxulub impact and mass extinction at the Cretaceous-Paleogene boundary (K-PgB),
17	ecosystems haltingly recovered under unstable conditions. An early Danian (65.9 Ma)
18	perturbation of the carbon cycle known as Dan-C2, which includes two carbon isotopic excursions
19	(CIEs), has been ascribed to inputs of greenhouse gases through large-scale volcanism of the
20	Deccan Traps. However, the relationship between Dan-C2, volcanism and environmental and
21	climatic changes during the early Danian remains ambiguous. Based on stable isotopes, calcium
22	carbonate content, magnetic susceptibility and planktic foraminifera, we present a
23	paleoenvironmental, paleoclimatic and paleoceanographic reconstruction of the early Danian
24	from the Caravaca section, Spain, one of the most complete and continuous K-PgB sections

25 worldwide. The paleobiological response of planktic foraminifera suggests very volatile 26 environmental conditions during the first 230 kyr of the Danian, as reflected in the rapid 27 succession of opportunistic/generalist blooms and episodic high occurrences of aberrant 28 specimens. According to our age model, the Dan-C2 has been identified at the Caravaca section 29 from 65.92 to 65.74 Ma. No evidence of strong carbonate dissolution through ocean acidification 30 was observed in the Dan-C2 interval or the rest of the studied section, excluding the K-PgB clay 31 bed. We find that blooms of highly eutrophic Chiloguembelitria and increases in aberrant planktic 32 foraminifera coincided with a major early Danian eruptive episode of Deccan Traps (Ambelani 33 Formation), occurring before the Dan-C2. Conversely, during both Dan-C2 CIEs, less 34 opportunistic taxa thrived, indicating changes in the upper part of the water column. This study 35 demonstrates that the relationship between marine biota and climate change was very complex 36 and rapidly changing during the early Danian. In addition, we propose that the Deccan volcanism 37 had adverse effects on marine plankton, mostly through strong eutrophication, while an increased 38 water column stratification during the Dan-C2 event resulted in a transient boost in the recovery 39 of ecosystems.

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#### 41 HIGHLIGHTS

42 Rapid changes among planktic foraminifera within the first 230 kyr of the Paleocene

43 Low carbonate dissolution in the Danian suggests low rates of volcanic CO<sub>2</sub> emissions

44 Evidence of the Dan-C2 event between 65.92 and 65.74 Ma

45 Dan-C2 CIEs associated with reorganization of planktic foraminiferal assemblages

46

#### 47 **KEYWORDS**

48 Deccan volcanism; Dan-C2; acme-stage; mass extinction; western Tethys

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## 52 1. Introduction

53 The Cretaceous-Paleogene boundary (K-PgB) is marked by one of the most devastating 54 geological events that has occurred on Earth (Alvarez et al., 1980; Smit and Hertogen, 1980) 55 caused by the impact of a ~10 km-diameter asteroid at the Yucatan Peninsula, Mexico, known as 56 the Chicxulub asteroid (Hildebrand et al., 1991). It is widely understood that the asteroid impact 57 caused a series of catastrophic environmental effects, including the blockage of solar radiation 58 leading to a cold and dark "impact winter", ocean acidification, and pollution by toxic heavy 59 metals, resulting in one of the greatest biotic crises on Earth (Kring, 2007; Premović, 2009; 60 Schulte et al., 2010; Vellekoop et al., 2014, 2016; Gulick et al., 2019; Henehan et al., 2019; Gibbs 61 et al., 2020). The environmental effects were lethal in the pelagic realm and caused the decimation 62 of calcareous plankton at the K-PgB (Smit, 1982; Arenillas et al., 2000a,b; Bown, 2005). Multiple 63 lines of evidence have pointed to the Chicxulub impact as the main cause of the K-PgB mass 64 extinction (e.g. Smit, 1999; Arenillas et al., 2006; Schulte et al., 2010; Lowery et al., 2018; 65 Henehan et al., 2019). Nonetheless, recent advances in radiometric dating constrain the eruptive 66 phases of Deccan Traps volcanism (in India), as well as the Chicxulub impact, to a period of only 67 a few hundred thousand years during magnetochron C29r (Chenet et al., 2007; Renne et al., 2015; 68 Schoene et al. 2015, 2019; Sprain et al., 2019) thereby hindering a clear distinction between the 69 specific roles of volcanism and impact in the K-PgB mass extinction. Consequently, these issues 70 remain a topic of intense debate 40 years since the impact hypothesis was first proposed (Alvarez 71 et al., 1980; Hull et al., 2020; Keller et al., 2020).

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Discrepancies in the age of the K-PgB and its stratigraphic position within the Deccan Traps, as well as uncertainty regarding the eruptive rates of its main phases, result in two models of Deccan Traps eruptions, and a controversy about the role of Deccan volcanism in the K-PgB mass extinction and early Danian climate change (Burgess, 2019; Hull et al., 2020; Keller et al., 2020). Based on <sup>40</sup>Ar/<sup>39</sup>Ar dating, and volcano-stratigraphic and biostratigraphic evidence, it has been argued that the most voluminous Deccan eruptions occurred during the early Danian, corresponding to the emplacement of the Poladpur, Ambenali and Mahabaleswar Formations of the Wai subgroup (Jay and Widdowson, 2008; Renne et al., 2015; Richards et al., 2015; Sprain et al., 2019). However, based on U/Pb dating of the Deccan Traps Formations, it has been proposed that the volcanic phase with the highest eruptive rate (Poladpur Formation) occurred in the latest Maastrichtian, only tens of thousands of years prior the K-PgB (Schoene et al., 2015, 2019, 2021).

84

85 Although geochemical signatures ascribed to Deccan volcanism, such as <sup>187</sup>Os/<sup>188</sup>Os 86 excursions and Hg enrichments, have been recognized prior to the KPB (Robinson et al., 2009; 87 Font et al., 2016,2018; Keller et al., 2020), several paleo-ecological and paleoclimate studies (e.g. 88 Thibault and Gardin, 2010; Thibault et al., 2016; Hull et al., 2020; Gilabert et al., 2021) have 89 shown that the influence of the Deccan volcanism during the latest Maastrichtian did not 90 contribute to the K-PgB mass extinction (although this is disputed by some; e.g. Keller et al., 91 2020 and references therein). A broad temporal coincidence also exists between post-K-PgB 92 Deccan volcanism and the first Danian hyperthermal event, known as Dan-C2 (Quillévéré et al., 93 2008), which has led some to speculate that the two are mechanistically linked (e.g. Coccioni et 94 al., 2010; Punekar et al., 2014a; Krahl et al., 2020). However, others have suggested that the Dan-95 C2 event could be astronomically controlled (Quillévéré et al., 2008; Barnet et al., 2019; Sinnesael 96 et al., 2019) and recent models of the CO<sub>2</sub> emission rates of Deccan volcanism suggest that 97 outgassing from Deccan volcanism alone was incapable of driving the magnitude of climate 98 change observed during the early Danian (Hull et al., 2020; Fendley et al., 2020).

99

During the earliest Danian, planktic foraminiferal and calcareous nannoplankton assemblages were characterized by low diversity, a high single-species dominance, rapid evolutionary turnovers, and blooms of smaller generalist or opportunist taxa that could thrive under eutrophic and unstable conditions (Romein, 1977; Smit, 1982; Huber et al., 2002; Lamolda et al., 2005; Arenillas et al., 2006; Jiang et al., 2010; Jones et al., 2019; Lowery et al., 2020). Recently, it has been proposed that non-calcareous algal and microbial communities bloomed in the open ocean in the short-term aftermath of the Chicxulub impact (Bralower et al., 2020). According to Bralower et al. (2020), these microbial blooms probably contributed to rapid
ecosystem recovery by removing nutrients and providing a food source for higher trophic orders,
enhancing pelagic ocean habitability as evidenced by the rapid recovery of planktic foraminifera
and calcareous nannoplankton after the impact.

111

112 The main planktic foraminiferal indicators of enhanced environmental stress across the K-113 PgB are the guembelitriid blooms (Kroon and Nederbragt 1990; Keller and Pardo, 2004; Pardo 114 and Keller, 2008; Ashckenazi-Polivoda et al., 2014; Punekar et al., 2014a,b) and the increases in 115 aberrant planktic foraminifera tests (Gerstel et al., 1986; Coccioni and Luciani, 2006; Arenillas 116 et al., 2018). Worldwide, blooms of Guembelitria and its descendant Chiloguembelitria have been 117 reported well above the K-PgB (Arenillas et al., 2018), and thus appear genetically disconnected 118 from the Chicxulub impact. However, the blooms did occur within the temporal range of Deccan 119 volcanism in the Danian which points to a potential cause and effect relationship (Keller et al., 120 2012; Punekar et al., 2014a,b; Arenillas et al., 2018). Similarly, an increase in aberrant 121 foraminifera tests after the K-PgB has been shown to continue locally at the El Kef and Aïn 122 Settara sections (Tunisia) for several hundreds of thousands of years after the K-PgB (Arenillas 123 et al., 2018), suggesting the persistence of stressed conditions.

124

125 Except for the immediate aftermath of the Chicxulub impact, the climatic and environmental 126 changes that occurred during the first thousand years of the Danian leading up to the Dan-C2 127 event have not been exhaustively examined (e.g., Quillévéré et al., 2008; Barnet et al., 2019). To 128 improve our understanding of the complex paleobiological changes that took place during the 129 early Danian, and their potential relationship with the Deccan volcanism and the Dan-C2 event, 130 we carried out a detailed analysis of the first ~750 kyr of the Danian at the Caravaca section (SE 131 Spain, western Tethys). We took a multi-proxy approach: quantitative, diversity, taphonomic 132 (fragmentation index) and teratological (percentage of aberrant specimens) analyses of planktic 133 foraminifera, as well as bulk geochemical (stable C- and O-isotopes, CaCO<sub>3</sub> content) and 134 magnetic susceptibility measurements. Caravaca is a well-known section for its excellent exposure, completeness and continuity (Smit, 1982, 2004; Molina et al., 2009), and provides an
exceptional opportunity to evaluate paleoclimatic, paleoceanographic and paleobiological
changes during the early Danian.

138

### 139 **2. Material and methods**

140

141 We revisited the Caravaca section, which is located in the Barranco del Gredero (38°04'36" 142 N, 1°52'42" W), southwest of Caravaca de la Cruz, SE Spain (Fig. 1). The Danian part of this 143 section consists mostly of hemipelagic marly limestones (Fig. 2), although it starts with the well-144 known K-PgB clay bed (Smit, 1982, 2004), consisting of a 1-2 mm-thick red air-fall layer and a 145 6 cm-thick dark clay bed, that is almost black in the lowermost 1.5 cm (Fig. 1E). This section was 146 chosen as an auxiliary section of the Global Boundary Stratotype Section and Point (GSSP) for 147 the base of the Danian Stage (Molina et al., 2009), as it represents one of the most continuous and 148 complete K-PgB sections worldwide (Smit and Hertogen, 1980; Smit and Romein, 1985). 149 Previous studies have focused on the K-PgB clay bed and the first one or two meters of the 150 lowermost Danian, with a significantly lower resolution above this interval (Smit, 1982, 2004; 151 Canudo et al., 1991; Coccioni and Galeotti, 1994; Kaiho and Lamolda, 1999; Arz et al., 2000; 152 Lamolda et al., 2005; Vellekoop et al., 2018; Sepúlveda et al., 2019). In contrast, we sampled the 153 first 820 cm of the Danian at high resolution, taking samples every 1-5 cm over the first 200 cm, 154 and every 25-30 cm across the rest of the section.







Fig. 1. A) Paleogeographical location map of Caravaca and other localities worldwide: DSDP Site 516,
ODP Sites 1262 and 1049, and Gubbio (Contessa Highway section). B) Detailed paleogeographical map
of the Western Tethys, with the star indicating the position of Caravaca (modified from Andeweg, 2002).
C) Caravaca section overview. D) Detail of the Cretaceous-Paleogene transition. E) Rock fragment of the
first 3 centimeters of the K-PgB clay bed at Caravaca, which includes the 1-2 mm thick ejecta-rich air-fall
layer and the basal part of dark clay bed.

# 165 2.1. Micropaleontological methods

166

For micropaleontological analyses, a total of 46 samples were disaggregated in  $H_2O_2$  for 3-4 hours. These samples were washed and sieved under running water; the size fraction >63 microns was collected, and the residue was oven-dried at 50 °C for 24 h. Representative splits of ca. 300 individuals per sample were studied for quantitative analyses, classifying the specimens at species level. Representative specimens from Caravaca were photographed with a JEOL JSM 6400 SEM (scanning electron microscope) at the Microscopy Service of the Universidad de Zaragoza (Spain).

175 For the Danian, we have used the planktic foraminiferal zonation of Arenillas et al. (2004), 176 which was updated by Metsana-Oussaid et al. (2019). In Fig. 2, this biozonation has been 177 compared with the more standardized zonation of Berggren and Pearson (2005), which was 178 revised by Wade et al. (2011). Although the taxonomy used by the authors differs, the close 179 correspondence of biozones and subbiozones is illustrated in Fig. 2. The stratigraphic distribution 180 of Danian planktic foraminiferal species across the Caravaca section is also illustrated in Fig. 2. 181 SEM photographs of index-species and other relevant Danian species are displayed in Fig. 3.







- 185 Fig. 2. Stratigraphic column and distribution of the Danian species at Caravaca. Gb. = Guembelitria; Chg. =
- 186 Chiloguembelitria; Pc. = Pseudocaucasina; Pg. Palaeoglobigerina; Pv.= Parvularugoglobigerina; W.=
- 187 *Woodringina; Ch.* = *Chiloguembelina; E.* = *Eoglobigerina; T.* = *Trochoguembelitria; G.* = *Globanomalina;*
- 188 *P*.= *Parasubbotina; Pr*.= *Praemurica; Gc*.= *Globoconusa; S*.= *Subbotina.*
- 189



192 Fig. 3. SEM photographs of Pseudocaucasina antecessor (A-D); Guembelitria cretacea (E); 193 Chiloguembelitria hofkeri (F); Chiloguembelitria danica (G); Trochoguembelitria alabamensis (H); 194 Palaeoglobigerina alticonusa (I); Chiloguembelina midwayensis (J); Parvularugoglobigerina

longiapertura (K); Parvularugoglobigerina eugubina (L); Woodringina hornerstownensis (M);
Chiloguembelina taurica (N); Eoglobigerina simplicissima (O); Eoglobigerina edita (P); Eoglobigerina
pentagona (Q); Parasubbotina pseudobulloides (R); Subbotina triloculinoides (S); Globanomalina
archeocompressa (T); Praemurica taurica (U); Praemurica inconstans (V); Globanomalina compressa
(W). White bar scales = 100 microns.

200

201 To identify the planktic foraminiferal acme-stages (PFAS) proposed by Arenillas et al. 202 (2006) for the lower Danian, we used quantitative data (Tables S1, S2) and the PAST software 203 (v4.0.3, Hammer et al., 2001) for R-mode cluster analyses using the well-known Bray-Curtis 204 index. We chose The Bray-Curtis similarity index since it is a more appropriate index for 205 abundance data (i.e. species assemblages) than other distance measures, such as the common 206 Euclidian distance (Beals, 1984; Ricotta and Podani, 2017). The Euclidean distance can lead to 207 misleading results when species abundance data contains zeros (i.e. absences of certain taxa) as 208 this method places more weight on the abundance differences between samples than on the 209 similarities in the assemblage of species (Legendre and Gallagher, 2001). In contrast, in the Bray-210 Curtis index common and scarce species have relatively similar weights, which means that the 211 assemblage (as opposed to the absolute abundances) becomes the more significant control on 212 distance (Ricotta and Podani, 2017). The criteria for distinguishing the boundaries between PFAS 213 are clear: PFAS-1 is characterized by a dominance of triserial guembelitriids (*Guembelitria*), 214 PFAS-2 by the tiny trochospiral parvularugoglobigerinids (Parvularugoglobigerina and 215 Palaeoglobigerina), and PFAS-3 by biserial Woodringina and Chiloguembelina. All three acme-216 stages, first recognized in the Spanish sections of Zumaia (Arenillas et al., 1998) and Agost 217 (Molina et al., 1998), have been identified in lower Danian sections worldwide (Arenillas et al., 218 2006, 2016; Gallala et al., 2009; Lowery et al., 2018; Renne et al., 2018).

219

To reconstruct paleo-environmental change, we have used several planktic foraminiferal proxies: paleoecological preferences of species, the abundance of aberrant specimens, fragmentation index, diversity indices and the planktic/benthic ratio. Early Danian planktic

223 foraminiferal paleoecology has previously been interpreted based on the isotopic signatures of 224 each species (e.g. Olsson et al., 1999; Aze et al., 2011; Birch et al., 2012). In order to discriminate 225 between normal and abnormal specimens, we followed the compendium of aberrant 226 morphologies of Arenillas et al. (2018) for early Danian planktic foraminifera. The fragmentation 227 index (ratio of broken vs. complete foraminifers) was calculated following the method of Berger 228 et al. (1982), and used to identify changes in carbonate preservation state. Benthic foraminifers 229 were picked to evaluate potential dissolution processes by calculating the planktic/benthic (P/B) 230 ratio (% planktic foraminifera of the total number of foraminifera), but they were not 231 taxonomically classified.

232

233 2.2 Geochemical and geophysical methods.

234

235 The inorganic and organic carbonate content was measured using duplicate subsamples from 236 70 samples that were weighed into ceramic boats, one of which was roasted in air at 420°C for 12 237 hours to remove organic carbon. The total carbon content (TC) of the unroasted subsample, and 238 the Total Inorganic Carbon (TIC) of the roasted subsample, were determined using a Strohlein 239 Coulomat 702, in the Department of Earth Sciences of the University of Oxford. The difference 240 between the amount of carbon determined in unroasted and pre-roasted samples provided an 241 estimate of Total Organic Carbon (TOC). Assuming the inorganic carbon content is all associated 242 with CaCO<sub>3</sub> allows the estimation of CaCO<sub>3</sub> content using the equation from Stax and Stein (1993):  $CaCO_3\% = TIC * 8.33$ . Reproducibility of %C using this method is typically better than 243 244 0.1%.

245

246 Measurements of stable carbon and oxygen isotope ratios ( $\delta^{13}$ C,  $\delta^{18}$ O) were performed on 247 homogenized bulk powdered sediment from the same 70 samples. Samples were analyzed in the 248 Department of Earth Sciences of the University of Oxford using a GasBench device attached to a 249 ThermoFisher Delta V Advantage gas source isotope ratio mass spectrometer. Oxygen and 250 carbon-isotopes are reported using the standard delta notation ( $\delta^{18}$ O,  $\delta^{13}$ C) in parts per mil (‰)

251	on the Vienna PeeDee Belemnite (VPDB) scale. Calibration of samples to the VPDB scale was
252	achieved using multiple analyses of an in-house standard, NOCZ, which has average values on
253	the VPDB scale of -1.90‰ for $\delta^{18}$ O and 2.18‰ for $\delta^{13}$ C. For $\delta^{18}$ O, NOCZ has been calibrated to
254	the VPDB scale by comparison with analyses of NBS-19 and NBS-18, which were assigned $\delta^{18}O$
255	values of -2.20‰ and -23.01‰ respectively. For $\delta^{13}$ C, NOCZ has been calibrated to the VPDB
256	scale by comparison with analyses of NBS-19, which was assigned a value of 1.95‰. Repeated
257	analyses of in-house standards suggest a reproducibility ( $\pm 1\sigma$ ) of <0.1 for both $\delta^{13}C$ and $\delta^{18}O$ .
258	
259	The magnetic susceptibility (MS) of 70 samples was measured at the University of Zaragoza,
260	Spain, with a Spinning Specimen Magnetic Susceptibility Anisotropy Meter KLY-35
261	Kappabridge. Samples were crushed in an agate mortar and measured in cylindrical plastic boxes
262	of 10 cm <sup>3</sup> in volume. MS values are reported relative to mass (m <sup>3</sup> /kg).
263	
264	3. Results
265	
266	3.1. Biostratigraphy and age model
267	
268	At the Caravaca section, a total of 49 species and 14 genera of Danian planktic foraminifera
269	(including species of the genus Guembelitria) have been identified. Relative abundances of each
270	species are shown in the Supplementary Table 1. Seven subbiozones have been identified: Mh.
271	holmdelensis and Pv. longiapertura Subzones (of the G. cretacea Zone), Pv. sabina and E.
272	simplicissima Subzones (of the Pv. eugubina Zone), and E. cf. trivialis, S. triloculinoides and G.
273	compressa (part) Subzones (of the P. pseudobulloides Zone). The stratigraphic interval studied
274	corresponds to P $\alpha$ , P1a, P1b and (part of) P1c of Berggren and Pearson (2005) and Wade et al.

275 (2011). At Caravaca, the bases of these subbiozones are at 0, 3, 22, 42, 107, 332 and 655 cm,

276 respectively, above the K-PgB (Fig. 2).

278 To establish the age model at the Caravaca section, we linearly interpolated between the K-279 PgB, the top of the K-PgB dark clay bed, the C29r/C29n magnetic reversal and the C29n/C28r magnetic reversal. Based on the <sup>40</sup>Ar/<sup>39</sup>Ar calibrations of Sprain et al. (2018), we have assigned 280 281 an age of 66.052 Ma to the K-PgB, 65.724 Ma to the C29r/C29n reversal and 65.075 Ma to the 282 C29n/C28r reversal. Based on cosmic <sup>3</sup>He sedimentation rates, Mukhopadhyay et al. (2001) 283 estimated a duration for deposition of the K-PgB dark clay bed of ~10 kyr. At Caravaca, the top 284 of the K-PgB dark clay bed is ~6 cm above the K-PgB, and, according to Smit (1982) and Groot 285 et al. (1989), the C29r/C29n and C29n/C28r reversals are at 5.1 m and 9.8 m above the K-PgB 286 respectively. Consequently, the average sedimentation rates at Caravaca are  $\sim 0.6$  cm/kyr for the 287 K-PgB dark clay bed, 1.58 cm/kyr for the Danian part of C29r and 0.72 cm/kyr for C29n. In total, 288 the studied section spans approximately the first 760 kyr of the Danian. According to this age 289 model, the bases of Mh. holmdelensis, Pv. longiapertura, Pv. sabina, E. simplicissima, E. cf. 290 trivialis, S. triloculinoides and G. compressa Subzones occurred at 0, 5, 20, 33, 75, 219, and 528 291 kyr after the K-PgB, respectively. It is remarkable that *Pseudocaucasina antecessor* (Arenillas 292 and Arz, 2017) has been identified at Caravaca for the first time. Its Lowest Occurrence Data 293 (LOD) is at 1.5 cm above the K-PgB, i.e. 2.5 kyr after the K-PgB. The LODs of its most direct 294 evolutionary descendants, Parvularugoglobigerina longiapertura and Palaeoglobigerina 295 alticonusa, are at 3 cm above the K-PgB, i.e. 5 kyr after the K-PgB.

296

297 *3.2. Acme-stratigraphy* 

298

Planktic foraminiferal assemblages identified in the lower Danian of the Caravaca section
are characterized by low diversities and high consecutive dominances of single taxon groups,
corresponding to the succession of the three acme-stages PFAS of Arenillas et al. (2006). This is
confirmed by the cluster analysis performed here (Fig. 4).

303

304 PFAS-1 spans the first 5 cm of the lower Danian of the Caravaca section (from the K-PgB
305 to the lowermost part of *Pv. longiapertura* Subzone), i.e. the first 8 kyr after the K-PgB boundary

according to our age model. PFAS-1 is dominated by triserial taxa, mainly *Guembelitria* and, to
a lesser extent, its descendant *Chiloguembelitria*. *Guembelitria* is the only Cretaceous genus that
increased its abundance after the K-PgB. In addition, we have identified a bloom of *Pseudocaucasina antecessor*, which starts within the PFAS-1 and ends at the lowermost part of
PFAS-2. (Fig. 5, Table S1).

311

312 PFAS-2 is placed at 5 to 55 cm above the K-PgB at Caravaca (from the lowermost part of 313 Pv. longiapertura Subzone to the middle part of the E. simplicissima Subzone), i.e. between 8 314 and 41 kyr after the K-PgB. This acme-stage is dominated by parvularugoglobigerinids, i.e. 315 Parvularugoglobigerina and Palaeoglobigerina (the first evolutionary radiation of Danian 316 species), comprising between 50 and 80% of the assemblages. The LODs of Woodringina and 317 Chiloguembelina, the first Danian biserial taxa, occurred within PFAS-2, but their combined 318 relative abundance never exceeds 3% except for the upper part of PFAS-2. Around 38 cm above 319 the K-PgB (upper part of the Pv. sabina Subzone), biserial taxa show a sharp increase, but they 320 do not exceed the parvularugoglobigerinids in abundance.

321

322 PFAS-3 has been recognized from 55 cm above the K-PgB to the top of the studied section 323 (from the lower part of the E. simplicissima Subzone to the lower part of the G. compressa 324 Subzone), i.e. between 41 kyr and at least 756 kyr after the K-PgB. The planktic foraminiferal 325 assemblages in PFAS-3 comprise mostly biserial taxa, i.e. the genera Woodringina and 326 Chiloguembelina, and especially the species Woodringina hornerstownensis (30.7% on average). 327 Although PFAS-3 assemblages are dominated by biserial taxa throughout, several substages can 328 be identified on the basis of changes in relative abundances of some other taxa. One of the most 329 striking features within PFAS-3 at Caravaca is the occurrence of three successive blooms of the 330 opportunist triserial Chiloguembelitria reaching maxima abundances of 48.5, 28.6, and 12.3% 331 respectively. These Chiloguembelitria blooms are successively less intense and alternate in time 332 with remarkable increases in the combined abundance of genera resulting from the second Danian 333 evolutionary radiation, including Eoglobigerina, Parasubbotina, Globanomalina, Praemurica,

and *Subbotina*, or "other genera" for short. The latter group reaches maxima abundance values of 43.1, 49.5 and 48.4% between each *Chiloguembelitria* bloom (Fig. 5). Cluster analyses strongly support the further division of PFAS-3 into 7 shorter substages following the alternation of major groups (Fig. 4), with each substage named as PFAS-3 plus a suffix:  $\alpha$ , T1, O1, T2, O2, T3, O3. Stratigraphic and temporal boundaries of each stage and substage, with average relative abundances of major groups are listed in Table 1.



341

Fig. 4. A) Agglomerative clustering based on the unweighted paired group method with the arithmetic mean
(UPGMA) and the Bray-Curtis similarity index. Oth/Tr\*. = Clusters with ambivalent affinity. B)
Stratigraphically constrained dendrogram.

345

Table 1. Stratigraphic height and age of PFAS and relative abundance of major groups

PFAS	Height from K-PgB (cm)		Age from K-PgB (kyr)		Major groups relative abundance (%)			
	Base	Тор	Base	Тор	Triserial	Parvul.	Biserial	Others

PFAS-3O3	625	820*	487	756*	2.3%	0.0%	56.9%	40.8%
PFAS-3T3	475	625	311	487	8.8%	0.0%	59.5%	31.7%
PFAS-3O2	357	475	235	311	6.6%	0.0%	53.6%	39.8%
PFAS-3T2	282	357	187	235	21.4%	0.0%	58.8%	19.8%
PFAS-301	232	282	155	187	2.4%	0.0%	55.6%	41.9%
PFAS-3T1	107	232	75	155	45.2%	0.0%	49.0%	5.8%
PFAS-3a	55	107	41	75	0.8%	24.7%	67.0%	7.5%
PFAS-2	5	55	8.3	41	9.7%	83.4%	6.8%	0.1%
PFAS-1	K-PgB	5	0	8.3	76.1%	23.1%	0.9%	0.0%

348 Table 1. Stratigraphic position and calibrated age of planktic foraminiferal acme-stages (PFAS) at

349 Caravaca, and relative abundances of the major planktic foraminiferal groups. \* = Top of the studied

- 350 section. Parvul.=Parvularugoglobigerinids
- 351

352 PFAS-3 $\alpha$  is characterized almost exclusively by biserial *Woodringina* and *Chiloguembelina*, 353 but mostly by *Woodringina* (Fig. 5). The LODs of *Eoglobigerina*, *Parasubbotina*, 354 *Globanomalina*, *Praemurica* and *Trochoguembelitria* are at 55-85 cm above the K-PgB, forming 355 the second evolutionary radiation of Danian species (Fig. 2 and Fig. 5). The Highest Occurrence 356 Data (HOD) of *Palaeoglobigerina* and *Parvularugoglobigerina* are recognized towards the top 357 of PFAS-3 $\alpha$ , as these species were completely replaced by the incoming species of the second 358 Danian evolutionary radiation.

359

PFAS-3T (1-3) are characterized by subsequent blooms of triserial *Chiloguembelitria*. PFAS-3T1 witnessed the LOD of the genus *Globoconusa*, which occupies the same ecological niche as *Guembelitria* and *Chiloguembelitria* (see Olsson et al., 1999), although during each *Chiloguembelitria* bloom the relative abundance of *Globoconusa* remains extremely low with values <1% (0.1, 0.2, and 0.9%).

365

366 PFAS-3O (1-3) refers to the substages characterized by the higher relative abundance of the
367 "other genera" combination. The genera *Globanomalina* (15.2%) and *Praemurica* (9.9%) are the

- 368 most abundant genera during each PFAS-30. It is noteworthy that the alternations between PFAS-
- 369 3T and PFAS-3O occurred rapidly, especially between the first three alternations: T1-O1, O1-T2,





Fig. 5. Relative abundance of the Danian planktic foraminiferal genera and major groups at Caravaca.
Biozones: a = (Wade et al., 2011); b = (Arenillas et al., 2004).

375

3/6 3.3. Diversity indic	es.
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377

378 Standard diversity indices are summarized in Table 2 and shown in full in Table S2. We have 379 calculated the average diversity values for PFAS-1-2 and for each substage of PFAS-3. The 380 diversity indices provide evidence for rapid and abrupt environmental changes from PFAS-1 to 381 the end of PFAS-3T1 (suggesting less resilient planktic foraminiferal assemblages), and more 382 stable environmental conditions from PFAS-3O1 onwards (suggesting more resilient 383 assemblages). However, it is noteworthy that the rapid evolutionary radiations which occurred 384 during the PFAS-2 and PFAS-3 $\alpha$  intervals (Figs. 2 and Fig. 5) highly influenced the values of the 385 diversity indices. Conversely, from the base of PFAS-3O1 to the top of the section, diversity

- 386 indices show more gradual changes, showing slightly higher average diversity values in PFAS-
- 387 3O substages than in the PFAS-3T substages (see Table 2).
- 388

Relative abundance of aberrant forms (%) Main diversity indices PFAS S  $\mathbf{H}'$ Е Triserial Parvul. **Biserial** Others  $1/\lambda$ Total PFAS-3O3 0.0% 23 2.42 6.86 0.48 0.0% 1.3% 1.3% 2.6% PFAS-3T3 23 2.37 6.91 0.47 0.4% 0.0% 1.9% 1.4% 3.7% PFAS-3O2 26 2.55 8.09 0.49 0.2% 0.0% 2.2% 1.7% 4.1% PFAS-3T2 24 2.31 6.60 0.42 2.8% 0.0% 5.1% 2.2% 10.0% PFAS-301 25 2.54 8.46 0.52 0.2% 0.0% 2.9% 2.4% 5.5% 6.9% PFAS-3T1 18 1.95 4.86 0.41 0.0% 5.1% 0.5% 12.5% 20 0.0% 0.7% PFAS- $3\alpha$ 1.88 4.24 0.33 1.3% 4.6% 6.6% PFAS-2 15 1.71 3.95 0.40 1.0% 9.0% 0.7% 0.7% 11.5% 6 0.0% PFAS-1 1.26 3.13 0.64 17.3% 3.7% 0.0% 21.0%

Table 2. Diversity indices and relative abundance of aberrant forms of major groups in each PFAS

390

Table 2. Average values of main diversity indices for each planktic foraminiferal acme-stage (PFAS), and
 relative abundance of the aberrant forms in total, and in each major group. S = Species richness; H '=

393 Shanon-Weaver index;  $1/\lambda$ .= Inverse Simpson index; E = Evenness; Parvul. = Parvularugoglobigerinids

394

395 *3.4. Aberrant index* 

396

We have found abnormal specimens of *Guembelitria* and almost every incoming Danian species (Supplementary Table S3), whereas reworked Cretaceous specimens within Danian sediments display almost no aberrations. According to the terminology of Arenillas et al. (2018), the most common aberrant morphologies identified at Caravaca are: 1) chamber abnormalities: aberrant chamber shapes, reduced chamber sizes and overdeveloped chamber sizes; 2) an abnormal ultimate chamber: aberrant shape, anomalous position and bulla-like chamber; 3) 403 multiple ultimate chambers: double or twinned ultimate chambers, and a proliferation of 404 chambers; 4) distortion in test coiling; 5) abnormal tests. Some examples of these aberrations are 405 illustrated in Fig. 6. The aberrant forms of planktic foraminifera are mainly abundant within the 406 first 357 cm (~230 kyr) of the Danian (i.e. from the K-PgB to the top of PFAS-3T2), close to the 407 base of the *S. triloculinoides* Subzone (Fig. 7).

408





410
411 Fig. 6. Examples of different aberrant morphologies within the studied specimens; (A) overdeveloped
412 chamber size; (B) protuberant chamber; (C) aberrant shape; (D) overdeveloped chamber size + anomalous
413 position; (E) proliferation of chambers; (F) aberrant chamber shape; (G) abnormal ultimate chamber; (H)
414 bulla-like chamber; (I) distortion in test coiling and reduced ultimate chamber size; (J) additional chamber;
415 (K) abnormal test; (L) abnormal ultimate chamber; (M) bulla-like chamber; (N) chamber in anomalous
416 position; (O) abnormal test; (P) ultimate chambers in anomalous position and distortion in test coiling; (Q)
417 proliferation of chambers; (R) overdeveloped ultimate chamber with aberrant shape.

418

PFAS-1 and PFAS-2 are characterized by high relative abundances of aberrant specimens
(Fig. 7 and Table 2). The species most commonly displaying aberrant forms are either *G. cretacea*(14.4%) or *Ps. antecessor* (3.5%) for PFAS-1, and *Pv. longiapertura* (5.8%) for PFAS-2. The

422	aberrant specimens of PFAS-3 $\alpha$ mainly belong to biserial taxa, especially to Woodringina
423	hornerstownensis (3.1%). The average aberrant index during PFAS-3 $\alpha$ (6.6%) is significantly
424	lower than during PFAS-2 (11.5%), while during PFAS-3T1 it reaches 12.5%, which is similar
425	to that reached previously in PFAS-2. Triserial and biserial aberrant forms are dominant during
426	PFAS-3T substages, especially for Chg. danica and W. hornerstownensis species. During PFAS3-
427	T1, T2 and T3, aberrant forms of Chg. danica represent, respectively, 5.2, 1.6 and 0.3% on
428	average of total planktic foraminiferal specimens, and 3.3, 2.5 and 1.0% for W. hornerstownensis.
429	Each triserial bloom, i.e. PFAS-1, PFAS-3T1, PFAS3-T2 and to lesser extent PFAS3-T3, displays
430	a transient increase of the aberrant index (Fig 7 and Table 2). Conversely, during the blooming
431	episodes of the "other genera", i.e. PFAS-301, PFAS-302 and PFAS-303, there are fewer
432	aberrant specimens. The most common biserial species with aberrant forms within PFAS-30
433	substages are W. hornerstownensis with respectively 1.8, 1.2 and 0.8% on average, and the most
434	common aberrant specimens of the "other genera" belong to the species Globanomalina
435	archeocompressa (0.55%) for PFAS3-O1, Praemurica taurica (0.4%) for PFAS3-O2 and Pr.
436	inconstans (0.3%) for PFAS3-O3.



439 Fig. 7. Comparison of the quantitative results for the major groups of planktic foraminifera. Parvularugoglo.
440 = parvularugoglobigerinids.

442 3.6. Carbonate preservation and magnetic susceptibility

444 At Caravaca, the CaCO<sub>3</sub> content ranges from 15.5 to 88.4 % with a mean value of 73.78% 445 (n = 70). The lowest CaCO<sub>3</sub> content has been identified in the K-PgB clay bed (0-6 cm, between 446 15.53 and 39.7%, and 28% on average). From 6 to 26 cm above the K-PgB, the CaCO<sub>3</sub> content 447 increases sharply to 70%, and from 26 to 115 cm the average CaCO<sub>3</sub> content is 83%. There is a 448 decrease in the average CaCO<sub>3</sub> content from 115 to 520 cm above the K-PgB (71%) with two 449 relatively low values identified at 245 (60.5%) and 430 (65.5%) cm above the K-PgB (Fig. 8C). 450 Finally, from 520 cm to the top of section, the average CaCO<sub>3</sub> content is relatively stable at 80% 451 on average.

452

453 Apparent dissolution features on tests, such as abrasion marks, broken and/or isolated 454 chambers, or corroded walls, have been identified, but they are not abundant. The planktic 455 for a miniferal preservation is moderate to good in most of the samples, with the exception of those 456 from the dark K-PgB clay bed. We consider that samples with planktic foraminiferal 457 fragmentation ratio or fragmentation index (FI) >40% represent intervals of strong dissolution 458 (e.g., Kucera et al., 1997; Gilabert et al., 2021). Values of planktic foraminiferal fragmentation 459 vary between 7 and 45% across the Caravaca section, with an average fragmentation of 18.3% (n 460 = 46). The average fragmentation values are high (45%) in the K-PgB dark clay bed, moderate 461 (20%) between 6 and 520 cm, and low (13%) from 520 cm to the top of the section (Fig. 8D).

462

463 At Caravaca, the P/B ratio (Fig. 8F) ranges between 11 and 100%, with three distinct 464 intervals: in the dark K-PgB clay bed P/B ratios range between 11 and 47%; from 6 to 26 cm 465 above the K-PgB, they are between 72 and 89%; and from 30 cm to the top of the studied section, 466 they are between 95 and 100%. Benthic foraminifera are more resistant to fragmentation and 467 dissolution than planktic foraminifera, and the P/B ratio is expected to decrease with increasing 468 dissolution intensity (Kucera et al., 1997). The significant, negative correlation r = -0.75 p < 0.01469 between FI and the P/B suggests that higher dissolution of planktic foraminifera is related to lower 470 P/B values, which are limited to the K-PgB clay bed.

472 Magnetic susceptibility (MS) oscillates between  $1.67 \times 10^{-8}$  and  $1.23 \times 10^{-7}$  m<sup>3</sup>/kg across the 473 Caravaca section, with a mean value of  $3.51 \times 10^{-8}$  m<sup>3</sup>/kg. MS values are within the standard range 474 of values for lithified marine samples containing typical paramagnetic minerals (Ellwood et al., 475 2008). MS values increase between 115 and 520 cm above the K-PgB, with maxima at 245 and 476 430 cm, mirroring the CaCO<sub>3</sub> curve (Figs. 8C and 8E). The strong negative correlation between 477 MS and CaCO<sub>3</sub> content suggests variations in the original detrital influx or variations in the flux 478 of carbonate, causing variations in the concentration of paramagnetic minerals.



471





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480

484 3.7. Stable isotopes (bulk carbonate  $\delta^{13}C$  and  $\delta^{18}O$ )

At Caravaca,  $\delta^{13}$ C and  $\delta^{18}$ O values for bulk carbonate show a moderate degree of correlation 486 487 (r = 0.62, p < 0.01) and evolve in parallel in some intervals. However, values are comparable with 488 those previously reported for the lower Danian at the Caravaca section (Kaiho et al., 1999; Sosa-489 Montes de Oca et al., 2016; Sepúlveda et al., 2019) and other sections worldwide (see compilation 490 in Hull et al., 2020), suggesting little influence of diagenesis. Only in the K-PgB clay bed do 491 %CaCO<sub>3</sub> and stable isotope values exhibit a significant correlation. In the rest of the studied section, the correlation between %CaCO<sub>3</sub> and  $\delta^{18}$ O or  $\delta^{13}$ C values is poor to very poor, with r = 492 0.54 (p < 0.01) between %CaCO<sub>3</sub> and  $\delta^{13}$ C, and r = 0.32 (p < 0.01) between %CaCO<sub>3</sub> and  $\delta^{18}$ O. 493 494 This lack of significant correlation suggests that the lithology exerts very little control on the 495 stable isotope values.

496

497 The  $\delta^{13}$ C values vary between -0.78‰ and +1.92‰ at Caravaca (Fig. 8A) with the lowest  $\delta^{13}$ C values registered within the K-PgB dark clay bed, ranging from -0.78‰ to 0.20‰. From 6 498 499 to 57 cm above the K-PgB,  $\delta^{13}$ C increases to the highest values of the section at ~1.92‰. Between the maximum  $\delta^{13}$ C value at 57 cm to 180 cm,  $\delta^{13}$ C displays very small oscillations between 1.92 500 501 and 1.71‰. From 180 cm to the top of the section,  $\delta^{13}$ C broadly displays a clear overarching trend 502 to lower values, with two negative carbon isotopic excursions (CIEs) superimposed upon this 503 trend. The first (CIE-1) has a minimum value of 0.93‰ at 245 cm and the second (CIE-2) a 504 minimum value of 0.69‰, at 460 cm above the K-PgB.

505

506 The  $\delta^{18}$ O values are broadly invarient across much of the Caravaca section, except for three 507 distinct negative excursions (Fig. 8B), the first at the K-PgB and the other two coinciding 508 approximately with the CIEs described above.  $\delta^{18}$ O values are the lowest of the whole section 509 within the K-PgB clay bed ranging from -3.77‰ to -2.82‰. The other two minima in  $\delta^{18}$ O occur 510 at 200 cm above the K-PgB (slightly below the CIE-1), and 460 cm, coincident with CIE-2.

511

### 512 4. Recognizing the Dan-C2 event at Caravaca

514 The Dan-C2 event was first recognized in the NW Atlantic (ODP 1049), and SE Atlantic 515 (DSDP 527 and 528) and defined as a pair of major, fairly symmetrical, negative excursions in 516  $\delta^{13}$ C and  $\delta^{18}$ O (Fig. 9), associated with decreased carbonate content and increased clay content 517 and magnetic susceptibility values (Quillévéré et al., 2008). At Caravaca, the stratigraphic interval 518 corresponding to Dan-C2, as defined in some Atlantic and Tethyan sections (Fig. 9), is recorded 519 between 200 and 490 cm above the K-PgB, from the middle part of E. cf. trivialis (P1a) Subzone 520 to the middle part of S. triloculinoides (P1b) Subzone, i.e. between 130 and 315 kyr after the K-521 PgB according to our age model. The CIEs identified in Caravaca (CIE-1 and CIE-2) are within 522 this stratigraphic interval and consequently they are correlated with the two characteristic negative 523 excursions of Dan-C2 defined elsewhere (Quillévéré et al., 2008; Coccioni et al., 2010). Using 524 our age model, the peak minimum values of CIE-1 and CIE-2 occur 158 and 295 kyr, respectively, 525 after the K-PgB and each CIE has a duration of ~40 kyr. In addition, both CIEs are associated 526 with the lowest CaCO<sub>3</sub> content and the highest MS values. According to our age model, the entire 527 Dan-C2 event lasted approximately 185 kyr, ending around the C29r/C29n magnetic reversal 528 (Fig. 9 and Fig. 10A). The small discrepancies in the assigned ages of Dan-C2 between Caravaca 529 and elsewhere (Fig. 9) are probably related to differences and uncertainties in the age models 530 and/or variations in the local sedimentation rates between tie points that are not represented by 531 linear interpolation. Nevertheless, all of different records suggest a broadly consistent age for 532 Dan-C2 and a termination of the event near the C29r/C29n reversal.



- 542 impact
- 543

Worldwide, the K-PgB dark clay bed was deposited above the Chicxulub-linked air-fall layer under conditions of global climatic warming and alterations in oceanic productivity and acidity (D'Hondt et al. 1998; Coxall et al. 2006; Kawaragi et al., 2009; Birch et al. 2016; Henehan et al., 2016, 2019). At Caravaca, as in most continuous marine sections, the K-PgB dark clay bed is characterized by very low values in %CaCO<sub>3</sub>,  $\delta^{18}$ O and  $\delta^{13}$ C (Figs. 8A, 8B and Figs. 10D, 10E) (see Schulte et al., 2010, and references therein). In addition, the highest planktic foraminiferal 550 fragmentation index values at Caravaca occur within the K-PgB dark clay bed. These geochemical 551 and preservational changes have been directly related to the decimation of pelagic marine 552 calcifiers at the K-PgB (Smit, 1982; Bown, 2005), and subsequent ocean acidification (Alegret et 553 al., 2012; Henehan et al., 2019). At Caravaca, the P/B ratio across the K-PgB dark clay bed is 554 very low, around 30% in comparison to the P/B ratio values-for most of the Danian (Table S2). 555 This decreased P/B ratio is more compatible with the sudden extinction of planktic foraminifera 556 at the K-PgB than with rapid paleobathymetric changes, as previously shown by Alegret et al. 557 (2003). Therefore, the K-PgB dark clay bed records a brief interval of time in which the 558 ecosystems collapsed and the oceans acidified (D'Hondt et al., 1998; Arenillas et al., 2006, 2018; 559 Kring, 2007; Henehan et al., 2019). Biological recovery, however, was relatively quick and 560 oceanic productivity was rapidly re-established after the K-PgB (Sepúlveda et al., 2009, 2019; 561 Lowery et al., 2018; Gibbs et al., 2020). Productivity may have been controlled by blooms in the 562 non-calcareous algal and microbial communities in the open ocean after the K-PgB event, which 563 potentially provided a food supply for higher trophic levels such as calcareous plankton (Bralower 564 et al., 2020).

565

566 The acme of the stress tolerant and opportunistic genus Guembelitria during PFAS-1 is 567 recorded within the K-PgB dark clay bed, immediately above the air-fall layer. 568 Contemporaneously, the start of an acme of the opportunist calcareous dinocyst Thoracosphaera 569 (Fig. 10B) has been reported above the K-PgB at Caravaca (Lamolda et al., 2005, 2016) and in 570 many other Tethyan sections (Romein, 1977; Smit, 1982; Pospichal, 1996; Gardin, 2002; 571 Lamolda et al., 2005, 2016; Fornaciari et al., 2007; Thibault et al., 2018). The highest planktic 572 foraminiferal aberrant index (21%) of the whole section occurs in PFAS-1 (Fig. 10C); a value 573 which is comparable to other Tethyan and North Pacific localities (Gerstel et al., 1986; Coccioni 574 and Luciani, 2006; Arenillas et al., 2018), suggesting stressed conditions on a global scale. 575 Detrimental environmental effects such as eutrophication, warming, ocean acidification, low 576 oxygenation, and the remobilization of pollutants and toxic heavy metals all occurred potentially 577 in the aftermath of Chicxulub impact and can be attributed to it (Smit, 1999; Coccioni and Luciani, 578 2006; Ballent and Carignano, 2008; Omaña et al., 2012; Arenillas et al., 2018; Henehan et al.,
579 2019). Nonetheless, a minor contribution by Deccan volcanism to some of these environmental
580 changes cannot be entirely ruled out.

581

582 5.2. PFAS-2: recovery and the first radiation of planktic foraminifera

583

584 Within PFAS-2, an initial recovery of planktic foraminiferal assemblages and a first 585 evolutionary radiation took place. This first evolutionary radiation was mostly related to the 586 evolution and proliferation of the tiny trochospiral species belonging to Parvularugoglobigerina 587 and Palaeoglobigerina genera. Recently, Arenillas and Arz (2017) proposed that 588 parvularugoglobigerinids originated from a benthic foraminifer that evolved into a planktic form, 589 such as Ps. antecessor. Both Parvularugoglobigerina and Palaeoglobigerina genera radiated and 590 proliferated during PFAS-2, which, together with the continued dominance of Thoracosphaera 591 (Lamolda et al., 2005, 2016) and the very high (11.5%) abundance of aberrant specimens, 592 suggests that conditions remained unstable and stressed throughout PFAS-2.

593

594 At Caravaca, a rapid rebound (< 20 kyr) of the carbonate preservation state between PFAS-595 1 and PFAS-2 is supported by the rapid increase in %CaCO3 and P/B ratios, and a decrease in the 596 fragmentation index and MS values (Fig. 8). According to Henehan et al. (2016, 2019), the initial 597 surface ocean acidification after the K-PgB, together with the extinction of calcifying organisms, 598 would have led to a transient reduction in the marine alkalinity sink. With the return of marine 599 calcifiers, the excess of alkalinity in seawater was removed leading to a rapid rebound and 600 overshoot in surface ocean pH within 40 kyr after the K-PgB. This process likely explains the rapid increase in %CaCO<sub>3</sub> during PFAS-2. The  $\delta^{13}$ C values sharply increase through PFAS-2, 601 602 returning to relatively stable background values similar to those recorded in the uppermost 603 Maastrichtian of Caravaca (e.g. Kaiho et al., 1999; Sosa-Montes de Oca et al., 2016; Gilabert et al., 2021). This trend in  $\delta^{13}$ C also suggests that there was a rapid return of oceanic productivity 604 605 (Sepúlveda et al., 2019).

#### 607 5.3. Onset of PFAS-3 (PFAS-3 $\alpha$ ) and the second radiation of planktic foraminifera

608

609 The biserial taxa Woodringina and Chiloguembelina proliferated throughout PFAS-3 but 610 especially during PFAS-3a, when Woodringina was the dominant taxon (Fig. 5). A second 611 radiation of Danian planktic foraminifera took place within PFAS- $3\alpha$  with the first appearance of 612 several incoming Danian genera including Eoglobigerina, Globanomalina, Parasubbotina, 613 Praemurica and Trochoguembelitria. Most of these new genera seem to have occupied 614 thermocline and subthermocline depths (see Olsson et al., 1999; Aze et al., 2011 and references 615 therein) thus suggesting an initial reoccupation of deeper depth habitats and an incipient but 616 increased water column stratification. The dominance of Woodringina during PFAS-3a suggests 617 relatively warm conditions (see Olsson et al., 1999 and references therein) during this stage. A 618 similar bloom of biserial taxa at Gubbio has been interpreted as an overall reduction in 619 oxygenation of the mixed layer in the oceans (Coccioni et al., 2010) similar to conditions during 620 blooms of biserial taxa during the late Maastrichtian (Pardo and Keller, 2008).

621

622 During PFAS-3a, Thoracosphaera was still the dominant calcareous nannoplankton genera 623 although Braarudosphaera started to replace it as the dominant taxon, at least in the western 624 Tethys (Romein 1977; Smit, 1982; Lamolda et al., 2016). Braarudosphaera species are thought 625 to have been abundant in the lower photic zone under low temperature, low salinity, and eutrophic 626 conditions, and therefore Braarudosphaera species are typically considered opportunists and are 627 associated with episodes of environmental stress (Bukry, 1974; Cunha and Shimabukuro, 1997; 628 Kelly et al., 2003; Lamolda et al., 2005, 2016; Jones et al., 2019). The appearance of incoming 629 species of planktic foraminifera and nannofossils characteristic of deeper water depth habitats 630 suggests a first step for the recolonization of deeper niches. However, it is well-known that the 631 entire reoccupation of the deeper ocean niches by planktic foraminifera, as well as the rebound of 632 diversity levels comparable to pre-KBP levels, took several million years (Aze et al., 2011; Birch

633 et al., 2012, 2016; Lowery and Fraas, 2019). Within PFAS-3α, carbonate parameters (%CaCO<sub>3</sub>

and the fragmentation index) are similar to those in PFAS-2.

635

## 636 5.4. Chiloguembelitria blooms during PFAS-3 and the Dan-C2 event

637

638 Triserial guembelitriid blooms like those recorded during PFAS-3T substages are the most 639 common planktic foraminiferal indicators of high environmental stress (Kroon and Nederbragt 640 1990; Coccioni and Luciani, 2006; Pardo and Keller, 2008; Punekar et al., 2014a,b; Arenillas et 641 al., 2018). In addition, during each triserial bloom identified at Caravaca, a rise in the aberrant 642 index has been identified (Fig. 10B and 10C). Danian triserial guembelitriid blooms have been 643 commonly ascribed to the effect of Deccan volcanism, and linked to the Dan-C2 event (Punekar 644 et al., 2014a; Keller et al., 2016). At Caravaca, the two main Chiloguembelitria blooms (PFAS-645 3T1 and T2) are related to strong increases in the aberrant index (Fig. 7), suggesting higher stress 646 conditions, similar to those reported in Tunisian sections (Arenillas et al., 2018).

647

A triserial guembelitriid acme, such as the bloom of *Guembelitria* (here *Chiloguembelitria*) during PFAS-3T1, has frequently been reported at other Tethyan localities (Arenillas et al., 2000a, b, 2018; Keller, 2003; Coccioni et al., 2010; Punekar et al., 2014a,b), always above the onset of the biserial acme of PFAS-3 (here PFAS-3 $\alpha$ ). In addition, this acme has been reported in the Gulf of Mexico (Arz et al., 2001; Abramovich et al., 2011), in the North Pacific (Smit and Romein, 1985), in the Western North Atlantic (Mateo et al., 2016), and in the Parathetys (Punekar et al., 2016). Therefore, PFAS-3T1 probably characterizes a global response to environmental stress.

655

During PFAS3T-1, the low-oxygenated sub-thermocline dweller *Chiloguembelina* (Boersma
and Premoli Silva, 1989; Olsson et al., 1999; Aze et al., 2011; Luciani et al., 2020) became more
abundant than during PFAS-3α, suggesting the progressive reoccupation of the deeper-most
niches initiated in PFAS-3α. *Chiloguembelina* stabilized in abundance through the section (Fig.

5 and Fig. 10B) suggesting that the oxygen minimum zone did not show major changes during the studied interval. Nevertheless, near the end of PFAS3-T1 (around the onset of Dan-C2), the *Braarudosphaera* bloom was followed by an acme of *Neobiscutum* species (Fig. 10B; Romein, 1977; Gardin and Monechi, 1998, Gardin, 2002; Lamolda et al., 2016; Thibault et al., 2018), suggesting rapidly changing ecological conditions in the upper part of the water column during the beginning of the Dan-C2 event.

666

667 Conversely to what may be expected, the minima of both Dan-C2 CIEs occurred during 668 PFAS-3O substages instead of during triserial blooms. This lack of coincidence between the 669 evidence for C-cycle perturbation and biotic stress was also noted in Italy (Coccioni et al., 2010) 670 and can also be observed in the data reported by Punekar et al. (2014a) from Israel, Egypt and the 671 USA. Since bulk  $\delta^{18}$ O and  $\delta^{13}$ C mostly reflect changes in the upper part of the water column it 672 can be assumed that transient surface water warming occurred during both CIEs. No evidence of 673 bottom water warming has been detected in the Dan-C2 interval, based on benthic foraminiferal 674  $\delta^{18}$ O (Fig. 9; see Barnet et al., 2019 and references therein). Instead, the warming indicated by 675 the bulk oxygen isotope data may have led to a rapid thermal stratification of the upper part of 676 the water column that caused a stronger thermal gradient between the near-surface waters and the 677 thermocline, creating more differentiated ecological niches. The increase of the deeper water 678 dwelling species (such as Chiloguembelina, Eoglobigerina, Globanomalina and Parasubbotina) 679 during both CIEs agrees with transient but enhanced ocean stratification (Fig. 5 and Figs. 10B, 680 10E). The overall planktic foraminiferal assemblage response during Dan-C2 interval is 681 represented by an alternation between the triserial blooms (PFAS-3T1, 2, 3) and increases in other 682 genera (PFAS3-O1, 2) which are progressively less abrupt through time (Fig. 8 and Fig. 10B). 683 These ecological alternations suggest rapidly changing conditions but also a slightly more 684 resilient and stable ocean, compared to the very earliest Danian. During the Dan-C2 event, CaCO3 685 dissolution due to ocean acidification has been invoked (see Coccioni et al., 2010; Krahl et al., 686 2020 and references therein) but, at Caravaca, the dissolution-sensitive parameters (fragmentation 687 index, CaCO<sub>3</sub> content and P/B ratio) do not show evidence of strong dissolution, although we

688 note that there are slight decreases in the %CaCO<sub>3</sub> content and slight increases in the 689 fragmentation index within the Dan-C2 interval, roughly coincident with the two CIEs.

690

# 691 5.5. End of PFAS-3: Shift to a more stable ocean

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The stabilization of the planktic foraminiferal assemblages at Caravaca seems to have been completed in PFAS-3O3, although there are some differences in comparison to the previous PFAS-3O substages. During PFAS-3O3, *Praemurica* significantly increases in abundance, and the triserial *Chiloguembelitria* is replaced by the trochospiral guembelitriid *Globoconusa* (Fig. 5, Fig. 10B and Table S2). The combined abundances of the *Chiloguembelitria* and *Globoconusa* genera is lower than the abundance reached by *Chiloguembelitria* in any of the preceding acme stages and substages, suggesting lower environmental stress conditions from PFAS-3O3 onwards.

700

701 During PFAS-3O3 newly incoming nannoplankton taxa become dominant, including the r-702 selected opportunist Futyana petalosa and the first K-strategists Cruciplacolithus and 703 Coccolithius (Romein, 1977; Gardin, 2002; Thibault et al., 2018; Jiang et al., 2019). 704 Cruciplacolithus and Coccolithius are generally reported as oligotrophic taxa (Jiang et al., 2010, 705 2019), suggesting that the upper ocean waters at Caravaca became more oligotrophic during 706 PFAS-3O3. Oligotrophic conditions at Caravaca are also supported by the very low abundance 707 of Chiloguembelitria and by the rise of the Praemurica lineage, within which planktic 708 foraminifera species first acquire symbionts during the Paleocene (Norris, 1996; Birch et al., 709 2012). In addition, carbonate content, magnetic susceptibility, fragmentation index, the P/B ratio 710 and isotopic proxies show only minor oscillations in PFAS-3O3, suggesting relatively stable 711 conditions (Fig. 8).





Fig. 10. Synthesis of the results obtained in this study: A) Planktic foraminiferal biozonations: a = Wade et al. (2011); b = Arenillas et al. (2004); B) Relative abundances of Danian planktic foraminiferal genera, and correlation of the planktic foraminiferal acme-stages (PFAS) and calcareous nannoplankton blooms (see references in the text); C) Relative abundance of planktic foraminiferal aberrant specimens; D) Main carbonate sensitivity parameters; E) Bulk carbonate stable isotopes.

#### 719 6. What was the environmental impact of the Deccan Traps during the earliest Danian?

720

721 For several decades, identifying the role of the Deccan Traps (India) in environmental and 722 climatic change across the K-PgB has been difficult, mainly due to the uncertainties associated 723 with radioisotopic dating (e.g., Courtillot et al., 1986; Vandamme et al., 1991; Chenet et al., 724 2007). However, recent improvements in radiometric methods have led to refined estimates of 725 the duration of Deccan volcanism, at less than 1 Myr (Schoene et al., 2019; Sprain et al., 2019), 726 and allowed for a more robust correlation between volcanism and climate change (e.g., Barnet et 727 al., 2018; Hernandez Nava et al., 2021). Whether the major eruptive episodes of the Deccan Traps 728 occurred before or after the K-PgB remains a topic of intense debate, since the timing has profound implications for the role of the Deccan Traps in the K-PgB mass extinction (Burgess,
2019; Hull et al., 2020; Keller et al., 2020). This controversy over the role of volcanism in the
extinction event is mostly related to uncertainty over the stratigraphic position of the K-PgB
within the Deccan Traps sequence.

733

734 Based on  ${}^{40}\text{Ar}/{}^{39}\text{Ar}$  dating, Sprain et al. (2018) reported an age of  $66.052 \pm 0.008/0.043$  for 735 the K-PgB, supporting the hypothesis that the most voluminous formations of the Deccan Traps, 736 which belong to the Wai subgroup (Poladpur, Ambenali and Mahabaleswar), are placed above 737 the K-PgB (Jay and Widdowson, 2008; Renne et al., 2015; Richards et al., 2015; Sprain et al., 738 2019). According to the eruptive model of Sprain et al. (2019), the K-PgB is near the base of the 739 Poladpur Formation, and thus the most voluminous eruptive episodes of the Deccan Traps may 740 be early Danian in age. However, based on U-Pb dating, Clyde et al. (2016) reported an age of 741  $66.021 \pm 0.24/0.039$  Ma for the K-PgB, supporting the emplacement of the Poladpur Formation 742 prior to the K-PgB (Schoene et al., 2015, 2019, 2021; Kasbohm et al., 2021; Fig. 11). According 743 to the eruptive model of Schoene et al. (2019), the emplacement of the Poladpur Formation, 744 (which achieved the highest eruption rate of Deccan volcanism) occurred in the latest 745 Maastrichtian, preceding the K-PgB by only some tens of thousands of years. Schoene et al. 746 (2021) recalculated the eruptive volumes of the model of Sprain et al. (2019) in terms of eruptive 747 rate (Fig. 11), which clearly distinguishes between the mega-pulse eruptive model of Schoene et 748 al. (2019) and the quasi-continuous eruptive model of Sprain et al. (2019).



751 Fig. 11. Age models and eruption rates for the Deccan Traps based on Fig. 4 of Schoene et al. (2021), 752 compared with the age of key early Danian paleontological and climatic events as determined in the 753 Caravaca section. The yellow band shows the most probable stratigraphic range for the K-PgB position 754 within the Deccan Traps, considering both the range of error on U-Pb and <sup>40</sup>Ar/<sup>39</sup>Ar dates. Our preferred 755 age model for Caravaca is anchored at the K-PgB using an <sup>40</sup>Ar/<sup>39</sup>Ar age of 66.052 Ma, but we also show 756 how the relationship between Deccan volcanism and the PFAS-3T1, 3O1 and 3T2 and Dan-C2 events could 757 vary by changing the age of the K-PgB to that derived from U-Pb (66.021 Ma). Error margins for both 758 radiometric techniques are also shown and incorporated into the age estimates of early Danian events. E.r. 759 = eruption rate; CI =Confidence interval (1) = (Schoene et al., 2019); (2) = (Sprain et al., 2019); (3) = 760 (Clyde et al., 2016); (4) = (Sprain et al., 2018).

The Rajahmundry Traps (RT, SE India) bear witness to the scale of early Danian Deccan volcanism, extending ~1000 km from the erupting center and forming the longest lava flows on Earth (Self et al., 2008; Keller et al., 2011, 2012). The exposed flows of the RT have been dated as early Danian in age (Keller et al., 2008; Fendley et al., 2020) and have been geochemically assigned to the Ambenali and Mahabaleshwar Formations in the Western Ghats (Baksi, 2001; Self et al., 2008). The age of the RT is also compatible with the recently published radiometric

768 ages of the Ambenali and Mahabaleshwar Formations (Schoene et al., 2019; Sprain et al., 2019). 769 However, the eruption and outgassing rates in each phase of Deccan volcanism are considered 770 potentially more important for climate change than the volume of basalt erupted (Self et al., 2006; 771 Gertsch et al., 2011; Hernandez Nava et al., 2021). Recent estimations of the amount and rates of 772 CO<sub>2</sub> release from Deccan volcanism predict only minor increases in atmospheric CO<sub>2</sub> compared 773 to the background Cretaceous-Paleogene atmospheric reservoir (Self et al., 2006; Schmidt et al., 774 2016; Steinthorsdottir et al., 2016; Henehan et al., 2016; Fendley et al., 2020). Modelling of 775 different climate sensitivities and volcanic outgassing scenarios suggest that Deccan volcanism 776 alone was insufficient to have driven warming during the early Danian (Hull et al., 2020; Fendley 777 et al., 2020). Conversely, the influence of volcanic  $SO_2$  and halogen emissions could have been 778 profound at times of flood basalt emplacement (Self et al., 2006). Although volcanic aerosols 779 have a short residence time in the troposphere of  $\sim 1$  week, flood basalt activity could have 780 provided a semi-continuous SO<sub>2</sub> supply (Self et al., 2006), which may have resulted in cooling 781 and/or ocean acidification on geologic time scales (Gertsch et al., 2011; Courtillot and Fluteau, 782 2014).

783

784 Our data show no indications of ocean acidification or temperature changes in the first 50 785 kyr of the Danian, except within the K-PgB clay bed, and thus the effect of volcanic outgassing 786 directly following the Chicxulub impact (Renne et al., 2015; Sprain et al., 2019) may have been 787 obscured by the extinction of calcifying plankton (Henehan et al., 2019). Alternatively, our data 788 could indicate that there was no significant Deccan volcanic activity during the earliest Danian. 789 This would be in accordance with Schoene et al. (2019), who suggested that the Poladpur eruptive 790 pulse occurred in the latest Maastrichtian and was followed by a period of volcanic inactivity in 791 the Deccan for about 100 kyr, extending into the earliest Danian. The return to higher stressed 792 environmental conditions at Caravaca occurred ~70 kyr after the K-PgB, as evidenced by the 793 Chiloguembelitria bloom in PFAS-3T1, which is broadly coincident with the eruption of the 794 Ambenali Formation (Schoene et al., 2019; Sprain et al., 2019). Recent pH estimates, based on 795 boron isotopes, show initial surface water acidification after the K-PgB, followed by a rapid rebound and overshoot in surface ocean pH within 40 kyr, before pH values returned to pre-K-PgB background levels after ~120 kyr (Henehan et al., 2019). Previous ocean pH estimates predicted a rise in pH due to the extinction of calcifying organisms and the consequent transient reduction in the marine alkalinity sink (Caldeira et al., 1990; Henehan et al., 2016). The latter is supported by observations of improved deep-sea carbonate preservation during the time of the pH overshoot (e.g. Minoletti et al., 2005; Alegret and Thomas, 2013; Tobin et al., 2017) and by our own data (see section 5.1 and 5.2).

803

804 At Caravaca, the aberrant index follows a declining trend during the first  $\sim 70$  kyr of the 805 Danian, from PFAS-1 to PFAS-3a, suggesting a reduction in the impact of the different 806 environmental stressors with a rapid recovery of productivity. Conversely, during the following 807  $\sim$ 80 kyr (in PFAS-3T1), the opportunistic *Chiloguembelitria* bloomed at a global scale, and the 808 aberrant index increased again (see section 5.4), suggesting a new episode of environmental 809 stress. Commonly, guembelitriid blooms are reported from shallow marine areas, often near 810 volcanic provinces with a high nutrient flux (Pardo and Keller, 2008). However, these 811 opportunistic taxa radiated and proliferated in the pelagic realm after the K-PgB (Olsson et al., 812 1999; Arenillas et al., 2000a,b; Keller, 2003), suggesting that, during the early Danian, their 813 ecological preferences of high nutrient availability were met in the open ocean. Although the 814 uncertainties in the temporal correlation between Deccan Traps volcanic phases, the K-PgB, and 815 climatic and paleobiological events are still significant, our data suggest that the 816 Chiloguembelitria bloom (PFAS-3T1) is coeval with the emplacement of the Ambenali 817 Formation (Fig. 11).

818

The emplacement of the Ambenali formation and the apparent co-occurrence with the Dan-C2 event in the upper part of C29r (Fig. 10A and 10D), has led some authors to link these two events mechanistically (Coccioni et al., 2010; Krahl et al., 2020; Punekar et al., 2014a). However, the Dan-C2 event was not associated with bottom water warming (Quillévéré et al., 2008; Coccioni et al., 2010; Barnet et al., 2019; Krahl et al., 2020) which raises questions as to whether 824 this event can be considered a 'true' hyperthermal event or not (see discussion in Barnet et al., 825 2019). Ocean acidification has also been invoked during Dan-C2 (Coccioni et al., 2010; Krahl et 826 al., 2020) but no convincing evidence has been identified during the Dan-C2 interval at Caravaca 827 nor in other sections to support this (see Barnet et al., 2019). The exact mechanisms that drove 828 the Dan-C2 event are poorly resolved, and several hypotheses have been put forward: pulses of 829 massive Deccan volcanic eruptions (Krahl et al., 2020); a combination of Deccan volcanism with 830 an orbital configuration which perturbed the carbon cycle (e.g. Coccioni et al., 2010; Barnet et 831 al., 2019; Sinnesael et al., 2019); and/or passive degassing of  $CO_2$  due to the interaction of 832 intrusive magma bodies with the crust (see Sprain et al., 2019; Fendley et al., 2020). The quasi-833 continuous eruptive model suggests that the Ambenali Formation represents a relatively long 834 volcanic episode (Renne et al., 2015; Sprain et al., 2019). Although this scenario does not provide 835 a relatively rapid trigger mechanism for environmental change, it still permits a hypothesis that 836 the Deccan volcanism could add to emissions of  $CO_2$  by passive degassing over longer timescales, 837 contributing, albeit perhaps in a minor way, to early Danian climate change and the Dan-C2 event. 838 Conversely, the hypothesis of a shorter (<100 kyr), more intense, volcanic pulse (Schoene et al., 839 2019) is not supported by geochemical evidence from marine records (Hull et al., 2020), and 840 would imply a minimal impact of volcanic activity on climate. Regardless of the eruptive model 841 favoured, and considering the estimated duration for the Ambelani eruptive episode, volcanic 842 outgassing from the Deccan Traps by itself was probably insufficient to drive significant warming 843 during the early Danian (Fendley et al., 2020; Hull et al., 2020), although it may have exacerbated 844 environmental stress. Our age model allows us to suggest that the Ambenali phase coincided with 845 the first Dan-C2 CIE, but not with the second CIE (Fig. 11). If Deccan volcanism was not the 846 cause of the second CIE, then it seems that other factors, such as orbital forcing, are required to 847 fully explain the Dan-C2 event.

848

849 Since the proposed warming for the Dan-C2 event is only observed in bulk and planktic 850 foraminiferal isotopic records (Fig. 9), the increase in temperature probably only affected surface 851 ocean waters. A transient but enhanced thermal stratification of the upper part of the water column 852 may have resulted in more differentiated ecological niches. This is supported by our data at 853 Caravaca, which show rapid turnovers of planktic foraminiferal assemblages in PFAS-3O1 and 854 O2 coinciding approximately with both Dan-C2 CIEs. Remarkably, continental records of the 855 first ~700 kyr of the Danian have documented increases in ecologic diversification of plants and 856 mammals (suggested by increased species richness and taxonomic composition), coinciding with 857 the warmer intervals of the early Danian (Lyson et al., 2019; Chiarenza et al., 2020). Therefore, 858 if the Dan-C2 event influenced both marine and terrestrial ecosystems (driven by Deccan 859 volcanism, orbital forcing, or a combination of those and additional mechanisms), it did not cause 860 harmful environmental effects, but instead may have temporarily boosted the recovery of 861 ecosystems.

862

## 863 6. Conclusions

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865 High-resolution planktic foraminiferal, geochemical and paleomagnetic analyses of the first 866 ~750 kyr of the Danian at Caravaca (Spain, western Tethys) were carried out. Planktic 867 foraminiferal assemblages after the Cretaceous-Paleogene boundary (K-PgB) are characterized 868 by a rapid succession of planktic foraminiferal acme-stages (PFAS). The first acme is of the 869 triserial Guembelitria (PFAS-1), the second of the tiny trochospiral Parvularugoglobigerina-870 Palaeoglobigerina (PFAS-2), and the third of the biserial Woodringina-Chiloguembelina (PFAS-871 3). Within PFAS-3, seven shorter substages are distinguished: PFAS-3 $\alpha$  during the maximal 872 bloom of biserials, PFAS-3T1, T2 and T3 for blooms of the triserial Chiloguembelitria, and 873 PFAS-301, O2, O3 for increases in abundance of the trochospiral genera Eoglobigerina, 874 Praemurica, Globanomalina and Parasubbotina. Triserial blooms and a high abundance of 875 aberrant forms are striking evidence of enhanced environmental stress, occurring especially 876 within PFAS-1 and PFAS-3T substages, during the first 230 kyr of the Danian.

877

878 On the basis of  $\delta^{18}$ O and  $\delta^{13}$ C, CaCO<sub>3</sub> content and magnetic susceptibility, the Dan-C2 event 879 has been identified for the first time at the Caravaca section. The two carbon isotopic excursions 880 (CIEs) that characterize the Dan-C2 event have been linked to surface water warming that caused 881 enhanced thermal stratification. During both CIEs, the PFAS-3T assemblages were rapidly 882 replaced by the PFAS-3O assemblages containing less opportunistic taxa, probably because the 883 water depth habitats became more differentiated. High carbonate dissolution is restricted to the 884 K-PgB clay bed, while the Dan-C2 event and the rest of the section show no evidence of 885 significant carbonate dissolution episodes. Although there are still uncertainties in radiometric 886 dating, our data suggest that the first and largest Chiloguembelitria bloom (PFAS-3T1) coincided 887 with the emplacement of the Ambenali Formation of the Deccan Traps. Conversely, the Dan-C2 888 event was decoupled from both the Ambenali eruptive pulse and the *Chiloguembelitria* bloom, 889 starting long after these events had occurred. This suggests that volcanic outgassing of CO<sub>2</sub> was 890 insufficient to drive warming, which only occurred only when the effects of volcanic CO<sub>2</sub> were 891 combined with other factors such as a specific orbital configuration. More high-resolution 892 multidisciplinary studies are needed to fully assess the relationship between Deccan Traps 893 volcanism on early Danian climate, and its potential contribution in reshaping life on Earth after 894 the end-Cretaceous mass extinction.

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905 References

906	Abramovich, S., Keller, G., Berner, Z., Cymbalista, M., Rak, C., 2011. Maastrichtian Planktic
907	Foraminiferal Biostratigraphy and Paleoenvironment of Brazos River, Falls County,
908	Texas, U.S.A., in: Keller, G., Adatte, T. (Eds.), The End-Cretaceous Mass Extinction and
909	the Chicxulub Impact in Texas. SEPM (Society for Sedimentary Geology), Tulsa, pp. 123-
910	156. https://doi.org/10.2110/sepmsp.100.123.
911	Alegret, L., Thomas, E., 2013. Benthic foraminifera across the Cretaceous/Paleogene boundary
912	in the Southern Ocean (ODP Site 690): Diversity, food and carbonate saturation. Mar.
913	Micropaleontol. 105, 40-51. https://doi.org/10.1016/j.marmicro.2013.10.003.
914	Alegret, L., Molina, E., Thomas, E., 2003. Benthic foraminiferal turnover across the
915	Cretaceous/Paleogene boundary at Agost (southeastern Spain): Paleoenvironmental
916	inferences. Mar. Micropaleontol. 48, 251-279. https://doi.org/10.1016/S0377-
917	8398(03)00022-7.
918	Alegret, L., Thomas, E., Lohmann, K.C., 2012. End-Cretaceous marine mass extinction not
919	caused by productivity collapse. Proc. Natl. Acad. Sci. USA 109, 728-732.
920	https://doi.org/10.1073/pnas.1110601109.
921	Alvarez, L.W., Alvarez, W., Asaro, F., Michel, H. V., 1980. Extraterrestrial cause for the
922	Cretaceous-Tertiary extinction. Science 208, 1095-1108.
923	https://doi.org/10.1126/science.208.4448.1095.
924	Andeweg, B., 2002. Cenozoic tectonic evolution of the Iberian Peninsula: effects and causes of
925	changing stress fields. PhD thesis, Vrije Universiteit Amsterdam, Amsterdam, 192 pp.
926	Arenillas, I., Arz, J.A., 2017. Benthic origin and earliest evolution of the first planktonic
927	foraminifera after the Cretaceous/Palaeogene boundary mass extinction. Hist. Biol. 29, 25-
928	42. https://doi.org/10.1080/08912963.2015.1119133.

- Arenillas, I., Arz, J.A., Molina, E., 1998. El límite Cretácico/Terciario de Zumaya, Osinaga y
  Músquiz (Pirineos): control bioestratigráfico y cuantitativo de hiatos con foraminíferos
  planctónicos. Rev. Soc. Geol. Esp. 11(1-2), 127-138.
- 932 Arenillas, I., Arz, J.A., Molina, E., Dupuis, C., 2000a. The Cretaceous/Paleogene (K/P)
- 933 boundary at Aïn Settara, Tunisia: Sudden catastrophic mass extinction in planktic
- 934 foraminifera. J. Foraminifer. Res. 30, 202-218. https://doi.org/10.2113/0300202.
- Arenillas, I., Arz, J.A., Molina, E., Dupuis, C., 2000b. An independent test of planktic
- 936 for a for a turnover across the Cretaceous/Paleogene (K/P) boundary at El Kef, Tunisia:
- 937 Catastrophic mass extinction and possible survivorship. Micropaleontology 46, 31-49.
- 938 Arenillas, I., Arz, J.A., Molina, E., 2004. A new high-resolution planktic foraminiferal zonation
- and subzonation for the lower Danian. Lethaia 37, 79-95.
- 940 https://doi.org/10.1080/00241160310005097.
- 941 Arenillas, I., Arz, J.A., Grajales-Nishimura, J.M., Murillo-Muñetón, G., Alvarez, W., Camargo-
- 942 Zanoguera, A., Molina, E., Rosales-Domínguez, C., 2006. Chicxulub impact event is
- 943 Cretaceous/Paleogene boundary in age: New micropaleontological evidence. Earth Planet.
- 944 Sci. Lett. 249, 241-257. https://doi.org/10.1016/j.epsl.2006.07.020.
- 945 Arenillas, I., Arz, J.A., Grajales-Nishimura, J.M., Meléndez, A., Rojas-Consuegra, R., 2016.
- 946 The Chicxulub impact is synchronous with the planktonic foraminifera mass extinction at
- 947 the Cretaceous/Paleogene boundary: New evidence from the Moncada section, Cuba.
- 948 Geol. Acta 14, 35-51. https://doi.org/10.1344/GeologicaActa2016.14.1.4.
- 949 Arenillas, I., Arz, J.A., Gilabert, V., 2018. Blooms of aberrant planktic foraminifera across the
- 950 K/Pg boundary in the Western Tethys: Causes and evolutionary implications.
- 951 Paleobiology 44, 460-489. https://doi.org/10.1017/pab.2018.16.

952	Arz, J.A., Arenillas, I., Molina, E., Sepulveda, R., 2000. La estabilidad evolutiva de los
953	foraminíferos planctónicos en el Maastrichtiense superior y su extinción en el límite
954	Cretácico/Terciario de Caravaca, España. Rev. geol. Chile 27, 27-47.
955	http://dx.doi.org/10.4067/S0716-0208200000100003.
956	Arz, J.A., Alegret, L., Arenillas, I., Liesa, C., Molina, E., Soria, A.R., 2001. Extinción de
957	foraminíferos del límite Cretácico/Terciario en Coxquihui (México) y su relación con las
958	evidencias de impacto. Rev. Esp. Micropaleontol. 33(2), 221-236.
959	Ashckenazi-Polivoda, S., Rak, C., Almogi-Labin, A., Zsolt, B., Ovadia, O., Abramovich, S.,
960	2014. Paleoecology of the K-Pg mass extinction survivor Guembelitria (Cushman):
961	isotopic evidence from pristine foraminifera from Brazos River, Texas (Maastrichtian).

962 Paleobiology 40, 24-33. https://doi.org/10.1666/13317.

Aze, T., Ezard, T.H.G., Purvis, A., Coxall, H.K., Stewart, D.R.M., Wade, B.S., Pearson, P.N.,

964 2011. A phylogeny of Cenozoic macroperforate planktonic foraminifera from fossil data.

965 Biol. Rev. 86, 900-927. https://doi.org/10.1111/j.1469-185X.2011.00178.x.

- 966 Ballent, S.C., Carignano, A.P., 2008. Morphological abnormalities in Late Cretaceous and early
- 967 Paleocene foraminifer tests (northern Patagonia, Argentina). Mar. Micropaleontol. 67,

968 288-296. https://doi.org/10.1016/j.marmicro.2008.02.003.

- 969 Baksi, A.K., 2001. The Rajahmundry Traps, Andhra Pradesh: Evaluation of their petrogenesis
- 970 relative to the Deccan Traps. Proc. Indian Acad. Sci. Earth Planet. Sci. 110, 397-407.
- 971 https://doi.org/10.1007/BF02702903.
- 972 Barnet, J.S.K., Littler, K., Kroon, D., Leng, M.J., Westerhold, T., Röhl, U., Zachos, J.C., 2018.
- 973 A new high-resolution chronology for the late Maastrichtian warming event: Establishing
- 974 robust temporal links with the onset of Deccan volcanism. Geology 46, 147-150.
- 975 https://doi.org/10.1130/G39771.1.

976	Barnet, J.S.K., Littler, K., Westerhold, T., Kroon, D., Leng, M.J., Bailey, I., Röhl, U., Zachos,
977	J.C., 2019. A High-Fidelity Benthic Stable Isotope Record of Late Cretaceous-Early
978	Eocene Climate Change and Carbon-Cycling. Paleoceanogr. Paleoclimatology 34, 672-
979	691. https://doi.org/10.1029/2019PA003556.
980	Beals, E.W., 1984, Bray-Curtis ordination: An effective strategy for analysis of multivariate

- 981 ecological data. Adv. Ecol. Res. 14, 1-55. https://doi.org/10.1016/S0065-2504(08)60168982 3.
- Berger, W.H., Bonneau, M., Parker, F.L., Plateau, L., 1982. Foraminifera on the deep-sea floor:
  lysocline and dissolution rate. Oceanol. Acta 5, 249-258.
- Berggren, W.A., Pearson, P.N., 2005. A revised tropical to subtropical Paleogene planktonic
- 986 foraminiferal zonation. J. Foraminifer. Res. 35, 279-298. https://doi.org/10.2113/35.4.279.
- 987 Birch, H.S., Coxall, H.K., Pearson, P.N., 2012. Evolutionary ecology of Early Paleocene
- 988 planktonic foraminifera: size, depth habitat and symbiosis. Paleobiology 38, 374-390.
  989 https://doi.org/10.1666/11027.1.
- Birch, H.S., Coxall, H.K., Pearson, P.N., Kroon, D., Schmidt, D.N., 2016. Partial collapse of the
- marine carbon pump after the Cretaceous-Paleogene boundary. Geology 44, 287-290.
  https://doi.org/10.1130/G37581.1.
- Boersma, A., Premoli Silva, I., 1989. Atlantic Paleogene biserial heterohelicid foraminifera and
  oxygen minima. Paleoceanography 4, 271-286.
- Bown, P., 2005. Selective calcareous nannoplankton survivorship at the Cretaceous-Tertiary
  boundary. Geology 33, 653-656. https://doi.org/10.1130/G21566.1.
- Burgess, S., 2019. Deciphering mass extinction triggers. Science 363, 815-816.
- https://doi.org/10.1126/science.aaw0473.

- 999 Bralower, T.J., Cosmidis, J., Heaney, P.J., Kump, L.R., Morgan, J. V., Harper, D.T., Lyons,
- 1000 S.L., Freeman, K.H., Grice, K., Wendler, J.E., Zachos, J.C., Artemieva, N., Chen, S.A.,
- 1001 Gulick, S.P.S., House, C.H., Jones, H.L., Lowery, C.M., Nims, C., Schaefer, B., Thomas,
- 1002 E., Vajda, V., 2020. Origin of a global carbonate layer deposited in the aftermath of the
- 1003 Cretaceous-Paleogene boundary impact. Earth Planet. Sci. Lett. 548, 116476.
- 1004 https://doi.org/10.1016/j.epsl.2020.116476.
- 1005 Bukry, D., 1974. Coccoliths as paleosalinity indicators evidence from Black Sea, in: Degens,
- E.T., Ross. D.A. (Eds.), The Black Sea–Geology, Chemistry, and Biology. AAPG Memoir
  20, Tulsa, pp. 353-363.
- 1008 Caldeira, K., Rampino, M.R., Volk, T., Zachos, J.C., 1990. Biogeochemical modeling at mass
- 1009 extinction boundaries: Atmospheric carbon dioxide and ocean alkalinity at the K/T
- 1010 boundary, in: Kauffman E.G., Walliser O.H. (Eds), Extinction Events in Earth History.
- 1011 Lecture Notes in Earth Sciences, 30, Springer, Berlin, Heidelberg, pp. 333-345.
- 1012 https://doi.org/10.1007/bfb0011156.
- 1013 Canudo, J.I., Keller, G., Molina, E., 1991. Cretaceous/Tertiary boundary extinction pattern and
- 1014 faunal turnover at Agost and Caravaca, S.E. Spain. Mar. Micropaleontol. 17, 319-341.
- 1015 https://doi.org/10.1016/0377-8398(91)90019-3.
- 1016 Chenet, A.L., Quidelleur, X., Fluteau, F., Courtillot, V., Bajpai, S., 2007. <sup>40</sup>K-<sup>40</sup>Ar dating of the
- 1017 Main Deccan large igneous province: Further evidence of KTB age and short duration.
- 1018 Earth Planet. Sci. Lett. 263, 1-15. https://doi.org/10.1016/j.epsl.2007.07.011.
- 1019 Chiarenza, A.A., Farnsworth, A., Mannion, P.D., Lunt, D.J., Valdes, P.J., Morgan, J. V.,
- 1020 Allison, P.A., 2020. Asteroid impact, not volcanism, caused the end-Cretaceous dinosaur
- 1021 extinction. Proc. Natl. Acad. Sci. USA 117, 17084-17093.
- 1022 https://doi.org/10.1073/pnas.2006087117.

1023	Clyde, W.C., Ramezani, J., Johnson, K.R., Bowring, S.A., Jones, M.M., 2016. Direct high-
1024	precision U-Pb geochronology of the end-Cretaceous extinction and calibration of
1025	Paleocene astronomical timescales. Earth Planet. Sci. Lett. 452, 272-280.
1026	https://doi.org/10.1016/j.epsl.2016.07.041.
1027	Coccioni, R., Galeotti, S., 1994. K-T boundary extinction: Geologically instantaneous or
1028	gradual event? Evidence from deep-sea benthic foraminifera. Geology 22, 779-782.
1029	https://doi.org/10.1130/0091-7613(1994)022<0779:KTBEGI>2.3.CO;2.

- 1031 Morphological abnormalities induced by impact-related extreme environmental stress?, in:
- 1032 Cockell, C., Gilmour, I., Koeberl, C. (Eds.), Biological Processes Associated with Impact
- 1033 Events. Springer-Verlag, Berlin, pp. 179-196. https://doi.org/10.1007/3-540-25736-5\_8
- 1034 Coccioni, R., Frontalini, F., Bancalà, G., Fornaciari, E., Jovane, L., Sprovieri, M., 2010. The

1035 Dan-C2 hyperthermal event at Gubbio (Italy): Global implications, environmental effects,

and cause(s). Earth Planet. Sci. Lett. 297, 298-305.

- 1037 https://doi.org/10.1016/j.epsl.2010.06.031.
- 1038 Courtillot, V., Besse, J., Vandamme, D., Montigny, R., Jaeger, J.-J., Cappetta, H., 1986. Deccan
- 1039 flood basalts at the Cretaceous/Tertiary boundary? Earth Planet. Sci. Lett. 80, 361–374.
- 1040 Courtillot, V., Fluteau, F., 2014. A review of the embedded time scales of flood basalt
- 1041 volcanism with special emphasis on dramatically short magmatic pulses, in: Keller, G.,
- 1042 Kerr, A.C. (Eds.), Volcanism, Impacts, and Mass Extinctions: Causes and Effects. Geol.
- 1043 Soc. Am. Spec. Pap. 505, p. 301-317. https://doi.org/10.1130/2014.2505(15).
- 1044 Coxall, H.K., D'Hondt, S., Zachos, J.C., 2006. Pelagic evolution and environmental recovery

after the Cretaceous-Paleogene mass extinction. Geology 34, 297-300.

1046 https://doi.org/10.1130/G21702.1.

- 1047 Cunha, A.S., Shimabukuro, S., 1997. *Braarudosphaera* blooms and anomalous enrichments of
   1048 *Nannoconus*: evidence from the Turonian South Atlantic, Santos Basin, Brazil. J.
   1049 Nannoplankt. Res. 19, 51-55.
- D'Hondt, S., Donaghay, P., Zachos, J.C., Luttenberg, D., Lindinger, M., 1998. Organic carbon
  fluxes and ecological recovery from the Cretaceous-Tertiary mass extinction. Science 282,
  276-279. https://doi.org/10.1126/science.282.5387.276.
- 1053 Ellwood, B.B., Tomkin, J.H., Ratcliffe, K.T., Wright, M., Kafafy, A.M., 2008. High-resolution
- 1054 magnetic susceptibility and geochemistry for the Cenomanian/Turonian boundary GSSP
- 1055 with correlation to time equivalent core. Palaeogeogr. Palaeoclimatol. Palaeoecol. 261,
- 1056 105-126. https://doi.org/10.1016/j.palaeo.2008.01.005.
- 1057 Fendley, I.M., Sprain, C.J., Renne, P.R., Arenillas, I., Arz, J.A., Gilabert, V., Self, S.,
- 1058 Vanderkluysen, L., Pande, K., Smit, J., Mittal, T., 2020. No Cretaceous-Paleogene
- 1059 boundary in exposed Rajahmundry Traps: A refined chronology of the longest Deccan
- 1060 lava flows from <sup>40</sup>Ar/<sup>39</sup>Ar dates, magnetostratigraphy, and biostratigraphy. Geochemistry,

1061 Geophys. Geosystems 21, 1-20. https://doi.org/10.1029/2020gc009149.

1062 Font, E., Adatte, T., Sial, A.N., de Lacerda, L.D., Keller, G., Punekar, J., 2016. Mercury

anomaly, Deccan volcanism, and the end-Cretaceous mass extinction. Geology 44, 171-

- 1064 174. <u>https://doi.org/10.1130/G37451.1</u>.
- 1065 Font, E., Adatte, T., Andrade, M., Keller, G., Mbabi Bitchong, A., Carvallo, C., Ferreira, J.,
- 1066 Diogo, Z., Mirão, J., 2018. Deccan volcanism induced high-stress environment during the
- 1067 Cretaceous–Paleogene transition at Zumaia, Spain: Evidence from magnetic,
- 1068 mineralogical and biostratigraphic records. Earth Planet. Sci. Lett. 484, 53-66.
- 1069 https://doi.org/10.1016/j.epsl.2017.11.055.

1071	Fornaciari, E., Giusberti, L., Luciani, V., Tateo, F., Agnini, C., Backman, J., Oddone, M., Rio,
1072	D., 2007. An expanded Cretaceous-Tertiary transition in a pelagic setting of the Southern
1073	Alps (central-western Tethys). Palaeogeogr. Palaeoclimatol. Palaeoecol. 255, 98-131.
1074	https://doi.org/10.1016/j.palaeo.2007.02.044.
1075	Gallala, N., Zaghbib-Turki, D., Arenillas, I., Arz, J.A., Molina, E., 2009. Catastrophic mass
1076	extinction and assemblage evolution in planktic foraminifera across the
1077	Cretaceous/Paleogene (K/Pg) boundary at Bidart (SW France). Mar. Micropaleontol. 72,
1078	196-209. https://doi.org/10.1016/j.marmicro.2009.05.001.
1079	Gardin, S., 2002. Late Maastrichtian to early Danian calcareous nannofossils at Elles
1080	(Northwest Tunisia). A tale of one million years across the K-T boundary. Palaeogeogr.
1081	Palaeoclimatol. Palaeoecol. 178, 211-231. https://doi.org/10.1016/S0031-0182(01)00397-
1082	2.
1083	Gardin, S., Monechi, S., 1998. Palaeoecological change in middle to low latitude calcareous
1084	nannoplankton at the Cretaceous/Tertiary boundary. Bull. Soc. géol. Fr. 169, 709-723.
1085	Gerstel, J., Thunell, R.C., Zachos, J.C., Arthur, M.A., 1986. The Cretaceous/Tertiary boundary
1086	event in the North Pacific: Planktonic foraminiferal results from Deep Sea Drilling Project
1087	Site 577, Shatsky Rise. Paleoceanography 1, 97-117.
1088	https://doi.org/10.1029/PA001i002p00097.
1089	Gertsch, B., Keller, G., Adatte, T., Garg, R., Prasad, V., Berner, Z., Fleitmann, D., 2011.

- 1090 Environmental effects of Deccan volcanism across the Cretaceous-Tertiary transition in
- 1091 Meghalaya, India. Earth Planet. Sci. Lett. 310, 272-285.
- 1092 https://doi.org/10.1016/j.epsl.2011.08.015.
- 1093 Gibbs, S.J., Bown, P.R., Ward, B.A., Alvarez, S.A., Kim, H., Archontikis, O.A., Sauterey, B.,
- 1094 Poulton, A.J., Wilson, J., Ridgwell, A., 2020. Algal plankton turn to hunting to survive

- 1095 and recover from end-Cretaceous impact darkness. Sci. Adv. 6, eabc9123.
- 1096 https://doi.org/10.1126/sciadv.abc9123.
- 1097 Gilabert, V., Arz, J.A., Arenillas, I., Robinson, S.A., 2021. Influence of the Latest Maastrichtian
- 1098 Warming Event on planktic foraminiferal assemblages and ocean carbonate saturation at
- 1099 Caravaca, Spain. Cretac. Res. 125, 104844. https://doi.org/10.1016/j.cretres.2021.104844
- 1100 Groot, J.J., de Jonge, R.B.G., Langereis, C.G., ten Kate, W.G.H.Z., Smit, J., 1989.
- Magnetostratigraphy of the Cretaceous-Tertiary boundary at Agost (Spain). Earth Planet.
  Sci. Lett. 94, 385-397. https://doi.org/10.1016/0012-821X(89)90155-6.
- 1103 Gulick, S.P.S., Bralower, T.J., Ormö, J., Hall, B., Grice, K., Schaefer, B., Lyons, S., Freeman,
- 1104 K.H., Morgan, J. V., Artemieva, N., Kaskes, P., De Graaff, S.J., Whalen, M.T., Collins,
- 1105 G.S., Tikoo, S.M., Verhagen, C., Christeson, G.L., Claeys, P., Coolen, M.J.L., Goderis, S.,
- 1106 Goto, K., Grieve, R.A.F., McCall, N., Osinski, G.R., Rae, A.S.P., Riller, U., Smit, J.,
- 1107 Vajda, V., Wittmann, A., 2019. The first day of the Cenozoic. Proc. Natl. Acad. Sci. USA
- 1108 116, 19342-19351. https://doi.org/10.1073/pnas.1909479116.
- 1109 Hammer, Ø., Harper, D.A.T., Ryan, P.D., 2001. PAST: Paleontological statistics software
- 1110 package for education and data analysis. Paleontol. Electron. 4 (1), 9.
- Henehan, M.J., Hull, P.M., Penman, D.E., Rae, J.W.B., Schmidt, D.N., 2016. Biogeochemical
  significance of pelagic ecosystem function: An end-Cretaceous case study. Philos. Trans.
- 1113 R. Soc. B Biol. Sci. 371. https://doi.org/10.1098/rstb.2015.0510.
- 1114 Henehan, M.J., Ridgwell, A., Thomas, E., Zhang, S., Alegret, L., Schmidt, D.N., Rae, J.W.B.,
- 1115 Witts, J.D., Landman, N.H., Greene, S.E., Huber, B.T., Super, J.R., Planavsky, N.J., Hull,
- 1116 P.M., 2019. Rapid ocean acidification and protracted Earth system recovery followed the
- 1117 end-Cretaceous Chicxulub impact. Proc. Natl. Acad. Sci. USA 116, 22500-22504.
- 1118 https://doi.org/10.1073/pnas.1905989116.

1119	Hernandez Nava, A., Black, B.A., Gibson, S.A., Bodnar, R.J., Renne, P.R., Vanderkluysen, L.,
1120	2021. Reconciling early Deccan Traps CO <sub>2</sub> outgassing and pre-KPB global climate. Proc.
1121	Natl. Acad. Sci. USA 118, e2007797118. https://doi.org/10.1073/pnas.2007797118.
1122	Hildebrand, A.R., Penfield, G.T., Kring, D.A., Pilkington, M., Antonio, C.Z., Boynton, W. V.,
1123	1991. Chicxulub crater: A possible Cretaceous/Tertiary boundary impact crater on the
1124	Yucatan peninsula, Mexico. Geology 19, 867-871. https://doi.org/10.1130/0091-

- 1125 7613(1991)019<0867:CCAPCT>2.3.CO;2.
- 1126 Huber, B.T., MacLeod, K.G., Norris, R.D., 2002. Abrupt extinction and subsequent reworking

1127 of Cretaceous planktonic foraminifera across the Cretaceous-Tertiary boundary: Evidence

from the subtropical North Atlantic. Geol. Soc. Am. Spec. Pap. 356, 277-289.

1129 https://doi.org/10.1130/0-8137-2356-6.277.

- 1130 Hull, P.M., Bornemann, A., Penman, D.E., Henehan, M.J., Norris, R.D., Wilson, P.A., Blum,
- 1131 P., Alegret, L., Batenburg, S.J., Bown, P.R., Bralower, T.J., Cournede, C., Deutsch, A.,
- 1132 Donner, B., Friedrich, O., Jehle, S., Kim, H., Kroon, D., Lippert, P.C., Loroch, D.,
- 1133 Moebius, I., Moriya, K., Peppe, D.J., Ravizza, G.E., Röhl, U., Schueth, J.D., Sepúlveda, J.,
- 1134 Sexton, P.F., Sibert, E.C., Śliwińska, K.K., Summons, R.E., Thomas, E., Westerhold, T.,
- 1135 Whiteside, J.H., Yamaguchi, T., Zachos, J.C., 2020. On impact and volcanism across the
- 1136 Cretaceous-Paleogene boundary. Science 367, 266-272.
- 1137 https://doi.org/10.1126/science.aay5055.

1138 Jay, A.E., Widdowson, M., 2008. Stratigraphy, structure and volcanology of the SE Deccan

- 1139 continental flood basalt province: Implications for eruptive extent and volumes. J. Geol.
- 1140 Soc. London. 165, 177-188. https://doi.org/10.1144/0016-76492006-062.
- 1141 Jiang, S., Bralower, T.J., Patzkowsky, M.E., Kump, L.R., Schueth, J.D., 2010. Geographic
- 1142 controls on nannoplankton extinction across the Cretaceous/Palaeogene boundary. Nat.
- 1143 Geosci. 3, 280-285. https://doi.org/10.1038/ngeo775.

- 1144 Jiang, S., Chen, X., Bernaola, G., 2019. Environmental controls on calcareous nannoplankton
- 1145 response to the Cretaceous/Paleogene mass extinction in the Tethys realm. Palaeogeogr.

1146 Palaeoclimatol. Palaeoecol. 515, 134-142. https://doi.org/10.1016/j.palaeo.2017.12.044.

- Jones, H.L., Lowery, C.M., Bralower, T.J., 2019. Delayed calcareous nannoplankton boom-bust
  successions in the earliest Paleocene Chicxulub (Mexico) impact crater. Geology 47, 753-
- 1149 756. https://doi.org/10.1130/G46143.1.
- 1150 Kaiho, K., Lamolda, M.A., 1999. Catastrophic extinction of planktonic foraminifera at the
- 1151 Cretaceous-Tertiary boundary evidenced by stable isotopes and foraminiferal abundance at
- 1152 Caravaca, Spain. Geology 27, 355-358. https://doi.org/10.1130/0091-
- 1153 7613(1999)027<0355:CEOPFA>2.3.CO;2.
- 1154 Kaiho, K., Kajiwara, Y., Tazaki, K., Ueshima, M., Takeda, N., Kawahata, H., Arinobu, T.,
- 1155 Ishiwatari, R., Hirai, A., Lamolda, M.A., 1999. Oceanic primary productivity and
- dissolved oxygen levels at the Cretaceous/Tertiary boundary: Their decrease, subsequent
- 1157 warming, and recovery. Paleoceanography 14, 511-524.
- 1158 https://doi.org/10.1029/1999PA900022.
- 1159 Kasbohm, J., Schoene, B., Burgess, S., 2021. Radiometric constraints on the timing, tempo, and
- 1160 effects of large igneous province emplacement, in: Ernst, R.E., Dickson, J., Bekker, A.
- 1161 (Eds.), Large Igneous Provinces: A Driver of Global Environmental and Biotic Changes.
- 1162 American Geophysical Union. Carbon in Earth's Interior published by John Wiley & Sons,
- 1163 Inc., pp. 27-82. https://doi.org/https://doi.org/10.1002/9781119507444.ch2.
- 1164 Kawaragi, K., Sekine, Y., Kadono, T., Sugita, S., Ohno, S., Ishibashi, K., Kurosawa, K., Matsui,
- 1165 T., Ikeda, S., 2009. Direct measurements of chemical composition of shock-induced gases
- 1166 from calcite: an intense global warming after the Chicxulub impact due to the indirect
- 1167 greenhouse effect of carbon monoxide. Earth Planet. Sci. Lett. 282, 56-64.
- 1168 https://doi.org/10.1016/j.epsl.2009.02.037.

- 1169 Keller, G., 2003. *Guembelitria* dominated late Maastrichtian planktic foraminiferal
- assemblages mimic early Danian in central Egypt. Mar. Micropaleontol. 47, 71-99.
- 1171 https://doi.org/10.1016/S0377-8398(02)00116-0.
- Keller, G., Pardo, A., 2004. Disaster opportunists Guembelitrinidae: Index for environmental
  catastrophes. Mar. Micropaleontol. 53, 83-116.
- 1174 https://doi.org/10.1016/j.marmicro.2004.04.012.
- 1175 Keller, G., Adatte, T., Gardin, S., Bartolini, A., Bajpai, S., 2008. Main Deccan volcanism phase
- 1176 ends near the K-T boundary: Evidence from the Krishna-Godavari Basin, SE India. Earth

1177 Planet. Sci. Lett. 268, 293-311. https://doi.org/10.1016/j.epsl.2008.01.015.

- 1178 Keller, G.K., Bhowmick, P.K., Upadhyay, H., Dave, A., Reddy, A.N., Jaiprakash, B.C., Adatte,
- 1179 T., 2011. Deccan volcanism linked to the Cretaceous-Tertiary boundary mass extinction:
- 1180 New evidence from ONGC wells in the Krishna-Godavari Basin. J. Geol. Soc. India 78,
- 1181 399-428. https://doi.org/10.1007/s12594-011-0107-3.
- 1182 Keller, G., Adatte, T., Bhowmick, P.K., Upadhyay, H., Dave, A., Reddy, A.N., Jaiprakash,
- 1183 B.C., 2012. Nature and timing of extinctions in Cretaceous-Tertiary planktic foraminifera
- 1184 preserved in Deccan intertrappean sediments of the Krishna-Godavari Basin, India. Earth
- 1185 Planet. Sci. Lett. 341-344, 211-221. https://doi.org/10.1016/j.epsl.2012.06.021.
- 1186 Keller, G., Punekar, J., Mateo, P., 2016. Upheavals during the Late Maastrichtian: Volcanism,
- 1187 climate and faunal events preceding the end-Cretaceous mass extinction. Palaeogeogr.
- 1188 Palaeoclimatol. Palaeoecol. 441, 137-151. https://doi.org/10.1016/j.palaeo.2015.06.034.
- 1189 Keller, G., Mateo, P., Monkenbusch, J., Thibault, N., Punekar, J., Spangenberg, J.E.,
- 1190 Abramovich, S., Ashckenazi-Polivoda, S., Schoene, B., Eddy, M.P., Samperton, K.M.,
- 1191 Khadri, S.F.R., Adatte, T., 2020. Mercury linked to Deccan Traps volcanism, climate

- change and the end-Cretaceous mass extinction. Glob. Planet. Change 194, 103312.
- 1193 https://doi.org/10.1016/j.gloplacha.2020.103312.
- Kelly, D.C., Norris, R.D., Zachos, J.C., 2003. Deciphering the paleoceanographic significance
  of Early Oligocene *Braarudosphaera* chalks in the South Atlantic. Mar. Micropaleontol.
- 1196 49, 49-63. https://doi.org/10.1016/S0377-8398(03)00027-6.
- 1197 Krahl, G., Bom, M.H.H., Kochhann, K.G.D., Souza, L. V., Savian, J.F., Fauth, G., 2020.
- 1198 Environmental changes occurred during the early Danian at the Rio Grande Rise, South
- 1199 Atlantic Ocean. Glob. Planet. Change 191, 103197.
- 1200 https://doi.org/10.1016/j.gloplacha.2020.103197.
- 1201 Kring, D.A., 2007. The Chicxulub impact event and its environmental consequences at the
- 1202 Cretaceous-Tertiary boundary. Palaeogeogr. Palaeoclimatol. Palaeoecol. 255, 4-21.
- 1203 https://doi.org/10.1016/j.palaeo.2007.02.037.
- 1204 Kroon, D., Nederbragt, A.J., 1990. Ecology and paleoecology of triserial planktic foraminifera.
  1205 Mar. Micropaleontol. 16, 25-38. https://doi.org/10.1016/0377-8398(90)90027-J.
- 1206 Kucera, M., Malmgren, B.A., Sturesson, U., 1997. Foraminiferal dissolution at shallow depths
- 1207 of the Walvis Ridge and Rio Grande Rise during the latest Cretaceous: Inferences for
- deep-water circulation in the South Atlantic. Palaeogeogr. Palaeoclimatol. Palaeoecol.
- 1209 129, 195-212. https://doi.org/10.1016/S0031-0182(96)00133-2.
- 1210 Lamolda, M.A., Melinte, M.C., Kaiho, K., 2005. Nannofloral extinction and survivorship across
- 1211 the K/T boundary at Caravaca, southeastern Spain. Palaeogeogr. Palaeoclimatol.
- 1212 Palaeoecol. 224, 27-52. https://doi.org/10.1016/j.palaeo.2005.03.030.
- 1213 Lamolda, M.A., Melinte-Dobrinescu, M.C., Kaiho, K., 2016. Calcareous nannoplankton
- 1214 assemblage changes linked to paleoenvironmental deterioration and recovery across the

- 1215 Cretaceous-Paleogene boundary in the Betic Cordillera (Agost, Spain). Palaeogeogr. 1216 Palaeoclimatol. Palaeoecol. 441, 438-452. https://doi.org/10.1016/j.palaeo.2015.10.003. 1217 Legendre, P., Gallagher, E.D., 2001. Ecologically meaningful transformations for ordination of 1218 species data. Oecologia 129, 271-280. https://doi.org/10.1007/s004420100716. 1219 Lowery, C.M., Fraass, A.J., 2019. Morphospace expansion paces taxonomic diversification 1220 after end Cretaceous mass extinction. Nat. Ecol. Evol. 3, 900-904. 1221 https://doi.org/10.1038/s41559-019-0835-0. 1222 Lowery, C.M., Bralower, T.J., Owens, J.D., Rodríguez-Tovar, F.J., Jones, H., Smit, J., Whalen, 1223 M.T., Claeys, P., Farley, K., Gulick, S.P.S., Morgan, J. V., Green, S., Chenot, E., 1224 Christeson, G.L., Cockell, C.S., Coolen, M.J.L., Ferrière, L., Gebhardt, C., Goto, K., 1225 Kring, D.A., Lofi, J., Ocampo-Torres, R., Perez-Cruz, L., Pickersgill, A.E., Poelchau, 1226 M.H., Rae, A.S.P., Rasmussen, C., Rebolledo-Vieyra, M., Riller, U., Sato, H., Tikoo, 1227 S.M., Tomioka, N., Urrutia-Fucugauchi, J., Vellekoop, J., Wittmann, A., Xiao, L., 1228 Yamaguchi, K.E., Zylberman, W., 2018. Rapid recovery of life at ground zero of the end-
  - 1229 Cretaceous mass extinction. Nature 558, 288-291. <u>https://doi.org/10.1038/s41586-018-</u>
  - <u>0163-6</u>.
  - 1231 Lowery, C.M., Bown, P.R., Fraass, A.J., Hull, P.M., 2020. Ecological Response of Plankton to

1232 Environmental Change: Thresholds for Extinction. Annu. Rev. Earth Planet. Sci. 48.

1233 https://doi.org/10.1146/annurev-earth-081619-052818.

- 1234 Luciani, V., D'Onofrio, R., Filippi, G., Moretti, S., 2020. Which was the habitat of early Eocene
- 1235 planktic foraminifer *Chiloguembelina*? Stable isotope paleobiology from the Atlantic
- 1236 Ocean and implication for paleoceanographic reconstructions. Glob. Planet. Change 191,
- 1237 103216. https://doi.org/10.1016/j.gloplacha.2020.103216.

- 1238 Lyson, T.R., Miller, I.M., Bercovici, A.D., Weissenburger, K., Fuentes, A.J., Clyde, W.C.,
- 1239 Hagadorn, J.W., Butrim, M.J., Johnson, K.R., Fleming, R.F., Barclay, R.S., Maccracken,
- 1240 S.A., Lloyd, B., Wilson, G.P., Krause, D.W., Chester, S.G.B., 2019. Exceptional
- 1241 continental record of biotic recovery after the Cretaceous–Paleogene mass extinction.
- 1242 Science 366, 977-983. https://doi.org/10.1126/science.aay2268.
- 1243 Mateo, P., Keller, G., Adatte, T., Spangenberg, J.E., 2016. Mass wasting and hiatuses during the
- 1244 Cretaceous-Tertiary transition in the North Atlantic: Relationship to the Chicxulub

1245 impact? Palaeogeogr. Palaeoclimatol. Palaeoecol. 441, 96-115.

1246 https://doi.org/10.1016/j.palaeo.2015.01.019.

- 1247 Metsana-Oussaid, F., Belhai, D., Arenillas, I., Arz, J.A., Gilabert, V., 2019. New sections of the
- 1248 Cretaceous-Paleogene transition in the southwestern Tethys (Médéa, northern Algeria):
- 1249 planktic foraminiferal biostratigraphy and biochronology. Arab. J. Geosci. 12.
- 1250 https://doi.org/10.1007/s12517-019-4402-4.
- 1251 Minoletti, F., De Rafelis, M., Renard, M., Gardin, S., Young, J., 2005. Changes in the pelagic
- 1252 fine fraction carbonate sedimentation during the Cretaceous-Paleocene transition:
- 1253 Contribution of the separation technique to the study of Bidart section. Palaeogeogr.
- 1254 Palaeoclimatol. Palaeoecol. 216, 119-137. https://doi.org/10.1016/j.palaeo.2004.10.006.
- Molina, E., Arenillas, I., Arz, J.A. 1998. Mass extinction in planktic foraminifera at the
  Cretaceous/Tertiary boundary in subtropical and temperate latitudes. Bull. Soc. géol. Fr.
  1257 169(3), 351-363.
- 1258 Molina, E., Alegret, L., Arenillas, I., Arz, J.A., Gallala, N., Grajales-Nishimura, J.M., Murillo-
- 1259 Muñetón, G., Zaghbib-Turki, D., 2009. The Global Boundary Stratotype Section and Point
- 1260 for the base of the Danian Stage (Paleocene, Paleogene, "Tertiary", Cenozoic): Auxiliary
- sections and correlation. Episodes 32, 84-95.
- 1262 https://doi.org/10.18814/epiiugs/2009/v32i2/002.

- 1263 Mukhopadhyay, S., Farley, KA., Montanari, A., 2001. A short duration of the Cretaceous-
- 1264 Tertiary boundary event: Evidence from extraterrestrial Helium<sup>3</sup>. Science 291, 1952-1955.
  1265 https://doi.org/10.1126/science.291.5510.1952
- 1266 Norris, R.D., 1996. Symbiosis as an evolutionary innovation in the radiation of Paleocene
- 1267 planktic foraminifera. Paleobiology 22, 461-480.
- 1268 https://doi.org/10.1017/S0094837300016468.
- 1269 Olsson, R.K., Berggren, W.A., Hemleben, C., Huber, B.T., 1999. Atlas of Paleocene Planktonic
- 1270 Foraminifera. Smithson. Contrib. to Paleobiol. 85, 1-252.
- 1271 https://doi.org/10.5479/si.00810266.85.1.
- 1272 Omaña, L., Alencáster, G., Hernández, J.R.T., Doncel, R.L., 2012. Morphological abnormalities
- 1273 and dwarfism in Maastrichtian foraminifera from the Cárdenas Formation, Valles-San
- 1274 Luis Potosí Platform, Mexico: Evidence of paleoenvironmental stress. Bol. Soc. Geol.
- 1275 Mex. 64, 305-318. https://doi.org/10.18268/BSGM2012v64n3a4.
- 1276 Pardo, A., Keller, G., 2008. Biotic effects of environmental catastrophes at the end of the
- 1277 Cretaceous and early Tertiary: *Guembelitria* and *Heterohelix* blooms. Cretac. Res. 29,
- 1278 1058-1073. https://doi.org/10.1016/j.cretres.2008.05.031.
- Pospichal, J.J., 1996. Calcareous nannoplankton mass extinction at the Cretaceous/Tertiary
  boundary: An update. Geol. Soc. Am. Spec. Pap. 307, 335-360. https://doi.org/10.1130/08137-2307-8.335.
- 1282 Premović, P.I., 2009. Experimental evidence for the global acidification of surface ocean at the
- 1283 Cretaceous-Palaeogene boundary: The biogenic calcite-poor spherule layers. Int. J.
- 1284 Astrobiol. 8, 193-206. https://doi.org/10.1017/S1473550409990139.

- Punekar, J., Mateo, P., Keller, G., 2014a. Effects of Deccan volcanism on paleoenvironment
  and planktic foraminifera: A global survey. Spec. Geol. Soc. Am. Spec. Pap. 505, 91-116.
  https://doi.org/10.1130/2014.2505(04).
- 1288 Punekar, J., Keller, G., Khozyem, H., Hamming, C., Adatte, T., Tantawy, A.A., Spangenberg,
- 1289 J.E., 2014b. Late Maastrichtian-early Danian high-stress environments and delayed
- recovery linked to Deccan volcanism. Cretac. Res. 49, 63-82.
- 1291 https://doi.org/10.1016/j.cretres.2014.01.002.
- 1292 Punekar, J., Keller, G., Khozyem, H.M., Adatte, T., Font, E., Spangenberg, J., 2016. A multi-
- 1293 proxy approach to decode the end-Cretaceous mass extinction. Palaeogeogr.
- 1294 Palaeoclimatol. Palaeoecol. 441, 116-136. https://doi.org/10.1016/j.palaeo.2015.08.025.
- 1295 Quillévéré, F., Norris, R.D., Kroon, D., Wilson, P.A., 2008. Transient ocean warming and shifts
- in carbon reservoirs during the early Danian. Earth Planet. Sci. Lett. 265, 600-615.
  https://doi.org/10.1016/j.epsl.2007.10.040.
- 1298 Renne, P.R., Sprain, C.J., Richards, M.A., Self, S., Vanderkluysen, L., Pande, K., 2015. State
- 1299 shift in Deccan volcanism at the Cretaceous-Paleogene boundary, possibly induced by

1300 impact. Science 350, 76-78. https://doi.org/10.1126/science.aac7549.

- 1301 Renne, P.R., Arenillas, I., Arz, J.A., Vajda, V., Gilabert, V., Bermúdez, H.D., 2018. Multi-
- 1302 proxy record of the Chicxulub impact at the Cretaceous- Paleogene boundary from
- 1303 Gorgonilla Island, Colombia. Geology 46, 547-550. <u>https://doi.org/10.1130/G40224.1</u>.
- 1304 Ricotta, C., Podani, J., 2017. On some properties of the Bray-Curtis dissimilarity and their
- ecological meaning. Ecol. Complex. 31, 201-205.
- 1306 https://doi.org/10.1016/j.ecocom.2017.07.003.
- 1307 Richards, M.A., Alvarez, W., Self, S., Karlstrom, L., Renne, P.R., Manga, M., Sprain, C.J.,
- 1308 Smit, J., Vanderkluysen, L., Gibson, S.A., 2015. Triggering of the largest Deccan

- 1309 eruptions by the Chicxulub impact. Bull. Geol. Soc. Am. 127, 1507-1520.
- 1310 https://doi.org/10.1130/B31167.1.
- 1311 Robinson, N., Ravizza, G., Coccioni, R., Peucker-Ehrenbrink, B., Norris, R., 2009. A high-
- resolution marine <sup>187</sup>Os/<sup>188</sup>Os record for the late Maastrichtian: Distinguishing the
- 1313 chemical fingerprints of Deccan volcanism and the K-P impact event. Earth Planet. Sci.
- 1314 Lett. 281, 159-168. https://doi.org/10.1016/j.epsl.2009.02.019.
- Romein, A.J.T., 1977. Calcareous nannofossils from the Cretaceous/Tertiary boundary interval
  in the Barranco del Gredero (Caravaca, Prov. Murcia, SE Spain). Proc. K. Ned. Akad.
- 1317 Wet. Ser. B 80, 256-279.
- 1318 Schmidt, A., Skeffington, R.A., Thordarson, T., Self, S., Forster, P.M., Rap, A., Ridgwell, A.,
- 1319 Fowler, D., Wilson, M., Mann, G.W., Wignall, P.B., Carslaw, K.S., 2016. Selective
- 1320 environmental stress from sulphur emitted by continental flood basalt eruptions. Nat.
- 1321 Geosci. 9, 77-82. https://doi.org/10.1038/ngeo2588.
- 1322 Schoene, B., Samperton, K.M., Eddy, M.P., Keller, G., Adatte, T., Bowring, S.A., Khadri,
- 1323 S.F.R., Gertsch, B., 2015. U-Pb geochronology of the Deccan Traps and relation to the
- end-Cretaceous mass extinction. Science 347, 182-184.
- 1325 https://doi.org/10.1126/science.aaa0118.
- 1326 Schoene, B., Eddy, M.P., Samperton, K.M., Keller, C.B., Keller, G., Adatte, T., Khadri, S.F.R.,
- 13272019. U-Pb constraints on pulsed eruption of the Deccan Traps across the end-Cretaceous
- 1328 mass extinction. Science 363, 862-866. https://doi.org/10.1126/science.aau2422.
- 1329 Schoene, B., Eddy, M. P., Keller, C. B., and Samperton, K. M., 2021. An evaluation of Deccan
- 1330 Traps eruption rates using geochronologic data: Geochronology 3, 181–198.
- 1331 https://doi.org/10.5194/gchron-3-181-2021.

- 1332 Schulte, P., Alegret, L., Arenillas, I., Arz, J.A., Barton, P.J., Bown, P.R., Bralower, T.J.,
- 1333 Christeson, G.L., Claeys, P., Cockell, C.S., Collins, G.S., Deutsch, A., Goldin, T.J., Goto,
- 1334 K., Grajales-Nishimura, J.M., Grieve, R.A.F., Gulick, S.P.S., Johnson, K.R., Kiessling,
- 1335 W., Koeberl, C., Kring, D.A., MacLeod, K.G., Matsui, T., Melosh, J., Montanari, A.,
- 1336 Morgan, J. V., Neal, C.R., Nichols, D.J., Norris, R.D., Pierazzo, E., Ravizza, G.,
- 1337 Rebolledo-Vieyra, M., Reimold, W.U., Robin, E., Salge, T., Speijer, R.P., Sweet, A.R.,
- 1338 Urrutia-Fucugauchi, J., Vajda, V., Whalen, M.T., Willumsen, P.S., 2010. The Chicxulub
- asteroid impact and mass extinction at the Cretaceous-Paleogene boundary. Science 327,
- 1340 1214-1218. https://doi.org/10.1126/science.1177265.
- 1341 Self, S., Widdowson, M., Thordarson, T., Jay, A.E., 2006. Volatile fluxes during flood basalt
- eruptions and potential effects on the global environment: A Deccan perspective. Earth
  Planet. Sci. Lett. 248, 518-532. https://doi.org/10.1016/j.epsl.2006.05.041.
- 1344 Self, S., Jay, A.E., Widdowson, M., Keszthelyi, L.P., 2008. Correlation of the Deccan and
- 1345 Rajahmundry Trap lavas: Are these the longest and largest lava flows on Earth? J.
- 1346 Volcanol. Geotherm. Res. 172, 3-19. https://doi.org/10.1016/j.jvolgeores.2006.11.012.
- 1347 Sepúlveda, J., Wendler, J.E., Summons, R.E., Hinrichs, K.U., 2009. Rapid resurgence of marine
- 1348 productivity after the Cretaceous-Paleogene mass extinction. Science 326, 129-132.
- 1349 https://doi.org/10.1126/science.1176233.
- 1350 Sepúlveda, J., Alegret, L., Thomas, E., Haddad, E., Cao, C., Summons, R.E., 2019. Stable
- 1351 Isotope Constraints on Marine Productivity Across the Cretaceous-Paleogene Mass
- 1352 Extinction. Paleoceanogr. Paleoclimatology 34, 1195-1217.
- 1353 https://doi.org/10.1029/2018PA003442.
- 1354 Sinnesael, M., Montanari, A., Coccioni, R., Frontalini, F., Gattacceca, J., Snoeck, C., Wegner,
- 1355 W., Koeberl, C., Morgan, L.E., De Winter, N.J., DePaolo, D.J., Claeys, P., 2019.
- 1356 Multiproxy Cretaceous-Paleogene boundary event stratigraphy: An Umbria-Marche

- 1357 basinwide perspective. Geol. Soc. Am. Spec. Pap. 542, 133-158.
- 1358 https://doi.org/10.1130/2019.2542(07).
- Smit, J., 1982. Extinction and evolution of planktonic foraminifera after a major impact at the
  Cretaceous/Tertiary boundary. Geol. Soc. Am. Spec. Pap. 190, 329-352.
- 1361 https://doi.org/10.1130/SPE190-p329.
- 1362 Smit, J., 1999. The global stratigraphy of the Cretaceous-Tertiary boundary impact ejecta.
- 1363 Annu. Rev. Earth Planet. Sci. 27, 75-113. https://doi.org/10.1146/annurev.earth.27.1.75.
- 1364 Smit, J., 2004. The section of the Barranco del Gredero (Caravaca, SE Spain): A crucial section
- 1365 for the Cretaceous/Tertiary boundary impact extinction hypothesis. J. Iber. Geol. 31, 179-
- 1366 191. https://doi.org/10.5209/JIGE.33967.
- Smit, J., Hertogen, J., 1980. An extraterrestrial event at the Cretaceous-Tertiary boundary.
  Nature 285, 198-200. https://doi.org/10.1038/285198a0.
- 1369 Smit, J., Romein, A.J.T., 1985. A sequence of events across the Cretaceous-Tertiary boundary.
- 1370 Earth Planet. Sci. Lett. 74, 155-170. https://doi.org/10.1016/0012-821X(85)90019-6.
- 1371 Sosa-Montes de Oca, C., Rodríguez-Tovar, F.J., Martínez-Ruiz, F., 2016. Geochemical and
- 1372 isotopic characterization of trace fossil infillings: New insights on tracemaker activity after
- 1373 the K/Pg impact event. Cretac. Res. 57, 391-401.
- 1374 https://doi.org/10.1016/j.cretres.2015.03.003.
- 1375 Sprain, C.J., Renne, P.R., Clemens, W.A., Wilson, G.P., 2018. Calibration of chron C29r: New
- 1376 high-precision geochronologic and paleomagnetic constraints from the Hell Creek region,
- 1377 Montana. Bull. Geol. Soc. Am. 130, 1615-1644. https://doi.org/10.1130/B31890.1.

- 1378 Sprain, C.J., Renne, P.R., Vanderkluysen, L., Pande, K., Self, S., Mittal, T., 2019. The eruptive
- 1379 tempo of Deccan volcanism in relation to the Cretaceous-Paleogene boundary. Science.

1380 363, 866-870. https://doi.org/10.1126/science.aav1446.

- 1381 Stax, R., Stein, R., 1993. Long-term changes in the accumulation of organic carbon in Neogene
- 1382 sediments, Ontong Java Plateau. Proc., Sci. results, ODP, Leg 130, Ontong Java Plateau
- 1383 130, 573-584. https://doi.org/10.2973/odp.proc.sr.130.039.1993.
- 1384 Steinthorsdottir, M., Vajda, V., Pole, M., 2016. Global trends of pCO<sub>2</sub> across the Cretaceous-
- 1385 Paleogene boundary supported by the first Southern Hemisphere stomatal proxy-based
- 1386 pCO<sub>2</sub> reconstruction. Palaeogeogr. Palaeoclimatol. Palaeoecol. 464, 143-152.
- 1387 https://doi.org/10.1016/j.palaeo.2016.04.033.
- Thibault, N., Gardin, S., 2010. The calcareous nannofossil response to the end-Cretaceous warm
  event in the Tropical Pacific. Palaeogeogr. Palaeoclimatol. Palaeoecol. 291, 239-252.
  https://doi.org/10.1016/j.palaeo.2010.02.036.
- 1391 Thibault, N., Galbrun, B., Gardin, S., Minoletti, F., Le Callonnec, L., 2016. The end-Cretaceous
- in the southwestern Tethys (Elles, Tunisia): orbital calibration of paleoenvironmental
- events before the mass extinction. Int. J. Earth Sci. 105, 771-795.
- 1394 https://doi.org/10.1007/s00531-015-1192-0.
- 1395 Thibault, N., Minoletti, F., Gardin, S., 2018. Offsets in the early Danian recovery phase in
- 1396 carbon isotopes: Evidence from the biometrics and phylogeny of the *Cruciplacolithus*
- 1397 lineage. Rev. Micropaleontol. 61, 207-221. https://doi.org/10.1016/j.revmic.2018.09.002.
- 1398 Tobin, T.S., Bitz, C.M., Archer, D., 2017. Modeling climatic effects of carbon dioxide
- emissions from Deccan Traps volcanic eruptions around the Cretaceous-Paleogene
- boundary. Palaeogeogr. Palaeoclimatol. Palaeoecol. 478, 139-148.
- 1401 https://doi.org/10.1016/j.palaeo.2016.05.028.

1402	Vandamme, D., Courtillot, V., Besse, J., Montigny, R., 1991. Paleomagnetism and age
1403	determinations of the Deccan traps (India): results of a Nagpur-Bombay traverse and
1404	review of earlier work. Rev. Geophys. Space Phys. 29, 159–190.
1405	Vellekoop, J., Sluijs, A., Smit, J., Schouten, S., Weijers, J.W.H., Sinninghe Damsté, J.S.,
1406	Brinkhuis, H., 2014. Rapid short-term cooling following the Chicxulub impact at the
1407	Cretaceous-Paleogene boundary. Proc. Natl. Acad. Sci. USA 111, 7537-7541.
1408	https://doi.org/10.1073/pnas.1319253111.
1409	Vellekoop, J., Esmeray-Senlet, S., Miller, K.G., Browning, J. V, Sluijs, A., van de
1410	Schootbrugge, B., Sinninghe Damsté, J.S., Brinkhuis, H., 2016. Evidence for Cretaceous-
1411	Paleogene boundary bolide "impact winter" conditions from New Jersey, USA. Geology
1412	44. https://doi.org/10.1130/G37961.1.
1413	Vellekoop, J., Woelders, L., van Helmond, N.A.G.M., Galeotti, S., Smit, J., Slomp, C.P.,
1414	Brinkhuis, H., Claeys, P., Speijer, R.P., 2018. Shelf hypoxia in response to global warming

after the Cretaceous-Paleogene boundary impact. Geology 46, 683-686. 1415

1416 https://doi.org/10.1130/G45000.1.

- 1417 Wade, B.S., Pearson, P.N., Berggren, W.A., Pälike, H., 2011. Review and revision of Cenozoic
- 1418 tropical planktonic foraminiferal biostratigraphy and calibration to the geomagnetic
- 1419 polarity and astronomical time scale. Earth-Science Rev. 104, 111-142.
- 1420 https://doi.org/10.1016/j.earscirev.2010.09.003.