



High-resolution Hg chemostratigraphy: A contribution to the distinction of chemical fingerprints of the Deccan volcanism and Cretaceous–Paleogene Boundary impact event



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ABSTRACT

There is a renewed interest in volcanism as the major trigger for dramatic climatic changes at the Cretaceous–Paleogene transition (KTB), which were accompanied by a decrease in biodiversity and mass extinction. We have used Hg contents as proxy for volcanic activity at the classical localities of Gubbio (Italy) and Stevns Klint (Denmark) where the KTB layer is easily recognizable, and at a near-complete succession exposed at the Bajada del Jagüel locality in the Neuquén Basin, Argentina. These three localities display similar $\delta^{13}\text{C}_{\text{carb}}$ trends with markedly negative excursion at the KTB layer. Bulk-rock oxygen isotopes yielded similar pathways across the KTB layers in these localities and, if considered near-primary, the negative $\delta^{18}\text{O}$ excursion at the KTB in Gubbio and Bajada del Jagüel suggest warming temperatures during this transition, whereas the negative excursion immediately followed by positive one at Stevns Klint points to a cycle of warm followed by colder climate. At Stevns Klint, Hg contents reach 250 ng g^{-1} within the KTB layer (Fiskeler Member) and 45 ng g^{-1} at 1.5 m above that, while within the Scaglia Rossa Formation at Gubbio, three Hg peaks across the KTB are observed, one of them within the KTB layer (5.3 ng g^{-1}). Hg shows several peaks across the KTB in the Neuquén Basin, with up to 400 ng g^{-1} in the Jagüel Formation. The phenomena that caused dramatic changes at the KTB probably expelled huge amounts of Hg into the atmosphere as recorded by these high Hg levels. A co-variation between Hg and Al_2O_3 in the studied sections suggest that Hg is adsorbed onto clays. Hg concentrations and also Hg isotopes are perhaps a powerful tool in the assessment of the role of volcanic activity during extreme climatic and biotic events, and in assessing the role of meteorite impact versus volcanism as the predominant cause of past global catastrophes and mass extinction.

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1. Introduction

It is well known that major geologic time boundaries in the Phanerozoic are marked by dramatic changes in the geological record, including biological extinction, fluctuations of sea-level and atmospheric composition. These changes are reflected in sharp change in biotic activity and included major mass extinction at the Permian–Triassic and

Cretaceous–Paleogene transitions, and also by the chemical composition of sediments.

It is well known that major catastrophic events concurrent to massive mass extinction occurred in the Phanerozoic. Magaritz (1989) observed that $\delta^{13}\text{C}$ values drop from a high level, reaching a minimum after the Ediacaran–Cambrian, Permian–Triassic and Cretaceous–Paleogene transitions, and then increase to a new level. He proposed that these changes reflect variations in the exogenic carbon cycle that may correlate with variations in the total biomass. Recently, it was proposed that the variation in Hg abundances may correlate with some major geologic time boundaries such as the Permian–Triassic (PTB) (Sanei et al.,

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2012) and the Cretaceous–Paleogene (KTB) transitions (Sial et al., 2013).

Since the early 1980s, after the work by Alvarez et al. (1980), the KTB mass extinction has been often ascribed to an impact of a huge meteorite that generated the large Chicxulub multi-ring crater in the Yucatán Peninsula, Mexico. Some studies, however, have raised the possibility that such a colossal impact may have predated the Cretaceous–Paleogene transition (e.g. Keller et al., 2004, 2009). However, Renne et al. (2013) proposed on the basis of ^{40}Ar – ^{39}Ar dating of tektites and bentonite beds that the KTB mass extinction and the Chicxulub impact were synchronous to within 32,000 years. Alternatively, the Deccan volcanism whose second phase, the world's longest lava flows (Self et al., 2008), has been correlated with the KTB at 65.5 Ma, spanning the magnetochrons 29r and 29n of the geomagnetic polarity time scale (Chenet et al., 2007; Hooper et al., 2010), has been regarded as responsible for this mass extinction (Keller et al., 2008). Rapid climate changes, sea-level fluctuations and volcanism during 100,000 years preceding the Cretaceous–Paleogene transition have also questioned the bolide impact theory as being solely responsible for the mass extinction (e.g. Keller, 2010, 2011; Gertsch et al., 2011 and references therein). The renewed interest in volcanism as the major trigger of the KTB crisis has stimulated Nascimento-Silva et al. (2011, 2013) and Sial et al. (2013) to initiate a preliminary investigation using Hg as a proxy of volcanism in the sedimentary record across this chronological time boundary. Major tools for this is the geochemical behavior of some elements and their isotopic ratios such as C, O and Sr isotopes, well-established tools in reconstructing past climate and sea water compositions.

Ocean acidification has been claimed as an additional factor contributing to the decline in biodiversity just before and during the KTB. Deccan eruptions likely caused acid rains in an extensive area of the globe, resulting in weathering and dissolution effect on the continental surface. Dissolution of foraminifera (Gertsch et al., 2011) and iron oxide minerals (bio- and detrital magnetite) dissolution are, perhaps, important evidence of the acidification of the oceans during this critical period (Font et al., 2014).

This report aims at extending and better understanding the use of Hg as a volcanogenic trace-element in identifying whether or not volcanism has played significant role in climatic reorganization during the KTB and on the most famous and persistent paleontological murder riddle. Hg concentrations and C-isotope chemostratigraphy, in parallel, are traced along the two classical KTB localities of Gubbio and Stevns Klint where the KTB clay layer is well preserved, and substantial evidence for a meteorite impact has been put forward by Alvarez et al. (1980). In addition, this study also focuses on the sedimentary transition within the Neuquén Basin, Argentina, a locality with a near-complete KTB stratigraphic record in South America, thus expanding the results of previous Hg investigations on the Yacoraite Formation (Salta Basin, Argentina) and Paraíba Basin, Brazil (Nascimento-Silva et al., 2011; Sial et al., 2013). Previous studies have not found the influence of meteoritic impact ejecta on the sedimentary sequences bracketing the KTB in South America (e.g. Barrio, 1990; Marquillas et al., 2003, 2007, 2011; Aberhan et al., 2007; Keller et al., 2007; Nascimento-Silva et al., 2013; Sial et al., 2013 and references therein). However, Scasso et al. (2005) have referred to the presence of tsunami deposits at the KTB in the Neuquén Basin.

We add new Hg analyses to the preliminary data for the KTB layer (Fiskeler Member) at Stevns Klint, previously reported (Sial et al., 2013). We also report preliminary Hg-isotope data for this KTB site, as a further approach to assessing the role of meteorite impact versus volcanism as the major cause of the catastrophe and concurrent mass extinction during the KTB.

2. The Cretaceous–Paleogene transition

Substantial evidence for a giant bolide impact on the Earth's surface coincident with the best-studied global biotic event, the Cretaceous–

Paleogene mass extinction event (~65.5 Ma), has been built up for over the last three decades (Schultze et al., 2010 and references therein). The importance of this impact (or impacts) as the sole extinction mechanism, however, is still a matter of debate.

Anomalously high iridium and platinum-group element concentrations in almost one hundred KTB sites worldwide (e.g. Alvarez et al., 1980; Claeys et al., 2002), presence of glass microspherules (Smit and Klaver, 1981; Smit, 1999) and shocked quartz (Bohor et al. 1984; Bohor, 1990; Morgan et al., 2006; Kamo et al., 2011) have been used to argue in favor of the bolide impact hypothesis as the mechanism responsible for the KTB mass extinction. Presence of high temperature/high pressure (HTP) phase of fullerenes (complexes C molecules) in deposits supposedly associated with events involving an impact of a large bolide (e.g., Chicxulub and Bedout craters; Becker et al., 2000a) provides further evidence for a crucial role of bolide impacts in the mass extinction. Parthasarathy et al. (2008) reported the presence of the natural toluene-soluble fullerene and the toluene-insoluble high-pressure/temperature phase of fullerene C_{60} in carbonaceous matter extracted from iridium-rich intertrappean sediments of the Anjar KTB site, India. The conditions of high-pressure/temperature regimes required for the formation of HTP fullerene phases can be created by an energetic impact event (Parthasarathy et al., 2008). This has led these authors to suggest that the occurrences of such phases in the KTB layer at Anjar could be linked to a contemporaneous bolide impact. The cage structure of fullerene has the unique ability to encapsulate and retain noble gases and this has allowed one to determine that Allende and Murchison meteorites and KTB fullerenes contain trapped noble gases (He, Ar, Xe), the isotopic composition of which can only be described as typical of extraterrestrial origin (e.g. Stevns Klint; Becker et al., 2000a, 2000b; Anjar, Parthasarathy et al., 2008). Denne et al. (2013) have reported a KTB massive deposit in deep waters of the Gulf of Mexico that substantiates widespread slope failure induced by the Chicxulub impact and that provides further evidence of a single impact coincident with the KTB mass extinction.

However, some Maastrichtian–Danian deposits in several basins around the globe show prominent geochemical and isotopic anomalies preceding the KTB. Analyses of stratigraphic variations of whole-rock elemental concentrations and stable isotopic compositions in some of these basins have led to detection of Ba anomalies far below the KTB in the Cauvery Basin (India), in Israel, in northeastern Mexico and in Guatemala (Ramkumar et al., 2004, 2005), and Hg anomalies across the KTB in the Yacoraite Basin, Argentina, and in the Paraíba Basin, Brazil (Sial et al., 2013). Stüben et al. (2005) reported the presence of bentonite layers and Pt and Pd-dominated PGE anomalies below and above the KTB in Mexico as an indication of volcanic activity. A complete shallow-marine succession with preserved KTB clay layer reported by Racki et al. (2011) at Lechówka near Chełm, southeastern Poland, shows anomalous amounts of iridium (9.8 ng g^{-1}), gold, nickel, and elevated Ir/Au ratios. These signals are consistent with a chondrite meteoritic origin, but a major positive iridium spike in Maastrichtian marls has been found 10 cm below the thin KTB clay layer at this location. Keller et al. (2004) also reported iridium enrichment in Maastrichtian sediments at about 20 cm below the KTB in the Yaxcopoil-1 core drilled within the Chicxulub crater, leading these authors to assume that the Chicxulub bolide impact predated the KTB mass extinction.

Multiple iridium-enriched layers have been observed at some KTB sections (e.g. McLean, 1985; Donovan et al., 1988; Graup and Spettel, 1989) in contrast to most reported KTB sites. Such a distribution of iridium anomalies implies episodic, iridium-delivering events over an extended period of time. In KTB sites where there is no evidence for a bolide impact (e.g. Lattengebirge, Bavarian Alps), the multiple iridium-enriched layers probably had a common volcano source, according to Graup and Spettel (1989). The iridium enrichment observed in airborne particles from the Kilauea volcano (Zoller et al., 1983) substantiates the hypothesis that volcanism could be the iridium source.

Investigating the vertical extent of the iridium anomaly at the Bottaccione Gorge, Gubbio, over 2 m below the KTB and 2 m above, Rocchia et al. (1990) observed that iridium concentrations stand above background over almost 3 m of section (~500,000 years) indicating a protracted duration of the source of iridium on top of which the main KTB anomaly proper stands.

Multidisciplinary investigations of sediments pertaining to the KTB exposed at the Brazos River in Falls County, Texas, discovered the existence of an impact spherule-rich sandstone layer (Adatte et al., 2011). Based on biostratigraphy and the $\delta^{13}\text{C}$ shift, the KTB at this site is located up to 1 m (50–100 ky) above this sandstone horizon and is inconsistent with a single catastrophic bolide impact on Yucatán and concurrent megatsunami deposits. The primary Chicxulub ejecta layer, according to these authors, lies about 40–65 cm below the sandstone horizon in a 3-cm-thick yellow clay layer that consists of altered impact glass interbedded in claystones.

Occurrences of pre-KTB geochemical anomalies have raised questions on the relative timing of biotic and isotopic events near this boundary and their relevance to the impact scenario (e.g., Kaminski and Malmgren, 1989; Keller, 2010), supporting the view of multi-causal stepwise extinction of biota across the KTB. Ramkumar et al. (2005, 2010) have interpreted Ba anomalies as well as two $^{87}\text{Sr}/^{86}\text{Sr}$ anomalies preceding the KTB exposed in the Cauvery Basin as the result of increased detrital influx caused by sea-level and climatic changes, and Deccan volcanism. Sial et al. (2013) suggested a possible link between the Hg anomalies across the KTB from their study areas in South America, and the Deccan volcanism. It seems that these geochemical records are in conformity with gradually increasing environmental stress and deterioration during the Maastrichtian, rather than to a sudden singular asteroid impact on a global scale. Carbonate clumped isotope paleothermometry reported by Tobin et al. (2014) demonstrated a changing environment over the last 300 ky of the Cretaceous, with ecosystems becoming more susceptible to an abrupt event like a bolide impact. Stinnesbeck et al. (2012) revisited several Maastrichtian sections in Latin America and verified that in all of them the disappearance of ammonites was completed prior to the end of the Maastrichtian, therefore seemingly independent of an asteroid impact at, or near, the end of the Cretaceous. This finding is in consonance with previous investigations on mass extinction across the KTB in Western European Tethys where the extinction of ammonite species and inoceramid bivalves could have resulted from a gradual event that predated the KTB (Marshall and Ward, 1996).

The López de Bertodano Formation, Seymour Island, Antarctic Peninsula, exhibits one of the most expanded known KTB sections (Tobin et al., 2012). Using magnetostratigraphy and isotopic data from carbonate-secreting macrofauna, these authors have presented a high-resolution, high-latitude paleotemperature record spanning the KTB interval. They have found two prominent warming events synchronous with the three main phases of Deccan Trap flood basalt volcanism. The onset of the second one is contemporaneous with a local extinction that predates the bolide impact. These findings have led Tobin et al. (2012) to interpret the KTB extinction as the sum of multiple, independent events, at least at high latitudes.

Climate warming in Antarctica, prior to the KTB, has been also recognized by Bowman et al. (2013), based on the *I. clavus* dramatically decrease in abundance, on the pollen and spore record and available $\delta^{18}\text{O}$ values from macrofossils. According to these authors, these records support the presence of ephemeral ice sheets on Antarctica during the Maastrichtian, highlighting the extreme sensitivity of this region to global climate change.

The importance of Deccan-related environmental changes leading to the end-Cretaceous mass extinction has been also explored by Font et al. (2011) who have applied a high-resolution environmental magnetic study to the Bidart (France) and Gubbio (Italy) KTB sections. Their results have shown that the KTB in these sections is identified by an abrupt positive shift of the magnetic susceptibility with an anomalous

interval of very low MS, containing an unknown Cl-bearing iron oxide similar to specular hematite, depicted just below the iridium-rich layer of the KTB. The grain-size and morphology of this Cl-iron oxide are typically in the range of hematitic dust currently transported by winds from the Sahara desert to Europe. Based on SEM-EDS analyses, Font et al. (2014) suggested that this Cl-rich iron hydroxide (aeolian akaganeite) generated by a heterogeneous reaction between HCl-rich volcanic gas and liquid–solid aerosols within buoyant atmospheric plumes formed above the newly emitted Deccan flood basalts and further transported to Bidart and Gubbio through the stratosphere.

The low MS interval observed corresponds to loss in biomagnetite and detrital magnetite, interpreted to result from reductive dissolution linked to ocean acidification (Font et al., 2014). These independent markers (aeolian akaganeite deposition and iron oxide dissolution) argue for important paleoenvironmental changes linked to Deccan phase 2 at the dawn of the KTB in the Atlantic and Tethysian realm. Their discovery suggests a global-scale phenomenon and may be regarded as a new benchmark for widespread effects of the Deccan volcanism and witnesses the nature and importance of the related atmospheric change.

The onset of a large igneous province (LIP) eruption often postdated a mass extinction event, except for the eruption of the Deccan Traps which coincided precisely with a mass extinction according to Courtillot and Renne (2003) and Wignall (2001). However, based on planktic foraminifera, the Deccan phase 2 may have started around 200,000 years before the KTB (in the CF1 and CF2 biozones) and ended at the KTB (Keller et al., 2012). An enormous amount of metals, including Hg, has been potentially transferred to the atmosphere by the Deccan eruptions, worsening the environmental conditions and leading to the huge KTB mass extinction, in a similar way to that proposed for the Siberian Traps that emitted 3.8×10^9 ton Hg (Sanei et al., 2012) at the Permian–Triassic boundary.

2.1. The studied sections

Investigations in the present report have used well-known sequences of Stevns Klint (Højerup and Fiskeler members at the Højerup section), Gubbio (Scaglia Rossa Formation at the Bottaccione Gorge section) and Neuquén Basin (Jagüel Formation at the Bajada del Jagüel section), in which previous studies had applied high-resolution biostratigraphy and environmental multi-proxies such as: (1) carbon-isotope stratigraphy; (2) sedimentology, microfacies analysis and bulk rock mineralogy; (3) clay mineralogy to infer paleoclimatic conditions; (4) platinum group elements (PGE), trace and major elements geochemistry; and (5) major and trace element geochemistry.

2.1.1. The Neuquén Basin, Patagonia, Argentina

The Neuquén Basin in the Patagonia of western Argentina, is located on the eastern side of the Andes in Argentina and central Chile, between 32° and 40°S latitude, and has a sedimentary record with a 6000 m of Mesozoic infill, ranging from the Upper Triassic to Miocene (Vergani et al., 1995; Legarreta and Uliana, 1999; Howell et al., 2005; Fig. 1). In this basin, the Cretaceous–Paleogene transition was recorded in the Upper Cretaceous–Lower Paleocene Malargüe Group that rests discordantly on the Late Cretaceous Neuquén Group and comprises the Allen, Jagüel, Roca and El Carrizo formations in the eastern part of the basin (Legarreta et al., 1989; Casadio, 1998; Heredia and Salgado, 1999).

Detailed investigations on this basin have used multiple environmental proxies to understand major geological events at the Cretaceous–Paleogene transition, including biostratigraphy, microfacies analysis, bulk rock/clay mineralogy, isotope stratigraphy, trace (including PGE) and major element chemistry (e.g. Barrio, 1990; Howell et al., 2005; Scasso et al., 2005; Aberhan et al., 2007; Keller et al., 2007; Musso et al., 2012).

Whereas the Maastrichtian succession in this foreland basin is transgressive, a regressive trend was recorded in the Paleocene (Scasso et al.,

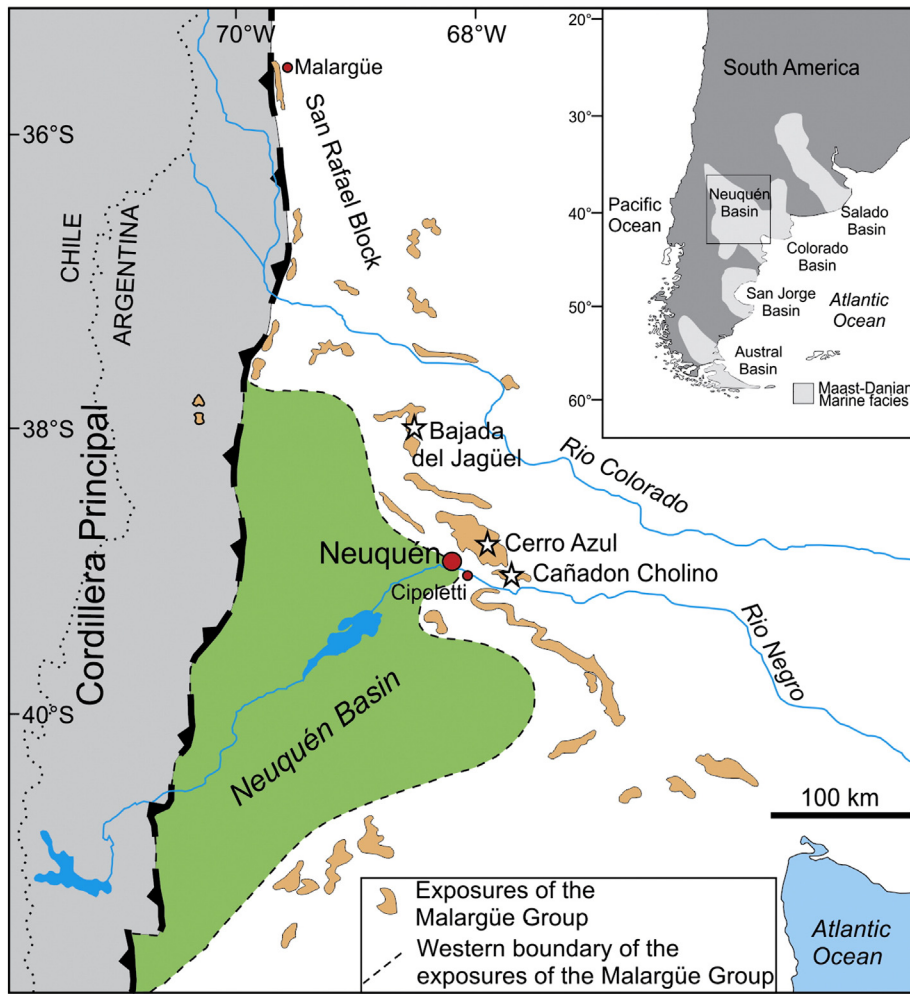


Fig. 1. (a) Map showing the location of the Maastrichtian–Danian Neuquén Basin; (b) Outcrops of rocks of the Malargüe Group and the eastern extension of this basin with location of the study sections cited in the text (modified from Aguirre-Urreta et al., 2011).

2005; Aguirre-Urreta et al., 2011). The Jagüel Formation, a 90-m thick succession, encompasses the KTB (Uliana and Dellapié, 1981; Uliana and Biddle, 1988) and consists of monotonous marine mudstones deposited in a mid- to outer-shelf environment, and is covered by bioclastic limestones of the Danian Roca Formation (Fig. 2a,b). An interval bracketing the KTB has been recognized on the basis of foraminifers and calcareous nannofossils (Bertels, 1970; Nañez and Concheyro, 1997), ostracods (Bertels, 1975), and palynomorphs (Papú et al., 1999). The position of the KTB is constrained to a single, thin coarse-

grained sandstone bed (Palamarczuk and Habib, 2001; Palamarczuk et al., 2002; Scasso et al., 2005) in the upper half of the Jagüel Formation.

Scasso et al. (2005) have described in detail this sandstone bed in which calcareous nannofossils have confirmed its Danian age. This 15–25 cm thick unconsolidated to slightly consolidated muddy sandstone (Fig. 2b) occurs within the homogeneous shelf mudstone sequence of the Jagüel Formation and contains abundant volcanogenic plagioclase (dated at 66 ± 0.5 Ma by the $^{40}\text{Ar}/^{39}\text{Ar}$ method) volcanic lithics, broken shells and shark teeth (Palamarczuk and Habib, 2001; Palamarczuk

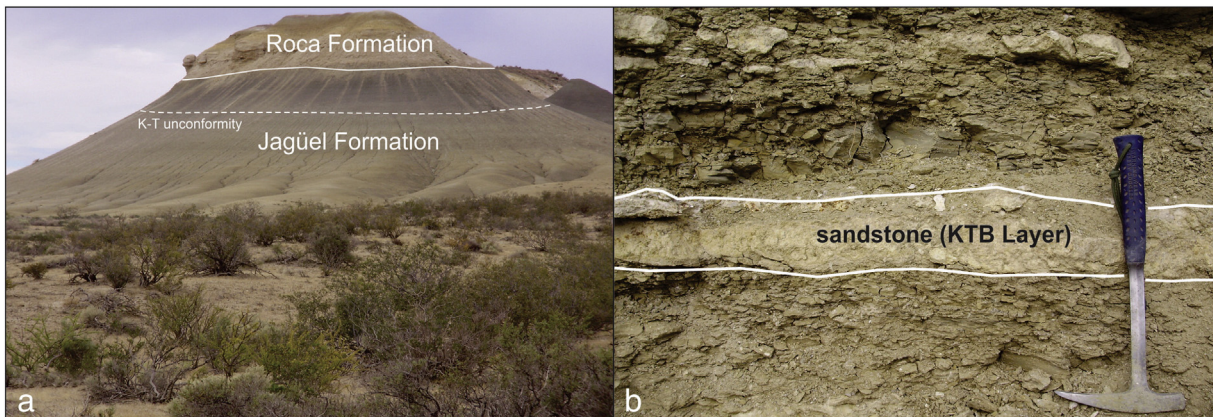


Fig. 2. Some field aspects of the Neuquén Basin: (a) Bajada del Jagüel section; (b) Cañadon Cholino section.

et al., 2002). Although this sandstone displays markedly pinch and swell structure, it can be traced laterally for 5 km in the area. According to Scasso et al. (2005), it shows abundant rip-up clasts, erosional base, coarse-grain size, normal grading and hummocky cross-bedding, and has been assumed to represent a tsunami deposit in a shelf environment, related to the Chicxulub bolide impact in Mexico. However, no spherule, shocked quartz or enrichment of meteoritic component have been found by these authors in this layer, or in the mudstones immediately above. Besides, PGE concentrations in this boundary bed are mostly below the detection limit and below the values for the average continental crust (Scasso et al., 2005).

Keller et al. (2007), however, contended that the KTB in this basin is marked by an erosional surface that marks a hiatus at the base of the 15–25 cm thick volcanoclastic sandstone described by Scasso et al. (2005). According to Keller et al. (2007), this sandstone contains diverse planktic foraminiferal zone P1c assemblages and nannofossils of zone NP1b immediately above it and, therefore, its deposition occurred at about 500 ky after the KTB hiatus. No evidence of the Chicxulub impact or related tsunami deposition was reported by these authors. Aberhan et al. (2007) found no sign of the Danian planktic foraminiferal zone P0 neither impact tracer in the KTB sandstone layer at the Bajada del Jagüel section, deducing that this section is slightly incomplete (missing about 100 ky).

In this study, we have examined and sampled the same section studied by Keller et al. (2007) and Aberhan et al. (2007) at Bajada del Jagüel (38° 06'11" S, 68° 23' 24"W; Fig. 2a), for which a large amount of paleontological, sedimentological and geochemical data with precise biostratigraphic control have been reported. We have also examined and sampled two other sections in this basin, one at the Cerro Azul (38° 50'51"S, 67° 52' 07"W) and another at the Cañadon Cholino (38° 53'16" S, 67° 41'25" W; Fig. 2b), along a river canyon, both located to the east of Cipoletti town. The KTB is within a 90 m-thick succession of monotonous, bioturbated, middle to outer shelf mudstones of the Jagüel Formation (Fig. 2a). At the Bajada del Jagüel and Cañadon Cholino sections, the KTB tuffaceous, coarse-grained sandstone layer described above, is easily recognized (Fig. 2a, b). The Jagüel Formation is overlain by shallow-marine bioclastic limestones of the Danian Roca Formation.

2.1.2. The Danish Basin, Stevns Klint, Denmark

The Danish Basin exposes at the Stevns Klint coastal cliffs, Denmark, the most famous and best exposed KTB section in the world, with an exceptional KTB layer immediately beneath a pronounced topographic overhang separating the lowermost Danian Cerithium Limestone Member from the overlying lower Danian bryozoan limestone of the Stevns Klint Formation (Fig. 3).

The Stevns Klint coastal cliffs (Fig. 4a), located ca 45 km south of Copenhagen, on the east coast of the island of Sjælland, represent the longest, best preserved, largest lateral extensive (except for Hell Creek, Eastern Montana, United States; see Johnson et al., 2002) and complete KTB site in the world (~14.5 km long). The exposed 45 m-thick succession displays the stratigraphic evolution from the latest Cretaceous, across the KTB and into the early Paleogene (Surlyk, 1997; and references therein). This classical KTB site is one of the three discovery localities of the famous iridium anomaly (the other two localities are Gubbio, Italy, and Woodside Creek, New Zealand), of about 160 times the background (Alvarez et al., 1980; Hansen et al., 1988; Schmitz et al., 1988), taken as evidence of extraterrestrial origin and basis for the asteroid impact hypothesis of Alvarez et al. (1980). Therefore, this is a key locality in the ongoing debate about mass extinction in general and, in particular, for the KTB, besides being the type locality of the Danian Stage together with the succession exposed in the nearby Faxø quarry.

The most detailed stratigraphic studies of this area are those accomplished by Surlyk (1997), Surlyk et al. (2006) and Lauridsen et al. (2012). At the Højerup coastal cliff section, one can see outcrops of the Maastrichtian Møns Klint Formation (Sigerslev and Højerup members; Surlyk et al., 2013; Hansen and Surlyk, 2014), overlain by the Lower Danian Rødvig Formation (Fiskeler, Cerithium members) which in turn is covered by the lower to middle Danian Korsnæb Member of the Stevns Klint Formation (Surlyk et al., 2006). The Danian is represented by bryozoan limestone mounds outlined by thick black flint bands which illustrate the geometry, dimensions and architecture of one of the finest, ancient cool-water carbonate mound complexes in the world, formed shortly after the KTB mass extinction (Surlyk, 1997; Hart et al., 2004; Machalski and

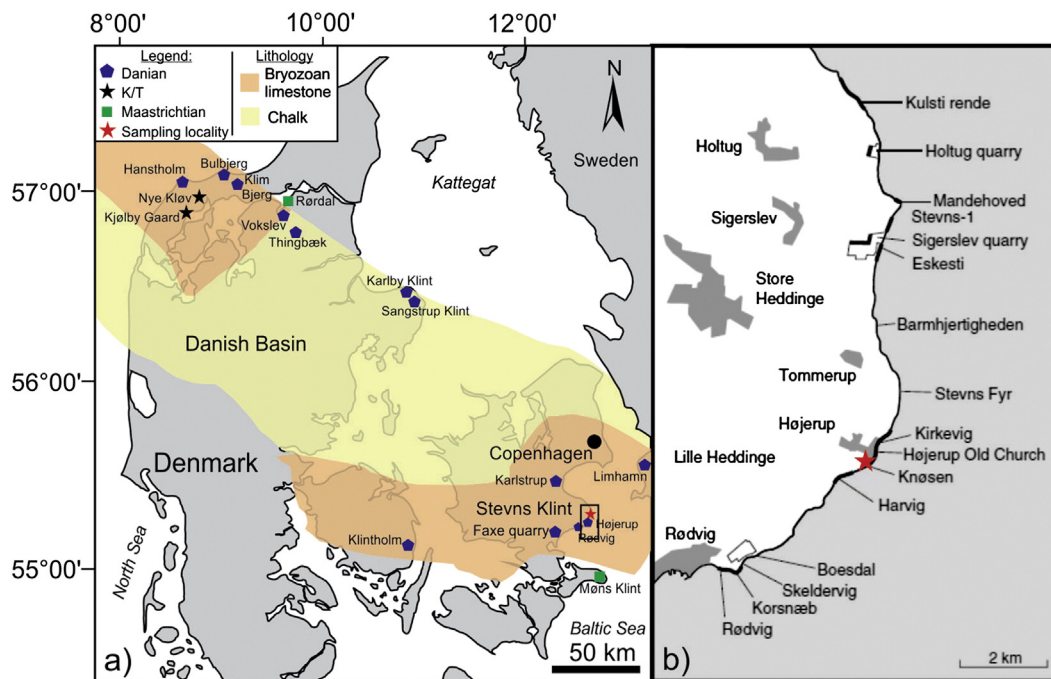


Fig. 3. (a) Map of Denmark with major structural elements (based on Surlyk et al., 2006; Lauridsen et al., 2012; modified from Sial et al., 2013) showing location of sampled site at Stevns Klint; (b) Key localities along the 14 km long cliff are shown in addition to towns and villages (grey shading) in the area (modified from Hansen and Surlyk, 2014).

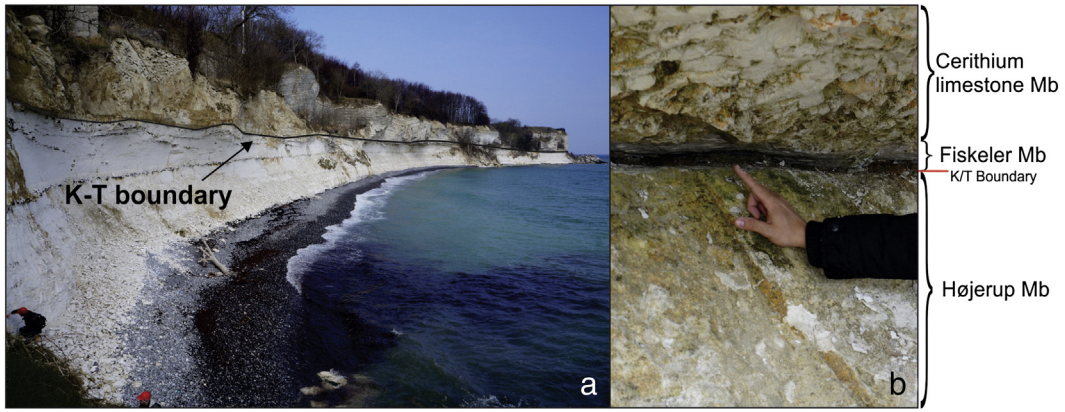


Fig. 4. Field aspects of Stevns Klint, Danish Basin, Denmark: (a) Panoramic view of the Højerup section at Stevns Klint showing the position of the KTB; (b) A closer view of the KTB layer (Fiskeler Member) in the Højerup section.

Heinberg, 2005; Surlyk et al., 2006; Lauridsen et al., 2012 and references therein). In summary, Stevns Klint is a world-class KTB site to achieve description of the mass extinction and recovery of the fauna and one of the best localities to document changes in sea level, surface and bottom sea-water temperature, ecological and

evolutionary conditions and changes in the last phases of the Late Cretaceous greenhouse world.

Relatively high concentrations of metals such as Ni, Co, and Zn, besides Ir, are found in the KTB layer at Stevns Klint (Christensen et al., 1973; Schmitz, 1985; Elliott, 1993). Studies conducted by Frei and Frei

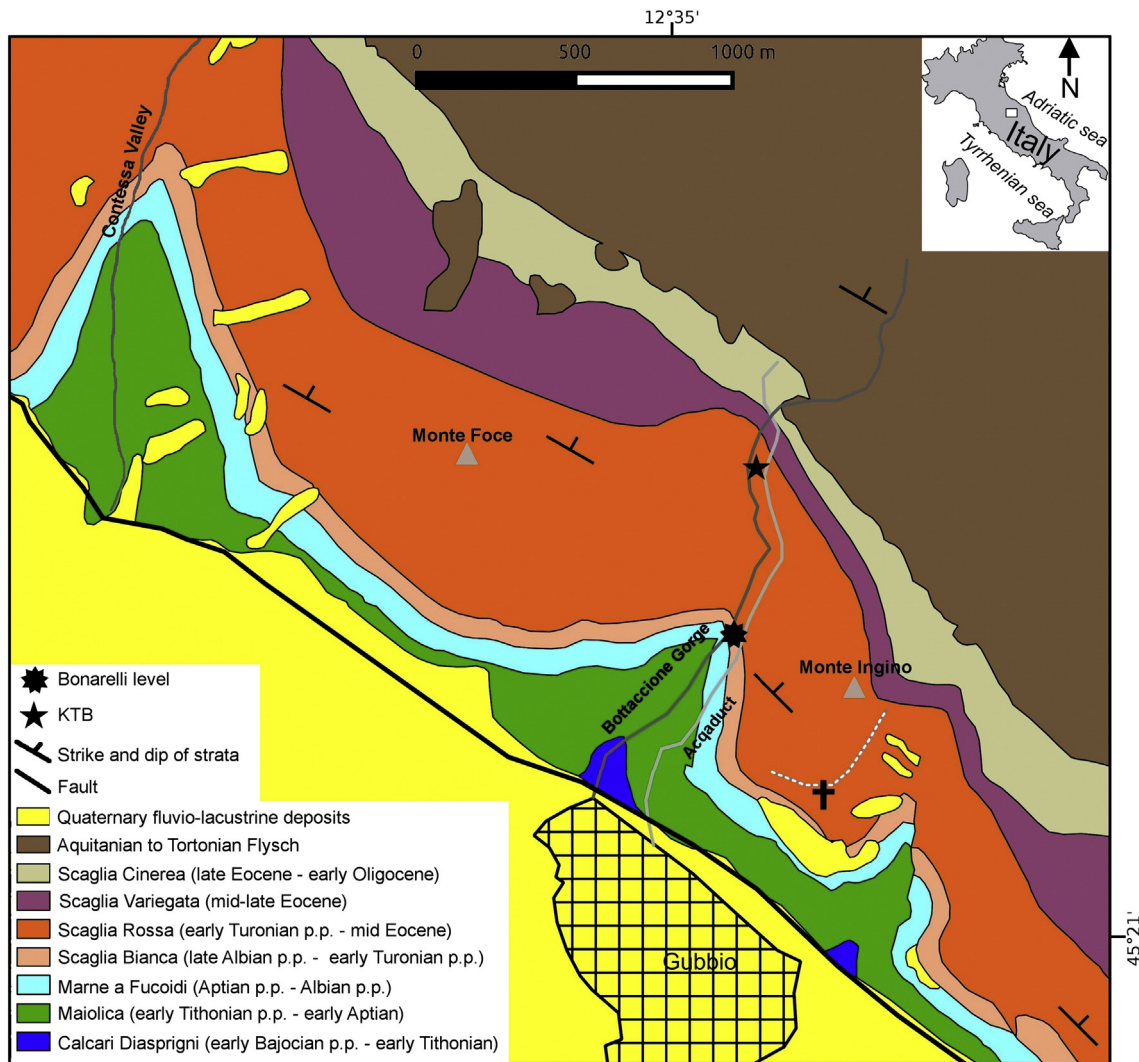


Fig. 5. Summary geological map of the surrounding Gubbio town, where it is shown the spatial distribution of Jurassic to Middle Cretaceous rocks, Scaglia Group rocks, and location in plant view of the Bonarelli level and the KTB (modified from Alvarez et al., 1977).

(2002) included Os, Sr, Nd and Pb isotope data obtained from a profile across the KTB layer (Fiskeler Member, informally known as Fish Clay, about 5 to 10 cm thick, separating the underlying Maastrichtian chalk from the overlying Danian limestone sequence; Fig. 4). The behavior of these isotopes in the Fish Clay layer, according to Frei and Frei (2002), supports that PGEs originated from global input of cosmogenic material into the ocean derived from a likely chondritic impactor.

2.1.3. The Umbria–Marchean succession, Central Apennines, Italy

The Gubbio Mountains form a NW–SE chain, about 13 km long and 3 km wide. Structurally this chain displays a NNE–SSW-trending anticlinorium of Mesozoic–Cenozoic rocks whose core is Jurassic in age. It shows the same orientation as the Central Apennines and is cross-cut lengthwise by a fault system whose major fault is known as Gubbio Fault that has dropped the southwestern limb of the anticlinorium, which has been filled with Quaternary fluvio-lacustrine sediments (Boncio et al., 2000; Mirabella et al., 2004).

The two better known KTB sections in this basin are those in the Bottacione Gorge (Figs. 5, 6a, b) and in the Contessa Highway, running through a parallel valley. The Bottacione Gorge begins just outside the Gubbio town and cuts through pelagic sedimentary rocks, spanning from Middle Jurassic to Upper Eocene, including the Calcarei Diasprigni, Maiolica, Marne a Fucoidi formations and the Scaglia Group (Scaglia Bianca, Scaglia Rossa, Scaglia Variegata and Scaglia Cinerea formations), all belonging to the Umbria–Marche Basin.

The Cretaceous to upper Eocene marine sedimentary rocks have been subject of numerous biostratigraphical studies (e.g. Luterbacher and Premoli Silva, 1964; Premoli Silva and Sliter, 1994, and references therein) that represented the basis for magnetostratigraphical (e.g. Pialli, 1976; Alvarez et al., 1977; Napoleone et al., 1983) stratigraphical–sedimentological (Arthur and Fisher, 1977; Premoli Silva et al., 1977, 1988; Wonders, 1979, 1980; Monechi and Thierstein, 1985; Cresta et al., 1989; Coccioni et al., 1992; Cecca et al., 1994; Galeotti et al., 2000), and geochemical investigations (O, C and Sr isotopes; e.g. Jenkyns et al., 1994).

The Bottacione section has been proposed as the magnetostratigraphic standard for the Upper Cretaceous–Eocene interval. It became famous since the discovery of a 1 cm-thick clay layer at the KTB within the Scaglia Rossa Formation, enriched in iridium and interpreted by Alvarez et al. (1980) as evidence of an extraterrestrial impact. On the contrary of what is observed at Stevns Klint, the KTB layer at Gubbio has a very short lateral extent as the succession is tilted.

The Cenomanian–Turonian Bonarelli level, a regional stratigraphic marker which represents the second major oceanic anoxic event in the Cretaceous (OAE 2), occurs near the top of the Scaglia Bianca Formation (Fig. 5) and consists of a 1 m-thick cherty black shale, very rich in radiolarians, along with some fish remains and more than 23% organic

carbon (Arthur and Premoli Silva, 1982). The contact between the Scaglia Bianca and the overlying Scaglia Rossa formations is characterized by alternating gray and pink colored beds. The Scaglia Rossa Formation is about 320 m thick in this section, brackets the KTB and consists of predominantly pink to red pelagic limestones with cherty nodules and calcareous marls, with bedding thickness ranging from 10 to 20 cm. The formation contains an apparently continuous record across the Mesozoic–Tertiary transition. Based on abundant planktonic and benthic foraminifera and calcareous nannofossils, this formation has been dated as Upper Cretaceous to lower middle Eocene (Premoli Silva and Sliter, 1994; Alvarez, 2009 and references therein).

The KTB crops out on the eastern side of the main road and lies around 240 m above the base of the formation. The topmost of the Cretaceous in this section is represented by a 30 cm-thick whitish limestone bed (bleached, Fig. 6b, c), overlain by the 2 cm-thick dark clay KTB layer which correlates to the well-known KTB mass-extinction event (Sepkoski, 1996; Keller, 2011). The base of the Paleocene is defined on the occurrence of *Globigerina eugubina* foraminiferal Zone (= *Parvurugoglobigerina eugubina*: Luterbacher and Premoli Silva, 1964).

The KTB layer is iridium enriched but devoid of calcium carbonate (Alvarez et al., 1980) and falls in the reversal interval equated to Chron 29r (Alvarez et al., 1977). In this report, we have studied and sampled this section bracketing the KTB.

3. Analytical methods

Analyses of C and O isotopes of carbonates were performed at the Stable Isotope Laboratory (LABISE) of the Department of Geology, Federal University of Pernambuco, Brazil. Extraction of CO₂ gas from micro-drilled powder (1 mm drill bit was used, avoiding fractures, recrystallized portions and weathered surfaces) was performed in a conventional high vacuum extraction line after reaction with 100% orthophosphoric acid at 25 °C for one day (three days allowed, when dolomite was present). Released CO₂ was analyzed after cryogenic cleaning in an Isotech double inlet, triple-collector SIRA II or Thermofinnigan Delta V Advantage mass spectrometers and results are reported in δ notation in permil (‰) relative to the VPDB standard. The uncertainties of the isotope measurements were better than 0.1‰ for carbon and 0.2‰ for oxygen, based on multiple analyses of an internal laboratory standard (BSC). Alumina oxide analyses were performed on bulk samples by X-ray fluorescence at the LABISE, using a Rigaku RIX 3000 XRF unit equipped with a Rh tube and six analyser crystals.

Organic carbon isotopic values were obtained from the total organic carbon of insoluble residues of some samples from the Neuquén Basin only. After removing the outer layer of surface oxidation and any large veins, whole rock samples were crushed into powder. Insoluble

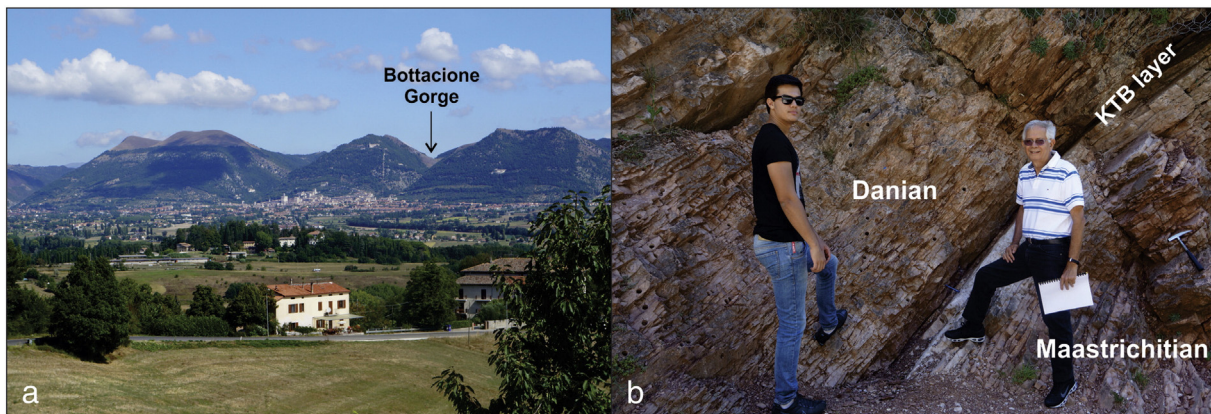


Fig. 6. (a) Gubbio town and location of the Bottacione Gorge; (b) KTB layer (1–2 cm thick) within the Scaglia Rossa Formation at the Bottacione Gorge; A. Sial (father) and A. Sial (son) for scale.

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 30 days to dissolve all carbonate minerals. Care was taken to ensure that acid was added and acidification continued until there was absolutely no visible carbonate dissolution so that the analyses would not be affected by contamination from residual inorganic carbon. The insoluble residues were then rinsed with DI water, dried and loaded into tin capsules for isotopic analysis. $\delta^{13}\text{C}_{\text{org}}$ values were obtained using a COSTECH elemental combustion system linked to a Delta V Advantage at the LABISE.

Mercury concentrations were determined at the LABOMAR, Federal University of Ceará, Brazil. Homogenized 0.5 to 1.0 g samples of sediments, dried at 60 °C to constant weight, were digested with an acid mixture (50% aqua regia solution), and heated at 70 °C for 1 h, in a thermal-kinetic reactor (“cold finger”). Glass and plastic ware were decontaminated by immersion for 1 day in (10% v/v) Extran solution (MERCK), 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 Hg²⁺ reduction with SnCl₂. All samples were analyzed in duplicates, showing reproducibility within 9.5%. A certified reference material (NIST 2702, Canada) was simultaneously 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. Concentration values were not corrected for the recoveries found in the certified material.

4. C and O isotopes composition of the studied sections

Chemostratigraphic correlation based on bulk sediment carbon isotopes is increasingly used to facilitate high-resolution correlation over large distances as we know that at times of rapid environmental change, chemostratigraphy backed by biostratigraphy can enable correlation at much higher resolution than biostratigraphy alone (Brenchley et al., 2003). However, complications arise from a multitude of possible influences from local differences in biological, diagenetic and physical-chemical factors on individual $\delta^{13}\text{C}$ records that mask the global signal (Wendler, 2013). In the present study, it is possible to assess global versus local contribution in the $\delta^{13}\text{C}$ record by comparing C and O-isotope pathways across the KTB from sections far apart from each other, such as in the Neuquén Basin, Stevns Klint and Gubbio.

Prior to the interpretation of the C- and O-isotope data generated in this study, we reviewed the more frequent criteria for screening samples that have not undergone post-depositional alteration, as discussed below. The most effective parameter to screen primary values is the Mn/Sr ratio, because Sr is preferentially removed during recrystallization of meta-stable carbonate phases, while Mn becomes enriched during formation of late-stage ferroan calcite cement (Ripperdan et al., 1992; Kaufman et al., 1993; Jacobsen and Kaufman, 1999). Limestones are considered to be unaltered only when Mn/Sr < 1.5 and $\delta^{18}\text{O} > -10\%$ VPDB according to Fölling and Frimmel (2002). The Mn/Sr ratios for all samples in this study are < 1.5 and $\delta^{18}\text{O}$ values for all samples are > -10% VPDB.

It is known that correlation between C and O isotopes is observed in material that has been altered by burial diagenesis, but it is also known that covariance between these two isotopes cannot be used for unequivocal identification of meteoric late diagenesis. The reason is that covariance is also observed in modern calcareous skeletons and in pristine fossils where it reflects simultaneous depletion in $\delta^{13}\text{C}$ and $\delta^{18}\text{O}$ due to kinetic fractionation effects related to growth rate (McConnaughey, 1989; Wendler et al., 2013) or to vital effects related

to pH (Spero et al., 1997; Bijma et al., 1999; Zeebe, 1999; Adkins et al., 2003; Zeebe, 2007).

In the Neuquén Basin, we have collected fourteen closely-spaced samples perpendicular to the strike of the strata in a section at the Bajada del Jagüel, thirteen samples in a section at Cerro Azul, and eight samples in a section along the river canyon at Cañadón Cholino, not far from the Cipoletti town (Fig. 1). C and O isotopes were analyzed in these samples and results are shown in Table 1 and plotted in Figs. 7, 8 and 9.

High resolution $\delta^{13}\text{C}_{\text{carb}}$ and $\delta^{18}\text{O}$ pathways for Maastrichtian samples from the Bajada del Jagüel section (Fig. 7) show little fluctuation and are followed by a strong negative excursion at the KTB layer that, if primary in nature, records a warming event around the KTB. $\delta^{13}\text{C}_{\text{carb}}$ and $\delta^{18}\text{O}$ shift to lower values in the Danian samples but also show limited fluctuation. In a $\delta^{13}\text{C}_{\text{carb}}-\delta^{18}\text{O}$ plot (Fig. 7), Maastrichtian and Danian samples show a certain degree of co-variation. Fossil shells and bioclastic material are prominent in the analyzed marl to pelitic rocks at the Bajada del Jagüel section, likely with substantial influence on the bulk-rock isotope analyses and, therefore, the explanation for co-variation given above may be applicable.

The $\delta^{13}\text{C}_{\text{org}}$ curve for organic matter extracted from samples of this section (Fig. 7) exhibits a pathway with a minimum (~-26‰) in marl about 0.5 m below the KTB layer, a negative excursion at the KTB, a well-defined positive excursion to -23‰ immediately above the KTB, and a gradual and strong decrease to -26‰ upsection. The behavior

Table 1

C, O isotope analyses and Hg contents for three sections in the Jagüel Formation, Neuquén Basin, Argentina.

Sample	Height (cm)	$\delta^{13}\text{C}_{\text{carb}}$ ‰ VPDB	$\delta^{13}\text{C}_{\text{org}}$ ‰ VPDB	$\delta^{18}\text{O}$ ‰ VPDB	Hg (ng·g ⁻¹)
<i>(a) Bajada del Jagüel section</i>					
KT 12	100	0.4	-25.9	-6.2	11.04
KT 11	100	0.4	-25.0	-5.8	12.14
KT 10	35	-0.4	-	-7.1	11.87
KT 9	20	-0.3	-20.0	-6.7	12.85
KT 8	3	-0.3	-23.8	-6.3	8.57
KT 7	3	-2.2	-24.0	-10.6	8.55
KT 6	10	-1.0	-24.7	-8.2	10.19
KT 5	17	-0.6	-24.3	-6.8	15.80
KT 4	25	1.3	-24.2	-3.8	13.95
KT 3	35	1.1	-25.1	-3.7	15.80
KT 2	45	1.1	-25.5	-3.6	16.56
KT 1	55	0.6	-25.3	-3.6	15.81
KT 0	55	0.9	-24.7	-4.1	14.79
KT(-1)	0	0.7	-25.0	-4.8	12.53
<i>(b) Cerro Azul section</i>					
CAZ 13	150	-	-25.3	-	45.41
CAZ 12	150	-1.6	-24.9	-6.5	415.25
CAZ 11	50	-0.6	-24.9	-3.8	72.80
CAZ 10	50	0.1	-25.5	-4.9	160.57
CAZ 9	75	-	-24.6	-	332.53
CAZ 8	25	-0.2	-24.7	-3.9	99.68
CAZ 7	25	0.1	-25.8	-2.9	46.05
CAZ 6	50	0.3	-26.5	-2.8	18.70
CAZ 5b	0	-	-25.2	-	30.50
CAZ 5a	75	-0.1	-	-3.1	332.53
CAZ 4	75	-0.3	-24.2	-3.3	7.03
CAZ 3	75	-	-24.9	-	337.10
CAZ 2	75	-0.6	-24.2	-3.8	20.38
CAZ 1	0	-0.2	-24.6	-4.9	40.65
<i>(c) Cañadon Cholino section</i>					
CCL 8	200	-	-24.1	-	17.81
CCL 7	35	-	-24.6	-	16.61
CCL 6	200	-	-25.9	-	16.78
CCL 5	100	-	-24.7	-	5.56
CCL 4	200	-3.9	-24.7	-9.4	7.68
CCL 3	300	-0.6	-23.2	-5.5	6.45
CCL 2	300	-3.2	-24.1	-5.8	18.52
CCL 1	0	-0.4	-25.9	-4.7	21.12

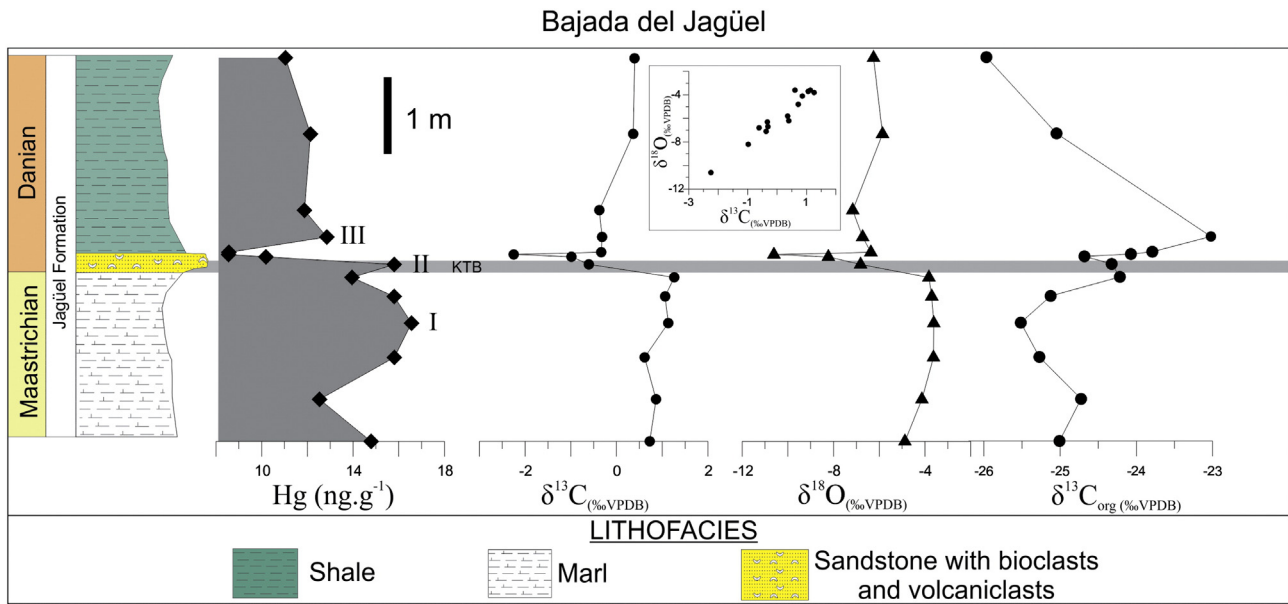


Fig. 7. $\delta^{13}\text{C}_{\text{carb}}$, $\delta^{18}\text{O}$, $\delta^{13}\text{C}_{\text{org}}$ and Hg, variation curves across the KTB at the Bajada del Jagüel section, Neuquén Basin, Argentina. Some co-variation in the $\delta^{13}\text{C}_{\text{carb}}-\delta^{18}\text{O}$ plot is also shown.

of $\delta^{13}\text{C}_{\text{org}}$ in this section somehow opposes that of $\delta^{13}\text{C}_{\text{carb}}$ as expected and is interpreted as a response to a gradual deterioration of the environment in the topmost Maastrichtian with gradual decrease of the organic productivity, followed by its gradual recovery early in the Danian. In the Cerro Azul section, $\delta^{13}\text{C}_{\text{carb}}$ and $\delta^{18}\text{O}$ stratigraphic curves (Fig. 8) show similar behavior to that of Bajada del Jagüel section but with less apparent co-variation. The behavior of $\delta^{13}\text{C}_{\text{carb}}$ versus $\delta^{13}\text{C}_{\text{org}}$ is similar to that observed in the Bajada del Jagüel section, the curves displaying opposite behavior around the KTB (Fig. 8). At the Cañadón Cholino section, the amount of inorganic carbon in the rocks analyzed is very low and only a $\delta^{13}\text{C}_{\text{org}}$ curve is displayed in Fig. 9 where it essentially replicates the behavior of the $\delta^{13}\text{C}_{\text{org}}$ curve observed in Fig. 8.

Kaminski and Malmgren (1989) have made C- and O-isotope measurements on KTB samples from two Danish localities—Stevns Klint

and Kjølbj Gaard. Results indicated a large positive $\delta^{18}\text{O}$ excursion coincident with the Ir anomaly in the Fiskeler Member, followed by large fluctuations in $\delta^{18}\text{O}$ values in the lowermost Danian while C isotopes have shown a negative $\delta^{13}\text{C}$ excursion across the KTB. Kaminski and Malmgren (1989) have observed that a shift toward more negative $\delta^{13}\text{C}$ values begins at 0.75 m (<10,000 years) below the KTB layer at Kjølbj Gaard while at Stevns Klint, the lowest $\delta^{13}\text{C}$ values are observed at an interval of low dinoflagellate abundance in the Danian. These $\delta^{13}\text{C}$ patterns have been ascribed to decrease of surface-water productivity that started several thousand years before the KTB event.

High resolution C- and O-isotope investigations at the Stevns Klint succession across the KTB have been also performed by Hart et al. (2004), Machalski and Heinberg (2005). In both studies, a $\delta^{13}\text{C}$ negative shift was found at the KTB (values drop from around +2‰ in the

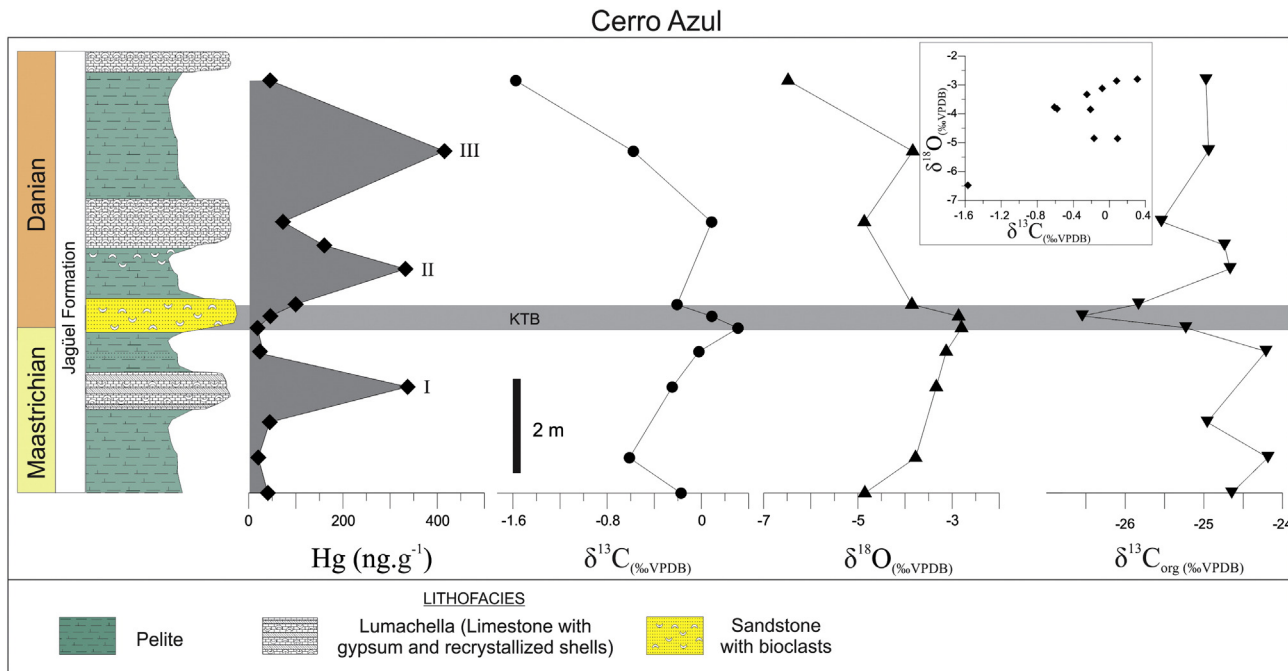


Fig. 8. $\delta^{13}\text{C}_{\text{carb}}$, $\delta^{18}\text{O}$, $\delta^{13}\text{C}_{\text{org}}$, Hg variation curves across the KTB at the Cerro Azul section across the KTB. No co-variation is apparent in a $\delta^{13}\text{C}_{\text{carb}}-\delta^{18}\text{O}$ plot.

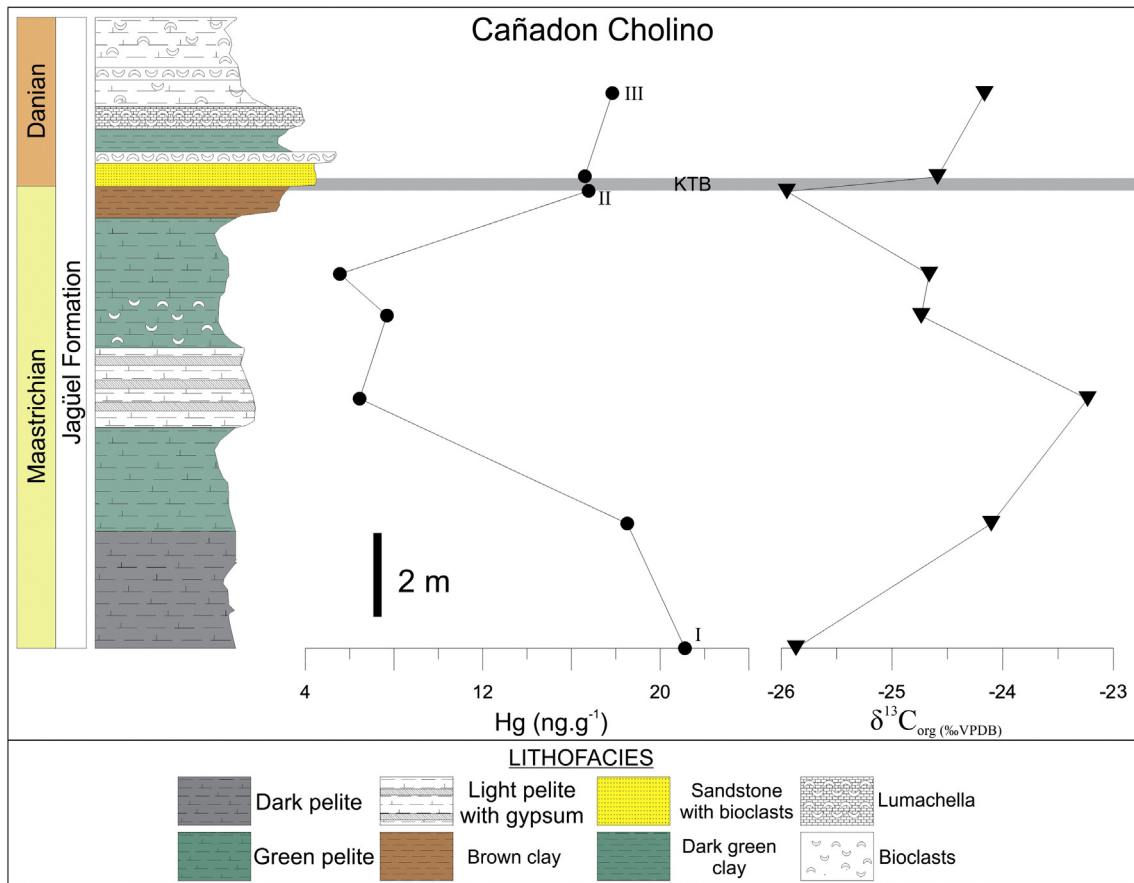


Fig. 9. Hg and $\delta^{13}\text{C}_{\text{org}}$ variation curves across the KTB at the Cañadon Cholino section.

Table 2

C, O isotopes and Hg contents across the KTB at Stevns Klint, Denmark. Stratigraphic heights are given in relation to the KTB layer.

Formation/member		Sample	Height (cm)	$\delta^{13}\text{C}$ (‰ VPDB)	$\delta^{18}\text{O}$ (‰ VPDB)	Hg (ng g^{-1})		
Danian Stevns Klint Formation	Korsnaeb Member	SKc	1050	1.4	-1.9	1.8		
		N7	990	1.19	-2.33	1.0		
		N6	960	1.38	-1.61	48.2		
		N5	930	1.28	-1.8	6.8		
		N4	900	1.23	-1.87	11.7		
		N3	870	1.16	-1.81	3.0		
		N2	840	1.28	-1.93	6.6		
		Danian Rødvig Formation	Cerithium Limestone	D3	830	1.4	-1.6	4.55
				D2	815	1.6	-1.5	4.51
			Member	N1	810	1.15	-1.87	4.5
No. 3	805			1.7	-1.5	2		
Fish Clay (Fiskeler Member; KTB)	FC-1			0			127.72	
	FCD		803	1.9	-0.5	67.9		
	FCC		802	1.7	-0.9	257.94		
Maastrichtian Møns Klint Formation	Gray Chalk (Højerup Member)		FCB	801	1.3	-1.7	194.53	
		FCA	800	1.3	-1.7	108.76		
		FK-1	800	1.61	-1.41			
		FK-2	800	1.56	-1.72			
		M2	785	1.8	-1.4	9.09		
		N(-1)	770	0.71	-3.61			
	White Chalk (Sigerslev Member)	M1A	770	1.7	-2	0.88		
		No. 2	760	1.8	-1.7	2.3		
		N(-2)	740	1.71	-1.59	<1.26		
		N(-3)	685	1.94	-1.61	4.6		
N(-4)	660	1.93	-1.74	6.5				
No. 1	0	1.7	-1.9	11.3				

uppermost Højerup Member to +1% at the Fiskeler Member; Table 2), while the $\delta^{18}\text{O}$ chemostratigraphic curve shows a negative anomaly ($\sim -1.5\%$ VPDB). In addition, Thibault et al. (2012) have shown very detailed C-isotope profiles in Stevns Klint and Rørdal localities, correlating them with nannofossil bio-events. Their study, as well as the one performed by Surlyk et al. (2013) have confirmed the $\delta^{13}\text{C}$ negative shift described above. In the present study, $\delta^{13}\text{C}$ and $\delta^{18}\text{O}$ for a high resolution section across the KTB at Højerup (Fig. 10) show the same behavior described in Hart et al. (2004). In a $\delta^{13}\text{C}$ - $\delta^{18}\text{O}$ plot, there is no clear co-variation.

O- and C-isotope measurements on topmost Maastrichtian and Paleocene limestone samples from the Bottaccione Gorge section and from the adjacent Contessa Highway section have been reported by Corfield et al. (1991). Their C-isotope data from the Bottaccione Gorge show $\delta^{13}\text{C}$ maxima at the Cenomanian/Turonian boundary and during the Paleocene, as well as a pronounced $\delta^{13}\text{C}$ minimum 7 m above the clay layer at the KTB. Their data from the Contessa Highway show a $\delta^{13}\text{C}$ minimum 9 m above the KTB clay and they have interpreted these minima as probable diagenetic artifacts. According to these authors, the Paleocene of the Bottaccione Gorge was affected by slumping or faulting and diagenetic overprinting, giving a spurious $\delta^{13}\text{C}$ carb recovery after the KTB. Their O-isotope data indicated an increase in $\delta^{18}\text{O}$ (possible cooling) in the Danian in both, the Bottaccione and Contessa Highway sections.

In the present study, twenty-eight samples have been collected, perpendicular to the strike of the strata, at the Bottaccione Gorge section, bracketing the KTB. These samples have been analyzed for C and O isotopes and results are shown in Table 3 and the C and O-isotope pathways are displayed in Fig. 11. Samples from the topmost Maastrichtian rocks in

Højerup Section

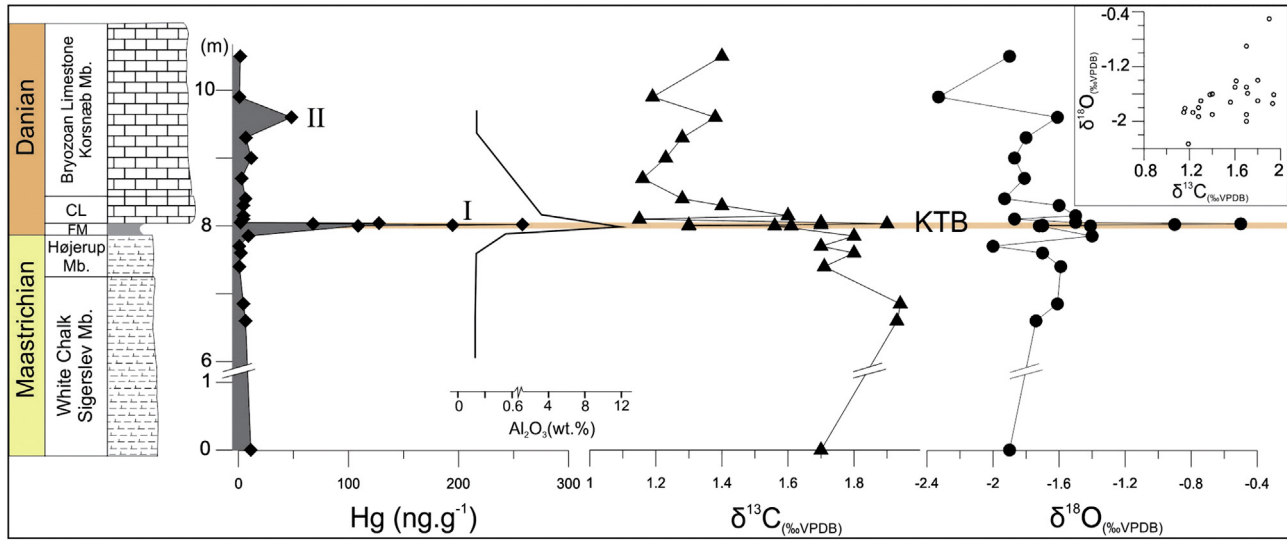


Fig. 10. Hg, Al_2O_3 , $\delta^{13}\text{C}$ and $\delta^{18}\text{O}$ variation curves across the KTB at Stevns Klint, Denmark. Al_2O_3 curve is from Schmitz et al. (1992) and $\delta^{13}\text{C}$ and $\delta^{18}\text{O}$ are from this study. No co-variation is apparent in a $\delta^{13}\text{C}$ – $\delta^{18}\text{O}$ plot.

this section show a high resolution $\delta^{13}\text{C}$ pathway with limited variation (from 2.32 to around 2‰) with a marked drop at the KTB, followed upsection by substantial gradual decrease of $\delta^{13}\text{C}$ values with a minimum at about 7 m above the KTB. The $\delta^{18}\text{O}$ stratigraphic profile, in turn, shows less fluctuation but displays a marked negative anomaly at the KTB indicative of warming conditions.

In a $\delta^{13}\text{C}$ – $\delta^{18}\text{O}$ plot (Fig. 11), Maastrichtian and Danian values group in two distinct areas in this diagram; as there is no clear correlation between C and O-isotope values we assume that these are near-primary isotope signals and that the negative $\delta^{18}\text{O}$ anomaly suggests warming in the KTB, followed by immediate cooling.

5. Mercury behavior across the Cretaceous–Paleogene boundary

Cataclysmic volcanoes have the potential of injecting enough Hg to the atmosphere to change global and regional Hg cycles. Volcanic emissions are a significant natural source of Hg to the atmosphere, with average annual emissions of about 90 tons, 60% of which from eruptions and 40% from degassing activities (Nriagu and Becker, 2003).

The association of higher contents of Hg with modern carbonates finely inter-layered with terrigenous sediments may have ultimately resulted from increased atmospheric Hg deposition related to

Table 3

C and O isotopes and Hg concentrations at the Botaccione Gorge section, Gubbio, Italy.

Sample	Height (cm)	Lithology	$\delta^{13}\text{C}_{\text{carb}}$ ‰VPDB	$\delta^{18}\text{O}$ ‰VPDB	Hg content (ng.g ⁻¹)
GP-28	120	Marl	1.26	−3.29	3.44
GP-27	70	Marl	1.27	−3.18	1.62
GP-26	100	Marl	1.38	−2.60	3.79
GP-25	100	Marl	1.18	2.36	1.51
GP-24	100	Marl	1.43	−2.53	1.55
GP-23	100	Marl	1.34	−2.55	1.83
GP-22	100	Marl	1.28	−2.82	0.87
GP-21	100	Marl	1.48	−2.36	0.42
GP-20	20	Marl	1.58	−2.04	0.68
GP-19	30 (clay)		1.77	−1.69	0.57
GP-18	30 (clay)	Marl (rhythmites; carbonate production decreases in the Danian)	2.02	−1.57	0.72
GP-17	20 (clay)		2.08	−1.99	0.44
GP-16	0.40		2.03	−1.79	0.75
GP-15	1 cm thick KTB layer	Greenish brown clay layer, about 170 m from the base of the Scaglia Rossa Formation			5.23
GP-14	4	Mudstone with planktic foraminifera	2.28	−2.83	0.22
GP-13	10	Mudstone with planktic foraminifera	2.32	−2.57	0.16
GP-12	10	Mudstone with planktic foraminifera	2.29	−2.68	0.35
GP-11	10	Mudstone with planktic foraminifera	2.27	−2.67	0.22
GP-10	30	Mudstone with planktic foraminifera	2.24	−2.85	0.37
GP-9	30	Mudstone with planktic foraminifera	2.17	−2.75	0.38
GP-8	100	Mudstone with planktic foraminifera	2.18	−2.80	0.47
GP-7	100	Mudstone with planktic foraminifera	2.07	−3.22	0.42
GP-6	100	Mudstone with planktic foraminifera	2.15	−2.82	0.88
GP-5	100	Mudstone with planktic foraminifera	2.32	−3.46	0.48
GP-4	100	Mudstone with planktic foraminifera	2.13	−2.57	0.97
GP-3	100	Mudstone with planktic foraminifera	2.15	−2.32	1.38
GP-2	100	Mudstone with planktic foraminifera	2.23	−2.39	1.75
GP-1	0.0	Mudstone with planktic foraminifera	2.29	−2.30	0.37

Bottaccione Section

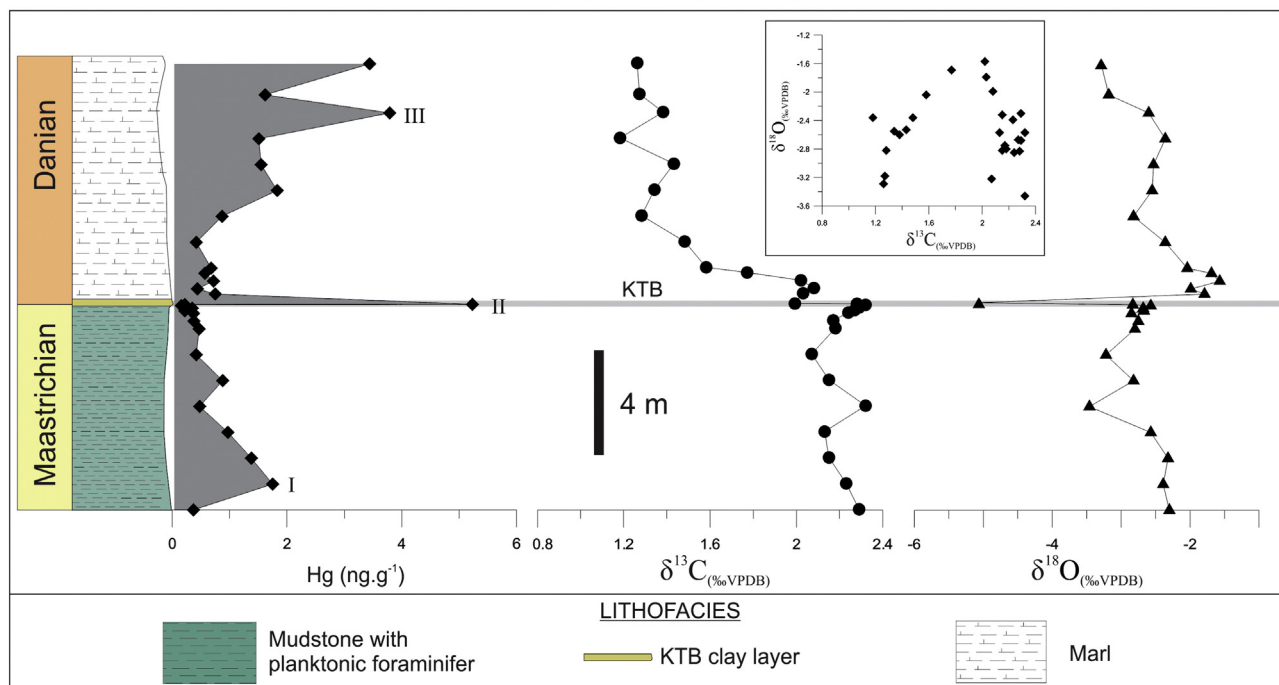


Fig. 11. $\delta^{13}\text{C}$, $\delta^{18}\text{O}$ and Hg variation curves across the KTB at the Bottaccione Gorge section, Gubbio, Italy. Some co-variation becomes apparent in a $\delta^{13}\text{C}_{\text{carb}}-\delta^{18}\text{O}$ plot.

volcanism (Roos-Barracough et al., 2002). Gaseous Hg from volcanic activity is transported far from their source reaching a regional and even global scale, contrary to most elements present in ash. As a result, many studies have reported synchronous Hg spikes in the sedimentary record associated with recent (e.g. Martínez-Cortizas et al., 1999; Roos-Barracough et al., 2002; Roos-Barracough and Shoty, 2003) and prehistoric volcanic activities (e.g. Sial et al., 2010; Nascimento-Silva et al., 2011, 2013; Sanei et al., 2012).

When reaching a surface environment that is depleted of organic scavenging capacity due to climate changes at the KTB, Hg^{+2} would be kept mostly in solution, readily adsorbed onto clays and transported to sedimentary basins. In summary, high levels of Hg associated to clay-bearing carbonates is in agreement with an increased flux of volcanic-derived Hg from the landmass into the marine realm.

Leaching of this Hg from the land surface, and subsequent transport into the oceans, could eventually have led to the known accumulation of Hg in argillaceous carbonates. For example, higher Hg accumulation rates were found to be much larger in sediments deposited after glacial maxima, when runoff is increased, than in sediment layers deposited before that, and it seems to represent a global phenomenon, as suggested by similar results obtained by the analysis of sediments in the Amazon region (Santos et al., 2001); Antarctica (Vandal et al., 1993) and Europe (Martínez-Cortizas et al., 1999).

Sanei et al. (2012) associated increasing atmospheric Hg deposition with catastrophic Siberian Traps volcanic eruptions, followed by the disruption of the organic fixation of Hg and consequently increasing dissolved mercury fluxes. Anomalous Hg contents have been observed across the KTB in Dolenja Vas, Slovenia, regarded as a probable result of sub-aerial volcanic activity (Palinkaš et al., 1996). High Hg contents were recorded across the KTB in the Paraíba Basin, Brazil, similar to peaks observed in Dolenja Vas, and much higher Hg values ($\sim 250 \text{ ng g}^{-1}$) were encountered across the KTB in the Yacoraite Formation, Salta Basin, Argentina (Sial et al., 2013) and have been ascribed to volcanism. Anomalous Hg contents across the KTB have been reported by Hildebrand and Boynton (1989) who have claimed this as an evidence for acidic rain they have deemed responsible for the associated mass extinction.

Besides a direct volcanic origin of Hg, higher Hg concentrations in the atmosphere could result from widespread reduction of biological activity that decreases or even shuts down the scavenging and biological fixation of Hg from the atmosphere, increasing the dissolved Hg fluxes. Under reduced bioproductivity, Hg is less likely captured by organic matter, one of the major Hg sinks (Sanei et al., 2012).

5.1. Mercury chemostratigraphy

Mercury concentrations were measured across the KTB in the Neuquén Basin in the same three sections analyzed for C and O isotopes. At the Bajada del Jagüel section, thirteen samples were analyzed (Table 1) and three mercury spikes are present (Fig. 7), one of them of about 17 ng g^{-1} Hg (labeled I) at about 65 cm below the KTB, a second one at the KTB (about 16 ng g^{-1} , labeled II) and a third one, less pronounced, of about 13 ng g^{-1} , around 15 cm above the KTB (labeled III).

At the Cerro Azul section mercury concentration was measured in thirteen samples, and also three mercury spikes with much higher concentrations than those at the Bajada del Jagüel section are evident. One of them (340 ng g^{-1} , labeled I) is about 1.5 m below the KTB, a second one of about 333 ng g^{-1} (labeled II) is situated about 1 m above the KTB and a third one of about 415 ng g^{-1} (labeled III) is about 3.5 m above the KTB. At the Cañadon Cholino section, the three highest values encountered in eight of the analyzed samples were of about 21, 17 and 18 ng g^{-1} , respectively at 8 m below KTB (labeled I), at the KTB (labeled II) and about 2 m above that (labeled III).

Results of Hg concentrations in eight samples from across the KTB at Stevns Klint, that were analyzed for Sr, Nd, Pb and Os isotopes by Frei and Frei (2002) are listed in Table 2. In Fig. 10, the Hg stratigraphic variation curve is compared with Al_2O_3 curve (data are from Schmitz et al., 1992) and with $\delta^{13}\text{C}$ and $\delta^{18}\text{O}$ stratigraphic variation curves. A peak of Hg of about 260 ng g^{-1} coincides with the $\delta^{18}\text{O}$ maximum value and $\delta^{13}\text{C}$ minimum value in the Fish Clay layer at the KTB that records an increase of continental weathering, subsequent clay generation and Hg fixation.

Table 4

Mercury contents (ng g⁻¹) for Cretaceous–Paleogene bulk samples from Paraíba Basin, Brazil, and Salta Basin, Argentina (modified from Sial et al., 2013).

Cretaceous–Paleogene transition (KTB) Carbonates																
(a) Paraíba Basin, Northeastern Brazil																
(a) Itamaracá drill hole, Paraíba Basin				(b) Olinda drill hole, Paraíba Basin				(c) Poty drill hole, Paraíba Basin				(d) Poty quarry section, Paraíba Basin				
Formation	Sample	Height (cm)	Hg	Formation	Sample	Height (m)	Hg	Formation	Sample	Height (m)	Hg	Formation	Sample	Height (cm)	Hg	
Maria Farinha	D 3240	32.4	2.91	Maria Farinha	D 3630	36.3	2.1	Maria Farinha	D 114	11.4	1.37	Maria Farinha	1-PO-01-PE 9,32	0	8.2	
	D 3270	32.7	2.97		D 3660	36.6	1.2		D 117	11.7	2.64		1-PO-01-PE 9,45	0.13	11.6	
	D 3300	33	3.35		D 3690	36.9	2.3		D 123	12.3	0.39		1-PO-01-PE 9,5	0.18	11.4	
	D 3360	33.6	4.38		D 3720	37.2	2.2		Gramame	D 126	12.6		0.73	1-PO-02-PE 9,6	0.28	14.3
	D 3390	33.9	3.85		D 3750	37.5	2.2			D 129	12.9		0.14	1-PO-01-PE 10,05	0.73	11.2
Gramame	D 3420	34.2	1.59	Gramame	D 3810	38.1	1.7	D 132	13.2	0.13	1-PO-01-PE 10,12	0.8	10.5			
	D 3450	34.5	1.06		D 3840	38.4	4.5	D 135	13.5	0.5	1-PO-01-PE 11,7	2.38	10.5			
	D 3480	34.8	0.84		D 3900	39	2.3	D 138	13.8	0.53	1-PO-01-PE 12,55	3.23	9.5			
	D 3510	35.1	2.6		D 3960	39.6	8.9	D 141	14.1	0.18	1-PO-01-PE 13,2	3.88	16.6			
	D 3540	35.4	1.5		D3990	39.90	11.5	D 144	14.4	0.46	1-PO-01-PE 14,75	5.43	9.4			
								D 171	17.1	0.16	1-PO-01-PE 14,85	5.53	13.0			
								D 174	17.4	0.27	1-PO-01-PE 15,2	5.88	6.3			
						D 177	17.7	0.42	1-PO-01-PE 17,32	8.0	8.0					
						D 201	20.1	0.12								
						D204	20.4	0.28								
						D207	20.7	0.25								
						D222	22.2	0.43								
						D225	22.5	0.17								
						D228	22.8	0.38								
						D276	27.6	0.48								
						D279	27.9	0.56								
						D282	28.2	0.46								
Cretaceous–Paleogene transition (KTB) bulk samples																
(b) Salta Basin, Argentina																
(e) Yacoraite Formation, Cabra Corral, Provincial road N. 47								(f) Yacoraite and Olmedo formations, Cabra Corral Dam, by the bridge								
Formation/Member	Sample	Height (cm)	Hg	Formation/Member	Sample	Height (m)	Hg									
Maastrichtian Yacoraite Formation (Juramento Member)	E-37	0	7.41	Danian Olmedo Formation	Yacor-24	40.1	10.4									
	E-36 (all)	80	34.52		Yacor 23	23.5	3.66									
	E-34	150	4.07		Yacor 17	2.65	1.51									
Maastrichtian Yacoraite Formation (Alemania Member)	E-31	360	1.54	Maastrichtian Yacoraite Formation (Alemania Member)	Yacor 16	14.85	1.51									
	E-30	610	13.93		Yacor-12	6.5	19.07									
	E-29	570	6.06		Yacor-11	5.3	9.8									
	E-28	180	75.22		Yacor-9	29.1	14.26									
	E-27	180	75.22		Yacor-5	14.15	1.51									
	C-908	15	248.23		Yacor-3	11.8	18.66									
	E-26	520	6.92		Yacor-1	0	12.87									
	E-25	40	7.39													
	C-604	1100	139.97													
	C-602	80	147.97													
	C-405	2130	12.28													
C-319	370	199.64														
C-318	60	70.85														
Maastrichtian Yacoraite Formation (Güemes Member)	C-315	540	3													
	C-313	225	39.6													
	C-108	420	35.3													
	K-14-E	170	18.64													
Maastrichtian Yacoraite Fm. (Amblayo Member)	K-G	190	12.42													
	K-I	180	7.44													
	K-J	190	8.36													
	K-14-K	90	38.61													

Mercury concentrations were analyzed in twenty-eight samples from the Bottaccione Gorge, Italy (Table 3) and shown in Fig. 11. Three mercury peaks are observed, a small one (around 2 ng g⁻¹, labeled I) at about 6 m below the KTB, the largest one at the KTB (around 6 ng g⁻¹, labeled II) which coincides with a minimum of $\delta^{13}\text{C}$ and $\delta^{18}\text{O}$, and a third one at about 7 m above the KTB (around 4 ng g⁻¹, labeled III).

Hg contents analyzed in bulk samples across the KTB in the Yacoraite Formation, Salta Basin, Argentina, and in the Paraíba Basin, northeastern Brazil (Sial et al., 2013) are listed in Table 4 for comparison with Hg results in the present report.

5.2. Mercury isotopes

There is a growing interest in using Hg stable isotope signatures as source tracers for Hg of natural and anthropogenic origins. The success of such an approach strongly depends on two factors, according to Sonke and Blum (2013): (a) different natural and anthropogenic Hg sources must have analytically discernable Hg isotope signatures and (b) processes responsible for transportation/transformation of emitted or discharged Hg into the environment must not obscure the original Hg source isotope signatures. Therefore, fractionation of Hg isotopes after release has to be either small relative to source differences or is predictable enough to be corrected for, allowing estimation of the source isotopic composition (Sonke and Blum, 2013).

According to previous studies, post-emission or post-discharge Hg transformations may not significantly modify the isotopic composition of particulate Hg (Foucher et al., 2009; Sonke et al., 2010; Estrade et al., 2011; Gehrke et al., 2011; Chen et al., 2012; Sun et al., 2013). If Hg transformations do shift original source isotope signatures, then under certain conditions the shift may prove to be constant (Laffont et al., 2009, 2011; Gehrke et al., 2011). However, if multiple Hg transformations modify the Hg isotope signatures in unpredictable ways, then source tracing will become very difficult (Sonke and Blum, 2013).

Isotopic composition of Hg is potentially one way to differentiate volcanogenic from meteoritic Hg. Characterization of the isotope composition of mantle-derived Hg is difficult because samples usually contain Hg that is either not entirely mantle-derived and/or because Hg has already undergone fractionation during chemical and phase transformations (Bergquist and Blum, 2009). Sherman et al. (2010) suggested that mantle-derived Hg might be isotopically heavier than crustal Hg, with $\delta^{202}\text{Hg}$ and $\Delta^{201}\text{Hg}$ values closer to 0‰. However, Zambardi et al. (2009) have found that $\delta^{202}\text{Hg}$ values from an active volcano in Italy range from -1.74‰ to -0.11‰ for gas and particulates, respectively.

Most natural samples that obtained Hg after it had been through cycling in the environment have $\delta^{202}\text{Hg}$ values significantly different from crust and mantle values (Bergquist and Blum, 2009). These authors observed that, under the light of the data available at that time, $\delta^{202}\text{Hg}$ varies in chondrites from around -1.0‰ to 0.0‰ while in volcanic rocks, a wider fluctuation could be observed, with values between -2.0 and 0.0‰.

It is known that volcanism followed major Neoproterozoic glaciations supplying CO₂ to the atmosphere that led to greenhouse effect and further cap carbonate deposition in the aftermath of glaciations. Hg concentration linked to such volcanism can potentially be used as a test of investigation on how volcanism could influence Hg background values, especially for the mass-independent fractionation of, for example, the odd Hg isotopes (¹⁹⁹Hg and ²⁰¹Hg). The photo-reduction processes may occur in the atmosphere during the transport of Hg and thus induce odd Hg isotope anomaly (Sherman et al., 2010; Gratz et al., 2010; Chen et al., 2012), which would be ultimately deposited and recorded in surface. Sial et al. (2010) reported Hg concentration levels in the Neoproterozoic cap carbonate in the Jacoca Formation, Sergipean Belt, northeastern Brazil (~280 ng g⁻¹) which are equivalent to levels reported for the KTB in the Yacoraite Formation (~250 ng g⁻¹; Sial et al., 2013) and for the KTB layer at Stevns Klint. This is important information to this discussion since volcanism, in absence of known

meteorite impact, was the main Hg supplier to the Earth's surface during Neoproterozoic cap carbonate formation.

Sial et al. (2014) reported preliminary Hg isotope analyses of samples from the KTB layer (Fiskeler Member) at Stevns Klint, Denmark with results reported in delta notation in permil (‰) relative to NIST SRM3133 Hg standard. Three among four $\delta^{202}\text{Hg}$ values measured yielded $\delta^{202}\text{Hg}$ values between -2.0 and -1.0‰. Two samples from the Jacoca cap carbonate also yielded values in this range. Interestingly, most of these samples showed small (but significantly higher than analytical precision of 0.04‰) positive $\Delta^{201}\text{Hg}$, in favor of long-term atmosphere transport prior to deposition. This points to a promising start of Hg isotopes in the identification of Hg source.

6. Discussion and conclusions

A complication for determining the origin of Hg in KTB sections arises from that meteorites may carry large amounts of Hg, and concurrently a much larger contribution from volcanic activity at the Deccan traps could have increased worldwide Hg contents across the KTB. Ozerova et al. (1973) reported an average of 6 ng g⁻¹ Hg in stone meteorites (abundance from 0 to 33 ng g⁻¹), about 500 ng g⁻¹ in carbonaceous chondrites, iron meteorites displaying the lowest Hg concentrations (generally 0.1 ng g⁻¹). Lauretta et al. (2001) reported bulk abundances of Hg of 294 ± 15 and 30 ± 1.5 ng g⁻¹ for the Murchison and Allende carbonaceous chondrites, respectively. Shima et al. (1974) have reported a bulk Hg analysis of 1330 ng g⁻¹ for the Parambu chondrite (state of Ceará, Brazil).

The nature of the main meteorite impactor is under debate but short-lived Cr isotopic compositions of sedimentary rocks from the KTB point to a Hg-poor carbonaceous chondrite type bolide (e.g. Shukolyukov and Lugmair, 1998). However, most studies on this matter, with the exception of that of Lauretta et al. (2001), have used analytical methodologies which are not advised for Hg analysis (see for example Randa et al., 2003). Analysis of samples from the Allende chondrite reviewed by Lauretta et al. (2001), for example, gave values ranging from 16.4 to 10,020 ng g⁻¹, whereas the same meteorite analyzed by Kumar et al. (2001) gave smaller concentrations from 16.4 to 17.8 ng g⁻¹. One reason for the large variability of Hg content reported for meteorites is the erroneous determination of Hg, mostly by radiochemical neutron activation, that ignored the presence of ⁷⁵Se in the released Hg fraction, resulting in an overestimation of the Hg content (Kumar et al., 2001). Bulk Hg analysis may also blur the actual Hg content due to contamination. For example, in a sequential thermal-desorption analysis of a sample of the Antarctica meteorite Y 82050, up to 62% of the Hg present was only released at temperature higher than 300 °C, suggesting strong contamination by Hg-like halogens, and this may be the case for all Antarctica meteorites (Kumar et al., 2001). Therefore, high Hg concentrations reported for meteorites should be taken with care.

Thermal analysis of Hg-bearing chondrites have shown that the major proportion of Hg is released at relatively high temperature (>340 °C) and it is synchronous with S releasing temperature. This strongly suggests that HgS is the major Hg compound in these meteorites. The same thermal behavior has been shown for some other meteorites (e.g. Jovanovic and Reed, 1976; Kumar and Goel, 1992) and, recently, reported by Komorowski et al. (2012) who first characterized the presence of Hg-Cu-bearing metal-sulfide in an unshocked H-3 chondrite, Tieschitz meteorite.

In this report, it is evident that in the two studied sections in Europe (Stevns Klint and Gubbio) a prominent Hg peak coincides with the KTB layer, being the peak at the Stevns Klint section almost 50 times larger than that at Gubbio. Both sections are within a similar dispersion radius in relation to the Chicxulub impact site in Mexico or to the Deccan traps in India, therefore similar Hg peaks would be expected.

Among possible explanations for this discrepancy, two may be invoked: (a) divergence in time lengths and/or factors governing Hg

deposition, and (b) preservation of Hg deposit from weathering and diagenesis. It is difficult to evaluate the time length involved in Hg transport and deposition, as the nature and extension of all variables controlling it are not known (e.g. mountain chains, air and marine currents, dispersion corridors along latitudinal zones). When reaching an environment which is depleted of organic scavenging capacity due to climate changes as at the KTB, Hg⁺² would be likely kept mostly in solution, readily adsorbed onto clays and transported to sedimentary basins.

The high levels of Hg associated to the Fiskeler Member at Stevns Klint may have resulted from increased flux of volcanic-derived Hg from the landmass into the marine realm. At the Gubbio KTB site, Hg accumulation may have resulted from a similar mechanism, but a more distal, deep marine (pelagic) setting probably determined a lower Hg concentration if compared to the more proximal Stevns Klint section. However, as the KTB layer is tilted at Gubbio, this could have also facilitated a gradual Hg washing out, resulting in a more modest remaining Hg peak.

One of the three Hg peaks observed at the Bajada del Jagüel section, Neuquén Basin, coincides with the KTB sandstone layer and is much smaller than the peak at the Fiskeler Member in Stevns Klint, but three times larger than that at the KTB in Gubbio. At the Cerro Azul section, Neuquén Basin, three large Hg peaks are evident, the one few cm above the KTB reaches about 330 ng g⁻¹, much larger than that at Bajada del Jagüel and Cañadon Cholino KTB sites that exhibit peaks around 16 ng g⁻¹. The Hg peak about 3 m above the KTB in the Cerro Azul section is the largest observed in this study (415 ng g⁻¹).

Likewise the studied sections in the Neuquén Basin, two sections in the Salta Basin, Argentina, bracketing the KTB also display contrasting sizes of Hg peaks. In one section, a Hg peak reaches 250 ng g⁻¹, a bit below the KTB, twenty times larger than the corresponding Hg peak in the other section (Sial et al., 2013).

Therefore, the contrast in Hg peak size between Gubbio and Stevns Klint sites, two far distant localities, is also observed within one single basin, as observed in the Neuquén and Salta basins. To find a plausible explanation for the Hg distribution pattern, it would be necessary to know the paleogeography of these basins in the Maastrichtian–Danian time interval, precise factors governing Hg accumulation/deposition before installation of any expressive diagenetic process, and to determine Hg dispersion factors.

Preliminary Hg isotope data for clays from the KTB layer in Stevns Klint (−2.0 to −1.0‰) reported by Sial et al. (2014) lie within the range for volcanogenic Hg as reported in previous studies (e.g. Bergquist and Blum, 2009), and the odd Hg isotope anomaly also sheds light on the long-range atmosphere transport of volcanic emission. However, one cannot assure, solely based on this isotopic similarity, that Hg is volcanogenic for two reasons: (a) limited Hg isotope data available and (b) processes involved as eruption, transportation and even deposition may have induced to Hg isotope fractionation and modification of the original Hg isotope signature. A more detailed study of the behavior of Hg isotopes in the KTB localities in this study is under way to confirm the volcanogenic origin suggested in the preliminary report of Sial et al. (2014). Conclusions stemming from this study attest to the influence of endogenous mechanisms in the KTB scenario, in superposition with the asteroid impact (or impacts) and assure that volcanism played a long-term influence in the concurrent mass extinction event.

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