Errata for Weppe et al., 2020

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Palembertina desplasi, an emended name for Palembertina deplasi Weppe, Blondel, Vianey-Liaud, Pélissié, and Orliac, 2020 (Artiodactyla, Cainotheriidae) from Palembert (Quercy, SW France)

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ABSTRACT

The cainotheriid (Artiodactyla) species Palembertina deplasi Weppe et al., 2020 from the Quercy area (Palembert, France; MP18-MP19) was erected with a misspelling in the species epithet. We propose a corrected replacement name: Palembertina desplasi.

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Keywords: new genus; new species; “Cainotherium plan”; Eocene; karstic infillings; occlusion


The cainotheriid species Palembertina deplasi Weppe et al., 2020 was erected and described by dental and mandibular material from the Palembert locality (Quercy, SW France) dated of the late Eocene (MP18-MP19; Weppe et al., 2020). Palembertina deplasi is characterized by intermediate morphology between the “robiacinid” and the “derived cainotheriid” types, and allows for a better
understanding of the evolution of the dental pattern of cainotheriids. This species was named for Claude Desplas, who helped to discover many fossil-yielding localities in the Quercy phosphorites. Unfortunately, the authors misspelled the name “Desplas” in the original publication and consequently the epithet of the new cainotheriid species dedicated to him in the article of Weppe et al. (2020). The name *Palembertina desplasi* is given here as a replacement name for *P. deplasi* Weppe et al., 2020. The holotype (left M2/, PAL 21) and paratypes (left dentary fragment with M/3, PAL 60; a left dentary fragment with P/4 and M/2, PAL 62; and a right isolated M/3, PAL 64) are maintained for the definition of the species.

REFERENCES

A new Cainotherioidea (Mammalia, Artiodactyla) from Palembert (Quercy, SW France): Phylogenetic relationships and evolutionary history of the dental pattern of Cainotheriidae

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ABSTRACT

Cainotheriidae are small artiodactyls restricted to Western Europe deposits from the late Eocene to the middle Miocene. From their first occurrence in the fossil record, cainotheriids show a highly derived molar morphology compared to other endemic European artiodactyls, called the “Cainotherium plan”, and the modalities of the emergence of this family are still poorly understood. Cainotherioid dental material from the Quercy area (Palembert, France; MP18-MP19) is described in this work and referred to Oxacron courtoisii and to a new “cainotherioid” species. The latter shows an intermediate morphology between the “robiacinid” and the “derived cainotheriid” types. This allows for a better understanding of the evolution of the dental pattern of cainotheriids, and identifies the enlargement and lingual migration of the paraconule of the upper molars as a key driver. A phylogenetic analysis, based on dental characters, retrieves the new taxon as the sister group to the clade including Cainotheriinae and Oxacroninae. The new taxon represents the earliest offshoot of Cainotheriidae.

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INTRODUCTION

Cainotheriidae are small artiodactyls documented from Western European deposits. They appeared abruptly in the fossil record at the end of the Eocene (MP18-MP19) and, unlike many endemic European ungulates that disappeared around the Eocene-Oligocene transition, they successfully crossed the Grande Coupure (Stehlin, 1910) and diversified during the Early Oligocene and at the beginning of the Miocene. They became extinct during the Middle Miocene. The morphology of their brachydont selenodont molars is, moreover, unique among mammals: named by Stehlin (1906) the “Cainotherium plan”, the cainotheriids pattern is characterized by the presence of a distolingual protocone on the upper molars, as well as by a large mediolingual cuspid on the lower molars (Weppe et al., 2020).

The family Cainotheriidae is composed of five genera: Oxacron Filhol, 1884, Paroxacron Hürzeler, 1936, Plesiomeryx Gervais, 1873, Caenomeryx Hürzeler, 1936, and Cainotherium Bravard, 1828 (Blondel, 2005). Hürzeler (1936) proposed to gather Oxacron and Paroxacron, mainly known from Late Eocene deposits, in the subfamily Oxacroninae Hürzeler, 1936, while including the other three genera, mainly Oligocene, in the Cainotheriinae Camp and VanDerHoof, 1940; these two subfamilies were each recovered as monophyletic by Weppe et al. (2020). Despite the great abundance of cainotheriid remains in collections, works on cainotheriid remain scarce, particularly for Eocene localities (Legendre, 1980; Legendre and Weidmann, 2000; Weppe et al., 2020). The last 50 years of excavations in Quercy area yielded abundant remains of cainotheriids in a rather precise chronological framework and allowed to clarify certain episodes in the evolutionary history of the group (Legendre, 1980; Hooker and Weidmann, 2000; Weppe et al., 2020). The last 50 years of excavations in Quercy area yielded abundant remains of cainotheriids in a rather precise chronological framework and allowed to clarify certain episodes in the evolutionary history of the group (Legendre, 1980; Blondel, 2005; Weppe et al., 2020). However, the modalities of the emergence of this family are still poorly understood. The first occurrence of cainotheriids in the fossil record is the species Oxacron courtoisii, documented and described by Gervais (1859) in the locality of La Débruge (Vaucluse, France) referred to the MP18 level. The dental morphology of this earliest species is derived compared to other endemic European artiodactyls and already shows the “Cainotherium plan”.

Because of their unique dental morphology, the phylogenetic position of Cainotheriidae within both endemic European artiodactyls and artiodactyls is still debated, and this family has been related to various endemic European families (Romer, 1966; Webb and Taylor, 1980; Gentry and Hooker, 1988) without reaching a consensus. Similarly, others have linked them to modern groups of artiodactyls such as ruminants (Geisler and Uhen, 2005; O’Leary and Gatesy, 2008; Lihoreau et al., 2015) or tylopods (Geisler and Uhen, 2003; Geisler et al., 2007; Thewissen et al., 2007). However, the recent phylogenetic study of Weppe et al. (2020) retrieved Cainotheriidae closely related to the endemic European families Mixtotheriidae, Anoplotheriidae, and Robiacinidae. According to their results, Robiacinidae is sister taxon to Cainotheriidae and forms with the latter the super-family Cainotherioida. This relationship, already proposed by Sudre (1969, 1977, 1978), is supported in part by a distal migration of the protocone on the upper molars. The family Robiacinidae is monogenic (Robiacina Sudre, 1969) and includes three species: Robiacina minuta Sudre, 1969, Robiacina quercyi Sudre, 1977, and Robiacina lavergnensis Sudre, 1977. The family is documented in the fossil record in MP16-MP17 levels. It presents a close dental morphology to Cainotheriidae, notably at the level of premolars and on the presence of a protocone subcentral and lingual (M3/) on the upper molars (Weppe et al., 2020). However, Cainotheriidae are dentally way more derived than Robiacinidae, and 15 synapomorphies define the cainotheriid node (Weppe et al., 2020). The earliest stages of the dental history of cainotheriids is therefore not yet documented by existing species.

In this work we describe unpublished cainotheriod dental material collected in the 1980’s in the locality of Palembert (Quercy area, Tarn-et-Garonne, France; Crochet et al., 1981) close to the village of Caylus. The fossils were found in piles of excavated rocks extracted from a former phosphate infillings exploitation. The mammalian assemblage dated this locality from MP18-MP19 levels. The new material is here referred to Oxacron courtoisii (Gervais, 1852) and to a new cainotheriod taxon Palembertina deplasi gen. nov. sp.
nov. showing an intermediate morphology between the "robiacinid" and the "derived cainotheriid" type. A phylogenetic analysis based on the taxon/characters matrix of Weppe et al. (2020), allows for discussing the phylogenetic affinities of this species and supports the proposed systematic framework. *Palembertina deplasi* gen. nov. sp. nov. provides a better understanding of the early evolutionary history of Cainotherioidea.

**MATERIAL AND METHODS**

**Institutional and Anatomical Abbreviations**

BMNH: British Museum of Natural History; PAL: specimen prefix for the Palembert locality in Montpellier University; MX/: upper molars; M/X: lower molars; PX/: upper premolars; P/X: lower molars.

**Material**

The material studied here comes from the locality of Palembert in the Quercy phosphorites area (South West of France) close to the village of Caylus (Tarn-et-Garonne, France), near to Malpérée and Rouchelou localities (Figure 1). The locality of Palembert is an area comprising various karstic excavations exploited at the end of the eighteenth century. The material was collected from excavated sediment piles mostly composed of clay, pisoliths, and calcareous fragments, disposed on one of the pits' edge. Fossils were sorted out after washing and sieving the sediment with a 0.7 mm sieve. The Palembert material consists of six cainotheriid specimens: a right dentary fragment with P/3-M/3 (PAL 59) and a right dentary fragment with M/2-M/3 (PAL 61), referred to *Oxacron courtosii*; a left M2/ (PAL 21), a right dentary fragment with M/3 (PAL 60), a right dentary fragment with P/4 and M/2 (PAL 62), and a right isolated M/3 (PAL 64), referred to *Palembertina deplasi* nov. gen. nov. sp. All these specimens are housed in the collection of Montpellier University. The age of the material described here was determined using biostratigraphic association of rodent species. The rodent assemblage documented in Palembert consists of the co-occurrence of *Patriotheridomys altus*? Vianey-Liaud, 1974, *Pseudoltinomys cuvieri* (Pomel, 1853), *Suevosciurus* indet. and *Blainvillimys civracensis* Vianey-Liaud and Ringeade, 1993, or *Blainvillimys rotundidens* Schlosser, 1884, pointing to a late Priabonian age. The scarcity and the partial preservation of the rodent material hindered the ability to precisely refer this locality to a Mammalian Paleogene (MP) reference level and the latter is here correlated with the MP18-MP19 levels.

**FIGURE 1.** Topographic map (géoportail; Da Silva Pires, 2008) and geographical localization of the Palembert locality, Quercy, France.
Measurements

Measurements (in millimetres) were taken using a mesuroscope Nikon 10 (binocular with measuring device). The general occlusal surface of the tooth is described by: L, the maximal mesiodistal length; and l, the maximal buccolingual width. The buccolingual aspect of the tooth is also expressed through two measurements: la, maximal width of mesial lobe; lp, maximal width of distal lobe. The height (h) corresponds to the maximal height of the crown (labial border for lower teeth and lingual border for upper teeth). L3 and l3, respectively, correspond to a maximal length and width of the third lobe (distal) of M/3. The variability of dental measures is estimated using the variation coefficient (V).

Phylogenetic Analysis

In order to test the phylogenetic position of *Palembertina* gen. nov. within Cainotherioidea, we performed a phylogenetic analysis based on the data matrix of Weppe et al. (2020). We modified five characters and included seven new ones. The character list and character processing (Appendix I), data matrix (Appendix II), and list of apomorphies (Appendix III) are provided online. Character coding is based on direct observations of specimens and on illustrations and descriptions from the following references: Gervais (1852); Gervais (1859); Stehlin (1906); Stehlin (1908); Matthew and Reed (1910); Hürzeler (1936); Berger (1959); Emry (1978); Sudre (1978); Gentry and Hooker (1988); Berthet (2003); Hooker and Weidmann (2000); Erfurt and Métais (2007); Hooker (2007); and Rincon et al. (2015). The matrix of characters was established with the software NDE 1.0 (Page, 2001). The phylogenetic analysis was performed with PAUP* version 4.0a166 (Swofford, 2020), using a heuristic search (>12 taxa) with TBR algorithm, and 1000 repetitions (10 trees retained by repetition; by stepwise addition).

Dental Nomenclature

The nomenclature of Boisserie et al. (2010; figure 2) and Weppe et al. (2020; figure 2) was used in this work to name crests/cristids and cusps, respectively. It is detailed and figured in Appendix IV.

SYSTEMATIC PALAEONTOLOGY

Class MAMMALIA Linnaeus, 1758
Order ARTIODACTYLA Owen, 1848
Super-family CAINOTHERIOIDEA Camp and VanDerHoof, 1940

**Family CAINOTHERIIDAE** Camp and VanDerHoof, 1940

**Emended diagnosis.** Cainotherioidea with a size ranging from small rabbit-sized species as *Romerolagus diazi* (e.g., *Palembertina*, *Oxacron*, *Paroxacron*) to slightly smaller than the extant tragulid ruminant *Tragulus javanicus* (e.g., *Cainotherium*). Complete postorbital bar; enlarged bullae and orbits. Morphology of the mandible variable within the family but tends to become massive and deepen distally in several genera. Presence of diastemata within the premolar row in the Eocene species but tend to be reduced or lost (*Cainotherium*) in the Oligocene and Miocene forms. Lower incisors with two endocristulids; lower canine incisiform. Mesial premolars narrow to large; P/4 tends to be molarized; P1/ and P2/ without mesial lobe; P3/ triangular and labially concave, with a lingual cusp; P4/ triangular to sub-rectangular. Crowns of cheek teeth moderately high. Lower molars with a large and individualized centrolingual metaconulid; short to long mediocristid; centroconulid and posttendocristulid (neotrigonid of Sudre, 1977) present; absence of hypoconulid on M/1 and M/2. Upper molars sub-triangular to quadrangular with a “W”-shaped ectoloph and crescent-shaped lingual cusps; parastyle large like other styles; mediostyle and distostyle small to large; protocone subcentral to distal and paracone mesiolingual; postparacristula present; lack of preprotocrista and paracone junction. Limbs tetradactyl with a reduction of lateral digits (II and V); forelimb shorter than hind limb; tail long.

**Remarks.** The justification for the erection of the new genus and species *Palembertina deplasi*, and its integration in the Cainotheriidae family, is provided in the following section and follows the results of phylogenetic analysis.

*Palembertina*, new genus

zoobank.org/3C5849FC-3895-4100-8494-0ADFC2F617F7

*Palembertina deplasi*, new species

zoobank.org/AA39B90C-5786-457F-97B8-10D64E21CA10

Figures 2E–K, 3C–D; Appendix V

*Oxacron? courtoisii*; Hooker and Weidmann 2000: 89 p., fig. 55j. (Left M1/ BMNH.30674, Bravard cat. no. 344, from La Débruge).

**Etymology.** *Palembertina* refers to the Palembert locality. The species is named for Claude Deplas, who helped to discover many localities and fossils in the Quercy phosphorites.
FIGURE 2. Material of the dentition of Oxacron courtoisii (A-D) and Palembertina deplasi gen. nov. sp. nov. (E-K). A, B, right dentary fragment with P3-M3 (PAL 59; mirror view); C, D, right dentary fragment with M2-M3 (PAL 61; mirror view); E, F, left dentary fragment with M3 (PAL 60); G, H, left dentary fragment with P4 and M2 (PAL 62); I, J, left M2/ (PAL 21); K, right isolated M3 (PAL 64; mirror view). In lingual view: J. In occlusal view: A, C, E, G, I, K. In buccal view: B, D, F, H. Scale bar equals 1 mm.
Holotype. Left M2/ (PAL 21), from Palembert (Tarn-et-Garonne, France).

Paratype. Left dentary fragment with M3 (PAL 60); left dentary fragment with P4 and M2 (PAL 62); right isolated M3 (PAL 64), from Palembert (Tarn-et-Garonne, France).

Diagnosis. Genus of Cainotherioidea, similar in size to Oxacron (400-600g). Upper molar with a centrolingual protocone without crest; parastyle similar to other styles; looped and distally inclined mesostyle; distostyle and mediostyle isolated and large; short and continuous premetacristula; long postparacristula; mesial cingulum pronounced. P4 with a talonid basin buccolingually compressed and a paraconid weakly lingually inclined. Lower molars with a small paraconid; strong lingual position of the protoconid; short medio cristid; talonid basin large with entoconid slightly more distal than hypoconid; large individualized metaconulid close to the metaconid; neotrigonid present; hypoconid absent on M1 and M2; large basin-shaped hypoconulid with a long and weakly marked posthypocristulid, and distal to the entoconid on M3. Mandible with short diastema between P2 and P3; horizontal ramus gracile, relatively shallow and height is rather constant between the premolar area and the last molar; strong incisura vasorum.

Differential diagnosis. The upper molars of Palembertina gen. nov. differs from other Cainotheriidae by the presence of a centrolingual protocone without crests, by isolated and larger distostyle and mediostyle, by a larger mesial and distal cingulum, and by the absence of a distally inclined mesostyle. It differs also by the presence of a smaller talonid basin on the P4. The lower molars of Palembertina gen. nov. are distinguished from those of other Cainotheriidae by a more lingual protoconid, a metaconulid closer to the metaconid with a shorter medio cristid, a larger talonid basin, a smaller centroconulid and shorter postendocentrocristulid, a less distal entoconid (close to the buccolingual position of hypoconid), and a basin-shaped hypoconulid more distal with a long posthypocristulid.

Palembertina gen. nov. differs from Robiacini dae on its upper molars by a stronger dilambdodonty, styles less crested, a looped mesostyle (postparacrista and premetacrista convex buccally), the presence of a postparacristula, the absence of preprotocrista, a shorter and continuous premetacristula, a longer preparacrista, the presence of a mediostyle and distostyle, and a larger mesial and distal cingulum. It differs also on its lower molars by an individual, larger and more distal metaconulid, the presence of a medio cristid and a neotrigonid, a slightly more distal entoconid (compared to buccolingual hypoconid position), and the absence of a hypoconulid on M1 and M2.

Material from Palembert. One left M2/ (PAL 21); one left mandibular fragment with M3 (PAL 60); one left mandibular fragment with M2 and P4 (PAL 62); one right isolated M3 (PAL 64). Palembert is located in the Quercy phosphorites area (South West of France) close to the village of Caylus (Tarn-et-Garonne, France).

Description. The upper dentition of Palembertina deplasi nov. gen. nov. sp. is only documented by the M2/ (PAL 21; Figures 2I–J, 3C). It bears five main cusps. The paraconule is mesiolingual and bears a preparacristula connecting the preparacristyla; the metaconule is large and presents a short and continuous premetacristula, as well as a postmetacristula joining the postmetacristyla; the protocone lacks a crest and is centrolingually positioned. The styles are also well-pronounced: the buccal styles are weakly crested with a loop-like and distally inclined mesostyle; the lingual styles, distostyle and mediostyle, are isolated and large. Finally, the M2/ has pronounced mesial and distal cingula.

The lower dentition of Palembertina deplasi nov. gen. nov. sp. is partly documented by a damaged P4 and M2, a complete, unworn M3, and a damaged M3. The crown of P4 (PAL 62; Figure 2G–H) is short and narrow and presents three main cusps: a large protoconule, metaconid, and paraconid. The protoconule and the metaconule are close to each other, the protoconule being more mesial than the metaconule. On the mesial part of the tooth, the paraconid is slightly inclined lingually. On the distal part, the talonid basin is buccolingually compressed and bears a small entoconid and a worn hypoconid. A postprotoconulid is visible between the protoconule and the hypoconid. The morphology of the lower molars is mainly based on M3 (PAL 60, PAL 64; Figures 2E–F, 2K, 3D), because of the poor preservation of the M2/ (PAL 62; Figure 2G–H). The lower molars present three lingual cusps: a mesial metaconule, a sub-central metaconulid, and a distal entoconid. On the trigonid, the paraconid is small, and the protoconule is very lingual. The talonid is large with a small centroconulid close to the metaconulid, and a postendocentrocristulid weakly expanded distolingually. The metaconulid is large, close to the metaconule, and the medio cristid is short. A large valley is present between the metaconulid and the entoconid. The entoconid has a loop-like shape and is slightly more distal than the hypoconid. M3 (PAL 60, PAL
**FIGURE 3.** Dental diagrams of M2/ and M3 in occlusal view. **A-B,** Robiacina; **C-D,** Palembertina deplasi gen. nov. sp. nov.; **E-F,** derived Cainotheriidae. In color, the diagnosis characters of interest: blue, protocone; light blue, paraconule; pink, metaconulid; red, entoconid; orange, hypoconulid; brown, neotrigonid (Sudre, 1977). Abbreviations: Centroulid., centroconulid; dist., distostyle; Entod., entoconid; Hypod., hypoconid; Hypoulid., hypoconulid; med., mediostyle; mes., mesostyle; Meta., metacone; Metad., metaconid; Metaul., metaconule; Metaulid., metaconulid; met., metastyle; Para., paracone; Parad., paraconid; Paraul., paraconule; par., parastyle; Proto., protocone; Protod., protoconid. Circles: cusps/cuspids/styles/conules/conulids; full lines: crests; dashed lines: valleys/fossa; dotted lines: cingula/cingulids. Arrows indicate the mesio-lingual side. For more detailed dental labelling, see APPENDIX IV. Scale bar equals 1 mm.
64; Figures 2E–F, 3D, Figure 2K) present a large hypoconulid, basin-shaped and distal to the entoconid. It connects to the rest of the talonid by the prehypocristid, which joins the posthypocristid and postentocristid. Moreover, the posthypocristid is weakly pronounced and extended lingually. Finally, the mesial cingulid is large.

The mandibular morphology of *Palombertina deplasi* nov. gen. nov. sp. is documented from the level of the second premolar (PAL 62; Figure 2G–H) mesially, to the base of the mandibular angle (PAL 60; Figure 2E–F) distally. The mandible (PAL 62) presents a small diastema between P/2 and P/3 and two mental foramina on the horizontal ramus, one below the mesial part of P/2 and one below the mesial part of P/4 (Figure 2H). The horizontal ramus is gracile, relatively shallow, and its height is rather constant between the premolar area and the last molar. A wide gap is present between the base of the ascending ramus and the distal margin of M/3 (Figure 2F). The angular apophysis is not preserved but seems to be medi ally inclined; it is mesially delimited by a shallow incisura vasorum (PAL 60).

Sub-family OXACRONINAE Hürzeler, 1936

Genus *Oxacron* Filhol, 1884

*Oxacron courtoisii* (Gervais, 1852)

Figure 2A–D; Appendix VI

*Cainotherium courtoisii*; Gervais, 1852: 80 p., plate XXXIV fig. 6 and plate XXXV fig. 4.

*Cainotherium courtoisii*; Gervais, 1859: 162 p., plate XXXIV fig. 6, plate XXXV fig. 4.

*Plesiomyx quinquedentatus*; Filhol, 1877: 430 p., fig. 314-316.

*Oxacron minimus*; Filhol, 1884: 64 p.

*Oxacron courtoisii* (Gervais, 1852); Stehlin, 1906: 677 p., fig. XCIV-XCVI.

**Lectotype.** Left mandibular fragment with P/3–M/3 (Gervais, 1852), from the lignites of La Débruge (MP18; Vaucouleuse, France).

**Paralectotype.** Left maxillary fragment with P4/-M3/ (Gervais, 1852), from the lignites of La Débruge (MP18; Vaucouleuse, France).

**Remarks.** Gervais (1852) erected the species *Cainotherium courtoisii*, and Stehlin (1906) subsequently placed it in the genus *Oxacron*. Gervais (1852) described and figured the species based on the small sized cainotheiids specimens from La Débruge but did not define a holotype. According to the International Code of Zoological Nomenclature, the name *Oxacron courtoisii* is available but not valid; therefore, we define here a lectotype and a paralectotype for *Oxacron courtoisii*, corresponding to the material from La Débruge originally figured by Gervais (1852): left mandibular fragment with P/3-M/3 (plate XXXV, fig. 4) for the lectotype; left maxillary fragment with P4/-M3/ (plate XXXIV, fig. 6) for the paralectotype.

**Emended Diagnosis.** Sole species of *Oxacron*, with diastemata short to long between P1–P2/ and P2–P3, and very short to absent between P1–P2 and C–P1. Premolars short; P1/, P/1 and P/2 caniniforms; P3/ with short mesial lobe and small protocone; P4/ triangular and narrow buccolingually; upper molars subtriangular, slightly distally inclined, with small paraconule. Mandible with a gracile and relatively shallow horizontal ramus of constant height between the premolar area and the last molar; coronoid process of the mandible high above the occlusal surface; mandibular condyle relatively long mediolaterally; angular apophysis slightly extended ventrally and elongated distally; strong incisura vasorum. Cranium without ethmoidal fissures; rostrum elongated and mediolaterally constricted; contact maxillofrontal large; sagittal crest lowly pronounced; choane large; basisphe noid apically rounded.

**Differential diagnosis.** *Oxacron* differs from Cainothy rerinae by the presence of subtriangular upper molars without crested styles, a shorter postparacristula, narrow premolars, a weaker postprotoconulid, a biradicate P1, diastemata between P2–P3 and P1–P2/, and a protocone more mesial than the paracone on P4/. It differs also from *Plesiomyx* and *Caenomeryx* species by smaller size, a smaller protocone on P3/, as well as the absence of a protocone on P2/. It differs from *Cainotherium* by smaller size, the absence of cingula/ids on pre molars, a triangular P3/, and the presence of a valley between the entoconid and the mesoconid on lower molars. *Oxacron* is distinguished from the other Oxacroninae *Paroxacron* by shorter premolars, a P1/, P/1, and P/2 caniniforms, a shorter mesial lobe on the P3/, a triangular P4/, as well as a smaller paraconule on the upper molars. Finally, *Oxacron* differs from *Palombertina* gen. nov. by many characters (see above).

**Material from Palombert.** One right dentary fragment with P/3-M/3 (PAL 59); one right dentary fragment with M/2-M/3 (PAL 61). Palombert is located in the Quercy phosphorites area (South West of France) close to the village of Caylus (Tarn-et-Garonne, France).

**Description.** *Oxacron courtoisii* from Palombert is only documented by its lower dentition with two fragmentary dentaries. According to the alveolar pattern of PAL 59 (Figure 2A–B), the P/1 is biradicate. Very short diastemata are present between
P/1-P/2 and P/2-P/3. P/3 and P/4 are short. The P/3 presents a strong protoconid and a short distal cristid lingually curved. The P/4 shows a protoconid more mesial than the metaconid. Its talonid basin is large with a small postprotoconulid and a small hypoconid. The entoconid is very weak.

The lower molars (PAL 59, PAL 61; Figure 2A–D) present the “Cainotherium plan” with three large lingual cusps: a mesial metaconid, a medial metaconulid, and a distal entoconid. The paraconid is small. The metaconulid is large, far from the metaconid and separated by a long mediocristid. The talonid is bucco-lingually constricted with a large centroconulid and a postendocentroconulid expanded distolingually. The entoconid is looped and more distal than the hypoconid. The M/3 presents a large hypoconulid without a posthypocristulid. The mesial cingulid is rather large.

The mandible (PAL 59) shows two mental foramina on the horizontal ramus, one below the mesial part of P/2 and one below the mesial part of P/4 (Figure 2B). The horizontal ramus is gracile and shallow, and its height is rather constant between the premolar area and the last molar. The angular apophysis, not preserved, is mesially delimited by a strong incisura vasorum (PAL 59, PAL 60; Figure 2B, 2D).

**RESULTS**

**Phylogenetic Analysis**

The cladistic analysis retained a single parsimonious tree (length = 183, Consistency Index [CI] = 0.48, Retention Index [RI] = 0.73, CI with only parsimony-informative characters = 0.45; Figure 4), reflecting a well-structured signal and relatively weak homoplasy. The topology is congruent with that of Weppe et al. (2020), the new taxon **Palembertina deplasi** nov. gen. nov. sp. finding its place within the monophyletic Cainotheriidae, as the sister taxon to the clade C (Oxacroninae, Cainotherinae). This position is supported by six non-ambiguous synapomorphies (BI = 4): a more distal (131-2; RI = 0.94) and lingual protocone (191; RI = 0.50) on upper molars and M3/, respectively, a sub-equal paracone and metacone on P3/ (661; RI = 1.00), the absence of protocone on P2/ (700; RI = 0.57), a small hypoconid on P/2-P/3 (740; RI = 0.75), and the presence of a postprotoconulid on the lower premolars (751; RI = 1.00). Robiacinidae (BI = 2) are characterized by the presence of a small paraconule (160; RI = 0.43) and a short preparacrista (271; RI = 1.00) on upper molars, as well as the presence of a metaconid on the P/3 (721; RI = 0.50).

Relationships outside Cainotherioidea are similar to that retrieved by Weppe et al. (2020). The Bremer index of some nodes is, however, higher in this new phylogenetic analysis.
DISCUSSION

Taxonomic Attribution of Palembert Cainotheriid Material

Two species of cainotheriid are described from the locality of Palembert: *Palembertina deplasi* gen. nov. sp. nov. (PAL 21, PAL 60, PAL 62, and PAL 64; Figures 2E–K, 3C–D) and *Oxa- crocn courtoisii* (PAL 59 and PAL 61; Figure 2A–D). These species are very similar in size (Appendix V, VI), but our decision to distinguish two species is based on numerous morphological differences: i) on the premolar proportion: the P/4 (PAL 62; Figure 2G) referred to *P. deplasi* gen. nov. sp. nov. presents a talonid basin strongly compressed buccolingually and a paraconid slightly inclined lingually, while the P/4 of *O. courtoisii* (PAL 59; Figure 2A) shows a large talonid basin and a paraconid strongly inclined lingually; ii) on the pattern structure of the upper molars: the material attributed to *P. deplasi* gen. nov. sp. nov. (PAL 21; Figures 2I–J, 3C) presents a paraconule large and lingual, a protocone without crest in a centrolingual position, as well as large and individualized lingual styles. In contrast, the upper molars of *O. courtoisii* (“Cainotherium plan”; Figure 3E), not documented in the material of Palembert but present in other Quercy localities, bear a small paraconule, a crested distal protocone, and small lingual styles; iii) on the pattern of the lower molars structure: *P. deplasi* gen. nov. sp. nov. (PAL 60, PAL 62 and PAL 64; Figures 2E–H, 2K, 3D) present molars with a trigonid and a talonid buccolingually large, a metaconulid in a mediolingual position, a neotrigonid weakly pronounced, and a crested basin-shaped hypoconulid. By opposition, PAL 59 and PAL 61 (Figure 2A–D) referred to *O. courtoisii* show molars with a compressed buccolingually trigonid and talonid, a more distal metaconulid, a neotrigonid with a large centroconulid and a long postendocentroconulid, as well as a hypoconulid without cristid (“Cainotherium plan”; Figure 3F); iv) on the man-
The mandible of the distal margin of M/3, while the mandibles of between the base of the ascending ramus and the incisura vasorum slightly marked and a wide gap visible. The number of specimens of P. deplasi gen. nov. sp. nov. is, however, small, and it is unfortunately not possible to discuss the intra-specific variability of this species. Intraspecific variation in the different cainotheriid species is generally manifested at the level of the length of the diastemata, the height of the mandibular corpus, and the degree of incisura vasorum marking.

Referral of the mandibles PAL 59 and PAL 61 (Figure 2A–D) to the species Oxacron courtoisi (Figure 2B, 2D) present a strongly marked incisura vasorum and a faintly visible gap. The number of specimens of P. deplasi gen. nov. sp. nov. is, however, small, and it is unfortunately not possible to discuss the intra-specific variability of this species. Intraspecific variation in the different cainotheriid species is generally manifested at the level of the length of the diastemata, the height of the mandibular corpus, and the degree of incisura vasorum marking.

This study describes a new cainotheriid taxon: Palembertina deplasi gen. nov. sp. The association between the isolated upper molar PAL 21 (Figures 2I–J, 3C) and the lower dentition (mandibles PAL 60, PAL 62, and the isolated M/3 PAL 64; Figures 2E–H, 2K, 3D) and the attribution to the same species has been validated by testing their occlusion (see Figure 5B, as paraconule and metaconulid). Palembertina gen. nov. presents a unique combination of characters, showing an intermediate morphology between the “robiacinid” (Figure 3A–B) and the “derived cainotheriid” pattern (“Cainotherium plan”; Figure 3E–F). It shows both: i) plesiomorphic characters retrieved the Robiacinidae: the protocone is centrolingual on the upper molars; on the lower molars, the talonid is large, the hypoconulid is in basin-shaped with a long posthypocristulid, and the entoconid is lowly distal; and on the P/4, the talonid basin is buccolingually compressed; ii) derived characters of the Cainotheriidae: on the upper molars, the dilambdodonty is strong, the parastyle size is similar to other styles, lingual styles (mediostyle and distostyle), and a postparacristula are present, and a preprotocrista and paraconule junction is absent; on the lower molars, the metaconulid is large, individualized and more distal, a neotrigonid is present, and a hypoconulid on M/1-M/2 is absent (Weppe et al., 2020; this study). We propose here to include Palembertina gen. nov. within the family Cainotheriidae. This contrasts with the definition of Stehlin (1906) who defined the Cainotheriidae by the presence of a distal protocone on the upper molars (“Cainotherium plan”; Figure 3E), which is not the case in Palembertina gen. nov. Indeed, according to our cladistic analysis, the character state “distal protocone” (132; RI = 0.94) is acquired at the node C, which could have been a good candidate as the Cainotheriidae node. However, we decided to include Palembertina gen. nov. within Cainotheriidae based on the shared character of nine non-ambiguous synapomorphies (see above; BI = 4), among which some characteristics are unique to this family: on upper molars, the presence of a medistyle and distostyle; on lower molars, the presence of a large and individualized metaconulid, of a mediocristid, and of a neotrigonid. Palembertina gen. nov. therefore, represents the first offshoot of the Cainotheriidae family and differs from the other members of the family by i) retaining the plesiomorphic state for the six non-ambiguous synapomorphies defining the clade C (see above and phylogenetic analysis), gathering Oxacroninae and Cainotheriinae and ii) showing unique features (three autapomorphies), such as the presence large lingual styles (mediostyle and distostyle) on the upper molars, unique within artiodactyls. These characters altogether (see also Systematic Palaeontology) led us to erect Palembertina deplasi as a new genus and a new species.

Early Evolutionary History of Cainotheriidae

Cainotheriidae appeared abruptly in the fossil record in the middle levels of the Priabonian with Oxacron courtoisi (Erfurt and Métais, 2007; Weppe et al., 2020). The specimens of Palembertina are morphologically and biometrically similar to those described by Gervais (1852; plate XXXV, fig. 4; lectotype), Hürzeler (1936; fig. 67), and Legendre (1980) (Escamps (MP19; Quercy)), and referred to as Oxacron courtoisi. They are also similar to those from other Quercy localities: Rosière 1–4 (MP 19), Sindou D (MP 19), Tabarly (MP 20) and Pecarel (MP 20) referred to this species (Remy et al., 1987).

This study describes a new cainotheriid taxon: Palembertina deplasi gen. nov. sp. The association between the isolated upper molar PAL 21 (Figures 2I–J, 3C) and the lower dentition (mandibles PAL 60, PAL 62, and the isolated M/3 PAL 64; Figures 2E–H, 2K, 3D) and the attribution to the same species has been validated by testing their occlusion (see Figure 5B, as paraconule and metaconulid). Palembertina gen. nov. presents a unique combination of characters, showing an intermediate morphology between the “robiacinid” (Figure 3A–B) and the “derived cainotheriid” pattern (“Cainotherium plan”; Figure 3E–F). It shows both: i) plesiomorphic characters retrieved the Robiacinidae: the protocone is centrolingual on the upper molars; on the lower molars, the talonid is large, the hypoconulid is in basin-shaped with a long posthypocristulid, and the entoconid is lowly distal; and on the P/4, the talonid basin is buccolingually compressed; ii) derived characters of the Cainotheriidae: on the upper molars, the dilambdodonty is strong, the parastyle size is similar to other styles, lingual styles (mediostyle and distostyle), and a postparacristula are present, and a preprotocrista and paraconule junction is absent; on the lower molars, the metaconulid is large, individualized and more distal, a neotrigonid is present, and a hypoconulid on M/1-M/2 is absent (Weppe et al., 2020; this study). We propose here to include Palembertina gen. nov. within the family Cainotheriidae. This contrasts with the definition of Stehlin (1906) who defined the Cainotheriidae by the presence of a distal protocone on the upper molars (“Cainotherium plan”; Figure 3E), which is not the case in Palembertina gen. nov. Indeed, according to our cladistic analysis, the character state “distal protocone” (132; RI = 0.94) is acquired at the node C, which could have been a good candidate as the Cainotheriidae node. However, we decided to include Palembertina gen. nov. within Cainotheriidae based on the shared character of nine non-ambiguous synapomorphies (see above; BI = 4), among which some characteristics are unique to this family: on upper molars, the presence of a medistyle and distostyle; on lower molars, the presence of a large and individualized metaconulid, of a mediocristid, and of a neotrigonid. Palembertina gen. nov. therefore, represents the first offshoot of the Cainotheriidae family and differs from the other members of the family by i) retaining the plesiomorphic state for the six non-ambiguous synapomorphies defining the clade C (see above and phylogenetic analysis), gathering Oxacroninae and Cainotheriinae and ii) showing unique features (three autapomorphies), such as the presence large lingual styles (mediostyle and distostyle) on the upper molars, unique within artiodactyls. These characters altogether (see also Systematic Palaeontology) led us to erect Palembertina deplasi as a new genus and a new species.
lingual cusps instead of the two generally observed in Artiodactyla (metaconid and entoconid). The “Cainotherium plan” therefore implies a rearrangement of upper molar cusps and an increase of the number of cuspids on lower molars.

Different hypotheses of cusps/cuspids homology have been proposed to explain the “Cainotherium plan” (for a general review see Weppe et al., 2020). According to Stehlin (1910) and Hürzeler (1936), the cainotheriid morphology of the upper molars results from a distal migration of the protocone and a lingual migration of the paraconule. The peculiar pattern of the lower molars, according to Sudre (1977) and Weppe et al. (2020), results from the development and distal migration of a supernumerary cuspid (metaconulid; Weppe et al., 2020) originating from the metaconid.

The dental morphology of the molars of the Robiacinidae (Figure 3A–B), sister group to Cainotheriidae, allows for a polarization of the dental characters of the molars and provides the start of an answer to understand the modalities of the cainotheriid pattern evolution (Figures 5, 6). Compared to other Palaeogene artiodactyls, the protocone is displaced distally on the upper molars (centrolingual), and the lower molars bear a small metaconulid in contact with the metaconid (this character is also present in Anoplotheriidae; Weppe et al., 2020 and this study). However, a great morphological gap (Figure 3) remains between Robiacinidae and derived Cainotheriidae (Cainotheriinae and Oxacroninae; “Cainotherium plan”). The new genus Palembertina described in this study documents an early stage of the evolutionary history of the cainotheriid dental pattern. Indeed, the upper molars of Palembertina deplasi nov. gen. nov. sp. (Figure 3C–D) present a plesiomorphic pattern with a centrolingual location of the protocone, similar to Robiacinidae, while the lower molars present a more derived morphology similar to derived Cainotheriidae with, notably, a large metaconulid (see taxonomic attribution of palembert cainotheriid material for other characters). The “Cainotherium plan” of derived Cainotheriidae (Figure 3E–F) is then formed by a more distal migration of the protocone and metaconulid on the upper and lower molars.

Concurrently to the “Cainotherium plan”, the Cainotheriidae also present a peculiar pattern of occlusion, directly linked to cusp/cuspids size and location (Figure 5). Robiacina shows rather triangular outlines of the upper molars related to the small size of the paraconule (Figures 3A, 5A) while the Palembertina gen. nov. and more derived cainotheriid show more quadrangular upper molars with a bigger, more lingual, paraconule (Figures 3C, 3E, 5B–C). In occlusion (Figure 5), the paraconule is interposed between the two mesio-lingual cusps of the lower molars (metaconid and metaconulid), and changes in size and position of these structures are related. The lingual development of the paraconule in Palembertina gen. nov. corresponds with the enlargement and distal displacement of the metaconulid on lower molars (Figure 5B). This tendency goes on to an extreme degree in more derived cainotheriids, accompanied by the fully distal migration of the protocone.
and metaconulid on the upper and lower molars (Figure 5C). The famous “Cainotherium plan” therefore results from a succession of dental changes (Figure 6); we agree with Sudre (1977) that the paraconule can be regarded as a key feature in the achievement of this pattern (Figure 5).

CONCLUSIONS

This study presents the first description of a new cainotherioid, *Palembertina deplasi* nov. gen. nov. sp. from the locality of Palembert (Quercy, France). It documents an early stage in the evolutionary history of the dental pattern of cainotheriids and shows an intermediate morphology between the “robiacinid” type and “derived cainotheriid” type (“Cainotherium plan”). The dental morphology of *Palembertina* gen. nov. allows us to understand better the evolution of the dental pattern of cainotheriids, defining the development and the lingual migration of the paraconule of the upper molars as a key feature of its formation.

Our phylogenetic analysis, based mainly on the matrix of Weppe et al. (2020), places *Palembertina* gen. nov. as the sister group to the clade including Cainotheriinae and Oxacroninae. Because of the number and non homoplastic nature of synapomorphies supporting their node, we consider *Palembertina* gen. nov. as an early offshoot of the Cainotheriidae family.
ACKNOWLEDGEMENTS

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APPENDICES

Appendix I-III are available online in a zipped file at https://palaeo-electronica.org/content/2020/3216-new-species-from-palembert.

APPENDIX I.

List of Dental Characters. (A) for Ordered characters; (NA) for Unordered characters.

APPENDIX II.

Taxa/Characters Matrix from Dental Material (Nexus Format).

APPENDIX III.

List of Apomorphy. --> for Ambiguous synapomorphy; ==> for Non-ambiguous synapomorphy
APPENDIX IV.

Dental diagrams of derived Cainotheriidae dentition according to dental nomenclature of Boisserie et al. (2010) and Weppe et al. (2020). For upper dentition, the term crista refers to crests originating from main cusps, the term cristula refers to crests originating from conules, the term cristyla refers to crests originating from styles. For lower dentition, the term cristid refers to crests originating from main cuspids, the term cristulid refers to crests originating from conulids. A, B, M2/; C, D, M3/.

Abbreviations: Centro-ulid, centroconulid; dist., distostyle; endometac, endometacrista; endoparac, endoparacrysta; Ento-d, entoconid; Hypo-ulid, hypoconulid; mediocid, mediocristid; med., mediostyle; mes., mesostyle; Meta, metacone; Meta-ulid, metaconulid; Meta-ulid, metaconulid; med., metaostyle; Para., paracone; Parul., paraconulid; par., parastyle; postcentro-ulid, postcentrocristulid; postdistostyle, postdistocristyla; postectocristyla, postectocristyla; postentocristyla, postentocristyla; posthypocristid, posthypocristid; postmetacristid, postmetacristid; postparacristid, postparacristid; postparacyla, postparacyla; precentro-ulid, precentrocristulid; predistostyle, predistocristyla; preentocristyla, preentocristyla; prehypocristid, prehypocristid; premetacristid, premetacristid; preparacristid, preparacristid; preparacyla, preparacyla; preproto-cristid, preproto-cristid; Proto-d, protoconid. Circles: cusps/styles/conules/conulids; full lines: crest; dashed lines: valleys/fossa; dotted lines: cingula/cingulids; bold full lines: crown contour. Arrows indicate the mesio-lingual side. Scale bar equals 1 mm.
APPENDIX V.

Table with measurements (mm) and statistical data of the permanent teeth of *Palembertina deplasi* gen. nov. sp. nov. from Palembert. N, number of specimens; L/lp/la/h/L3 and l3, see Methods; LV, limit values; M ± SE, Mean ± Standard Error; σ, standard deviation; V, coefficient of variation.

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## APPENDIX VI.

Table with measurements (mm) and statistical data of the permanent teeth of *Oxacron courtoisii* from Palembert. N, number of specimens; L/lp/la/h/L3 and l3, see Methods; LV, limit values; M ± SE, Mean ± Standard Error; σ, standard deviation; V, coefficient of variation.

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