- 1 Mercury enrichment and Hg isotopes in CretaceousePaleogene boundary successions:
- 2 Links to volcanism and palaeoenvironmental impacts
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24 Abstract

- We investigate the use of Hg as a proxy for volcanism by studying four distal and two proximal
- sections in relation to the Deccan volcanic center, straddling the CretaceousePaleogene (KPg)
- boundary at (a) Højerup (Denmark), Bottaccione and Padriciano (Italy), (b) Meghalaya and Jhilmili
- 28 (India), and (c) Bajada del Jagüel (Argentina). Hg sequestration by organic matter results in constant
- 29 Hg/TOC ratio and linear correlation between Hg content of the sediments and total organic carbon

(TOC). Elevated Hg concentrations that deviate from this linear relationship represent most likely true 30 Hg anomalies and these notable Hg/TOC spikes (all TOC <1%) are found in the Meghalaya, 31 Bottaccione and Højerup sections within the CF2 planktic foraminiferal biozone (spike I), at the KPg 32 boundary (spike II), and within the P1a planktic foraminiferal subzone (spike III). Spike III occurs also 33 in the Jhilmili section. No clear correlation between Hg/TOC and Al₂O₃ exists in any of the studied 34 sections. The Hg anomalies probably result from strong volcanic episodes of the Deccan phase-2 35 (started 250 kyr before the KPg boundary and lasted for 750 kyr) that exhaled sulfuric aerosols, carbon 36 dioxide and other toxic agents which reached a critical threshold, represented in true Hg enrichments in 37 the paleoenviron- ments. The possibility that Hg enrichments resulted from anoxia scavenging on the 38 seafloor and penetration downward into sediments is not supported in the stratigraphic record of Mo/Al 39 ratios redox proxy. Hg isotopes were analyzed in samples from all KPg boundary sections in this study 40 and from Bidart, France, the latter for comparison. Hg isotopes yielded δ^{202} Hg values ranging from 1 to 41 2% and Δ^{201} Hg signatures from 0 to 0.05% (spike II in Højerup, Bottaccione and Meghalaya KPg 42 boundary layers) consistent with volcanic emission of Hg (0 to 2‰). The δ^{202} Hg in spike I in 43 Meghalaya and Padriciano and spike III in Jhilmili is consistent with volcanic emission of Hg. Two 44 samples from Bajada del Jagüel and four from Bidart, however, display isotope signals compatible with 45 volcanic emission/chondrite Hg. The results of three other samples are characteristic for reworked 46 sediment, soil and/or peat. Most of the data show small positive Δ^{201} Hg, in favor of long-term 47 atmospheric transport prior to deposition, supporting a volcanic origin for the Hg. The present study 48 49 broadens, therefore, the potential use of Hg as stratigraphic marker and, moreover, confirms that in the critical KPg transition, Hg was enriched in paleoenvironments at three distinct stages during the 50 Deccan phase-2. 51

Cretaceous-Paleogene boundary, Chemostratigraphy, Total organic carbon, Molybdenum, Hg 53 isotopes 54

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1. Introduction

56 Volcanic emissions have the potential of injecting large amounts of Hg into the atmosphere and, 57 therefore, contribute with a sig- nificant natural input of Hg to the atmosphere (e.g. Schuster et al., 58 2002; Pyle and Mather, 2003). Many studies have reported Hg enrichments in sedimentary records that 59 are synchronous with modern (e.g. Martínez-Cortizas et al., 1999; Roos-Barraclough et al., 2002; 60 Roos-Barraclough and Shotyk, 2003) and prehistoric volcanic events (e.g. Palinkas et al., 1996; Sial et 61 al., 2010, 2013, 2014; Nascimento-Silva et al., 2011, 2013; Sanei et al., 2012; Grasby et al., 2013, 62 2015a, 2015b; Percival et al., 2015; Font et al., 2016). In contrast to most elements present in ash, Hg 63 derived from volcanic activity is mainly in gaseous form (Hg⁰), can be transported far in the 64 atmosphere, reaching even global-scale distribution prior to deposition in terrestrial and marine 65 environments and has a long atmospheric residence time (1e2 years). Through oxidation in the 66 atmosphere, Hg⁰ forms reactive Hg^{b2}, soluble in water and, there- fore, enriched in the rain (e.g. 67 Schroeder and Munthe, 1998; Witt et al., 2008 and references therein). Organic matter and clay min-68 erals scavenge Hg in the marine environment and fix it in bottom sediments on the sea floor (Grasby et 69 70 al., 2015a; Percival et al., 2015). If Hg reaches an environment with low organic scavenging capacity on the surface, then Hg^{b2} remains in solution and is, even-tually, adsorbed onto clays and transported 71 from land to sea. Therefore, high levels of Hg associated with argillaceous sediments can be explained 72 by increased flux of volcanogenic Hg from conti- nents into the oceans. Higher Hg accumulation rates 73 are typical and more pronounced in sediments deposited after glacial maxima, when runoff is 74 increased, compared to sediment layers deposited before such events. This peculiarity is perhaps a 75 phenomenon that can be observed globally as similar results have been observed in the Amazon region 76

(Santos et al., 2001), Antarctica (Vandal et al., 1993) and Europe (Martínez-Cortizas et al., 1999). 77 Biotic or abiotic reduction of Hg ²b to Hg⁰ (g) limits the scavenging and biological fixation of Hg from 78 79 the atmosphere and, in consequence, allows Hg re-emission and higher concentration in the atmosphere (Percival et al., 2015). Under reduced bioproductivity, Hg availability stops being captured by organic 80 matter which is one of the major Hg sinks (Sanei et al., 2012). A strong linear correlation between Hg 81 and total organic matter (TOC) contents in sediments and an apparent affinity of Hg for terrestrial 82 organic matter has been recognized (Outridge et al., 2007; Stern et al., 2009; Sanei et al., 2014) but the 83 importance of terrestrial versus aquatic retention of Hg is still an open question (Percival et al., 2015). 84 Increased Hg concentrations in sediments that deviate from a linear relationship between Hg and TOC 85 86 represent true Hg anomalies. On the other hand, Hg can be adsor- bed onto clays (Krupp, 1988). In this case one would expect a covariation between Hg and Al2O3, as for example observed in some sections 87 88 across the KPg boundary (e.g. Sial et al., 2013). It has also been suggested that Hg emission rates can be significantly enhanced in periods of large igneous province (LIP) volcanic activity when the marine 89 90 buffering control on Hg can be overwhelmed and thus generating Hg spikes in sediments (Sanei et al., 2012; Grasby et al., 2013, 2015a; Font et al., 2016). There- fore, Hg anomalies could serve as proxy for 91 periods of extensive volcanic activities when LIPs, for example, could have released toxic quantities of 92 Hg into environment. The Hg flux to the sedimentary realm accompanying large amounts of magmatic 93 sulfur and other toxic metals released by LIPs (e.g. Deccan Traps, Callegaro et al., 2014) can provide 94 missing links between the terrestrial and ma-rine records of biota extinctions as suggested by Grasby et 95 al. (2015a). Sanei et al. (2012) related enhanced atmospheric Hg depositions at the Permiane Triassic 96 transition to catastrophic volcanic erup- tions of the Siberian Trap LIP event, followed by the 97 discontinuity of the organic Hg fixation that lead to an increased dissolved Hg flux. The 98 Permiane Triassic biotic crisis, as well as the extinction events in the late Capitanian and at the 99 Smithian/Spathian boundary in NW Pangea, has been linked to enhanced Hg deposition (Grasby et al., 100

2013, 2015a, 2015b). In addition, the end-Pliensbachian extinction and Toarcian oceanic anoxic event 101 102 (OAE) are probably related to LIP activities (KarooeFerrar), according to Percival et al. (2015). Acidic rain has been deemed responsible for the mass extinction associated to the KPg boundary (Hsü and 103 McKenzie, 1985) and Hg enrichments in KPg boundary sedimentary rocks have been regarded as an 104 evidence for this type of rain (Hildebrand and Boynton, 1989). Anomalous Hg concentrations in 105 deposits spanning the KPg boundary in Dolenja Vas, Slovenia, likely resulted from sub-aerial volcanic 106 activity (Palinkas et al., 1996). Subsequent work suggested a possible connection between enhanced Hg 107 concentrations in sedimentary rocks across the KPg boundary and Deccan Traps events (Nascimento-108 Silva et al., 2011, 2013; Sial et al., 2013, 2014) and enabled a distinction of chemical fingerprints 109 110 related to the Deccan volcanism from an impact event at the KPg boundary (Sial et al., 2014). 111 However, these studies have not taken the mutual relationship between TOC content and Hg concentrations into account, and this precludes an evaluation on how lithological variation (organic 112 matter-poor versus organic matter-rich sediments) was related to the reported Hg anomalies. 113 Post-emission or post-discharge Hg transformations may not drastically modify the isotopic 114 composition of particulate Hg ac- cording to several authors (e.g. Foucher et al., 2009; Sonke et al., 115 2010; Estrade et al., 2011; Gehrke et al., 2011; Chen et al., 2012; Sun et al., 2013). When Hg 116 transformations lead to change of Hg isotope signatures, this change tends to be constant under certain 117 118 conditions (Laffont et al., 2009, 2011; Gehrke et al., 2011). Multiple Hg transformations may modify the Hg isotope signatures in un- predictable way, making difficult to trace the Hg source (Sonke and 119 Blum, 2013). The Hg isotopic composition may help in the distinction be- tween volcanogenic and 120 meteoritic Hg. It is not simple to precise the isotope composition of mantle-derived Hg because usually 121 it is not entirely mantle-derived and/or because it has undergone fractionation during chemical/phase 122 transformation at near surface regions (Bergquist and Blum, 2009). Mantle-derived Hg probably has 123 $d^{202}Hg$ values close to 0% according to Sherman et al. (2010), but Zambardi et al. (2009) have 124

reported values from 1.74% to 0.11% for gas and particulates, respectively, from an active volcano in 125 Italy. Generally, geogenic Hg does not have significant mass independent fractionation (MIF) of odd 126 Hg isotopes (D²⁰¹Hg or D¹⁹⁹Hg) (Blum et al., 2014). On the other hand, Hg in most nat- ural samples 127 displays d²⁰²Hg values significantly different from crust and mantle values after it has undergone 128 cycling in the sur- face environment (Bergquist and Blum, 2009). Interestingly, these samples 129 commonly have significant MIF signatures as a result of photoreactions (or evaporation), with 130 generally negative D²⁰¹Hg for surface soils, sediments and land plants, while positive D²⁰¹Hg values 131 are observed for precipitation, aquatic organisms and at-mospheric particles (Blum et al., 2014). Hg 132 isotopes are potentially able to discriminate natural from anthropogenic Hg sources (Sonke and Blum, 133 134 2013). 2. The Deccan province and the KPg boundary event 135 Continuous advances in radiometric age dating have gradually confirmed the ties between the 136 Kalkarindji, Viluy, Siberian, Central Atlantic magmatic province, and Deccan volcanism to the Ear-137 lyeMiddle Cambrian (Jourdan et al., 2014), end-Devonian, end-Permian, end-Triassic, and end-138 139 Cretaceous mass extinctions, respectively (Keller and Kerr, 2014). In the Deccan lavas which flooded the Indian subcontinent (Raja Rao et al., 1999), three phases of continental flood basalts are known 140 (e.g. Chenet et al., 2009) with a total of 6% volume erupted in phase-1 (C30neC31n), while 80% was 141 142 erupted in phase-2 (C29r), and 14% in phase-3 (C29n). An about 850,000 years quiescence of Deccan volcanism separated Deccan phase-1 from phase-2. Phase-2, the longest lava flows in the Phanerozoic 143 (Self et al., 2008) started at 66.288 ± 0.027 Ma (UePb zircon dating; Schoene et al., 2015), preceding 144 the KPg boundary (65.968 ± 0.085 Ma; Renne et al., 2013) with around 250,000 years, encompassing 145 the age interval of the CF2 and CF1 planktic fora- minifers biozones. Deccan phase-2 ended at 500,000 146 years after the KPg boundary ($65.552 \pm 0.026 \text{ Ma}$). About 1.1 million km³ of lavas were erupted 147

(Schoene et al., 2015). Deccan Traps eruptions have been blamed responsible for the mass extinction
documented in the intertrappean beds in India (e.g. McLean, 1985; Keller et al., 2011, 2012, and
references therein).
It is known that volcanic gases (e.g. CO2 and SO2) from a single eruptive event, and not the total
volume erupted, may cause devastating impacts on Earth's environment and life (Svensen et al., 2009;
Bond and Wignall, 2014). The Deccan Traps contain magmatic sulfur concentrations as high as 1900
ppm (Callegaro et al., 2014), much higher than in basalts from some other similarly sized igneous
provinces(e.g.Parana-Etendeka). Therefore, eruptions likely caused globally extensive acidic rains
before and during the KPg boundary, resulting in weathering and dissolution effects on land.
Dissolution of foraminifers (e.g. Gertsch et al., 2011) and of iron oxide minerals (bio- and detrital
magnetite) is may be considered as important evidence of acidification of oceans during this critical
period (Font et al., 2014). Punekar et al. (2014) investigated the environmental/biological effects of the
Deccan Traps eruptions using C and O isotopes and demonstrated that the global high-stress
environment prior to the KPg boundary is coeval with the Deccan eruption phase-2 in C29r. These
authors have also concluded that the Deccan phase 3 correlates with the delayed Danian biotic recovery
in C29n. Keller et al. (2016) reviewed the chronology and geographic distribution of the three Deccan
phases in India. It is not yet possible to precisely date the Deccan flood basalts associated with mass
extinction. Therefore, one cannot yet quantify the effects of volcanism and asteroid impact on the KPg
boundary mass extinction. In the present study, we use Hg, as tracer in order to investigate the
influence of distal volcanism, in worldwide chemostratigraphic profiles across the KPg boundary with
the overall aim to contribute to solve this problem.

3. Study aims

172 volcanogenic signals in marine sediments in sections straddling the boundary from different continents to confirm the global extent of Hg loading related to the Deccan Traps volcanism, and (b) examining 173 Hg isotopes from these sites to help telling apart influence of volcanism from bolide impact 174 fingerprints in KPg events. By examining these Hg stratigraphic patterns, we investigate worldwide 175 fingerprints of Deccan erup- tions in climate change. 176 4. Study areas 177 Four distal KPg boundary sections relative to the Deccan Traps volcanic center (Denmark, Italy and 178 Argentina) and two proximal sections (India) are the focus of this study (Fig. 1). They represent well-179 documented sites in the Northern Hemisphere (Højerup at Stevns Klint, Bottaccione at Gubbio and 180 Padriciano near Trieste), and in the Southern Hemisphere (Meghalaya, Jhilmili and Bajada del Jagüel). 181 Mercury chemostratigraphical data from the Højerup, Bottaccione, and Bajada del Jagüel sections are 182 found in Sial et al. (2014), who have not reported TOC% values alongside with Hg concentrations. 183 This lack of TOC data has limited the further use of the depicted Hg spikes as proxy for volcanism. No 184 Hg chemo- stratigraphy has been so far reported for the Padriciano section, a classical shallow-marine 185 carbonate platform (Trieste Karst, Italy), for Meghalaya, South Shillong Plateau (NE India) and for the 186 187 Jhilmili section (central India). The last two sections are located at proximal and very proximal 188 distances to the Deccan volcanic province, respectively, and are of paramount importance for the assessment of Hg as fingerprint of eruptions in the period of the end-Cretaceous mass extinction. 189 In selecting sections for this study, we paid much attention to stratigraphic completeness of the 190 sedimentary successions and the geographic distribution of the sites. It is known that the sections at 191 Højerup (Stevns Klint), Bottaccione (Gubbio) and Meghalaya sections, based on the study of planktic 192 foraminiferal biozones, are likely complete. Stevns Klint is the best preserved and laterally the most 193

extensive KPg succession (UNESCO World Heritage Site, since June 2014). All foraminiferal biozones

The present study aims at: (a) broadening the knowledge and contrasting Hg spikes (Hg/TOC) as

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are present in the uppermost Maastrichtian to lowermost Danian succession at Højerup (Rasmussen et 195 196 al., 2005; Surlyk et al., 2006; Thibault et al., 2015). 197 Refined magnetostratigraphy, planktic foraminiferal and calcareous nannofossil biostratigraphies are available for the stratigraphic succession at Gubbio (e.g. Coccioni et al., 2010; Coccioni and Premoli 198 Silva, 2015). These authors have reported the presence of CF1, CF2 and CF3 for aminiferal biozones in 199 200 the uppermost Maastrichtian, encompassing the Scaglia Bianca and Scaglia Rossa formations up to the KPg boundary. In the Shillong Plateau, seven successive planktic foraminiferal biozones from across 201 the KPg boundary at the Um Sohrynkew section (Meghalaya) are present (Mukhopadhyay, 2008; Pal et 202 al., 2015) and these are in stratigraphic order CF4, CF3, CF2 and CF1 in the upper Maas-trichtian part 203 204 and the P0 and Pa zones and P1a subzone in the lower Danian part, representing a biostratigraphically continuous record across the KPg boundary. The Jhilmili intertrappean beds record the lower Danian 205 (planktic foraminifers P1a subzone; Keller et al., 2009) and represent the closest section to the Deccan 206 volcanic center. 207 In the Bajada del Jagüel section, Neuquén Basin, the CF1 biozone and the base of the Danian (P0, P1a, 208 P1b) seem to be missing (Keller et al., 2007; Pardo and Keller, 2008). The latest Maastrichtian as-209 sociation of planktic foraminifers is composed of species which are not age-diagnostic and, therefore, it 210 211 is not possible to precisely pinpoint the exact biozones at this site. The absence of typical species of the 212 CF1 zone led Keller et al. (2007) to tentatively assign the latest Maastrichtian foraminifers in the Bajada del Jagüel sec- tion to the CF2 zone. Typical Danian species occurring at the base of the Pa 213 zone (P. eugubina or P. longiapertura) seem to be absent. 214 215 Palamarczuk et al. (2002) have mentioned co-occurrence at Bajada del Jagüel section, of dinoflagellate (Senoniasphaera inornata and Damassadinium californicum) and foraminifers (Guembelitria cretacea 216 and Hedbergella montmouthensis) within the KPg boundary sandstone and in foraminiferal zone Pa (P. 217 eugubina) above, suggesting that this 20 cm-thick sandstone which marks the KPg transition lies within 218

Zone P0. Besides, according to Palamarczuk et al. (2002), the 50 cm interval directly above the sandstone con- tains planktic for aminiferal species typical of zones Pa through P1c. An $^{40}\mathrm{Ar}/^{39}\mathrm{Ar}$ plateau age of 66 ± 0.5 Ma for feldspar from the KPg boundary ash bed was reported by the same author. Habib and Saeedi (2007) have recognized an abundance of Manumiella seelandica (dinoflagellate) immediately below the KPg boundary and stated that the global distribution of this species makes it an excellent biostratigraphic marker. Fossils of M. seelandica have been reported from mid-latitude sites in Argentina (Palamarczuk and Habib, 2001; Palamarczuk et al., 2002). The pattern of magnetic susceptibility for the sedimentary rocks across the KPg boundary suggests that the section at Bajada del Jagüel is incomplete with respect to the very uppermost Maastrichtian, although the time span of interest may be as short as 40 kyr (Nañez et al., 2002). Aberhan et al. (2007) found neither a sign of the Danian planktic foraminiferal zone P0 nor impact specific trace element concentrations in the KPg boundary sandstone layer in the Bajada del Jagüel section, and concluded that this succession lacks about 100 kyr of a sedimentary depositional period. The disagreement about how much of sedimentary record is missing in the Bajada del Jagüel section can be somehow addressed and tested by comparing the episodic distribution of Hg/TOC anomalies with those in more complete sections as done here. 4.1. The Danish Basin, Stevns Klint, Denmark The Stevns Klint (coastal cliffs) is located about 45 km south of Copenhagen, Denmark, on the island of Sjælland (Figs. 1, 2). The sediments, deposited in the Danish Basin, represent one of the most complete KPg boundary section worldwide. They exhibit a KPg boundary layer beneath a topographic

overhang separating the lowermost Danian Cerithium Limestone Member from the over-lying lower

Danian bryozoan limestone of the Stevns Klint For- mation (Fig. 8; Surlyk, 1997; Surlyk et al., 2006,

2013). The distinct boundary clay (Fiskeler Member) varies in thickness and is mainly around 5 cm,

but up to 40 cm at Kulstirenden, in the northernmost part of the cliff, including here the so-called 'red

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layer' just above its base (Hart et al., 2004). There is clear evidence for shallowing in the latest Maastrichtian before the KPg boundary, and for sea-water temperature fluctuations (Surlyk, 1997; Hart et al., 2004; Thibault et al., 2015). The 45 m-thick succession exposed at Stevns Klint exhibits the stratigraphic evolution of the Danish Basin from the latest Cretaceous, across the KPg boundary into the early Paleogene. The most comprehensive stratigraphic studies of these sections are those of Surlyk (1997), Rasmussen et al. (2005); Surlyk et al. (2006; 2013). At the Højerup coastal cliff section, the Maastrichtian Møns Klint Formation (Sigerslev and Højerup members; Surlyk et al., 2013; Hansen and Surlyk, 2014) is overlain by the lower Danian Rødvig Formation (Fiskeler and Cerithium Limestone members) which in turn is covered by the lower to middle Danian Korsnæb Member of the Stevns Klint Formation (Surlyk et al., 2006). The Cerithium Limestone is diachronous and becomes gradually younger from the southern part of Stevns Klint towards the northern part. A hiatus including all the P. eugubina zone is present at the Fiske-lereCerithium Limestone transition in the northern part of the cliff but it is absent in the southern part (Rasmussen et al., 2005). Danian bryozoan limestone mounds, outlined by black flint bands, formed shortly after the KPg boundary mass extinction (Lauridsen et al., 2012 and references therein). 4.2. The Umbria-Marchean succession, Central Apennines, Italy The Gubbio Mountains are part of the UmbriaeMarche Basin with two well-known KPg boundary sections (Figs. 1, 2 and 8B). These occur in two parallel valleys, the Bottaccione Gorge and the Contessa Highway. The former begins just outside the town of Gubbio and cuts through pelagic Middle Jurassic to upper Eocene sedimentary rocks, including the Calcari Diasprigni, Maiolica, Marne a

Fucoidi formations and the Scaglia Group (Scaglia Bianca, Scaglia Rossa, Scaglia Variegata and

Scaglia Cinerea formations). The Bottaccione section represents the magnetostratigraphic standard for

the Upper CretaceouseEocene interval (e.g. Galeotti et al., 2015) and contains a 1 cm-thick clay layer at

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the KPg boundary (Fig. 2) within the Scaglia Rossa Formation, enriched in Ir (Alvarez et al., 1980). In 267 contrast to Stevns Klint, the KPg boundary layer at Gubbio has a very short lateral extent as the 268 269 succession is tilted (Fig. 2B). Alternation of gray and pink colored beds marks the contact between the Scaglia Bianca and the 270 overlying Scaglia Rossa for- mations. The latter brackets the KPg boundary and consists of 271 predominantly pink to red pelagic limestones with cherty nodules and calcareous marls, with bedding 272 thicknesses between 10 and 20 cm. This Upper Cretaceous to lower middle Eocene succession 273 represents an apparently continuous stratigraphic record across the KPg boundary and based on 274 abundant planktic and benthic fora- minifers and calcareous nannofossils, it has been dated as Late 275 276 Cretaceous to early middle Eocene (Premoli Silva and Sliter, 1994; Alvarez, 2009; Coccioni and 277 Premoli Silva, 2015 and references therein). The KPg boundary crops out on the eastern side of the main road and lies around 240 m above the base 278 279 of the Scaglia Rossa Forma- tion (Figs. 2B, 8B). The topmost Cretaceous in this section is rep-resented by a 30e40 cm-thick whitish limestone bed, overlain by the 2 cm-thick dark clay KPg boundary layer 280 which correlates to the well-known KPg boundary mass-extinction event (Premoli Silva and Sliter, 281 1994). The lowermost Paleocene is correlated with the occurrence of Globigerina eugubina 282 foraminiferal zone (=Parvorugoglobigerina eugubina; Luterbacher and Premoli Silva, 1964). 283 284 4.3. The Trieste Karst, NW Adriatic platform, Italy 285 The Karst region including the Trieste area of Italy and a portion of Slovenia is part of the Adriatic 286 platform and is characterized by Maastrichtian to Danian peritidal carbonate deposits (Fig. 3). In Italy, 287 the thickest succession of this shallow carbonate platform is located at Padriciano (Fig. 3). In the 288 Trieste Karst, the KPg boundary is well-documented in some sections at Padriciano (Pugliese et al., 289 1995, 2000; Tewari et al., 2007). The interval including the KPg boundary is characterized by presence 290

of several peritidal cycles consisting of basal breccias, lagoonal limestone, and stromatolitic limestone 291 292 with typical fenestral fabric. The KPg boundary in the carbonate platform of the Karst region of Slovenia was first identified by 293 Drobne et al. (1989). The main criteria they used to identify the KPg boundary was by the occur-rence 294 of certain biota, particularly by the disappearance of Creta-ceous taxa and appearance of Paleogene 295 ones, by identification of foraminiferal zone SBZ1 and of magnetic polarity Ch29r, the presence of and 296 Ir anomaly and by the strong negative shift of δ^{13} C values across the KPg boundary (Hansen et al., 297 1995; Marton et al., 1995; Ogorelec et al., 1995; Hansen and Toft, 1996; Palinkas et al., 1996). The 298 same criteria have been adopted for locating the KPg boundary in the Trieste Karst and, in addition, the 299 presence of microtektites at the top of a Maastrichtian breccia near the village of Padriciano was also 300 301 regarded as evidence for the record of the KPg boundary in this region (Gregoric et al., 1998). 302 The peritidal shallow carbonate succession at Padriciano is represented by the Liburnia Formation (Fig. 3) which contains a Maastrichtian breccia (50 cm-thick, Fig. 2), another breccia (20 cm-thick) of dark 303 limestone with rudist shell fragments and foramin- ifers (Tewari et al., 2007) about 80 cm below the 304 305 upper breccia (Fig. 2). The KPg boundary is situated at the base of the upper breccia, the latter overlain 306 by a succession of light gray Danian limestone with Microcodium that is overlain by stromatolitic limestone also with Microcodium. A change in the foraminifer species from Miliolidae, Rhapydionina 307 308 liburnica, Fleuyriana adriatica to the occurrence of Microcodium and Bangiana hanseni is present 309 immediately below the upper breccia (Caffau et al., 1998). This brown limestone, with gastropod 310 fragments and Microcodium, is overlain by gray stromatolitic limestone capped or penetrated by Microcodium. Further peritidal cycles of gray limestone overlain by stromatolitic limestone with 311 Microcodium, forming thick units, follow up-section. Within lagoonal limestones of these cycles, some 312 Danian opportunistic and r-strategist foraminifers taxa (e.g. Bangiana hanseni and small Miliolidae) 313 appear (Pugliese et al., 2000; Tewari et al., 2007) and characterize the Danian larger benthic 314

foraminiferal zone 1 (SBZ 1; Serra-Kiel et al., 1998) which corresponds to the planktic foramineral P0,

Pa and P1 biozones of Berggren et al. (1995).

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4.4. The Neuquén Basin, Patagonia, Argentina

The Neuquén Basin in Patagonia of western Argentina is located between 32 and 40 S latitude on 319 320 the eastern side of the Andes in Argentina and central Chile. This foreland basin comprises Upper Triassic to Miocene deposits (Vergani et al., 1995; Legarreta and Uliana, 1999; Howell et al., 2005). 321 The KPg transition is situated in the Upper Cretaceouselower Paleocene Malargüe Group that com-322 prises the Allen, Jagüel, Roca and El Carrizo formations and depos- ited unconformably overlying the 323 Upper Cretaceous Neuquén Group (Legarreta et al., 1989; Casadío, 1998; Heredía and Salgado, 1999). 324 The KPg transition in this basin has been extensively investigated by biostratigraphy, microfacies 325 analysis, bulk rock/clay mineralogy, isotope stratigraphy, and trace and major element chemistry (e.g. 326 Howell et al., 2005; Scasso et al., 2005; Aberhan et al., 2007; Keller et al., 2007; Musso et al., 2012). 327 These and further results show that the Maastrichtian succession was deposited during trans- gression. 328 A regressive trend was recorded in the Paleocene (Scasso et al., 2005; Aguirre-Urreta et al., 2011). The 329 Jagüel Formation en- compasses the KPg boundary (e.g. Uliana and Biddle, 1988) and consists of 90-m 330 thick, monotonous marine mudstones deposited in a mid-to outer-shelf environment, covered by 331 332 bioclastic marine limestones of the Danian Roca Formation (Fig. 4). The KPg boundary interval has been identified on the basis of palynomorphs (Papú et al., 1999), ostracods (Bertels, 1975), calcareous 333 nannofossils and foraminifers (Nan~ez and Concheyro, 1997). The position of the KPg transition is 334 constrained to a single, thin coarse-grained sandstone bed (Palamarczuk and Habib, 2001; Palamarczuk 335 et al., 2002; Scasso et al., 2005) in the upper half of the Jagüel Formation, and Danian age above has 336 been confirmed by calcareous nannofossils (Scasso et al., 2005). This 15e25 cm thick non to slightly 337 lithified tuffaceous sandstone can be traced laterally for 5 km in the area. It occurs within the 338

homogeneous shelf mudstone of the Jagüel Formation and contains abundant volcanogenic plagioclase volcanic lithics, shell detritus and shark teeth (Palamarczuk et al., 2002). This KPg tran- sition sandstone shows abundant rip-up clasts, erosional base, coarse-grain size, normal grading and hummocky cross-bedding, and has been interpreted as a tsunami deposit in a shelf environ- ment, related to the Chicxulub bolide impact in Mexico (Scasso et al., 2005). Spherule, shocked quartz or enrichment of meteoritic com- ponents has not been reported from this layer or the mudstones immediately above. Keller et al. (2007) suggest that the KPg boundary is marked by erosional-based sandstone that signifies a hiatus. This sandstone, previously interpreted as a mark of the KPg transition, contains diverse planktic foraminiferal zone P1c assemblages and nanno- fossils of zone NP1b immediately above (Keller et al., 2007). These authors suggested that its deposition occurred at about 500 kyr after the KPg hiatus.

4.5. Meghalaya, South Shillong Plateau, NE India

The Um Sohrynkew section is situated in northeastern India, north of Bangladesh and in a distance of about 800e1000 km from the Deccan volcanic province (Figs. 5, 6 and 7). It is exposed along the Um Sohrynkew River in Meghalaya, next to Therria village, East Khasi Hills District, south Shillong Plateau, in the eastern Himalayas (Fig. 7). The succession comprises the most complete marine KPg succession known from India, and possibly worldwide (Gertsch et al., 2011 and references therein). The section displays strong evidence of mass extinction patterns (e.g. planktic foraminifers and larger ammonoids), a preserved KPg boundary layer, as well as the first appearance of Danian foraminifers, and indications for sea- level change (Keller et al., 2008, 2009; Tewari et al., 2010a, 2010b; Gertsch et al., 2011 and references therein). The KPg boundary at Um Sohrynkew (Fig. 5) is marked by a thin red clay layer enriched in Ir and other platinum group elements (Pandey, 1990; Bhandari et al., 1993, 1994; Garg et al., 2006) with abundant subangular quartz grains in a brown matrix (Gertsch et al. 2011). This

section consists of a continuous CampanianeEocene succession characteristic of coastal, estuarine and nearshore envi- ronments (Nagappa, 1959; Krishnan, 1968; Banerji, 1981; Tewari et al., 2010a, 2010b) with marine shelf sediments including thick sandstone layers, marl, shale and carbonates.

Based on the distribution of zonal indices, Mukhopadhyay (2008) and Pal et al. (2015) have recognized seven successive planktic foraminiferal zones across the KPg boundary at the Um Sohrynkew River section. These zones are, in stratigraphic order, CF4, CF3, CF2 and CF1 in the upper Maastrichtian part and Zone P0, Zone Pa and Subzone P1a in the lower Danian part, thus, representing a biostratigraphically continuous succession across the KPg boundary. We focus only on 2.5 m of the MaastrichtianeDanian interval of the Um Sohrynkew section, bracketing the KPg boundary. Twenty-five samples have been stratigraphically collected at an interval of 10 cm: seven samples from greenish sandstone and greenish gray clayey marl of the Maastrichtian Mahadeo Formation, one sample from the rust-red-colored sandyesilty KPg boundary layer (about 2 cm thick) and the rest of samples from the Danian Langpar For- mation whose basal portion is formed by a 10 cm-thick bioturbated sandstone overlain by dark to light gray shale to marl (Fig. 5).

4.6. Jhilmili intertrappean sediments, Madhya Pradesh, Central India

The Deccan intertrappean sediments have been regarded for long time as terrestrial deposits, Maastrichtian in age, until the work by Keller et al. (2009) who studied the sedimentology, microfacies, biostratigraphy and, C- and O-isotope chemo- stratigraphy of a thick intertrappean succession next to the village of Jhilmili (22°02'044" N, 79°09'034" E), Chhindwara District of Madhya Pradesh, central India. This succession consists of sedi- ments with early Danian planktic foraminifera sandwiched between two basalt flow horizons (the lower and upper trap basalts correspond, respectively, to C29r and the C29r/C29n transition) (Keller et al., 2009) and is located near the main road that links Seoni and Chhindwara, exposed in a shallow-river valley. Its lithological types comprise claystones, paleosols,

siltstones, clayeyesilty marlstones to limestones and can be subdivided into three different units 387 388 summarized below. The 6 m thick lowermost unit (unit 1), deposited in terrestrial and palustrine environments, overlies the 389 strongly weathered lower basalt trap. It consists of red clayey siltstone with carbonate nodules and 390 coarser quartz grains, becoming less abundant to- wards the top. The unit is overlain by another about 391 70 cm thick lacustrine-brackish deposit (unit 2), that consists of pink, reddish to yellow laminated 392 clays, containing marly limestones in its lower part, and with abundant volcanic glass shards. The top 393 of this unit is formed by coarse-grained nodular limestone that was probably deposited in fresh to 394 brackish-marine water deduced from the common freshwater and rare brackish-water ostracods along 395 396 with rare planktic foraminifers (Keller et al., 2009). Unit 3 is similar to the unit 2, consisting of red clayey siltstones, with clay and carbonate clasts 397 interbedded with fine-sand layers. Glass spherules and shards are present as well as a carbonate- nodule 398 layer. A 1 cm-thick black charcoal layer separates this unit from the overlying basaltic flow which is 399 strongly weathered. Immediately below this upper basalt trap, a 30 cm-thick tempestite has been 400 401 recognized. In this intertrappean succession, Keller et al. (2009) have iden-tified a thin aquatic interval of fresh 402 water ponds and lakes deposits overlain by shallow coastal marine sediments with brackish marine 403 404 ostracods and lower Danian zone P1a planktic foraminifers, very close to the KPg boundary. Their 405 most important contributions were: (a) discovery of environmental changes preserved within the intertrappean sediments at Jhilmili and (b) a marine incursion pointing to the existence of a seaway at 406 least 800 km long, across India, from the west through the NarmadaeTapti rift valley (Shukla and 407 Srivastava, 2008; Keller et al., 2009, Fig. 6). 408

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5. Methods

5.1. Analysis of total organic carbon (TOC), molybdenum and aluminum 411 All analyzed samples in this study were decarbonated using 6 M HCl prior to TOC analyses. Acid 412 remains were subsequently removed by a repeated procedure of rinsing, centrifugation, and decantation 413 of supernatant liquids. Remaining fractions were weighed out again after 24 h freeze drying at 414 temperatures of 54 C. The CaCO3 concentration was calculated by the weight difference of the 415 aliquots before, and after the acid treatment. The total organic carbon was measured on 160 mg of 416 decarbonated aliquot using a Carbon-Sulfur-Determinator (Eltra CS 500) at the Department of 417 Geosciences and Natural Resources Management, University of Copenhagen. The sample was 418 combusted for 90 s in a ceramic boat at a temperature of ~1350 C with a catalytic oxidation process. 419 The resulting CO2 signal was electronically linearized and integrated, and subsequently used to 420 calculate the TOC. The reproducibility, obtained from multiple analyses of the in-house reference 421 material SKK-9 (TOC 1/4 6.88%), is generally better than 0.1% (1 sd). 422 Molybdenum and aluminum were analyzed by ICPeMS at the Geological Survey (GEUS) in 423 Copenhagen, Denmark after sample digestion with 6N HCl, dissolving all carbonates, but probably 424 leaving some silicate detrital material back. These analyses are, therefore, incomplete whole-rock 425 426 analyses and the reported aluminum concentrations are regarded as minimum-rock concentrations. 427 Molybdenum has been measured over two different iso-topes (95Mo and 98Mo) and here the total Mo 428 concentrations are calculated from the 95Mo peak. 429 5.2. Analysis of mercury 430 Mercury concentrations were determined in homogenized powdered samples at the LABOMAR in the 431 Federal University of Cearà, Brazil, using the method described by Sial et al. (2014, p. 105) "glass and 432 plastic ware were decontaminated by immersion for 1 day in (10% v/v) Extran solution (MERCK), 433 434 followed by immersion for 2 days in diluted HCl (5% v/v) and final rinsing with Milli-Q water. All

chemical reagents used were of the least analytical grade. Cold Vapor Atomic Fluorescence Spectrophotometry, using a Millenium Merlin PSA spectrophotometer, was used for Hg determination, after Hg2b reduction with SnCl2. All samples were analyzed in duplicates, showing reproducibility within 9.5%. A certified reference material (NIST 2702, Canada) was simulta- neously analyzed to evaluate Hg determination accuracy. Such analysis showed a precision of 4%, as indicated by the relative standard deviation of three replicates, and presented Hg recovery of 98.8± 6.2%. The Hg detection limit estimated as 3 times the standard deviation of reagent blanks was 1.26 ng.g-1. In all cases, blank signals were lower than 0.5% of sample analysis. The con- centration values were not corrected for recoveries found in the certified material powder, following Sial et al. (2014, p. 105)." 5.3. Analysis of carbon isotopes Inorganic and organic δ^{13} C analyses of carbonates were per- formed at the Stable Isotope Laboratory (LABISE) of the Department of Geology, Federal University of Pernambuco, Brazil. Extraction of CO₂ gas was performed using a conventional high vacuum extraction line after reaction with 100% orthophosphoric acid at 25 °C for one day. Released CO₂ was analyzed in a Thermofinnigan Delta V Advantage mass spectrometer and results are reported in δ notation in permil (‰) VPDB. δ¹³Corg was analyzed from total organic carbon of insoluble residues of samples from all the studied sections. After whole-rock samples were crushed into powder "insoluble residues for organic carbon isotope analysis were obtained by acidifying these whole- rock powders in increasing concentrations (0.4, 0.6 and 0.8 mol l⁻¹) of H₃PO₄ for three days to dissolve all carbonate minerals. Care was taken to ensure that acid was added and acidification continued until there was absolutely no visible car-bonate dissolution so that the analyses would not be affected by contamination from residual inorganic carbon. The insoluble resi- dues were then rinsed with DI water, dried and loaded into tin capsules for isotopic

analysis (Sial et al., 2014, p. 105)." δ^{13} C org was measured using continuous flow elemental analysis

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COSTECH elemental combustion system (com- bined analytical and sampling error of $\pm 0.2\%$). 460 461 5.4. Analysis of Hg isotopes 462 Mercury isotope ratios were determined on the new MC-ICP-MS (Nu-Plasma II, Nu Instruments) 463 equipped with sixteen Faraday cups at the State Key Laboratory of Environmental Geochemistry, Insti-464 tute of Geochemistry, Chinese Academy of Sciences, China, following a method similar to Huang et al. 465 (2015). A continuous flow cold-vapor generation system (CV) was coupled with an Ari- dus II 466 desolvation unit (CETAC Technologies, U.S.) for Hg and Tl introduction, respectively. The 467 468 instrumental baseline was measured by de-focusing before each sample and standard. Both the internal standard method and the standard-sample bracketing technique were used to correct for instrumental 469 mass bias (Chen et al., 2010; Jiskra et al., 2012). 470 Mass-dependent fractionation (MDF) of Hg isotopes is reported in delta notation, δ , which is the permil 471 (‰) deviation relative to the SRM 3133 standard. Mass independent fractionation (MIF) of Hg isotopes 472 473 is reported using upper-case delta notation, Δ , which is the deviation of the measured isotope ratio from 474 the theoretical ratio predicted by MDF. The analytical quality was controlled by repeated measurement of standard materials. Long-term analysis gave average values of -0.54 \pm 0.10%, -0.02 \pm 0.04% and -475 $0.04\pm0.04\%$ for δ 202 Hg, Δ^{199} Hg and Δ^{201} Hg of UM-Almaden Hg (2SD, n = 21), and of -1.22 \pm 476 0.16%, $0.06 \pm 0.09\%$ and $0.03 \pm 0.10\%$ for those of Fluka Hg (2SD, n 1/4 13), respectively, in 477 accordance with the published results (Bergquist and Blum, 2009; Chen et al., 2010; Chen et al., 2012; 478 Jiskra et al., 2012). The 2SD of the isotopic compositions of the UM-Almaden were considered as the 479

analytical uncertainty for the isotopic compositions of samples. When the uncertainty of the replicate

isotopic measurements of one sample was larger than the 2SD of the UM-Almaden, the un-certainty

was applied to the sample (Huang et al., 2015).

isotope-ratio mass spectrometry with a Delta V Advantage mass spectrometer interfaced with a

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6. Results and data interpretation 484

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6.1. Carbon-isotope chemostratigraphy and total organic carbon (TOC) The covariation between carbonate and sedimentary organic C- isotope records can help establishing whether variations in the δ^{13} Ccarb record reflect changes in the isotopic composition of the ancient dissolved inorganic carbon pool (e.g. Oehlert and Swart, 2014, and references therein). Covariant $d\delta^{13}$ Ccarb and δ^{13} Corg records evidence that both the carbonate and organic matter were originally produced in the surface waters of the ocean and have retained their original δ^{13} C composition (e.g. Korte and Kozur, 2010; Meyer et al., 2013) as it is believed that no secondary processes are able to shift δ^{13} Ccarb and δ^{13} Corg in the same direction at the same rate (Knoll et al., 1986). Decoupled δ^{13} Ccarb and δ^{13} Corg records indicate diagenetic alteration (e.g. Grotzinger et al., 2011; Meyer et al., 2013) or that local syn-sedimentary processes have intro-duced noise in the δ^{13} Corg record (Maloof et al., 2010). Global δ^{13} C records of bulk sediment comparing C-isotope pathways across the KPg boundary from far apart sections at Højerup, Bottaccione and Bajada del Jagüel localities (Figs. 8 and 9), were previously published by Sial et al. (2014). New δ^{13} C carb data are presented here for the sections in the Meghalaya, Jhilmili and Padriciano (Tables 1 and 3; Figs. 8 and 10) and δ^{13} C org for the six sections under consideration (Tables 1e3, Figs. 8, 9, 10). Twenty-five samples were collected stratigraphically at centi- meter scale, perpendicular to the strike of the strata, from a 2.5 m section across the KPg boundary at the Um Sohrynkew River in Meghalaya (Table 3; Fig. 10A). The lowermost meter in this section is composed of greenish glauconitic sandstone and greenish-gray shale belonging to the Maastrichtian Mahadeo Formation, below the yellowish brown-reddish KPg boundary clay layer (samples KT- 1 through KT-7 in Table 3 lie immediately

below the KPg boundary clay layer). Light to medium-gray and dark-gray to light-medium gray shale 507 508 (samples KT-8 through KT-25; Table 3) were collected from the lowermost portion of the Danian 509 Langpar Formation, immediately above the KPg boundary layer. Twenty-four samples were collected from a 7.3 m section o intertrappean beds (Table 3; Fig. 10B) 510 between two basaltic flow layers in a shallow river valley next to the village of Jhilmili, Chhindwara 511 512 District of Madhya Pradesh. The lowermost layer in this section is composed of ferricrete paleosol with calcitic veins which is covered by a hard sediment layer within paleosol with clastic grains and calcitic 513 cement. This is followed upward by fine- grained, brick red with yellowish tint soft ferricrete, and then 514 by comparably harder grayish brown claystone with laminar features. Towards the top, this section 515 contains again fine-grained, brick red soft ferricrete with yellowish tint, which is covered by a greenish 516 clay layer with calcitic veins. Eighteen samples of carbonates from the Liburnia Formation have been 517 collected from a representative section across the KPg boundary at a quarry at Padriciano in the Trieste 518 519 Karst (Table 1; Fig. 8C). Among the six sections in this study, the three from Europe and the one from Bajada del Jagüel display 520 isotopically lighter carbonate composition immediately above the KPg boundary, portrayed by a shift 521 to lighter δ^{13} C values (Figs. 8 and 9). This picture could be partially a consequence of an increase of 522 volcanic CO₂ in the atmosphere (80% of the Deccan phase 2 eruptions took place within the time lapse 523 recorded in the sections examined) which may have accounted also for the large negative δ^{13} C 524 525 excursion at or just below the KPg boundary. Another important factor driving this negative excursion 526 is the collapse of bioproductivity at the boundary, with a gradual recovery in the early Danian. An abrupt decrease of δ^{13} C carb at the KPg boundary is noticeable at the Højerup, Bottaccione and 527 Meghalaya sections. At the Bajada del Jagüel and Padriciano localities, there is a strong decrease of 528 δ^{13} C values that predate (but very close) the KPg transition. In Jhilmili, the end-Cretaceous is 529 represented by basalt in the studied section and the δ^{13} C pathway shows a prominent minimum (-12‰) 530

at about 1 m above the lower basaltic flow, and upsection this curve gradually shows less negative 531 532 values, predominantly from -4 to -3%. 533 The total organic carbon content in the studied sections, expressed as percent TOC, are <1.0% (Tables 1e3), with the majority of samples yielding values < 0.2%. Among the complete sections in this study, 534 the highest TOC values were recorded in the KPg boundary layer in the Højerup section (~0.9%) and in 535 the uppermost Maastrichtian and Danian portions of the Meghalaya section (0.3e0.6%), while much 536 lower TOC values are observed in the whole Bottaccione section (0.01-0.09%). At Højerup, the TOC 537 stratigraphic variation curve shows a very monotonous pattern with almost no oscillation except a 538 prominent positive shift at the KPg boundary layer (Fig. 8A). At Gubbio, however, the TOC strati-539 540 graphic pathway shows vigorous oscillations with a negative shift about 4 m below the KPg boundary (within the CF2 biozone), a small negative excursion at the KPg boundary and a prominent positive 541 shift about 8 m above the KPg boundary (within the P1a biozone). At Meghalaya, the TOC 542 543 stratigraphic variation curve shows two marked positive excursions, about one meter below the KPg boundary (CF2 biozone) and another at the KPg boundary and in lowermost Danian (P0 biozone). 544 At the Padriciano section, the TOC values are all <0.3%, with highest values found in the Maastrichtian 545 portion of the section and a positive shift in the KPg boundary breccia. In Bajada del Jagüel, TOC 546 values range between 0.3 (Maastrichtian) and 0.8% (Danian), and a strong shift to values as low as 547 548 0.01% is the sandstone that marks the KPg transition (Table 2). At the Jhilmili section, the TOC stratigraphic pathway shows vigorous oscillations in the lower half of the section, followed by a strong 549 positive shift in the upper half of this curve. 550 The δ^{13} C org curve for organic matter from the Højerup section displays a gradual decrease of values 551 from -23% towards the KPg boundary layer with a minimum of -27% at the KPg boundary (Fig. 8A). 552 This is followed by a strong increase to -22% upsection and again by a negative shift to -27% and a 553 positive shift to -22‰. At the Bottaccione section (Fig. 8B), the δ^{13} Corg curve shows a less 554

pronounced decrease towards to the KPg boundary within the CF2eCF1 biozones, a discrete decrease at 555 the KPg boundary, and two positive shifts within the P1a biozone. The Meghalaya section shows a 556 different pattern (Fig. 10A) with no decrease of δ^{13} Corg towards the KPg boundary, contrary to what 557 was observed at Højerup and Bottaccione. At the KPg boundary, the δ^{13} Corg pathway shows a 558 maximum (-21‰) within the P0 biozone, followed upsection by a monotonous variation curve within 559 the P1a (1) subzone and a small increase to -24% within the P1a (2) subzone. A covariance record 560 between δ^{13} Ccarb and δ^{13} Corg at the Højerup and Bottaccione sections is depicted in Fig. 8A and B, 561 providing evi- dence that the carbonate and organic matter were produced in the surface waters of the 562 ocean and have probably retained their original respective δ^{13} C composition. At Meghalaya, the 563 δ^{13} Ccarb and δ^{13} Corg curves display opposite behaviors around the KPg boundary, with large positive 564 excursion of the δ^{13} Corg curve. 565 The Padriciano section is characterized by a δ^{13} Corg curve (Fig. 8C) similar to that observed at 566 Højerup, with a gradual decrease from -21 to -23% from one meter below the KePg transition to the 567 KPg transition, reaching a minimum about 0.5 m above this transition, and followed upsection by a 568 positive shift. The δ^{13} C org curve for Bajada del Jagüel section exhibits a minimum (~-26%) about 0.5 569 m below the KPg boundary (within the CF2 biozone), a negative excursion at the boundary, a positive 570 excursion (-23‰) immediately above the boundary, and a gradual and strong decrease to -26‰ 571 upsection (Fig. 9). For the Danian Jhilmili section, the δ^{13} Corg shows two positive shifts within the P1a 572 573 biozone with a strong negative shift at the top, close to the upper basalt flow. Except for Meghalaya, the behavior of the δ^{13} Corg stratigraphic variation curves for these sections may possibly be explained 574 by a response to growing deterioration of the environment at the end of the Maastrichtian with gradual 575 decrease of organic productivity, followed by its gradual recovery early in the Danian. 576

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6.2. Mercury concentrations, Hg/TOC and Mo/Al ratios

Mercury concentrations for stratigraphically collected samples from Højerup, Bottaccione (Table 1) 579 580 and Bajada del Jagüel (Table 2) sections are available in Sial et al. (2014). The Hg variation pattern for the Højerup section displays an Hg enhancement (260 ng.g⁻¹) that coincides with a minimum value of 581 δ¹³C in the Fiskeler Member at the KPg boundary (Fig. 8A). A second Hg enrichment (48 ng.g⁻¹) is 582 seen 1.6 m above the KPg boundary layer. In the Bottaccione section, three Hg enhancements (Fig. 8B) 583 were observed by Sial et al. (2014), a small one (2 ng.g⁻¹) at about 6 m below the KPg boundary, the 584 largest one at the KPg boundary (6 ng.g⁻¹) which coincides with a minimum of δ^{13} C, and a third one at 585 about 7 m above the KPg boundary (4 ng.g⁻¹). In the Hg strati- graphic variation curve for the Bajada 586 del Jagüel section (Fig. 9), three Hg enhancements are present, one at about 65 cm below the KPg 587 boundary (17 ng.g⁻¹Hg), a second one at this boundary (16 ng.g⁻¹) and a third one (13 ng.g⁻¹) around 15 588 cm above that. 589 590 Hg concentrations were measured in eighteen bulk samples from Padriciano (Table 1), twenty-five from Meghalaya and eigh- teen from Jhilmili (Table 3) and the corresponding Hg variation patterns are 591 shown in Figs. 8C, 10A and B. In Meghalaya, an Hg enrichment is seen at about 0.5 m below the KPg 592 boundary layer (Fig. 10A), preceding the prominent negative δ^{13} C that marks the KPg boundary. Two 593 prominent Hg enrichments are present in the intertrappean Danian sediments of the Jhilmili section. In 594 the Padriciano section, the Hg stratigraphic variation pathway shows a prominent Hg enrichment (183 595 ng.g⁻¹) at about 40 cm below the KPg boundary, followed by a second Hg enrichment (51 ng.g⁻¹) at 596 about 25 cm above the KPg boundary and a more discrete one (31 ng.g⁻¹; Fig. 9C) at about 2.40 m 597 above the KPg boundary. 598 599 In order to screen for Hg enrichments in these sections that represent true Hg loading to the environment, Hg concentrations have been normalized to corresponding TOC values following the 600 approach recommended by Grasby et al. (2015a, 2015b) and Percival et al. (2015). However, in 601 normalizing Hg concentrations to TOC, caution should be taken in using Hg-TOC pairs in which TOC 602

concentrations are <0.2% because too low Hg concentrations may produce, in some cases, unrealistic 603 604 or even false Hg/TOC spikes. Absolute Hg concentrations, TOC, Hg/TOC and Mo/Al ratios for all the studied sections are listed in 605 Tables 1e3 and are plotted in Figs. 8Ae10B. In most cases, elevated Hg concentrations, after 606 normalization to TOC, have generated a Hg/TOC spike. At Meghalaya, Bottaccione and Højerup, one 607 Hg/TOC spike is observed below the KPg boundary, within the age interval of the planktic 608 foraminiferal CF2 biozone (labeled I; Figs. 8A, B and 10A). This spike in the Bot-taccione section is 609 located about 4 m below the KPg boundary and is somewhat smaller than in the Meghalaya section. At 610 Padriciano, a large Hg/TOC spike is observed below the KPg boundary and, perhaps, can be correlated 611 612 with the spike described above, but presence of planktic foraminifers typical of the CF2eCF1 biozones has not been observed and, therefore, do not support such a correlation (Fig. 8C). 613 A second Hg/TOC spike is present at the KPg boundary at Højerup, Bottaccione, and Meghalaya 614 615 (labeled II in Figs. 8A, B and 10A). Similar to the Hg/TOC spike I, the size of spike II is variable from place to place, being larger at Højerup. At the Padriciano section (Fig. 8C), this spike is absent as the 616 KPg boundary is marked by an unconformity. Two large Hg/TOC spikes are present in the Jhilmili 617 section within the planktic foraminiferal subzone P1a (labeled III and IV; Fig. 10B). At Højerup, a 618 Hg/TOC spike is seen about 1e1.5 m above the KPg boundary (labeled III, Fig. 8A), about 4 m above 619 620 the KPg boundary at Bottaccione and about 0.5 m above at Meghalaya. At the Bajada del Jagüel section, a large Hg/TOC spike is observed in the sandstone layer that marks 621 the unconformity at the KPg boundary (Fig. 9). The presence of volcanoclastic sedimentary grains in 622 623 the sandstone is an evidence of local volcanism adjacent to the Neuqueñ Basin that may have contributed to enhancing the Hg/ TOC ratio background prior to the KPg boundary. This positive Hg/ 624 TOC excursion may correspond to the Hg/TOC enrichment (spike II) recorded in the Højerup and 625 Bottaccione sections. However, the paucity of the record of the CF1, Po, Pa and P1a planktic forami-626

niferal biozone interval, as pointed out by Keller et al. (2007), attest to a depositional or erosional 627 hiatus of about 700 kyr bracketing the KPg boundary. Therefore, this Hg enrichment is possibly related 628 629 to local volcanism next to this basin and the absence of spikes I and III could be justified by this depositional/erosional hiatus. 630 Covariation between Hg and Al₂O₃ (mostly associated with clays) is observed in some sections across 631 the KPg boundary and this led to the suggestion that Hg was probably adsorbed onto continental clays 632 with subsequent transport and deposition into the sea (e.g. Sial et al., 2013). However, there is no clear 633 correlation between Al₂O₃ and Hg/TOC (Tables 1-3), except in the KPg boundary layer at Højerup 634 (Fiskeler Member). 635 636 Molybdenum in sedimentary rocks has been suggested as an important elemental proxy for anoxia (e.g. Bond et al., 2015; Grasby et al., 2015a). Based on results from the Cariaco Basin, north central coast of 637 Venezuela, Lyons et al. (2003) demonstrated a high cor- relation between TOC content and Mo/Al 638 639 ratio in euxinic sediments and, therefore, Mo/Al trends may allow calculating the TOC composition. According to Wilde et al. (2004), however, this method demands refinement and no universal 640 elemental proxy for determining TOC has been established so far. Mo/Al ratios are useful for 641 discerning the original compositions of ancient rocks that were subjected to later diagenetic, low-grade 642 metamorphism or weathering. 643 644 Molybdenum concentrations, analyzed in almost all samples in this study, have been normalized to 645 corresponding Al values (Figs. 8Ae10B). At Højerup, low values of Mo/Al ratio characterize the topmost Maastrichtian (0.50 cm below the KPg boundary) and no correlation is observed between this 646 ratio and TOC. Within the Fiskeler Member clays, Mo/Al ratios are enhanced, while δ^{13} C org changes 647 to higher values and a positive shift of TOC values is observed. In the Cerithium Limestone and Stevns 648 Klint Formation, in the lower Danian, no correlation between TOC and Mo/Al is apparent and Mo/Al 649 ratios are higher than in the Fiskeler Member. 650

At Bottaccione, Mo/Al ratios are low, with a monotonous strat- igraphic variation curve in the upper Maastrichtian, but a negative shift is seen about 0.50 cm below the KPg boundary, coinciding with vigorous positive-negative shifts in TOC values followed by a positive one of Mo/Al at the KPg boundary. At Meghalaya, Mo/Al ratios are low in the upper Maastrichtian, but likewise Højerup and Bottaccione, it shifts to lower values within 20 cm below the KPg boundary, opposite to TOC values which exhibit higher values. In the Bajada del Jagüel section, Mo/Al ratios are also low, but with little variation and no particular correlation with TOC is apparent. In the Danian Jhilmili section, Mo/Al ratios are low, and in the lower half of this section no correlation is seen with TOC but in the upper half of the section there is a weak correlation with TOC values. In the Bottaccione, Padriciano, Meghalaya, and Bajada del Jagüel sections, minimum Mo/Al ratios do not coincide with δ^{13} C org peaks. The Højerup section is the only one which departs from this rule. Lowrie et al. (1990) investigated the origin of the whitish limestone beds at about 20e50 cm below the KPg boundary in the Bottaccione and Contessa sections at Gubbio. They believed that these white beds were deposited under the same conditions as the underlying pink beds of the Scaglia Rosa Formation whose whit- ening is related to removal of Fe2b ions by downward infiltration of reducing waters resulting from large quantity of organic matter produced by the extinction at the KPg boundary. However, Abrajevitch et al. (2015) demonstrated that a process of downward percolation of organicrich fluids is unlikely at Bidart and Gubbio and the stratigraphic record of Mo/Al redox proxy in the studied KPg sections here suggests that Hg/TOC spikes are likely not related to decrease in oxygenation. Therefore, anoxia was not the main cause for anomalous Hg/TOC observed across the KPg boundary.

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6.3. Hg isotopes

Mercury isotope analyses for the Hg spikes in the complete Højerup (Fiskeler Member), Bottaccione 674 675 (Scaglia Rossa Formation) and Meghalaya sections, besides Jhilmili, Padriciano and Bajada del Jagüel 676 sections are listed in Table 4, with results reported in delta notation in permil (%) relative to NIST SRM 3133 Hg standard. Six analyses from a section at Bidart, part of the Basque basin, France, one of 677 the most complete KPg boundary successions known (Bonté et al., 1984; Galbrun and Gardin, 2004; 678 Font et al., 2014, 2016), were included for comparison and are plotted alongside the analyses for all 679 sections in this study in Fig. 11. In this Figure, δ^{202} Hg (MDF) for all analyzed samples were plotted 680 against corresponding Δ^{201} Hg (MIF) values and the ranges for volcanic emission and 681 chondrite/volcanic emission Hg are indicated. 682 Two samples from the Meghalaya section yielded δ^{202} Hg values of -1.61% (spike I) and -1.89% (spike 683 II) and Δ^{201} Hg close to 0.0%. Three among four of the analyzed samples (spike II) from the Fiskeler 684 Member at Højerup yielded δ^{202} Hg between -1.00 and -2.00% and two of them display positive Δ^{201} Hg 685 686 or negative but very close to 0.0%. One sample from the KPg boundary layer at the Bottaccione section in Gubbio (Spike II) yielded a δ^{202} Hg value of -1.28% and positive Δ^{201} Hg. In addition, Hg 687 isotopes were analyzed in six samples from Bidart which yielded δ^{202} Hg values from -0.25 to -2.66 and 688 all positive Δ^{201} Hg values (Fig. 11). A sample from the KPg boundary layer from which Font et al. 689 (2016) determined a true Hg spike (corresponding to spike II in this study) yielded a δ^{202} Hg value of -690 0.74 compatible with volcanic or chondrite source Hg. So, in the light of or current Hg isotope data, a 691 volcanic source for Hg in the KPg boundary (spike II) is likely at these classical KPg localities. 692 One sample from the Padriciano locality (spike I) yielded a δ^{202} Hg value of -1.38% and slightly 693 positive Δ^{201} Hg, and as in Meghalaya, Hg isotopes seem to support volcanic source for the spike I in 694 these two localities. One sample from spike III at the Jhilmili site showed a value of -1.01% and 695 negative Δ^{201} Hg while another sample (spike labeled IV) yielded a δ^{202} Hg value of -2.18‰ and a 696 Δ^{201} Hg value of -0.28%. While the Hg isotope signature for spike III seems to support a volcanic 697

origin, the one for spike IV suggests some post-depositional alteration. Among the twelve samples 698 from Jagüel Formation, nine show δ^{202} Hg between -1.00 and -2.00% and Δ^{201} Hg are all positive, 699 compatible with a volcanic source Hg. 700 Interestingly, most of the analyzed samples in this study show slightly, but significantly higher than the 701 analytical precision of 0.04%, positive Δ^{201} Hg signatures. Since most continental samples (soils, 702 sediments and land plants) usually display negative or close to zero Δ^{201} Hg and post-deposition and 703 diagenetic processes likely would not induce odd-MIF, the positive Δ^{201} Hg values observed in most 704 samples here indicate a long-term atmospheric transport prior to deposition. During transportation in 705 706 the atmosphere, photoreduction of gaseous oxidized Hg (GOM) would enrich odd isotopes in water 707 droplets and particles, for example by adsorption, thus triggering positive Δ^{201} Hg values in final deposition (Chen et al., 2012; Blum et al., 2014). Our results seem point to Hg iso-topes as a 708 709 promising way for identification of the Hg source.

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7. Discussion

712 From the Hg/TOC stratigraphic patterns in this study, one is tempted to outline a preliminary picture of 713 the global Hg loading to the environment during the 750 kyr of Deccan phase-2 volcanic eruptions. However, as organic matter content in the examined sections tend to be very low (<0.2%), reliable 714 715 Hg/TOC ratios are not always obtained and so these ratios should be looked with caution. 716 Three Hg/TOC spikes are depicted from the three examined complete sections. One of them observed 717 within the CF2 foraminiferal biozone in Meghalaya, Højerup and Bottaccione sections (spike-I), in agreement with increased Hg loading to the atmosphere by early stages of Deccan phase-2 eruptions, 718 between 250 and 160 Kyr before the KPg boundary. Too low TOC values at Højerup and Bottaccione 719 resulted at enrichment of the Hg spike I in these two sections. As there is no foraminifer biozone 720 control at Padriciano, we speculate that the prominent Hg enrichment observed at this section (182 721

ng.g-1 of Hg), about one meter below the KPg boundary, is coeval with the spike-I in the Meghalaya 722 section. An apparent Hg spike (16 ng.g⁻¹) at about one meter below the KPg boundary at Bajada del 723 Jagüel, recognized by Sial et al. (2014), disappears after TOC normalization. 724 725 It became evident from this study that a second Hg/TOC spike (spike-II) coincides with the KPg boundary layer at Højerup and Bottaccione. At Meghalaya, presence of an Hg enhancement in the 726 proximity of the KPg boundary becomes evident only after normalization of the Hg concentration to 727 TOC. In this case, a proximal section to the Deccan, the Hg loading was only detected after 728 normalization to TOC. Enrichment of Hg in the KPg boundary at Padriciano, if any, was probably 729 swept off by erosion that pre-ceded deposition of the breccia that marks this boundary. It is uncertain 730 whether the notable Hg/TOC spike in the KPg boundary at Bajada del Jagüel is correlated to spike II or 731 resulted from Hg loading from local volcanism within the Neuqueñ Basin. A third Hg/ TOC spike in 732 the Danian (spike III) at Jhilmili, a very proximal sec- tion at the Deccan volcanic center, Højerup, 733 Bottaccione and Meghalaya within the P1a foraminiferal biozone, seems to be a record of late stages of 734 Deccan phase-2 eruptions within the 220-500 kyr after the KPg boundary. 735 Around 66 Ma ago, the Højerup and Bottaccione sites were located at similar distances relative to both 736 the Deccan Traps and the Chicxulub impact site (Fig. 1), therefore similar-sized Hg peaks would be 737 expected. "Two explanations for the observed discrepancy may be offered: (a) divergence in time 738 739 lengths and/or factors governing Hg deposition, (b) preservation of Hg deposit from weathering and diagenesis. The high level of Hg recorded at the KPg boundary layer at Højerup may have resulted 740 from an increased flux of volcanic-derived Hg from the landmass into the marine realm. At 741 Bottaccione, Hg accumulation likely resulted from a similar mechanism, but a distal, deep marine 742 (pelagic) setting probably determined a lower Hg concentration if compared to the more proximal 743 Højerup section. However, as the KPg boundary layer has been tectonically tilted at Bottaccione, this 744 745 could have also facilitated a gradual Hg leaching, resulting in a more modest remaining Hg

enrichment" (Sial et al., 2014, p. 111e112). One cannot discard the possibility of acid rain leaching of 746 Hg in the aftermath of the KPg boundary event. It is difficult to estimate how much mountain chains, 747 dispersion corridors along latitudinal zones, winds and marine currents have affected the Hg transport 748 and deposition during the KPg transition. Due to climate changes, the environment at that time was 749 probably depleted of organic scav- enging capacity and, therefore, Hg⁺² was likely kept in solution, 750 readily adsorbed onto clays and transported to the sea/ocean. 751 Large differences in Hg peak magnitude, as observed between Højerup and Bottaccione, are also 752 observed in sections bracketing the KPg boundary within a single basin (e.g. Salta and Neuqueñ basins; 753 Sial et al., 2013, 2014). One cannot totally discard the pos-sibility that the Chicxulub asteroid impactor 754 may have carried a large amount of Hg. If Hg concentrations measured in the CI- chondrite Orgueil 755 were typical for the KPg boundary asteroid impactor, full release of highly volatile Hg would 756 correspond to a total injected mass of 10⁶-10⁷ metric tons of Hg according to Meier et al. (2015), that 757 is, 10^4 - 10^5 times higher than present annual anthropogenic emissions, implying a global Hg deposit of 758 several thousand ng Hg/cm². Small Hg peaks recorded at the KPg boundary have been attributed to 759 terrestrial response to the Chicxulub impact by Hildebrand and Boynton (1989) or to volcanic activity 760 pulses (Sial et al., 2013, 2014; Font et al., 2016). Meier et al. (2015) measured the concentration and 761 762 isotopic composition of Hg in meteorites and at some KPg boundary sites, including Højerup, Bidart 763 (France) and Teapot Dome sites (USA). At Teapot Dome, they found a huge, double-spiked Hg peak of ~1000 ng.g⁻¹, and much smaller Hg concentrations at Højerup (200 ng.g⁻¹) and Bidart (80 ng.g⁻¹). The 764 Hg isotopic composition in Hg spikes in sedi- mentary rocks spanning the KPg boundary can 765 potentially help constraining the Hg source. Meier et al. (2015) did not report Hg isotope compositions 766 measured in their study of meteorites and KPg boundary sites to allow further comparison with the Hg 767 isotope data reported in the present study. 768

The δ^{202} Hg data in clays from the KPg boundary layer at Højerup, from the KPg boundary clay layer in the Scaglia Rossa Formation at Bottaccione and from Meghalaya (spike II) besides from Hg in the spike I (Meghalaya and Padriciano) and spike III at Jhilmili lie within the range for volcanogenic Hg (0.00 to -2.00‰) as reported by Bergquist and Blum (2009). The small and positive Δ^{201} Hg (Table 4) also sheds light on the long-range atmospheric transport of volcanic emission.

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8. Conclusions

(a) Three Hg/TOC spikes present in the studied complete sections (Meghalaya, Højerup and 776 Bottaccione) are proposed to represent a record of the Deccan phase-2 eruptions. One of these spikes is 777 situated within the CF2 foraminiferal biozone (e.g. Meghalaya), within the 250 kyr (beginning of 778 Deccan phase-2) and 160 kyr (CF2-CF1biozones boundary) before the KPg boundary after carbon 779 dioxide, sulfuric aerosols and other toxic agents reached a critical threshold. The second spike, at the 780 KPg boundary, is also coeval to the Deccan phase-2 (e.g. Højerup and Bottaccione) and a third one, 781 within the P1a foraminiferal biozone in the lowermost Danian (e.g. Jhilmili), is likely related to late 782 Deccan phase-2 eruptions (within the 220-500 kyr interval after the KPg boundary). These three 783 784 periods of anomalous Hg deposition identified here suggest events of enhanced Hg deposition over 785 broad areas of the globe. 786 b) The possibility that Hg enhancements around the KPg boundary at Højerup, Bottaccione, and 787 Meghalaya, could be postdepostional, resulting from scavenging by anoxia on the seafloor and transported downward into the uppermost 50 cm layer, is not confirmed by the stratigraphic record of 788 Mo/Al redox proxy that does not support a decrease in oxygenation. 789 (c) Differences in the magnitude of the Hg concentrations among the complete studied sections are due 790 to either dif- ference in sedimentation rates, proximity to the continents, or to the partial leaching of Hg 791 792 during weathering and/or diagenesis.

(d) In selecting Hg enriched values that may represent true volcanic Hg loading to the environment, it 793 794 is a common practice to normalize Hg concentrations by the corresponding TOC % values. This 795 approach potentially confirms true Hg enrichments but inaccuracy of measurements may lead to highly variable or unrealistic Hg/TOC spikes in cases where extremely low TOC values (<0.2%) are used (e.g. 796 uppermost Danian portion of the Meghalaya section). 797 (e) The δ^{202} Hg data for boundary clays or from samples of other sedimentary rocks spanning the KPg 798 boundary lie within the range of volcanogenic Hg. Small and positive Δ^{201} Hg seems to support a long 799 range atmospheric transport of Hg. This indicates a promising start of using Hg isotopes in the 800 identification of the Hg source. 801 802 Despite these stimulating results, one cannot assure that Hg is volcanogenic solely based on isotopic similarity because of: (a) limited amount of Hg isotope data, and (b) processes involved as eruption, 803 transportation and even deposition may have lead to Hg isotope fractionation/modification of the 804 805 original Hg isotope signals. We hope this study contributes to the growing agreement that a single large asteroid/comet impact 806 could not have been the sole cause of the end-Cretaceous mass extinction, but rather a contributing 807 factor along with volcanism. 808 Acknowledgments 809 810 We thank Gilsa M. Santana and Vilma S. Bezerra for assistance with stable isotope analyses in the LABISE and to Ingra K.C. Belmino for the help with the Hg analyses at the LABOMAR. We acknowl-811 edge N. Thibault (Copenhagen) for discussions about stratigraphic details of Danish successions. We 812 813 are grateful to Prof. Finn Surlyk (University of Copenhagen) and to an anonymous reviewer whose 814 comments and suggestions on an earlier version of the manuscript greatly contributed to improve it. 815 MKP and VCT are grateful to the Brazilian Council for Scientific and Technological Development

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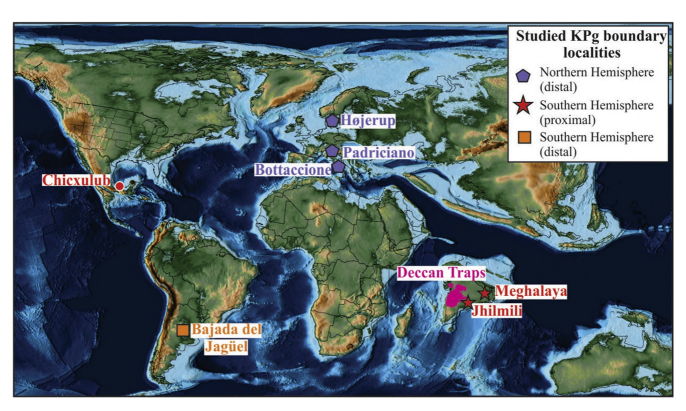


Fig. 1. Paleomap at 66 Ma showing paleogeography and location of the studied KPg boundary sections (based on Scotese, 2013). The location of the Chicxulub impact structure is indicated. Colored stars mark the three distinct types of studied KPg boundary sections in relation to the Deccan volcanic province center: (A) Northern Hemisphere (distal >5000 km); yellow; (B) Southern Hemisphere (proximal to very proximal; up to 1000 km); red; (C) Southern Hemisphere (distal > 5000 km); orange. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

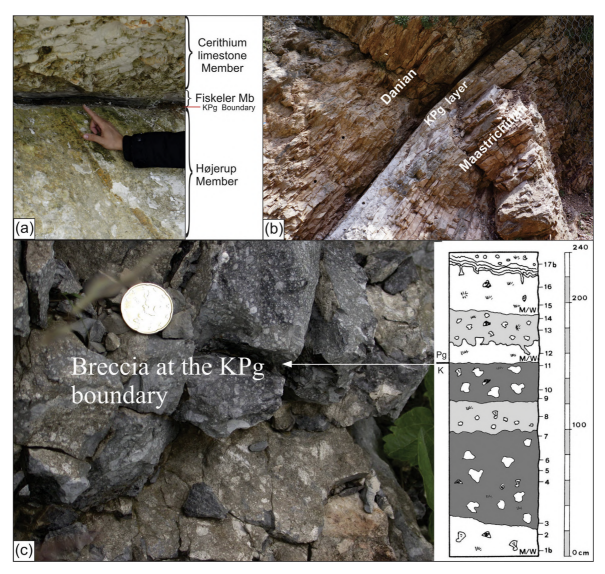


Fig. 2. (A) Closer view of the KPg boundary layer (Fiskeler Member) in the Højerup section, Stevns Klint, Denmark; (B) KPg boundary layer (1–2 cm thick) within the Scaglia Rossa Formation at the Bottaccione Gorge, near Gubbio, Italy; (C) Closer view of the upper breccia level at the KPg boundary in Padriciano, near Trieste, Italy.

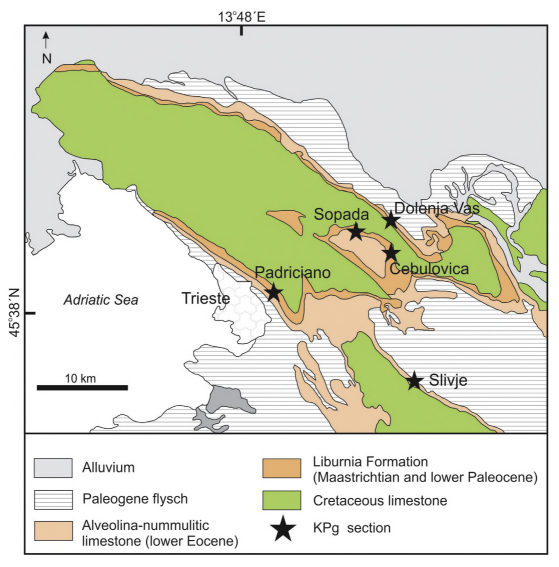


Fig. 3. Summary geological map showing five sites of studied KPg boundary sections in the Trieste Karst region of Slovenia and northeastern Italy (modified from Pugliese et al., 1995). Black stars mark the location of KPg boundary sections.

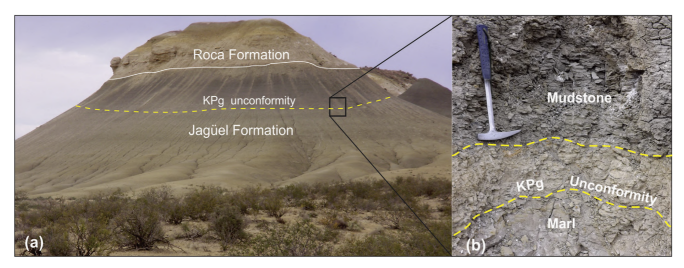


Fig. 4. Yellow volcanoclastic sandstone layer (20 cm thick) marking the KPg boundary in the Bajada del Jagüel section, Neuquén Basin, Argentina. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

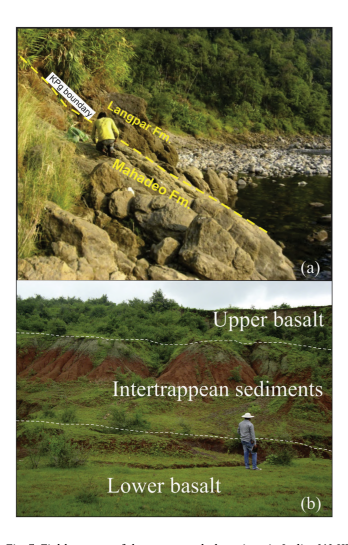


Fig. 5. Field aspects of the two sampled sections in India: (A) KPg boundary location between the Mahadeo and Langpar formations in an Um Sohrynkew River section, not far from Theria village, Shillong Plateau, northeastern India; (B) Jhilmili section showing the two basalt traps and sandwiched intertrappean sedimentary rocks.

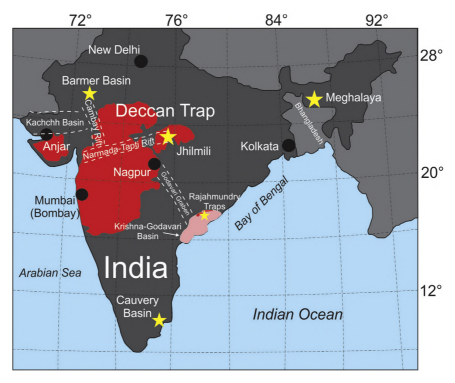


Fig. 6. Geographic map of India showing the location of well-known KPg boundary sections, marked by stars (Meghalaya, Jhilmili, Anjar, and Barmer, Krishna—Godavari and Cauvery basins). Also shown in this map is the Deccan volcanic province and the extent of the Narmada—Tapti seaway (modified from Cripps, 2002; Shukla and Srivastava, 2008; Keller et al., 2009).

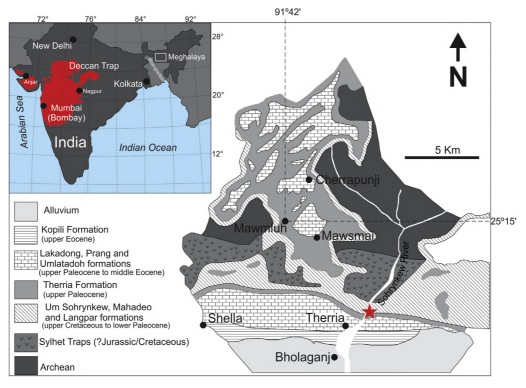
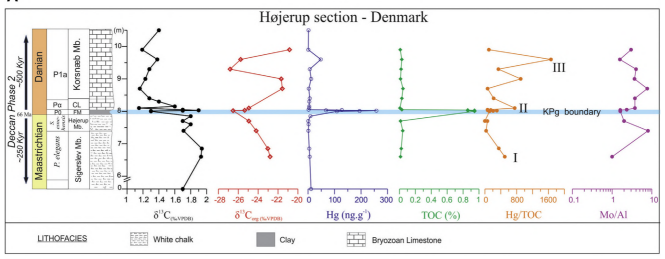
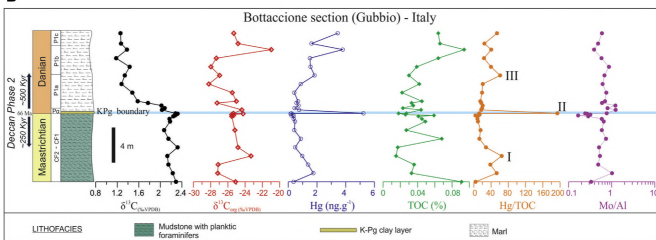


Fig. 7. Summary geological map of South Shillong Plateau, Meghalaya, northeastern India (modified from Tewari et al., 2010a, 2010b). Star indicates locality of the sampled section.





В



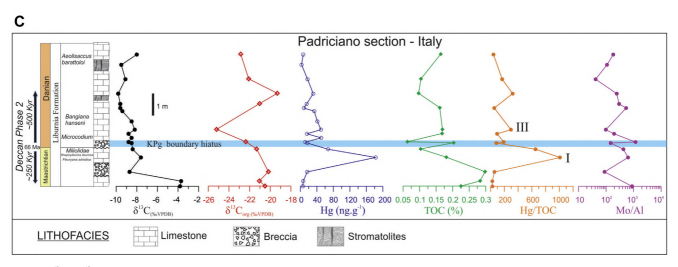


Fig. 8. (A) δ¹³C_{carb}, δ¹³C_{org}, Hg/TOC and Mo/Al variation patterns across the KPg boundary at Højerup, Stevns Klint (Maastrichtian planktic foraminiferal biostratigraphy from Surlyk et al., 2006, and Danian, from Rasmussen et al., 2005); (B) Bottaccione (Gubbio) section (planktic foraminiferal biostratigraphy from Coccioni et al., 2010; Coccioni and Premoli Silva, 2015) and (C) Padriciano section (foraminiferal biostratigraphy from Tewari et al., 2007), Italy.

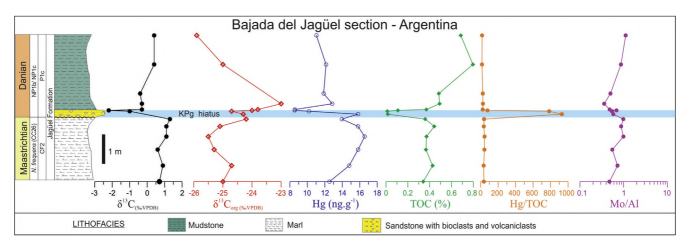


Fig. 9. δ¹³C_{carb}, δ¹³C_{carb}, δ¹³C_{org}, Hg/TOC and Mo/Al variation patterns across the KPg boundary in a section at Bajada del Jagüel, Neuquén Basin, Argentina (biostratigraphy according to Keller et al., 2007).

Table 1
C and O isotopes (% VPDB), total organic carbon (TOC %) and Hg (ng·g⁻¹) across the KPg boundary in sections at Højerup (Denmark), and Bottaccione and Padriciano (Italy).

(a) Højerup section, Stev	vns Klint, Denm	ark												
Formation	Member			Sample	Height (cm)	δ ¹³ C %	vPDB δ ¹³	Corg ‰ VPDI	B Hg (ng⋅g ⁻	1) TOC(%)	Hg/TOC	Mo (ppm)	Al (%)	Mo/Al
Danian	Korsnæb Mem	ber		SKc	1050	1.4		20.05	1.8	-	-	_	_	_
Stevns Klint Formation				N7 N6	990 960	1.19 1.38		20.85 25.76	1.0 48.2	0.010 0.029	100.0 1662.06	0.06 6 0.11	0.02 0.07	3.00 1.57
				N5	930	1.28		26.82	6.8	0.029	340.00		0.07	4.00
				N4	900	1.23		21.68	11.7	0.013	900.00		0.03	3.66
				N3	870	1.16	-2	21.57	3.0	0.044	68.18	0.23	0.03	7.66
				N2	840	1.28	_		6.6	0.030	220.00	0.15	0.04	3.75
Danian	Cerithium Lim	estone Me	mber	D3	830	1.4	_		4.55	-	_	-	_	_
Rødvig Formation				D2	815	1.6	_	24.04	4.51	-	750.00	- 0.15	-	_ 2.75
				N1 No. 3	810 805	1.15 1.7		24.94	4.5 2	0.006 0.030	66.66	0.15 0.14	0.04 0.06	3.75 2.33
	Fiskeler Memb	er (KPg ho	undary)	FC-1	804	_	_		127.72	0.859	148.68		1.14	1.60
	i ibheiri memb	er (ra g bo	undury)	FCD	803	1.9	-2	25.36	67.9	0.944	71.92		2.28	1.60
				FCC	802	1.7		26.52	257.94	0.859	300.27		2.28	1.60
				FCB	801	1.3	_		194.53	0.944	206.06	3.67	2.28	1.60
				FCA	800	1.3	_		108.76	0.944	115.21	3.67	2.28	1.60
Maastrichtian	Højerup Memb	oer (gray cl	nalk)	M2	785	1.8	_		9.09	_	-	_	_	-
Møns Klint Formation				N (-1)	770	0.71		24.94	-	0.014	-	0.02	0.01	2.00
				M1A	770 760	1.7	_		0.88	0.014	62.85	0.02	0.01	2.00
				No. 2 N (-2)	760 740	1.8 1.71	_	24.20	2.3 <1.26	- 0.041	- 30.73	0.08	0.01	8.00
				N (-2) N (-3)	685	1.71		24.20 23.03	<1.26 4.6	U.U4 I —	30./3 _	- 0.00	U.U I —	8.00 —
				N (-4)	660	1.93		22.80	6.5	0.015	433.3	0.01	0.01	1.00
	Sigerslev Mem	ber (white	chalk)	No. 1	0	1.7			11.3	-	-	-	_	-
(b) Bottaccione section,			,			-								
Lithology	-	Sample	Height	(cm) δ ¹	³ C _{carb} ‰	VPDB	δ ¹³ C _{org} %	vPDB I	Hg (ng⋅g ⁻¹)	TOC (%)	Hg/TOC	Mo (ppm)	Al (%)	Mo/Al
Marl	_	GP-28	1685.4		26		-25.32		3.44	0.064	53.75	0.19	0.32	0.59
		GP-27	1565.4		27		-24.82		1.62	0.066	24.54	0.15	0.31	0.48
		GP-26	1495.4		38		-20.97		3.79	0.094	40.31	0.16	0.41	0.39
		GP-25	1395.4	1.	18		-27.34	1	1.51	0.064	23.59	0.18	0.31	0.58
		GP-24	1295.4	1.	43		-27.97	1	1.55	0.039	39.74	0.19	0.22	0.86
		GP-23	1195.4	1.	34		-26.96		1.83	0.030	61.00	0.17	0.25	0.68
		GP-22	1095.4		28		-28.21		0.87	0.042	20.71	0.24	0.39	0.61
		GP-21	995.4		48		-25.49		0.42	0.022	19.09	0.15	0.20	0.75
Manil (almost a italia a a a al-		GP-20	895.4	1.58 1.77			-25.02			0.045			0.30	0.70
Marl (rhytmites; carbon		GP-19 GP-18	875.4		02		-27.25		0.57	0.033	17.27 20.57	0.34	0.57	0.91
decreases in the Dani	IdII)	GP-18 GP-17	845.4 815.4		02 08		_).72).44	0.035 0.023	19.13	0.39 0.12	0.32 0.15	1.21 0.80
		GP-16	795.4		03		-24.39		0.75	0.023	17.04	0.12	0.17	1.23
Greenish brown clay lay	/er	GP-15		oundary la		6.78	250		5.23	0.027	193.70	0.49	0.64	0.29
Mudstone with planktic		GP-14	794		28		-25.36		0.22	0.018	12.22	0.09	0.37	2.67
•		GP-13	790		32		-24.24		0.16	_	_	0.07	0.26	0.26
		GP-12	780	2.	29		-25.55	C	0.35	0.026	13.46	0.08	0.25	0.32
		GP-11	770		27		-25.35		0.22	0.059	3.72	0.05	0.30	0.16
		GP-10	760		24		-25.55		0.37	0.041	9.02	0.06	0.21	0.28
		GP-9	730		17		_		0.38	0.045	8.44	0.20	0.34	0.58
		GP-8 GP-7	700		18 07		- -25.19).47).42	0.049 0.027	9.59 15.55	0.17 0.23	0.26 0.39	0.65 0.58
		GP-7 GP-6	600 500		07 15		-25.19 -).42).88	0.027	15.55 12.94	0.23	0.39	0.58
		GP-5	400		32		_ _24.83).48	0.008	28.23	0.17	0.59	0.73
		GP-4	300		13		-23.32		0.97	0.017	64.66	0.23	0.43	0.53
		GP-3	200		15		-27.13		1.38	0.036	38.33	0.18	0.37	0.48
		GP-2	100		23		-27.18		1.75	0.033	53.03	0.58	0.57	1.01
		GP-1	0	2.	29		-25.11	C	0.37	0.091	4.06	0.26	0.78	0.33
(c) Padriciano section, It				. 12		. 42								
Stage Forma			ght (m)	δ ¹³ C %	o VPDB	_	rg ‰ VPDI			, 0,				Mo/Al
Danian Liburn		6.4		-8.0 9.5		-22.8	66	6.8	0.164				01	182
	VT 15 VT 14	5.9 5.2		-9.5 -9.1		- -22.0	10	2.9 17.5	- 0.104	- 169			01 08	108 41.62
	VT 14 VT 13	4.5		-9.1 -9.8		-22.0 -19.3		31.1	0.104				08 02	244.50
	VT 12	4.3		-9.6		-19.3		15.2	-	_			01	297
	VT 11	3.8		-9.6		-21.0	-	8.6	0.161	53			01	536
	VT 10	3.6		-9.4		_		33.6	-	-	_	_		-
	VT 9	3.1		-8.5		_		41.2	_	_	_	_		_
	VT 8	2.7		-8.2		-25.2	.0	49.9	0.169	295	5.26 1.	.96 0.	02	98.00
	VT 7	2.5		-8.8		_		15.9	0.168				01	196
	VT 6	2.3	5	-8.5		_		50.5	_	_	_	_		_
	VT 5	2.1	5	-8.8		-22.3	7	12.0	0.063	190	0.47 11.	.79 0.	01	1179.00
												(continu	ed on ne	xt page)

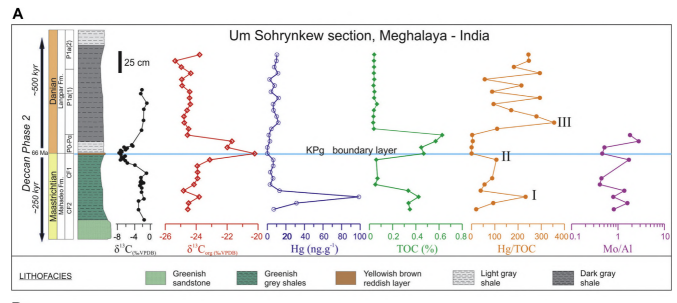
Table 1 (continued)

(c) Padriciano section, Italy											
Stage	Formation	Sample	Height (m)	δ^{13} C ‰ VPDB	δ ¹³ Corg ‰ VPDB	Hg $ng \cdot g^{-1}$	TOC (%)	Hg/TOC	Mo (ppm)	Al (%)	Mo/Al
KPg boundary		KPg breccia	2.1	-8.5	_	16.8	0.203	82.75	1.47	0.01	147
Maastrichtian		VT 4	1.8	-8.4	-21.32	67.3	0.103	653.39	8.52	0.02	426.00
		VT 3	1.4	-7.6	_	182.5	0.181	1008.28	25.16	0.04	629.00
		VT 2	0.7	-8.7	-20.18	16.7	0.298	56.04	0.88	0.01	88
		VT 1	0.25	-3.7	-21.05	6.9	0.283	24.38	1.17	0.00	0
		VT 0	0	-3.8	-20.51	5.9	0.225	26.22	8.76	0.01	876

Table 2
C- and O-isotope analyses (% VPDB), total organic carbon (TOC, %), Hg (ng·g⁻¹), Mo (ppm) and Al (%) in a KPg section at Bajada del Jagüel, Argentina.

(a) Bajada del Jagü	iel section, Neuquén Basin, Ai	gentina									
Formation	Lithology	Sample	Height (m)	$\delta^{13}C_{\text{$\%$ VPDB}}$	$\delta^{13}C_{org} \ _{WPDB}$	$Hg (ng \cdot g^{-1})$	TOC (%)	Hg/TOC	Mo (ppm)	Al (%)	Mo/Al
Jagüel Formation	Mudstone	KT 12	5.03	0.4	-25.9	11.04	0.685	16.11	1.92	1.72	1.11
		KT 11	4.03	0.4	-25.0	12.14	0.796	15.25	1.07	1.23	0.86
		KT10	3.03	-0.4	_	11.87	0.484	24.52	0.78	1.56	0.50
		KT9	2.68	-0.3	-23.0	12.85	0.491	26.17	0.55	1.53	0.35
		KT 8	2.48	-0.3	-23.8	8.57	0.370	23.16	0.61	1.27	0.48
	Sandstone with	KT7	2.45	-2.2	-24.0	8.55	0.109	78.44	0.75	1.09	0.68
	bioclasts and volcaniclasts	KT 6	2.42	-1	-24.7	10.19	0.013	783.84	0.57	1.05	0.54
		KT 5	2.32	-0.6	-24.3	15.80	0.017	929.41	0.75	1.29	0.58
	Marl	KT 4	2.15	1.3	-24.2	13.95	0.359	38.85	1.17	1.19	0.98
		KT 3	1.9	1.1	-25.1	15.8	0.442	35.74	1.36	1.55	0.87
		KT 2	1.55	1.1	-25.5	16.56	0.371	44.63	1.09	1.10	0.99
		KT 1	1.1	0.6	-25.3	15.81	0.366	43.19	0.59	1.07	0.55
		KT0	0.55	0.9	-24.7	14.79	0.427	34.63	0.74	1.02	0.72
		KT (-1)	0	0.7	-25.0	12.53	0.341	36.74	0.56	1.18	0.47

(a) Meghalaya section, Um	n Sohrynkew, northeas	tern India.										
Formation	Lithology	Sample	Height (cr	n) δ ¹³ C ‰ _V	PDB δ ¹³ Corg	‰ _{VPDB}	Hg ng∙g	g ⁻¹ TOC (%) Hg/TO	C Mo (pp	m) Al (%	(s) Mo/
Langpar (Danian)	Light gray shale	KT 25	10	0.1	-23.78	;	10.3	0.042	245.23			_
	Dark gray shale	KT 24	10	1.1	-25.36	i	9.8	0.039	251.28	3 –	-	_
			10	0.4	-24.96		7.4	0.040	185.00		_	_
			10	0.6	-24.35		11.5	0.039	294.87		_	_
			10	-0.1	-24.93		2.4	0.042	57.14		-	_
			10	0.4	-24.91		7.8	0.036	216.66		_	_
			10	-0.1	-24.42		3.9	0.044	88.63		_	
			10	-0.1	-24.43		12.1	0.041	295.12		_	_
			10	0.6	-24.38		6.2	0.065	95.38		_	_
			10 10	1.3 1.4	-24.59		6.7 10.0	0.039	171.79 285.71		_	_
			10	1.4	-24.79 -24.77		11.8	0.035 0.033	357.57		_	_
			10	-0.0	-24.77 -24.49		4.3	0.033	110.25		_	_
			10	-0.0 -0.1	-24.43		1.6	0.627	2.55		1.13	1.81
	Light gray shale		10	0.1	-21.69		2.8	0.571	4.90		1.05	2.79
	Light gruy shale		10	0.1	-21.99		0.0	0.447	0.00		1.39	0.51
			10	0.5	-20.22		0.0	0.468	0.00		1.16	0.45
KPg boundary layer	Yellowish brown to reddish layer		10	-8.13	-23.12		6.3	0.059	16.94		1.81	1.71
Mahadeo (Maastrichtian)	Greenish sandstone	KT8	10	-0.4	-23.93		6.3	_	_	_	_	_
,	and shale		10	0.4	-23.91		3.2	_	_	_	_	_
		KT6	10	-0.0	-23.91		6.2	0.070	88.57	0.66	1.50	0.44
		KT5	10	0.7	-24.14	ļ	2.9	0.052	55.76	0.59	1.45	0.40
		KT4	10	0.2	-24.82	!	13.1	0.335	39.10	1.67	1.22	1.36
		KT3	10	0.5	-23.81		98.6	0.424	232.54	1.21	1.53	0.79
		KT2 KT1	10 10	0.3 1.0	-24.49 -24.54		31.4 6.9	0.337 0.349	93.17 19.77		1.18 1.12	1.59 0.81
(b) Jhilmili Section, centra	l India											
	Lithology		Sample	Height (m)	δ ¹³ C ‰ vpns	$\delta^{13}C_{org}$	% VPDR	Hg ng∙g ⁻¹	TOC (%) I	Hg/TOC M	Io Al (%) Mo/
					VI DD						opm)`	
Upper Basalt Trap	2.5 m thick basaltic fl		JMR 20		_	-20.73		12.4	0.030	413.33 0		
Paleosol Section (Danian)	Greenish clay layer w	rith	JMR 19		-3.7	-28.27		15.3	0.093		.07 1.97	
	calcitic veins		JMR 18		-5.6	-25.44		12.3	0.060		.04 1.74	
	Soft ferricrete fine-gra			0.5	-3.3	-20.73		8.7	0.041	212.19 0		
	brick red with yellow	isn tint	JMR 16		-2.6	-23.33		100.2		3131.25 0		
				0.5	-2.6	- 24.45		19.2	0.036		.01 2.23	
	Claustone with lamin	an faatumaa	JMR 14		-3.20	-24.45		9.9	0.024 0.067		.03 2.22 .07 2.29	
	Claystone with lamin- grayish brown in colo		JMR 13 JMR 12		-3.5 -3.3	-25.53 -24.92		17.7 6.2	0.045	264.17 0 137.77 0	.07 2.23 .64 2.62	
	relatively harder	Λ,	-	0.4	-3.3 -4.7	-24.32 -23.26		14.8	0.043		.68 1.65	
	•	rained		0.0	-4.7 -4.1	-23.20 -		51.8			.36 3.07	
	Soft ferricrete, fine- grained, brick red with yellowish tint		JMR 09		-6.0	-23.38		14.5	0.035	414.28 0		
			JMR 08		-8.0	-25.31		11.8	0.056		.07 3.94	
				0.5	_	-20.33		19.0	0.062		.05 3.79	
	Hard sediment layer	within the	JMR 06		-11.0	-24.68		14.5	0.053		.60 2.03	
	paleosol with clasite g		JMR 05		-12.1	-24.41		24.1	0.074	325.67 0		
	Lowest layer of paleo	sol ferricrete	JMR 04	0.5	-3.4	-24.43		20.7	0.046	450.00 0	.28 2.7	0.10
	with calcitic veins		JMR 03		-9.9	-22.97		11.8	0.084	140.47 0	.03 3.88	0.00
Lower Basalt Trap	2.0 m thick basaltic fl		IMR 02	_	_			_	0.071 -	- 0	.04 1.48	0.02



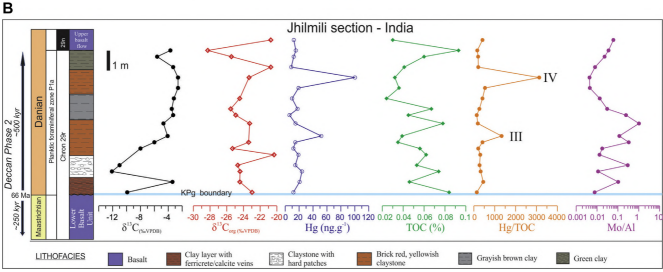


Fig. 10. δ¹³C_{carb}, δ¹³C_{org}, Hg/TOC and Mo/Al variation patterns across the KPg boundary at: (A) Um Sohrynkew River section (Meghalaya), south Shillong Plateau, northeastern India (δ¹³C_{carb} from Gertsch et al., 2011; planktic foraminifera biostratigraphy from Mukhopadhyay, 2008; Pal et al., 2015) and (B) Jhilmili section, Madhya Pradesh, central India (planktic foraminiferal biostratigraphy from Keller et al., 2009).

Table 4
Hg isotopes (% relative to NIST SRM 3133) in samples with Hg enrichments: spike I (Meghalaya, Padriciano); spike II (Højerup, Bottaccione, Meghalaya), and Bidart (France; for comparison); spike III (Jhilmili), besides samples from across the KPg boundary at Bajada del Jagüel.

Samples	δ^{199} Hg	δ^{200} Hg	δ^{201} Hg	δ^{202} Hg	Δ^{199} Hg	Δ^{200} Hg	Δ^{201} Hg		
Mass dependent fraction	ation (MDF)				Mass independent fractionation (MIF)				
(a) Fiskeler Member (KPg	g boundary layer), De	enmark							
FCA	-0.51	-1.06	-1.76	-2.34	0.08	0.11	-0.01		
FCB	-0.28	-0.74	-1.18	-1.62	0.13	0.07	0.04		
FCC	-0.29	-0.55	-0.86	-1.06	-0.02	-0.02	-0.06		
FCD	-0.43	-0.92	-1.44	-1.93	0.06	0.05	0.01		
(b) Bottaccione, Italy (KPs	g boundary layer)								
GP-15	-0.26	-0.59	-0.93	-1.28	0.06	0.05	0.03		
(c) Meghalaya, India									
KT-3	-0.35	-0.78	-1.16	-1.61	0.06	0.03	0.05		
KT-bulk	-0.64	-0.94	-1.58	-1.89	-0.16	0.01	-0.16		
(d) Jhilmili, India									
JM-16	-0.60	-0.57	-0.98	-1.01	-0.34	-0.06	-0.22		
JM-10	-1.01	-1.12	-1.92	-2.18	-0.46	-0.02	-0.28		
(e) Padriciano, Italy									
VT-3	-0.08	-0.69	-1.00	-1.38	0.27	0.00	0.04		
(f) Jagüel Formation, Neu	quén Basin, Argentir	าล							
K-T-12	0.07	-0.29	-0.29	-0.62	0.22	0.02	0.18		
K-T-11	-0.08	-0.67	-0.89	-1.41	0.27	0.04	0.17		
K-T-10	-0.14	-0.59	-0.77	-1.17	0.16	-0.01	0.10		
K-T-9	-0.13	-0.59	-0.76	-1.18	0.17	0.00	0.13		
K-T-8	-0.06	-0.59	-0.74	-1.15	0.13	-0.01	0.12		
K-T-7	-0.08	-0.52	-0.68	-1.00	0.17	-0.02	0.08		
K-T-6	-0.10	-0.48	-0.61	-0.87	0.12	-0.05	0.04		
K-T-5 (KPg layer)	-0.14	-0.57	-0.73	-1.05	0.13	-0.05	0.06		
K-T-4	-0.03	-0.36	-0.49	-0.77	0.17	0.02	0.09		
K-T-3	-0.12	-0.60	-0.88	-1.27	0.20	0.04	0.07		
K-T-2	-016	-0.61	-0.84	-1.22	0.15	0.01	0.08		
K-T-1	-0.12	-0.50	-0.68	-1.04	0.14	0.02	0.10		
(g) Bidart, France									
KPg boundary layer	-0.04	-0.34	-0.42	-0.74	-1.12	-0.15	0.03		
BI-9.28.5	-0.11	-0.57	-0.66	-1.44	-1.56	0.25	0.15		
BI-9.34.2	0.05	-0.11	-0.13	-0.31	-0.73	0.12	0.04		
BI-9.36.2	0.08	-0.10	-0.07	-0.25	-0.61	0.15	0.03		
BI-10.35.1	-0.33	-1.28	-1.68	-2.66	-4.30	0.33	0.06		
BI-11.7.3	0.13	-0.16	-0.04	-0.66	-0.73	0.30	0.17		

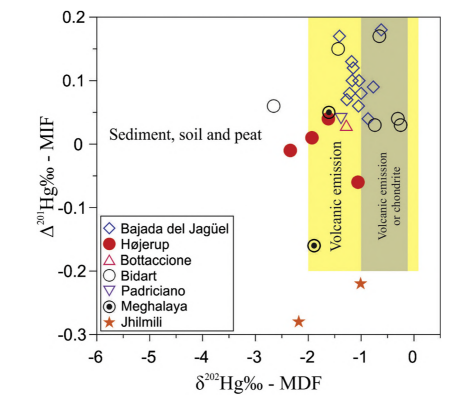


Fig. 11. In a δ^{202} Hg (MDF)— Δ^{201} Hg (MIF) plot, values most samples in this study lie within the range for volcanogenic Hg. Ranges for volcanogenic and chondritic Hg are from Bergquist and Blum (2009) and are shown as vertical bars.