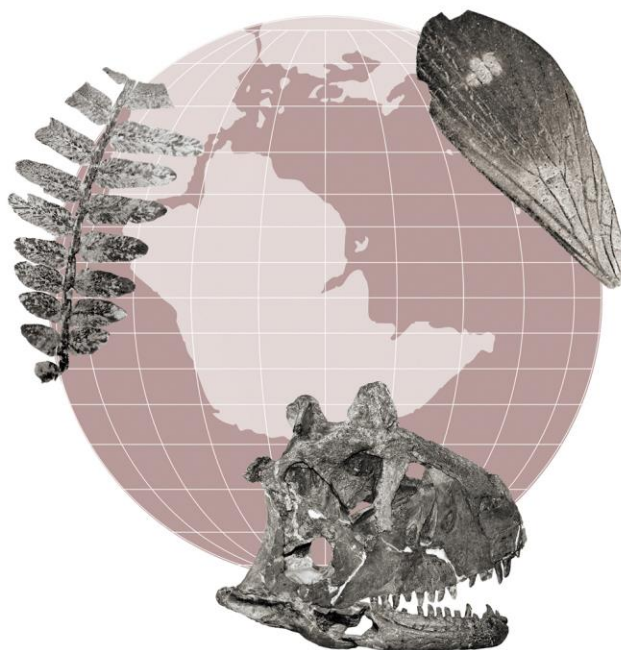




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1 **NEW CONTRIBUTIONS TO THE KNOWLEDGE OF ABELISAUROIDAE**  
2 **(DINOSAURIA, THEROPODA) FROM THE UPPER CRETACEOUS IBERO-**  
3 **ARMORICAN LANDMASS.**

4

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16

17 Running Header: ISASMENDI AND MALAFAIA: ABELISAUROID REMAINS FROM  
18 THE IBERIAN PENINSULA

19 Short Description: A revision of theropod remains from Ibero-Armorica, now identified  
20 as belonging to Abelisauridae, contributes to elucidating abelisaurid paleobiodiversity  
21 during the Late Cretaceous.

22

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24 **Abstract.** Abelisaurids were medium- to large-bodied theropod dinosaurs that inhabited  
25 Gondwana and Europe, even becoming the apex predators in the latest Cretaceous  
26 ecosystems of many of these areas. European abelisaurids remain elusive, but their  
27 remains are becoming increasingly abundant in the Late Cretaceous deposits of Ibero-  
28 Armorica. In this work, systematic, morphometric, and cladistic analyses of tooth  
29 samples from the Campanian–Maastrichtian deposits of three localities (Chera 2,  
30 Montrebei and Viso) from the Iberian Peninsula have allowed these elements to be  
31 reassigned to abelisaurids. The specimen from Chera 2 is assigned to *Arcovenator* sp.  
32 whereas teeth from Montrebei and Viso are classified as Abelisauridae indet. The latter  
33 represents the first confirmed abelisaurid remain from the Cretaceous of Portugal. The  
34 axial remains identified as belonging to *Rhabdodon* from the Laño site are here  
35 attributed to cf. *Arcovenator*. These findings indicate that abelisaurids were the only  
36 apex terrestrial predators and among the most abundant theropods in the Late  
37 Cretaceous faunas of the Ibero-Armorican landmass. The reclassification of mid- to  
38 large-sized isolated teeth from Ibero-Armorica as abelisaurids, rather than  
39 carcharodontosaurids or closely related forms, suggests that Abelisauridae had already  
40 become the dominant apex theropod lineage by the Cenomanian. The abelisauroid fossil  
41 record in Ibero-Armorica spans from the Albian to the latest Maastrichtian, indicating a  
42 complex and temporally extensive presence. Despite most of the specimens being  
43 fragmentary, the available evidence supports the persistence and diversification of  
44 abelisaurids across the Ibero-Armorican domain, with multiple evolutionary lineages  
45 arising either from a possible Albian stock or resulting from successive dispersals,  
46 followed by insular diversification throughout the Late Cretaceous.

47 **Keywords.** Dinosaur. Ceratosauria. Abelisaurid. Late Cretaceous. Iberian Peninsula.  
48 Paleobiogeography.

49 **Resumen.** NUEVOS APORTES AL CONOCIMIENTO DE ABELISAUROIDAE  
50 (THEROPODA, DINOSAURIA) DEL CRETÁCICO SUPERIOR  
51 IBEROARMÓRICANO. Los abelisáuridos fueron dinosaurios terópodos de tamaño  
52 medio a grande que habitaron Gondwana y Europa, llegando incluso a convertirse en los  
53 depredadores ápice en los ecosistemas del Cretácico final de muchas de estas regiones.  
54 Los restos europeos de abelisáuridos siguen siendo limitados, pero sus restos están  
55 comenzando a ser más abundantes en los depósitos del Cretácico Superior de Ibero-  
56 Armorica. En este trabajo, los análisis sistemáticos, morfométricos y cladísticos de  
57 muestras de dientes del Campaniense–Maastrichtiense de tres localidades (Chera 2,  
58 Montrebei y Viso) de la Península Ibérica han permitido reasignar estos elementos a  
59 Abelisauridae. El espécimen de Chera 2 se asigna a *Arcovenator* sp., mientras que los  
60 dientes de Montrebei y Viso se clasifican como Abelisauridae indet. Este último  
61 representa el primer resto confirmado de abelisáurido del Cretácico de Portugal. Los  
62 restos axiales identificados anteriormente como pertenecientes a *Rhabdodon*  
63 procedentes del yacimiento de Laño se atribuyen aquí a cf. *Arcovenator*. El registro  
64 fósil muestra que los abelisáuridos fueron los únicos depredadores terrestres dominantes  
65 y uno de los grupos de terópodos más abundantes en las faunas del Cretácico Superior  
66 de Ibero-Armórica. La reclasificación de dientes aislados de tamaño medio a grande  
67 como pertenecientes a abelisáuridos, en lugar de a carcarodontosáuridos u otros grupos  
68 cercanos, sugiere que Abelisauridae ya se había convertido en el grupo ápice de grandes  
69 depredadores hacia el Cenomaniense. El registro de abelisauroideos en Ibero-Armórica  
70 abarca desde el Albiense hasta el Maastrichtiense final, lo que indica una presencia  
71 compleja y prolongada en el tiempo de este grupo de terópodos. A pesar de la naturaleza  
72 fragmentaria de la mayoría de los restos, la evidencia apunta a la persistencia y  
73 diversificación de los abelisáuridos en el dominio Ibero-Armoricano, con múltiples

74 linajes evolutivos surgidos a partir de un posible stock albiense o de dispersiones  
75 sucesivas, que experimentaron una diversificación a lo largo del Cretácico Tardío en  
76 Europa.  
77 **Palabras clave.** Dinosaurio. Ceratosauria. Abelisáurido. Cretácico Tardío. Península  
78 ibérica. Paleobiogeografía.

79 ABELISAURIDAE comprises medium- to large-bodied (5–9 m in length) ceratosaurian  
80 theropods characterized by deep, heavily sculptured skulls with bony protuberances, and  
81 short, rounded snouts (Bonaparte, 1991; Wilson et al., 2003; Carrano and Sampson,  
82 2008; Canale et al., 2009; Pol and Rauhut, 2012; Cerroni et al., 2022; Amudeo-Plaza et  
83 al., 2023). They are recovered as a sister clade to the smaller-bodied Noasauridae within  
84 Abelisauroida (e.g., Novas, 1991; Bonaparte, 1991, 1996; Novas et al., 2013, Baiano et  
85 al., 2021). The oldest abelisaurid remains are from the Jurassic of South America and  
86 include an isolated abelisaurid tooth from the Late Jurassic of Uruguay (Soto et al.,  
87 2022) and a relatively complete skeleton of the putative early-branching abelisaurid  
88 *Eoabelisaurus mafi* from the Lower Jurassic of Patagonia (Pol and Rauhut, 2012).  
89 Despite being present throughout the Cretaceous of Europe and Gondwana, abelisaurids  
90 only became the dominant apex predators after carcharodontosaurids and spinosaurids  
91 declined and/or went extinct in the mid Cretaceous (Hendrickx et al., 2015a and  
92 references therein). However, they may have been more abundant than other inland  
93 predators (Sales et al., 2016). After the Cenomanian–Turonian transition, abelisaurids  
94 started to dominate the western European and Gondwanan landmasses (Candeiro and  
95 Martinelli, 2005; Carrano et al., 2012; Novas et al., 2013; Tortosa et al., 2014; Csiki-  
96 Sava et al., 2015; Hendrickx et al., 2015a).  
97 The European abelisaurid record has significantly increased over the last decades (e.g.,  
98 Allain and Pereda-Suberbiola, 2003; Buffetaut, 2005; Ósi et al., 2010; Ósi and  
99 Buffetaut, 2011; Tortosa et al., 2014; Csiki-Sava et al., 2015; Pérez-García et al., 2016;  
100 Isasmendi et al., 2022, 2024; Malafaia et al., 2025). The oldest definitive European  
101 abelisauroid is currently *Genusaurus sisteronis* from the Albien of Provence, southern  
102 France. Nevertheless, its phylogenetic position remains unresolved, having been  
103 recovered as a noasaurid (e.g., Carrano and Sampson, 2008), an early-branching

104 abelisaurid (e.g., Tortosa et al., 2014; Baiano et al., 2023), or a later branching  
105 abelisaurid more related to Furileosauria (Buffetaut et al., 2024; Buffetaut, 2025).  
106 *Caletodraco cottardi* is the second abelisaurid from the mid Cretaceous of France. It is  
107 based on a partial skeleton likely recovered from lower Cenomanian deposits in  
108 northwestern France and identified as a furileusaurian brachyrostran (Buffetaut et al.,  
109 2024). *Tarascosaurus salluvicus* was the first abelisaurid reported of lower Campanian  
110 deposits in southern France (Le Loeuff and Buffetaut, 1991) and despite its fragmentary  
111 nature and some authors considering it as a *nomen dubium* (Rauhut, 2003, Allain and  
112 Pereda-Suberbiola, 2003), this taxon was recovered amongst Abelisauridae in the  
113 phylogenetic analyses performed by Tortosa et al. (2014). Other material from the  
114 Iberian Peninsula, such as the Laño femora, have been compared with *Tarascosaurus*  
115 (Le Loeuff and Buffetaut, 1991), even though there is no conclusive evidence to support  
116 this attribution (Isasmendi et al., 2022; Malafaia et al., 2025). The best-known European  
117 abelisaurid is *Arcovenator escotae* from the upper Campanian of southern France,  
118 which has been recovered within Majungasaurinae (Tortosa et al., 2014). The holotype  
119 of *Arcovenator* is composed of a partial skeleton including cranial, axial and  
120 appendicular material. Additional isolated teeth and caudal vertebrae from other  
121 horizons and areas of the type locality have also been referred to this taxon.  
122 Furthermore, other skeletal elements from the Upper Cretaceous of France and the  
123 Iberian Peninsula have been assigned to this taxon (Tortosa et al., 2014; Pérez-García et  
124 al., 2016; Isasmendi et al., 2022; Malafaia et al., 2025). First regarded as Ceratosauria  
125 indet. by Carrano and Sampson (2008), *Betasuchus bredai*, from Maastrichtian strata of  
126 the Netherlands, was finally recovered as an abelisauroid more closely related to  
127 *Tarascosaurus* by Tortosa et al. (2014) and currently represents the latest record of the  
128 clade in Europe (Csiki-Sava et al., 2015). Other than abelisaurids, theropod remains

129 from the uppermost Cretaceous Ibero-Armorican mainly belong to small-sized  
130 coelurosaurians (e.g., Antunes and Sigogneau-Russell, 1991, 1992; Allain and Taquet,  
131 2000; Garcia et al., 2000; Laurent et al., 2002; Laurent, 2003; Ortega et al., 2015;  
132 Torices et al., 2015; Marmi et al., 2016; Puertólas-Pascual et al., 2018; Isasmendi et al.,  
133 2022, 2024; Santos Brilhante et al., 2022; Malafaia et al., 2023) whereas the majority of  
134 fossils from large bodied theropods have been either identified as indeterminate  
135 theropods (Antunes and Sigogneau-Russell, 1992; Laurent, 2003; Weishampel et al.,  
136 2004; Company, 2005; Ortega et al., 2015; Torices et al., 2015; Pérez-García et al.,  
137 2016), tetanurans (Carrano et al., 2012), or ornithomimosaur (Pereda-Suberbiola et al.,  
138 2000).

139 This study aims to review the large-bodied theropod material from several uppermost  
140 Cretaceous Iberian sites (Chera 2, Laño, Montebrei and Viso), which comprises isolated  
141 teeth and some axial elements, including a previously undescribed caudal vertebra.  
142 Aside from the systematic studies, morphometric and cladistic analyses were performed  
143 on the revised dental elements and the presence of Abelisauridae in Ibero-Armorica is  
144 evaluated.

145

146 **Institutional abbreviations.** **DPM**, Departamento de Paleontología de Madrid,  
147 Universidad Complutense de Madrid, Madrid, Spain; **IIPG**, Instituto de Investigación  
148 en Paleobiología y Geología, General Roca, Argentina; **MA**, Musée d'Angoulême,  
149 Angoulême, France; **MACN**, Museo Argentino de Ciencias Naturales 'Bernardino  
150 Rivadavia', Buenos Aires, Argentina; **MAU**, Museo Municipal "Argentino Urquiza",  
151 Neuquén, Argentina; **MCNA**, Museo de Ciencias Naturales de Álava/Arabako Natura  
152 Zientzien Museoa, Vitoria/Gasteiz, Spain; **MG**, Museu Geológico, Lisbon, Portugal;  
153 **MPCA**, Museo Provincial Carlos Ameghino, Cipolletti, Argentina; **MPCN**, Museo



154 Patagónico de Ciencias Naturales, General Roca, Argentina; **MPEF**, Museo  
155 Paleontológico Egidio Feruglio, Chubut, Argentina; **MPZ**, Museo de Ciencias  
156 Naturales de la Universidad de Zaragoza, Zaragoza, Spain; **UCPC**, Paleontology  
157 Collection of the University of Chicago, Chicago, USA; **UPUAM**, Unidad de  
158 Paleontología, Universidad Autónoma de Madrid, Madrid, Spain.  
159 **Anatomical abbreviations.** **cdl**, centrodiapophyseal lamina; **cv**, cervix; **dc**, distal  
160 carina; **dl**, dentine layer; **dt**, denticle; **fch**, facet for chevron; **iet**, irregular enamel-  
161 texture; **mc**, mesial carina; **ns**, neural spine; **pc**, pulp cavity; **poz**, postzygapophysis; **tp**,  
162 transverse process; **tu**, transverse undulation; **vg**, ventral groove.

163

## 164 **GEOGRAPHICAL AND GEOLOGICAL SETTINGS**

165 The skeletal elements here studied were recovered from four Iberian fossil sites namely,  
166 Chera 2, Laño, and Montebrei from Spain, and Viso from Portugal. The Chera 2 site is  
167 situated near the locality of Chera in the province of Valencia (Valencian Community),  
168 about 60 km west of the city of Valencia, in the eastern part of the Iberian Peninsula.  
169 The Laño site is located about 30 km south of the city of Vitoria/Gasteiz, in an  
170 abandoned sand quarry (the Laño quarry) between the towns of Laño and Albaina  
171 (County of Treviño), in the northern Iberian Peninsula. The Montrebei locality is  
172 located around 25 km southwest of the Tremp locality, east of the Congost de  
173 Montrebei, in the Alsamora municipality (Lleida, northeastern Iberian Peninsula).  
174 Finally, the locality of Viso is situated in the Montemor-o-Velho region (Coimbra), at  
175 km 20 of the railway (Sauvage 1897–1898), in the western part of the Iberian Peninsula  
176 (Fig. 1.1).  
177 Geologically, the Chera 2 site is located in the Chera Basin (southwestern Iberian  
178 Range), specifically in the Sierra Perenchiza Formation (Fm) (e.g., Company, 2005;

179 Company and Szentesi, 2012) (Fig. 1.1). Even though this lithostratigraphic unit reaches  
180 a maximum thickness of 300 m, it is only 50 m thick in the Chera area. It comprises  
181 lagoonal micritic and brecciated limestones at the base, which transition upwards into  
182 palustrine carbonates with evidence of pedogenic alterations towards the top (Platt,  
183 1989; Alonso et al., 1991; Company, 2005). This formation has been interpreted as a  
184 continental, shallow carbonate environment (Vilas et al., 1982) with shallow, low-  
185 salinity water bodies at the base (Platt and Wright, 1992) and restricted, ephemeral  
186 coastal carbonate lakes and ponds towards the top (Company, 2005). In this context, the  
187 Chera 2 site is thought to have formed in an isolated pond that developed in floodplains,  
188 where the fossils were transported after high-energy flows (Company, 2005). Peyrot et  
189 al. (2020) suggested a late Campanian–early Maastrichtian age for the Chera site based  
190 on palynological studies; however, Company et al. (2005) proposed a late Campanian  
191 age for it (Fig. 1.2).

192 The Laño site is located in the Northern Castilian Ramp of the Basque-Cantabrian  
193 Basin, specifically on the southern margin of the Miranda-Treviño Syncline (Pereda-  
194 Suberbiola et al., 2015; Corral et al., 2016) (Fig. 1.1). In the Laño quarry, the Sedano  
195 Fm crops out (Berreteaga, 2008; Corral et al., 2016). This formation mainly comprises  
196 siliciclastic deposits that can be divided into two units: (1) a lower unit consisting of  
197 marine marls and clays and (2) an upper unit composed of silty quartzarenites with  
198 interbedded dolomitized limestone layers (Floquet et al., 1982). The Sedano Fm  
199 represents a littoral environment, with a siliciclastic sequence in its upper part formed  
200 by deltaic aggradation in the subtidal and intertidal areas, prior to the progradation of  
201 the delta plain (Corral et al., 2016; Martín-Chivelet et al., 2019). At the Laño quarry, the  
202 Sedano Fm is 22 m thick and comprises two different units: (1) the basal unit, which  
203 begins with lag deposits of siliciclastic gravel overlain by different sandstone packs

204 with conglomeratic and erosive bases, and (2) the upper unit, which is mainly composed  
 205 of almost unconsolidated clayey sandstones. The fossils studied herein come from the  
 206 L1A and L2 vertebrate-bearing beds. Astibia et al. (1990) and Pereda-Suberbiola et al.  
 207 (2000) interpreted the Laño area as a braided river system, where channels, sandbars  
 208 and pools developed. Corral et al. (2016) dated the continental vertebrate site of Laño to  
 209 the late Campanian (chron C32n, 72–73.5 Ma) based on the combination of  
 210 lithostratigraphic and magnetostratigraphic analyses (Fig. 1.2).

211 The fossil locality of Montebrei is located in the South Pyrenean Basin, precisely in the  
 212 eastern Tresp Syncline (Fondevilla et al., 2016, 2019) (Fig. 1.1). The latter is the  
 213 largest of the various sub-basins formed by the compartmentalization of the synform  
 214 syntectonic sub-basins (Fondevilla et al., 2016). The uppermost Cretaceous and  
 215 Paleocene lithostratigraphic units crop out in the Tresp Syncline, encompassing  
 216 marine, transitional, and continental deposits. These include the middle Campanian–  
 217 Maastrichtian Arén Fm and the Maastrichtian–Paleocene Tresp Fm (Mey et al., 1968;  
 218 Nagtegaal et al., 1983; Mutti and Sgavetti, 1987). The latter overlays and laterally  
 219 transitions into the Arén Fm (Ardevol et al., 2000; Fondevilla et al., 2016). The Tresp  
 220 Fm has been informally divided into four units: the ‘Grey Garumnian’, the ‘Lower Red  
 221 Garumnian’, the ‘Vallcebre limestones and equivalents’, and the ‘Upper Red  
 222 Garumnian’ (Rosell et al., 2001; Perez-Pueyo et al., 2021). The Montebrei fossil locality  
 223 is situated within the ‘Grey Garumnian’, specifically in La Posa Formation (Torices  
 224 Hernández, 2002; Torices et al., 2015; Fondevilla et al., 2019). This unit consists of  
 225 grey marls and mudstones intercalated with sandstones, limestones, and occasional coal  
 226 beds, containing mixed marine and freshwater invertebrate faunas. It has been  
 227 interpreted as representing transitional environments such as lagoons, tidal mudflats,  
 228 and marshes (Eichenseer, 1988; Rosell et al., 2001; Díez-Canseco et al., 2014; Oms et

al., 2016). The ‘Grey Garumnian’ is dated to the latest Campanian–Maastrichtian (Díez-Canseco et al., 2014; Vicente et al., 2015, 2016; Fondevilla et al., 2016; Puertolas-Pascual et al., 2018) and the Montrebei fossil site dates more precisely to the ‘mid’ early Maastrichtian (C31r) (Fondevilla et al., 2019) (Fig. 1.2).

The vertebrate-bearing site of Viso is found in the Lusitanian Basin, either in the “Sandstones and Mudstones of Viso” of Barbosa et al. (2008) or Viso Fm of Malafaia et al. (2024) (Fig. 1.1). In the latter, a sandstone sequence is predominant, but mudstones, which have been dated to the Campanian–Maastrichtian, based on palynological studies, also crop out (Barbosa et al., 2008) (Fig. 1.2).

## **MATERIAL AND METHODS**

### **Geographic and stratigraphic context of the material**

The studied material consists of three isolated teeth from the Chera 2 site (CH 168), Montrebei (DPM-MON-T10 1) and Viso (MG 73), and two vertebrae from the Laño site (MCNA 8366 and 17433). MCNA 8366 comes from the L1A vertebrate-bearing bed, and MCNA 17433 was recovered from the L2 fossiliferous level. The specimens are in three separate institutions namely, the Universidad Complutense de Madrid of Spain (DPM-MON-T10 1), Museu Geológico of Lisbon, Portugal (MG 73), and the Arabako Natura Zientzien Museoa/Museo de Ciencias Naturales de Álava of Vitoria-Gasteiz, Spain (MCNA 8366 and 17433). The tooth CH 168 was studied based on photographs and a cast.

Historically, DPM-MON-T10 1 was considered as the tooth of an indeterminate theropod by Torices et al. (2015), before being assigned to an indeterminate Abelisauridae by Isasmendi et al. (2024). It should be noted that this specimen never received a proper description. The tooth MG 73 was first attributed to *Megalosaurus* sp.

254 by Sauvage (1897–1898) at the end of the 19<sup>th</sup> century, then to *Megalosaurus* cf.  
255 *pannoniensis* by Lapparent and Zbyszewski (1957), and recently to Theropoda indet. by  
256 Malafaia et al. (2024). The neural arch MCNA 8366 was originally assigned to  
257 *Rhabdodon* by Pereda-Suberbiola and Sanz (1999), while the vertebra MCNA 17433  
258 (previously labeled as MCNA 14077) was tentatively identified as Abelisauridae indet.  
259 by Isasmendi et al. (2021). Finally, the tooth CH 168 was referred to as Theropoda  
260 indet. by Company (2005), as ?Neoceratosauria indet. by Company et al. (2009) and as  
261 cf. *Arcovenator* by Isasmendi et al. (2022).

262 **Comparative methodology and terminology for the isolated teeth.** Unlike the  
263 postcranial remains, which was studied first-hand, the isolated teeth were studied under  
264 a stereo microscope (Nikon® SMZ645) and through first-hand observations. Crown-  
265 based measurements were taken using a digital caliper (Juning®), while denticle-based  
266 measurements were taken either under the stereo microscope or with the Imagej  
267 software (v.1.51j8). The study of the isolated teeth follows the anatomical, positional,  
268 directional and morphometric nomenclature proposed by Smith and Dodson (2003),  
269 Smith et al. (2005), and Hendrickx et al. (2015b).

270 **Measurements taken on the isolated teeth.** The following measurements were taken  
271 on the studied sample based on Currie et al. (1990), Smith et al. (2005), and Hendrickx  
272 et al.'s (2015b, 2020) methodology (see Supplementary material 1): crown base length  
273 (**CBL**), crown base width (**CBW**), mid-crown length (**MCL**), mid-crown width  
274 (**MCW**), crown height (**CH**), apical length (**AL**), extent of the mesiobasal denticles  
275 (**MDE**), length of the mesial serrated carina (**MSL**), distoapical denticle density (**DA**),  
276 distobasal denticle density (**DB**), distocentral denticle density (**DC**), mesioapical  
277 denticle density (**MA**), mesiobasal denticle density (**MB**), mesiocentral denticle density  
278 (**MC**), mesial denticle length (**MDL**), and distal denticle length (**DDL**). In addition, the

279 crown base ratio (**CBR**; as CBW/CBL), crown height ratio (**CHR**; as CH/CBL), mid-  
 280 crown ratio (**MCR**; as MCW/MCL), mesial angle of the crown (**CMA**), distal angle of  
 281 crown (**CDA**), average mesial denticle density (**MAVG**; as (MA+MC+MB)/ 3),  
 282 average distal denticle density (**DAVG**; as (DA+DC+DB)/ 3), and denticle size density  
 283 index (**DSDI**; as MC/DC) were calculated. The MCA and DCA were calculated using  
 284 the following equation (Smith, 2007; Serrano-Martínez et al., 2015):

$$285 \quad CMA = \arccos \frac{CBL^2 + AL^2 - CH^2}{2 \times CBL \times AL} \text{ and } CDA = \arccos \frac{CBL^2 + CH^2 - AL^2}{2 \times CBL \times AL}$$

286 **Linear discriminant analyses (LDA) on the isolated teeth.** The classification of the  
 287 isolated teeth was supported by several linear discriminant analyses performed using  
 288 PAST v.4.05 (Hammer et al., 2001). When performing the LDAs, variables expressed  
 289 as ratios (CBR, CHR and MCR) as well as the crown angles (CDA and CMA) were  
 290 excluded, as these are not independent but weighted variables (Hendrickx et al., 2015b).  
 291 Furthermore, to ensure that all variables were metric-based, measurements of denticle  
 292 length (MLD and DDL) were used instead of denticle densities, in accordance with the  
 293 approach of Hendrickx et al. (2020). Following Malafaia et al. (2025), the number of  
 294 flutes on the labial and lingual surfaces (**LAF** and **LIF**, respectively) were not included  
 295 in the analyses, as there are non-metric variables. Therefore, a total of 10 variables were  
 296 used in the performed LDAs, as Malafaia et al. (2025). These variables are: CBL, CBW,  
 297 CH, AL, MCL, MCW, MDE, MSL, MDL, and DDL. To perform the LDAs, the  
 298 database and a similar methodology proposed by Malafaia et al. (2025) were used (see  
 299 Supplementary material 1). Prior to conducting the morphometric analyses, the data  
 300 were normalized using a Log (x+1) transformation for all variables to avoid zero values.  
 301 In addition, as proposed by Young et al. (2019) and Hendrickx et al. (2020), an arbitrary  
 302 value of 100 denticles per five millimeters was used for those specimens with non-  
 303 denticulated carinae. Finally, specimens studied herein were not assigned to any

304 predefined group for the LDAs and were instead treated as belonging to an “unknown  
 305 taxon”.

306 Firstly, two linear discriminant analysis, one at the taxon level and another at the clade  
 307 level, were carried out using a database comprising 528 teeth with the data from  
 308 Hendrickx et al. (2020) (Database 1; Supplementary material 1) belonging to twenty-  
 309 four Cretaceous taxa (twenty-two genera, and two indeterminate family-level clades),  
 310 with the aim of minimizing potential noise (see Malafaia et al., 2025; Supplementary  
 311 material 1). This dataset gathers dental measurements from one non-abelisauroid  
 312 ceratosaurian (*Genyodectes*), nine abelisaurids (*Abelisaurus*, *Arcovenator*, *Aucasaurus*,  
 313 *Carnotaurus*, *Chenanisaurus*, ‘*Indosuchus*’, *Majungasaurus*, *Skorpiovenator*, and  
 314 *Abelisauridae* indet.), three spinosaurids (*Baryonyx*, *Suchomimus* and *Spinosaurinae*  
 315 indet.), one neovenatorid (*Neovenator*), two megaraptorans (*Australovenator* and  
 316 *Fukuiraptor*), five carcharodontosaurids (*Acrocanthosaurus*, *Carcharodontosaurus*,  
 317 *Eocarcharia*, *Giganotosaurus*, and *Mapusaurus*), and six tyrannosaurids  
 318 (*Albertosaurus*, *Alioramus*, *Daspletosaurus*, *Gorgosaurus*, *Tyrannosaurus* and  
 319 *Zhuchengtyrannus*).

320 Afterwards, two additional LDAs were performed, one at taxon level and another at  
 321 clade level, excluding the groups that plotted well-separated from the herein studied  
 322 teeth (Database 2; Supplementary material 1), as these were considered not closely  
 323 related. The excluded specimens belonged to all spinosaurids, as well as the  
 324 tyrannosaurids *Daspletosaurus*, *Tyrannosaurus* and *Zhuchengtyrannus*, and the  
 325 carcharodontosaurid *Carcharodontosaurus*.

326 Finally, a fifth LDA was performed with the third database (Database 3; Supplementary  
 327 material 1) compiled by Malafaia et al. (2025), which includes Ibero-Armorican  
 328 abelisaurids or specimens classified among *Abelisauridae*. These consist of the

329 indeterminate abelisaurids from the Western Tremp Syncline (South Pyrenean Basin)  
330 (Isasmendi et al., 2024), the abelisaurid teeth closely related to *Arcovenator* from Poyos  
331 (Malafaia et al., 2025), the specimens referred to cf. *Arcovenator* from Armuña (Pérez-  
332 García et al., 2016), and the *Arcovenator* specimens from Jas Neuf Sud and Laño  
333 (Tortosa et al., 2014; Hendrickx et al., 2020; Isasmendi et al., 2022). In this analysis, the  
334 isolated teeth of an indeterminate tetanuran from Iharkút (Ősi et al., 2010) were  
335 excluded from the dataset.

336 **Cladistic analysis.** A cladistic analysis was performed based on the dentition-based  
337 matrix (146 morphological dental characters) of Hendrickx et al. (2020), into which we  
338 scored the isolated teeth MG 73, CH 168, and DPM-MON-T10 1 (see Supplementary  
339 material 2). The positive constraints defined by Hendrickx et al. (2020) were applied in  
340 these analyses to recover a topology consistent with the most recent phylogenetic  
341 hypotheses. Additionally, to test the phylogenetic affinities of the studied specimens  
342 within the Abelisauridae clade, we constrained their placement and evaluated the  
343 additional steps required to achieve that position. The phylogenetic analysis was carried  
344 out using TNT 1.6 (Goloboff and Morales, 2023). The studied specimens were scored  
345 using Mesquite 3.7 (Maddison and Maddison, 2011) and subsequently imported into  
346 TNT. The search strategy follows the protocol used by Hendrickx et al. (2020),  
347 beginning with a combination of the tree-search algorithms, including Wagner trees,  
348 TBR branch swapping, sectorial searches, Ratchet (with the perturbation phase stopped  
349 after 20 substitutions), and Tree Fusing (5 rounds), until 100 hits of the same minimum  
350 tree length were achieved. The best trees recovered were subjected to a final round of  
351 TBR branch swapping. Zero-length branches in any of the recovered most parsimonious  
352 trees were collapsed. Consistency (CI) and retention (RI) indexes were obtained using  
353 the STATS.RUN command.



354

355

## SYSTEMATIC PALEONTOLOGY

356

Suborder THEROPODA Marsh, 1881

357

Infraorder CERATOSAURIA Marsh, 1884

358

Family ABELISAURIDAE Bonaparte and Novas, 1985

359

### **Abelisauridae indet.**

360

Figure 2

361 **Morphotype 1**

362 **Referred material.** DPM-MON-T10 1, one isolated mesial tooth (Fig. 2.1–2.7).

363 **Geographic occurrence.** Montrebei (Lleida, Spain).

364 **Stratigraphic occurrence.** ‘Grey Garumnian’ strata of the Tremp Formation; La Posa

365 Formation; ‘mid’ early Maastrichtian (C31r) (Torices Hernández, 2002; Torices et al.,

366 2015; Fondevilla et al., 2019).

367 **Description.** DPM-MON-T10 1 consists of a crown that lacks its basal portion and the

368 apex (Fig. 2.2–2.5). It is here interpreted as a mesial tooth, based on its symmetrical

369 ‘D’-shaped cross-section, relatively thick crown, and asymmetrical labial and lingual

370 surfaces (Hendrickx et al., 2015b, 2020). No denticles are complete, but the bases of the

371 mesial denticles and some distal denticle bases are preserved (Fig. 2.1). The enamel is

372 also largely missing, but a patch is preserved on the mesioapical part of the labial

373 surface (Fig. 2.6). The crown is ziphodont, not very compressed labiolingually and quite

374 distally curved, with its apicalmost part extending beyond the basodistalmost point of

375 the crown (Fig. 2.2–2.5). The mesial margin of the crown is strongly convex, and the

376 distal margin is concave in lateral view (Fig. 2.5). In distal view, the labial surface is

377 convex, and the lingual one is mostly straight, making the crown slightly lingually tilted  
378 (Fig. 2.4). The lingual and labial surface are convex, but the lingual surface becomes  
379 flatter close to the distal carina. The basalmost cross-section of the preserved crown is  
380 nearly subsymmetrical ‘D’-shaped, whereas the mid cross-section is lanceolate (Fig.  
381 2.7).

382 Both mesial and distal carinae are present and denticulated, but because the basal part of  
383 the crown is not preserved, their entire extension cannot be assessed. The mesial carina  
384 is mesiolingually oriented and straight; it is centered apically but slightly displaces  
385 lingually toward the base (Fig. 2.2). On the other hand, the distal carina is straight and  
386 strongly labially deflected (Fig. 2.4). The mesial denticle density is 10 denticles per five  
387 mm apically and at mid crown and there is no random variation of denticle-size.

388 The preserved enamel surface is slightly polished and exhibits micro-scratches. The  
389 enamel-texture is subtle, non-oriented, and irregular (Fig. 2.6).

## 390 **Morphotype 2**

391 **Referred material.** MG 73, one isolated lateral tooth (Fig. 2.8–2.12).

392 **Geographic occurrence.** Viso locality, region of Montemor-o-Velho, Coimbra, Portugal  
393 (Sauvage, 1897-1998; Malafaia et al., 2024).

394 **Stratigraphic occurrence.** Viso Formation (also known as Sandstones and Mudstones  
395 of Viso); Campanian–Maastrichtian (Barbosa et al., 2008; Malafaia et al., 2024).

396 **Description.** MG 73 consists of a lateral tooth crown that lacks the apex and a fragment  
397 of the root (Fig. 2.8–2.12). It is interpreted as a lateral tooth based on its symmetrical  
398 cross-section outline and low CBR value (see Hendrickx et al., 2015b, 2019, 2020). The  
399 enamel layer is eroded on its lingual surface (Fig. 2.11), and the mesial carina is almost  
400 entirely abraded (Fig. 2.8). The distal carina is also poorly preserved but the bases of the

401 denticles are visible along the carina and the basal denticles are relatively well  
402 preserved (Fig. 2.10–2.11).

403 This crown is relatively elongate, ziphodont, and strongly labiolingually flattened (Fig.  
404 2.8–2.12), with a CBR of 0.58 (CBL=12 mm and CBW=7 mm). The crown is distally  
405 curved, but it is not possible to determine whether the apex would have extended  
406 beyond its basodistalmost point (Fig. 2.11). In lateral view, the mesial margin is convex,  
407 while the distal margin is slightly concave, becoming straighter toward the base (Fig.  
408 2.11). In distal view, the lingual surface is slightly concave, and the labial surface is  
409 convex, suggesting that the apex was slightly tilted lingually (Fig. 2.10). There is no  
410 concave surface adjacent to the carinae, but the lingual surface becomes flat near the  
411 distal carina (Fig. 2.9). The basal cross-section of the crown is lanceolate (Malafaia et  
412 al., 2024) and the cross-section of the root is oval (Fig. 2.12). No basal constriction is  
413 present between crown and root.

414 The distal carina reaches the cervix and the distal denticles are present along the entire  
415 preserved length (Fig. 2.10). Denticles also appear to be present on the mesial carina,  
416 but only their bases are visible at mid-crown (Fig. 2.8). The full extent of the mesial  
417 carina cannot be determined (Malafaia et al., 2024). The distal carina is centrally located  
418 and slightly bowed whereas the mesial carina is more lingually positioned and straight  
419 (Fig. 2.8 and 2.10). There is no random variation in denticle size along the distal carina  
420 and the denticles decrease in size towards the base. Only the distal denticle densities  
421 could be measured. The DC value is 20 denticles per five millimeters, and the DB is  
422 22.5 denticles per five millimeters. The basal denticles are sub-quadrangular with  
423 slightly convex external margins and are separated by narrow interdenticular spaces.  
424 The denticles are poorly preserved at mid-crown, with only the bases of some visible.  
425 These are also separated by narrow interdenticular spaces and seem slightly longer

apicobasally than mesiodistally. However, because the external end is not preserved, this morphology cannot be confirmed. Interdenticular sulci are not visible adjacent to either the distal or mesial carinae. Some subtle transverse undulations are present on the enamel of the lingual surface of the crown (Fig. 2.9). The enamel texture is subtle, non-oriented, and irregular (Malafaia et al., 2024).

Genus ***Arcovenator*** Tortosa, Buffetaut, Vialle, Dutour, Turini and Cheylan, 2014

**Cf. *Arcovenator* sp.**

Figure 3–4

**Referred material.** MCNA 8366, an anterior dorsal vertebra; MCNA 17433, an anterior to middle caudal vertebra.

**Geographic occurrence.** Laño site, Treviño County, Spain.

**Stratigraphic occurrence.** Sedano Formation; upper Campanian (C32n) (Corral et al., 2016).

**Description.**

**Axial skeleton.**

**Dorsal vertebra (Fig. 3.1–3.4).** MCNA 8366 comprises the neural arch of an anterior dorsal vertebra (probably a D1 or D2, based on O’Connor, 2007). This neural arch is covered by an iron patina and lacks both prezygapophyses (Fig. 3.1–3.4). The neural spine and the left transverse process are badly damaged, with only their bases preserved. The right transverse process is laterally projected and horizontal but slightly inclined anteriorly, forming an angle of less than 20° relative to the horizontal plane (Fig. 3.1–3.4). It measures 46 mm in length and is developed in the anterior half of the vertebra. In dorsal view, the right transverse process expands anteroposteriorly towards the diapophysis, making it fan-shaped in this view (Fig. 3.4). The posterior edge of the

transverse process is straight, and its anterior margin is convex. In lateral view, the diapophysis is D-shaped and faces ventrolaterally (Fig. 3.2). The base of the neural spine is mediolaterally narrow anteriorly and broadens posteriorly. It extends posteriorly beyond the bases of the prezygapophyses, almost reaching the posterior margin of the postzygapophyses (Fig. 3.4). The postzygapophyses face ventrolaterally, are subcircular in outline, and appear to overhang the centrum (Fig. 3.1 and 3.2). The presence or absence of a hyposphyene-hypantrum articulation cannot be determined (Fig. 3.1).

**Caudal vertebrae (Fig. 3.5–3.8).** MCNA 17433 consists of the centrum and the base of the neural arch of an anterior to middle caudal vertebra (posterior to C5 due to its straighter anterior and posterior margins, based on Méndez, 2014), which are slightly mediolaterally compressed due to deformation (Fig. 3.5–3.8). The centrum and neural arch are almost fully fused, with the suture line being almost indiscernible (Fig. 3.7). The centrum is amphicoelous, spool-shaped, and slightly elongated (Fig. 3.5–3.8). MCNA 17433 measures 79 mm in anteroposterior length. Its anterior articular surface is 57 mm in height and 46 mm in width, while the posterior articular surface is 64 mm in height and 46 mm in width. Both the anterior and posterior articular surfaces are elliptical (Fig. 3.6 and 3.8). In lateral view, the anterior and posterior surfaces are straight, and the ventral margin is concave (Fig. 3.7). The centrum is medially compressed and exhibits relatively shallow pleurocentral depressions. However, the base of the neural arch is wider than the mid-centrum. The ventral surface of the centrum features an anteroposterior longitudinal groove, which is laterally delimited by a ridge on each side (Fig. 3.5). In the same view, the posterior end of the centrum exhibits quite prominent facets for articulation with the haemal arches, making the posterior end of the centrum more ventrally projected (Fig. 3.5).

474 The preserved portion of the neural arch includes the bases of the transverse processes  
475 (Fig. 3.5 and 3.7), which are centrally located and ventrally positioned in the neural  
476 arch. These processes are relatively thin and sheet-like and the preserved parts taper  
477 laterally. The ventral surface of the left transverse process exhibits a possible broad but  
478 not very protruding centrodiapophyseal lamina (Fig. 3.5 and 3.7).

479 ***Arcovenator* sp.**

480 Figure 4

481 **Referred material.** CH 168, one isolated lateral tooth.

482 **Geographic occurrence.** Chera, Valencia, Spain.

483 **Stratigraphic occurrence.** Sierra Perenchiza Formation; upper Campanian (Company,  
484 2005; Company et al., 2005).

485 **Description.** CH 168 preserves the crown and part of the root. It is interpreted as a  
486 lateral tooth based on its symmetrical cross-section outline and low CBR value (see  
487 Hendrickx et al., 2015b, 2019, 2020). The base of the crown is distally damaged, and  
488 fractures are present throughout the crown (Fig. 4). Parts of the carinae are also missing.  
489 An apically located elliptical wear facet is present on the labial surface (Fig. 4.5). The  
490 crown is ziphodont, labiolingually compressed, and distally curved. The apex extends  
491 slightly beyond the basodistalmost point of the crown (Fig. 4). In lateral view, the  
492 mesial margin is convex, while the distal margin is concave apically, becoming  
493 straighter toward the base (Fig. 4.3 and 4.5). In distal view, the lingual surface is  
494 straight basally and convex apically whereas the labial margin is convex basally and  
495 becomes straighter apically (Fig. 4.4). Both labial and lingual surfaces are convex, with  
496 no concave surfaces adjacent to any carinae. The lingual margin becomes more planar  
497 near the base of the distal carina. The basal cross-section of the crown is oval to

498 lanceolate (Fig. 4.6), and the mid cross-section is lenticular. The cross-section of the  
499 root is more oval, with no basal constriction between the root and the crown.  
500 The CBL measures 15.47 mm, whereas the CBW measures 9.88 mm, resulting in a  
501 CBR of 0.64. The CH is 30.02 mm, being the CHR of 1.94. The AL measures 33.52  
502 mm.

503 Both mesial and distal carinae are denticulated (Fig. 4.1 and 4.8). The mesial carina is  
504 straight, centered apically and mesiolingually displaced toward the base of the crown.  
505 The mesial carina does not reach the cervix but extends from the apex to approximately  
506 two-thirds of the crown (Fig. 4.2). The distal carina reaches the cervix, and it is  
507 sigmoidal in distal view and labially deflected (Fig. 4.4).

508 The denticles are largest at mid-crown on the mesial carina and gradually decrease in  
509 size basally (Fig. 4.2, 4.3 and 4.5). The wear facet prevents from determining whether  
510 there was a denticle-size discrepancy apically. Nevertheless, the distal denticles are  
511 larger apically (Fig. 4.3–4.5). There is no random variation in denticle size along the  
512 carinae. The denticles are almost equal in size or slightly larger on the mesial carina,  
513 with a DSDI value of 1. The mesial denticle density is 15 mm per 5 mm, whereas the  
514 distal denticle density is 12 denticles per 5 mm apically, 15 denticles per 5 mm at mid-  
515 crown, and 17 denticles per 5 mm at the base. The outline of the denticles is  
516 symmetrically to slightly asymmetrically convex and perpendicular to the carinae (Fig.  
517 4.1 and 4.8). The mesial denticles are mainly subquadrangular but become slightly  
518 apicobasally subrectangular apically (Fig. 4.1). The distal denticles, on the other hand,  
519 are mesiodistally subrectangular at mid-crown and basally, but almost subrectangular  
520 apically (Fig. 4.8). Subtle interdenticular sulci are present on both carinae. These are  
521 short, straight, and basally inclined.

522 A few transverse undulations are present on the crown, but they are particularly subtle.  
523 Subtle marginal undulations are similarly present on the labial surface of the crown,  
524 close to the distal carina. The enamel texture is non-oriented and irregular (Fig. 4.7).

525

## 526 **RESULTS**

### 527 **Morphometric analyses**

528 The discriminant analyses exhibit similar variance in their axes (Fig. 5 and 6). Axis 1  
529 accounts for 42.45–55.62 % of the total variance, while Axis 2 explains 20.54–30.06 %.

530 The reclassification rates obtained in the LDAs are relatively high, ranging from 63.42  
531 to 75.85 %, and are significantly higher using the clade-level databases (see

532 Supplementary material 1).

533 The classification of the specimens studied herein within Theropoda varies depending  
534 on the database and the linear discriminant analysis (Table 1). In LDA 1 (clade level,  
535 database 1), the tooth from Viso (MG 73) is classified as Abelisauridae (Jackknifed),  
536 the tooth from Chera 2 (CH 168) as a non-abelisauroid ceratosaurian (Jackknifed), and  
537 the Montrebei tooth (DPM-MON-T10 1) as Carcharodontosauridae (classification and  
538 Jackknifed). In LDA 2 (taxon level, database 1), MG 73 is classified as *Carnotaurus*  
539 (classification and Jackknifed), CH 168 as *Gorgosaurus* (Jackknifed), and DPM-MON-  
540 T10 1 as *Acrocanthosaurus* (classification and Jackknifed). In LDA 3 (clade level,  
541 database 2), MG 73 is classified as a megaraptoran (Jackknifed), CH 168 as a  
542 tyrannosaurid (classification and Jackknifed), and DPM-MON-T10 1 as a megaraptoran  
543 (Jackknifed). In LDA 4 (taxon level, database 2), MG 73 is again classified as  
544 *Carnotaurus* (classification and Jackknifed), CH 168 as *Gorgosaurus* (Jackknifed), and  
545 DPM-MON-T10 1 as *Aucasaurus* (classification and Jackknifed). Finally, the  
546 discriminant analysis performed using the Ibero-Armorican abelisaurid dental record



(Database 3) classifies MG 73 as *Arcovenator* (Jas Neuf Sud) (classification and Jackknifed), CH 168 as the indeterminate abelisauid teeth from Poyos, which are closely related to *Arcovenator* (Jackknifed), and DPM-MON-T10 1 as Abelisauridae indet. 1 from the Western Tremp Syncline (Jackknifed) (Table 1).

### **Cladistic analyses**

The cladistic analysis performed on the data matrix including MG 73 and using a constrained tree topology recovered a single most parsimonious tree (CI = 0.200; RI = 0.457; L = 1304). MG 73 is grouped with noasaurids within a well-resolved Abelisauroida clade (Fig. 7.1). The specimen from Viso shares with other abelisauroids the spacing between mid-crown denticles on the distal carina, which is less than one-third of the denticle width (state 0; character 107) and the presence of a smooth or irregular (non-oriented) enamel-texture (state 0; character 121). The only synapomorphy found for Noasauridae that can be verified in MG 73 is the average number of mid-crown denticles per five millimeters on the distal carina (character 89), which ranges between 16 and 29 (state 1), whereas in abelisauids it is usually lower, between 9 and 15 (state 2). Within Noasauridae, the specimen from Viso is positioned as the sister taxon to *Masiakasaurus*, based on the presence of tenuous transverse undulations on the crown (state 1; character 113). Forcing the specimen from Viso into the Abelisauridae clade recovered three most parsimonious trees, with the consensus tree (CI = 0.199; RI = 0.451) being four steps longer (L = 1311) than that obtained when treating the specimens as floating taxon. Interestingly, constraining MG 73 within the group comprising *Arcovenator*, *Majungasaurus*, and *Indosuchus* yielded a single most parsimonious tree (CI = 0.200; RI = 0.453) and did not require additional steps (L = 1307) relative to the analysis with the specimens as floating taxon. However, no synapomorphies were recovered for this group in the analysis.

572 The analysis performed for specimens CH 168 recovered five most parsimonious trees  
573 (CI = 0.200; RI = 0.453; L = 1307) and the consensus tree (CI = 0.199; RI = 0.452; L =  
574 1309) places this specimen within Coelurosauria, allied with the tyrannosauroids  
575 *Gorgosaurus*, *Raptorex*, *Alioramus*, *Tyrannosaurus*, and *Daspletosaurus* (Fig. 7.2). The  
576 analysis found the following tyrannosauroid synapomorphies present in the Chera 2  
577 specimen: (i) crown height between 1 and 6 cm (state 1; character 69); (ii) the denticles  
578 on the mesial carina at two-thirds height of the crown have a subquadrangular shape  
579 (state 1; character 95); and (iii) mesial and distal denticles of similar size ( $0.8 < \text{DSDI} <$   
580  $1.2$ ) (state 0; character 105). However, all these features are also shared with most  
581 abelisaurids (Hendrickx et al. 2020). Forcing the specimen from Chera into the  
582 Abelisauridae clade recovered three most parsimonious trees (CI = 0.199; RI = 0.451; L  
583 = 1310), with the consensus tree (CI = 0.199; RI = 0.449) being five steps longer (L =  
584 1314) than that obtained when treating the specimens as floating taxon. This analysis  
585 recovered the following abelisaurid synapomorphies present in CH 168: (i)  
586 subquadrangular shape of denticles at two-thirds height of the crown on mesial carina in  
587 lateral view (state 1; character 95) and (ii) horizontal subrectangular shape of mid-  
588 crown denticle on distal carina in lateral view (state 1; character 96).

589 Finally, the analysis performed for DPM-MONT-T10 1 recovered seven most  
590 parsimonious trees (CI = 0.200; RI = 0.454; L = 1305; consensus tree CI = 0.180; RI =  
591 0.376; L = 1454). The strict consensus tree shows a poor resolution, with most taxa  
592 placed in a large polytomy (Fig. 8). Only a few clades such as Megalosauroidea,  
593 Abelisauridae, Allosauroidea and some coelurosaurian groups are better resolved. The  
594 most parsimonious trees from this analysis placed the specimen from Montrebei within  
595 the Paraves clade, sometimes allied with troodontids and other times with avialan taxa.  
596 Forcing the specimen into the Abelisauridae clade recovered a single most parsimonious

tree (CI = 0.199; RI = 0.452) and required four additional steps (L = 1309) relative to the most parsimonious trees obtained when treating the specimens as floating taxon. On the other hand, constraining its position within a group comprising *Arcovenator*, *Majungasaurus*, and *Indosuchus* also yielded a single most parsimonious tree (CI = 0.199; RI = 0.451) and required only one further step (L = 1310) compared to the previous analysis.

## DISCUSSION

### Comparisons and identification of the isolated teeth

Almost every morphometric analyses support the attribution of the MG 73 specimen to Abelisauridae, as this tooth consistently grouped within Abelisauridae, *Arcovenator*, or *Carnotaurus* in the LDAs (Table 1). The classification of CH 168 and DPM-MON-T10 1 is more ambiguous, as neither was grouped within Abelisauridae in any of the first three LDAs performed. Indeed, the CH 168 specimen was classified as a tyrannosaurid, a non-abelisauroid ceratosaurian, or *Gorgosaurus*, while DPM-MON-T10 was grouped within Carcharodontosauridae, Megaraptora, or *Acrocanthosaurus*. Nevertheless, DPM-MON-T10 was also grouped with Abelisauridae indet. morphotype 1 from the Western Treppe Syncline or *Aucasaurus*. Therefore, the morphometric analyses also strongly suggest abelisaurid affinities for this specimen. Regarding the cladistic analyses, the results do not support direct attribution of the teeth to Abelisauridae. The Viso specimen was recovered within Abelisauroidae, but more closely related to Noasauridae. Nevertheless, forcing the specimen into Abelisauridae, in a group comprising *Arcovenator*, *Majungasaurus*, and *Indosuchus*, does not require any additional steps, suggesting that this hypothesis is as parsimonious as the position recovered in the unconstrained analysis. The Chera 2 specimen was allied with Tyrannosauroidae.

However, all features found as synapomorphies for this clade are also shared with most abelisaurids. Forcing the specimen into Abelisauridae requires five additional steps compared to the length of the consensus tree recovered on the unconstrained analysis and one extra step to place it within the group comprising *Arcovenator*, *Majungasaurus*, and *Indosuchus*. The analysis performed for the Montrebei specimen yielded very poor resolution, with most taxa placed in a large polytomy. The most parsimonious trees placed the specimen within the Paraves clade and forcing its position into Abelisauridae required four additional steps. The lack of European abelisaurid dental and mandibular elements, along with the limited representation of European tooth data in the matrix, may account for the poor resolution of the cladistic analyses.

#### **Abelisauridae indet.**

**Morphotype 1.** The isolated tooth DPM-MON-T10 1 from Montrebei shares with Abelisauridae a ziphodont crown with a ‘D’-shaped cross-section and the irregular, non-oriented enamel-texture (Hendrickx et al., 2019, 2020). Therefore, this tooth can be confidently assigned to Abelisauridae, which is also supported by the results of some morphometric analyses, despite the cladistic analysis not yielding sufficient resolution to test this attribution. Furthermore, the distal position of the apex and the profile of the distal margin are similar to the condition found in the distal teeth of the mesial dentition in abelisaurids, with the crown being more distally recurved (Hendrickx et al., 2020). Despite most abelisaurid taxa exhibiting a salinon-to J-shaped basal cross-section, specimen DPM-MON-T10 1 has a ‘D’-shaped cross-section, as observed in *Rahiolisaurus* and MPCN-PV 69 (Novas et al., 2010; Gianechini et al., 2015; Hendrickx et al., 2020). The Montrebei tooth is particularly thick, a condition similarly present in the mesialmost teeth of several abelisaurid taxa such as *Chenanisaurus*,

647 *Majungasaurus*, and *Rahiolisaurus* (Hendrickx et al., 2020) but differing from the more  
 648 labiolingually compressed mesial teeth of other abelisaurids such as the premaxillary  
 649 teeth of *Majungasaurus* (see Hendrickx et al., 2020). Although the crown lacks its  
 650 cervical portion, it seems to have been weakly elongated, as in all known members of  
 651 Abelisauridae (Hendrickx et al., 2020).  
 652 The mesiolingually oriented mesial carina of the Montrebei specimen resembles that of  
 653 the first premaxillary tooth of abelisaurids, as well as some teeth possibly belonging to  
 654 *Abelisaurus* (MPCA 1, 5 and 267; see Hendrickx et al., 2020), mesial abelisaurid  
 655 crowns from the Western Treppe Syncline (Abelisauridae indet. 1) and Poyos  
 656 (Abelisauridae indet. morphotype 1), along with the lateral teeth of *Arcovenator* and  
 657 closely related specimens from Poyos (Hendrickx and Mateus, 2014, Tortosa et al.,  
 658 2014; Hendrickx et al., 2020; Isasmendi et al., 2022, 2024; Malafaia et al., 2025). The  
 659 strongly lingually deflected distal carina of DPM-MON-T10 is a feature shared with the  
 660 mesial abelisaurid crown from the Western Treppe Syncline, the isolated mesial crown  
 661 IIPG-09, and the lateral crowns of *Arcovenator* (Hendrickx et al., 2020; Meso et al.,  
 662 2021; Isasmendi et al., 2022, 2024). Two distinctive features of the Montrebei crown  
 663 are the position of the apex, which apparently extended beyond its basodistalmost point,  
 664 and its concave distal profile. These features resemble the condition seen in  
 665 ‘*Indosuchus*’ and differ from the mesial dentition of *Chenanisaurus*, *Majungasaurus*,  
 666 *Skorpiovenator*, the mesial teeth closely related to *Arcovenator* from Poyos, and mesial  
 667 teeth from the Allen Formation in Patagonia (Smith, 2007; Hendrickx et al., 2020; Meso  
 668 et al., 2021; Malafaia et al., 2025). However, the distal maxillary teeth of ‘*Indosuchus*’  
 669 do not exhibit a distal profile as concave as that of the Montrebei specimen (see  
 670 Hendrickx et al., 2020; fig. 6.f). Therefore, in light of these considerations, DPM-MON-  
 671 T10 1 is here assigned to Abelisauridae indet.

672 **Morphotype 2.**

673 The isolated tooth MG 73 from Viso shares a combination of features with other  
674 members of Abelisauridae, namely a ziphodont crown, a straight or gently convex distal  
675 profile, a lanceolate basal cross-section, denticulated mesial and distal carinae (with the  
676 distal carina reaching the cervix), presence of large number of small denticles in the  
677 distal carina (c. 20 denticles per 5 mm), sub-quadrangular distal denticles, narrow  
678 interdenticular space, and an irregular, non-oriented enamel texture (see Hendrickx et  
679 al., 2020). Furthermore, the cladistic and morphometric analyses carried out herein  
680 further support its referral to Abelisauridae or, at least to Abelisauroidae.

681 An irregular enamel surface texture and the presence of transverse undulations are two  
682 dental features typically observed in Abelisauridae dentition (e.g., Canale et al., 2009;  
683 Longrich et al., 2017; Hendrickx et al., 2019, 2020; Meso et al., 2021; Isasmendi et al.,  
684 2022, 2024). Transverse undulations are also present in other Iberian abelisaurid teeth  
685 such as the three abelisaurid morphotypes from the latest Maastrichtian of the Western  
686 Tremp Syncline (South Pyrenean Basin) (Isasmendi et al., 2024) and the teeth of  
687 *Arcovenator* from the upper Campanian of Armuña (UPUAM 14044) and Laño  
688 (Isasmendi et al., 2022). Unlike MPZ 2004/8, MPZ 2017/804, and MPZ 2022/86 from  
689 the Maastrichtian of Western Tremp Syncline but similar to the lateral teeth of most  
690 Abelisauridae (Hendrickx et al., 2020), concave surfaces adjacent to the carinae are  
691 absent in MG 73.

692 MG 73 shares a moderately compressed crown (CBR of 0.58) with *Abelisaurus*,  
693 *Arcovenator*, *Aucasaurus*, and *Majungasaurus* (Hendrickx et al., 2020), as well as the  
694 Abelisauridae indet. morphotypes 1 and 3 from the Western Tremp Syncline (Isasmendi  
695 et al., 2024) and many teeth from Poyos (Malafaia et al., 2025).

Typically, the mesial and distal carinae in lateral abelisaurid crowns are centrally located on their respective mesial and distal margins (Hendrickx et al., 2020). However, in the MG 73 specimen, the mesial carina is lingually positioned and straight, resembling the condition present in the teeth referred to *Arcovenator* or a closely related taxon from Jas Neuf Sud, Laño, and Poyos (Tortosa et al., 2014; Hendrickx et al., 2020; Isasmendi et al., 2022; Malafaia et al., 2025). The distal carina of MG 73 is centrally located, as in most abelisaurids (Hendrickx et al., 2020), differing from the strongly deflected distal carinae of *Arcovenator* from Jas Neuf Sud, the largest teeth from Laño, and most of the *Abelisauridae* indet. lateral morphotypes from the Western Treppe Syncline (Tortosa et al., 2014; Hendrickx et al. 2020; Isasmendi et al., 2022, 2024). The distocentral denticle density of MG 73 is 20 denticles per five millimeters, which is slightly higher than the range typically found in most *Abelisauridae* taxa that have 9 to 15 denticles per five millimeters (Hendrickx et al., 2020). The sub-quadrangular basal distal denticles of MG 73 are similar in shape to those located at mid-crown in *Abelisaurus*, *Kryptops*, *Rugops*, some *Aucasaurus*, and most *Majungasaurus*, differing from the denticle shape present in *Arcovenator*, some teeth of *Aucasaurus*, and a few crowns of *Majungasaurus* (Hendrickx et al., 2020). Although many abelisaurids exhibit interdenticular sulci, MG 73 lacks them, similar to the condition present in the crowns of *Aucasaurus*, *Arcovenator*, *Rugops*, and UCPC 10, which either lack these sulci or have them restricted to the base of the distal denticles (Smith, 2007; Hendrickx et al., 2020). Therefore, despite MG 73 exhibiting many abelisaurid traits, some features are distinct from any previously described member of *Abelisauridae*, and hence, this crown is regarded as *Abelisauridae* indet.

***Arcovenator* sp.**

720 The isolated tooth CH 168 from Chera 2 has traits present in abelisaurid teeth, including  
721 an irregular and non-oriented enamel texture, CHR and CBR values, denticle shape,  
722 denticle density, and the DSDI value (see Hendrickx et al., 2020). Furthermore, one of  
723 the LDA classifies this specimen as *Arcovenator*. Nevertheless, the cladistic analyses do  
724 not clarify the phylogenetic affinities of the Chera 2 specimen. This tooth lacks some  
725 typical dental features found in most members of Abelisauridae (see Hendrickx et al.,  
726 2020), such as a straight or gently convex distal profile, a mesial carina extending to the  
727 cervix, or the characteristic arrangement of the carinae. The CBR is moderate for a  
728 lateral tooth, a feature shared with *Aucasaurus*, *Carnotaurus*, *Majungasaurus*, some  
729 teeth assigned to *Arcovenator* sp. from Laño, and some teeth from Poyos (Hendrickx et  
730 al., 2020; Isasmendi et al., 2022; Malafaia et al., 2025). Its CHR value is also moderate  
731 (1.94), similar to those of *Arcovenator*, *Chenanisaurus*, and some lateral teeth of  
732 *Majungasaurus*, among others (Hendrickx et al., 2020). Comparable CHR values are  
733 also found in teeth assigned to *Arcovenator* sp. from Laño and in lateral teeth belonging  
734 to an abelisaurid closely related to *Arcovenator* from Poyos (Isasmendi et al., 2022;  
735 Malafaia et al., 2025). Furthermore, among abelisaurid teeth, only those referred to  
736 *Arcovenator* from Armuña, Jas Neuf Sud, and Laño, as well as those closely related to  
737 *Arcovenator* from Poyos, exhibit an apex that extends beyond the basodistalmost point  
738 of the crown (Tortosa et al., 2014; Hendrickx et al., 2020; Isasmendi et al., 2022;  
739 Malafaia et al., 2015; this study).

740 The lateral crowns of abelisaurid theropods typically have mesial and distal carinae that  
741 extend along the midline of the crown from the apex to the root (Hendrickx et al. 2020),  
742 unlike the CH 168 tooth from Chera 2. Indeed, in this specimen the mesial carina is  
743 straight, displaces mesiolingually toward the base, and extends from the apex to  
744 approximately two-thirds of the crown. On the other hand, the distal carina reaches the



cervix but is sigmoidal and labially deflected. This condition more closely resembles that present in teeth referred to *Arcovenator* from the Jas Neuf Sud locality (Tortosa et al., 2014; Hendrickx et al., 2020), the largest teeth assigned to *Arcovenator* from Laño (Isasmendi et al., 2022), and the morphotype 2 abelisaurid teeth from Poyos (Malafaia et al., 2025).

Regarding the denticles, the DSDI value of the Chera 2 tooth is 1, similar to the teeth referred to *Arcovenator* from Jas Neuf Sud and Laño, as well as some specimens closely related to this taxon from Poyos , but differing from some *Arcovenator* teeth, in which the DSDI exceed 1.2 (Hendrickx et al., 2020; Isasmendi et al., 2022; Malafaia et al., 2025). The mesiocentral denticle density of CH 168 (MC of 15 denticles per 5 mm) is similar to the MC values of *Abelisaurus* and *Arcovenator*, while the distocentral denticle density of the specimen studied here (DC of 15 denticles per 5 mm) more closely resembles those of *Arcovenator* or *Aucasaurus*, whereas *Abelisaurus* teeth have lower DC values (Tortosa et al., 2014; Hendrickx et al., 2020).

The outline of the denticles in CH 168 is symmetrically to slightly asymmetrically convex as in *Arcovenator* teeth from Jas Neuf Sud and Laño, the teeth from Poyos, *Skorpiovenator*, and some abelisaurid teeth from the Cenomanian of northern Africa (Richet et al., 2013; Hendrickx et al., 2020; Isasmendi et al., 2022; Malafaia et al., 2025). The CH 168 tooth further resembles the teeth of *Arcovenator* in exhibiting subquadrangular or apicobasally subrectangular mesial denticles and mesiodistally subrectangular distal denticles (Hendrickx et al., 2020; Isasmendi et al., 2022; Malafaia et al., 2025). Therefore, based on the results of the morphometric analyses and the morphological similarities of the CH 168 tooth with other *Arcovenator* teeth, this specimen is assigned to *Arcovenator* sp.

770 **Postcranial remains.**

771 Despite their fragmentary nature, the Laño specimens exhibit several morphological  
772 features that support their attribution to Abelisauroidae. The presence in MCNA 17433  
773 of a centrodiapophyseal lamina on the transverse processes of the anterior and middle  
774 caudal vertebrae allows it to be confidently referred to this clade (Méndez, 2014,  
775 Tortosa et al., 2014), and Abelisauridae according to Baiano et al. (2023). In addition,  
776 the neural arch is also broader than the mid-centrum, which is a feature shared with  
777 *Masiakasaurus* and several abelisaurids (Baiano et al., 2023). In addition, the elliptical  
778 shape of the articular surfaces of the caudal vertebra, the presence of a ventral  
779 longitudinal groove on the centrum, and the lateral orientation of transverse process of  
780 the anterior dorsal and the anterior to middle caudal vertebra and the fan-shaped  
781 morphology of the transverse processes of the anterior dorsal vertebra allows these  
782 specimens to be referred to Majungasaurinae. Furthermore, since the presence of  
783 *Arcovenator* has previously been documented in Laño (Isasmendi et al., 2022), these  
784 remains are assigned to cf. *Arcovenator* sp., pending the discovery of additional  
785 diagnostic material.

786 The horizontally projected transverse processes seen in MCNA 8366 are also present in  
787 the anterior dorsals of *Majungasaurus* and *Sinraptor* (Currie and Zhao, 1993,  
788 O'Connor, 2007) and differ from those of *Allosaurus* and *Sigilmassasaurus* (Madsen,  
789 1976; Evers et al., 2015), in which the transverse processes are more ventrally  
790 projected. Furthermore, the fan-shaped transverse processes, which are laterally  
791 projected in MCNA 8366, are also seen in the anteriormost dorsal of *Majungasaurus*  
792 and *Sigilmassasaurus* (O'Connor, 2007; Evers et al., 2015). The morphology of the  
793 transverse process in the specimen from Laño, with an anterior convex and posterior  
794 straight edge that make the diapophysis project face laterally, is similar to that of

795 *Majungasaurus* (UA 8678; fig. 3 of O'Connor, 2007) although in this taxon the anterior  
796 convex edge is less pronounced. The posterior straight edge of the transverse processes  
797 of MCNA 8366 differs from the concave edges and posterolaterally facing diapophysis  
798 exhibited by the anteriormost dorsals of, for instance, *Carnotaurus* (MACN-CH 894),  
799 *Eoabelisaurus* (MPEF PV 3990), *Viavenator* (MAU-Pv-LI-530; fig. 7D of Filippi et al.,  
800 2018) and the indeterminate brachyrostran (MAU-Pv-LI-665; fig 6D of Méndez et al.,  
801 2022). This indicates that this condition could be a majungasaurine synapomorphy.  
802 Nevertheless, this hypothesis needs to be phylogenetically tested.

803 The ventral surface of the MCNA 17433 caudal vertebra exhibits a ventral longitudinal  
804 groove, a feature also present in *Allosaurus*, *Arcovenator*, *Aucasaurus*, *Ceratosaurus*,  
805 *Majungasaurus*, and *Viavenator*, as well as in many spinosaurids and megalosaurids  
806 (e.g., Madsen, 1976; Madsen and Welles, 2000; O'Connor, 2007; Benson, 2010;  
807 Méndez, 2014; Tortosa et al., 2014; Filippi et al., 2018; Malafaia et al., 2020). This  
808 contrasts with some abelisaurids, such as *Ekrixinatosaurus* and *Rajasaurus*, which  
809 exhibit a keeled ventral surface (Méndez, 2014). The presence of a centrodiapophyseal  
810 lamina on the ventral surface of the preserved transverse processes is a feature shared  
811 with Abelisauridae (e.g., *Arcovenator*, *Aucasaurus*, *Carnotaurus*, *Ekrixinatosaurus*,  
812 *Kurupi* and *Majungasaurus*) and *Masiakasaurus*, but not with *Allosaurus* or  
813 *Ceratosaurus* (Carrano et al., 2002; Coria et al., 2002; Calvo et al., 2004; O'Connor,  
814 2007; Méndez, 2014; Tortosa et al., 2014; Iori et al., 2021; Baiano et al., 2023).

815 Nevertheless, despite deformation, the bases of the transverse processes in MCNA  
816 17433 do not appear to be strongly dorsally directed. This differs from many  
817 abelisaurids such as *Aucasaurus*, *Carnotaurus*, *Ekrixinatosaurus* and *Skorpiovenator*,  
818 and is more similar to the condition observed in *Allosaurus*, *Arcovenator*, *Ceratosaurus*,  
819 *Majungasaurus*, and *Rahiolisaurus* (Madsen, 1976; Madsen and Welles, 2000;

820 O'Connor, 2007; Méndez, 2014; Baiano et al., 2023). The anterior and posterior  
821 articular facets of the MCNA 17433 caudal vertebra are close to elliptical in outline  
822 (note that although the centrum appears strongly mediolaterally compressed, this is  
823 likely a taphonomic artifact). This shape is more similar to that observed in  
824 *Arcovenator*, *Majungasaurus*, *Rajasaurus*, and *Sinraptor*, and differs from the more  
825 subcircular surfaces of, for instance, *Allosaurus*, *Aucasaurus*, *Carnotaurus*,  
826 *Ekrixinatosaurus*, *Ichthyovenator*, *Kurupi*, *Tyrannosaurus* or *Viavenator* (Currie and  
827 Zhao, 1993; Brochu, 2003; Wilson et al., 2003; O'Connor, 2007; Allain et al., 2012;  
828 Méndez, 2014; Tortosa et al., 2014; Filippi et al., 2018; Iori et al., 2021; Baiano et al.,  
829 2023).

### 830 **New contributions to the knowledge of European abelisaurids.**

831 In the Late Cretaceous European archipelago, skeletal remains of mid- to large-bodied  
832 theropods have been recovered in Central and Western Europe (e.g., Ósi et al., 2010;  
833 Ósi and Buffetaut, 2011; Tortosa et al., 2014; Isasmendi et al., 2022; Buffetaut et al.,  
834 2024). Among these remains, indeterminate tetanurans and abelisaurids have been  
835 identified. In the central European region, specifically at the Santonian site of Iharkút in  
836 Hungary and the Campanian site of Styria in Austria, teeth assigned to early-branching  
837 indeterminate tetanurans have been recovered (Ósi et al., 2010, 2012). Additionally, the  
838 Iharkút site has yielded further theropod remains referred to Abelisauridae (Ósi et al.,  
839 2010; Ósi and Buffetaut, 2011), indicating that two mid- to large-sized theropod clades  
840 coexisted in Central Europe during the Santonian. However, the scarce theropod  
841 remains in this area makes it challenging, at this moment, to provide a more accurate  
842 interpretation of the phylogenetic relationships of this fauna. Western Europe, on the  
843 other hand, appears to lack mid- to large-bodied early-branching tetanurans, with

844 Abelisauridae representing the main apex predator (e.g., Csiki-Sava et al., 2015;  
 845 Isasmendi et al., 2022; Buffetaut et al., 2024; Malafaia et al., 2025).  
 846 Abelisaurid theropods have historically been interpreted as biogeographical indicators  
 847 of Gondwanan affinities within Late Cretaceous European dinosaur assemblages  
 848 (Buffetaut et al., 1988; Le Loeuff and Buffetaut, 1991; Csiki-Sava et al., 2015). Current  
 849 evidence supports the persistence of multiple abelisauroid lineages within the European  
 850 archipelago up to the latest Cretaceous (Carrano and Sampson, 2008; Tortosa et al.,  
 851 2014; Buffetaut et al., 2024; Buffetaut, 2025). Based on the French fossil record,  
 852 Tortosa et al. (2014) proposed that small-bodied abelisaurids likely originated from an  
 853 Albian lineage of early-branching abelisaurids represented by *Genusaurus sisteronis*  
 854 from Provence, which subsequently diversified into distinct lineages. It should be noted  
 855 that this taxon has, however, been variably interpreted as a noasaurid (e.g., Carrano and  
 856 Sampson, 2008), an early-branching abelisaurid (e.g., Tortosa et al., 2014; Baiano et al.,  
 857 2023), or an abelisaurid more closely related to Furileosauria (Buffetaut et al., 2024;  
 858 Buffetaut, 2025).  
 859 The Cenomanian fossil record of Ibero-Armorica is scarce and currently consists of the  
 860 considerably incomplete *Caletodraco cottardi* from the early Cenomanian glauconitic  
 861 Chalk of Normandy (Buffetaut et al., 2024) (Fig. 9, loc. 1). This taxon, which is known  
 862 from ilia, some axial elements and a possible tooth, was interpreted as a furileusaurian  
 863 brachyrostran, potentially reflecting a more intricate evolutionary history of European  
 864 abelisaurids, with the presence of different abelisaurid lineages in Ibero-Armorica  
 865 (Buffetaut, 2024; Buffetaut et al., 2024; Malafaia et al., 2025). Other Cenomanian  
 866 abelisaurid remains from France and Spain include isolated teeth from Algora (Segovia,  
 867 Spain), “La Buzinie” (Charentes, France), and Limanes (Asturias, Spain). The isolated  
 868 theropod teeth from Algora (Utrillas Formation) (Fig. 9, loc. 4), were initially referred

869 to Carcharodontosauridae indet. (Torices et al., 2012) before being reassigned to cf.  
 870 Abelisauridae by Pérez-García et al. (2020). Additionally, a medium- to large-sized  
 871 caudal vertebra (ALG 192) from the same site was attributed to Theropoda indet.  
 872 (Pérez-García et al., 2020). Two other isolated teeth (MA BZN 1 and 2) from the “La  
 873 Buzinie” locality (Fig. 9, loc. 2) were similarly assigned to Carcharodontosauridae by  
 874 Vullo et al., 2007 before being reassigned to cf. Abelisauridae indet. by Malafaia et al.  
 875 (2025), based on their crown ornamentation and denticle densities. Another isolated  
 876 tooth from the Cenomanian La Manjoya Fm in the Limanes municipality (Fig. 9, loc. 3)  
 877 was also identified as Theropoda indet. closely related to Carcharodontosauridae (Ruiz-  
 878 Omeñaca et al., 2009). Malafaia et al. (2025) agreed on the morphometric affinities  
 879 suggested by Pérez-García et al. (2020) for the teeth from Algora, Charentes, and  
 880 Limanes. The Limanes tooth belongs to a mid-sized theropod. The apex of the crown  
 881 does not seem to have extended beyond its basodistalmost point (Ruiz-Omeñaca et al.,  
 882 2009; fig. 2a and b), both mesial and distal margins are convex in lateral view (Ruiz-  
 883 Omeñaca et al., 2009), the distal denticles are asymmetrically convex and apically  
 884 inclined, and the enamel exhibits an irregular texture, all features present in  
 885 Abelisauridae but absent in Carcharodontosauridae (Hendrickx et al., 2019, 2020). The  
 886 mesial carina is mesiolingually oriented (Ruiz-Omeñaca et al., 2009; fig. 2c), being  
 887 straight and centered apically but displaced lingually to the base, as in the first  
 888 premaxillary tooth of Abelisauridae, some teeth that may belong to *Abelisaurus* (MPCA  
 889 1, 5 and 267; see Hendrickx et al., 2020), Abelisauridae indet. morphotype 1 tooth from  
 890 the Western Tremp Syncline and Poyos, as well as in the lateral teeth of *Arcovenator*  
 891 and closely related specimens from Poyos (Hendrickx and Mateus, 2014, Tortosa et al.,  
 892 2014; Hendrickx et al., 2020; Isasmendi et al., 2022, 2024; Malafaia et al., 2025). Ruiz-  
 893 Omeñaca et al. (2009) indicate a mesial denticle density of 2 denticles/mm and a distal

894 denticle density of 2.22 denticle/mm (giving a DSDI value of 0.9). However, these  
895 measurements were taken from different positions along the crown, which compromises  
896 their comparability. The inferred MA is 14 denticles per five mm and the DA is 12.5  
897 denticles per five mm. Furthermore, an accurate DSDI value cannot be calculated  
898 because the distal carina is not preserved at mid-crown. Therefore, based on the  
899 combination of features shared with Abelisauridae, this tooth is here regarded as  
900 Abelisauridae indet. These findings suggest that carcharodontosaurians were already  
901 extinct in Ibero-Armorica by the Cenomanian and that abelisaurids had become the apex  
902 theropods in those ecosystems.

903 No abelisaurid or large-sized theropod remains that may belong to Abelisauridae have  
904 been recovered to date from the Turonian, Coniacian, or Santonian of the Ibero-  
905 Armorican landmass. In the Campanian–Maastrichtian Viso locality (Beira Litoral,  
906 Portugal) (Fig. 9, loc. 5), several theropod remains were reported and figured by  
907 Antunes and Sigogneau-Russell (1992) and Galton (1996). Among the Viso theropod  
908 material, tooth VI 1 (Antunes and Sigogneau-Russell, 1992; pl. I, fig. 5) was initially  
909 referred to as Theropoda indet. However, due to its considerable size, it may be  
910 attributable to Abelisauridae as the herein studied tooth from Viso (MG 73). The MG 73  
911 specimen had previously been identified as *Megalosaurus* sp. (Sauvage, 1897–1898),  
912 *Megalosaurus* cf. *pannoniensis* (Lapparent and Zbyszewski, 1957), and Theropoda  
913 indet. (Malafaia et al., 2024). Nonetheless, the morphological characteristics exhibited  
914 by this specimen warrant its reassignment to Abelisauridae indet., representing the first  
915 unequivocal abelisaurid remain from the Cretaceous of Portugal.

916 Theropod remains are relatively rare in lower Campanian deposits of Ibero-Armorica.  
917 At the Lambeau du Beausset locality in Var (France) (Fig. 9, loc. 6), the abelisaurid  
918 *Tarascosaurus* was erected based on fragmentary material (Le Loeuff and Buffetaut,

1991). This taxon has since been regarded as *nomen dubium* (Rauhut, 2003; Allain and Pereda-Suberbiola, 2003) or referred to *Abelisauroida incertae sedis* (Carrano and Sampson, 2008). Nevertheless, it was recovered as an early-branching member of *Abelisauridae* in the phylogenetic analyses performed by Tortosa et al. (2014). Additional material from the Iberian Peninsula, specifically femora from the Laño site (Fig. 9, loc. 11), has been compared to *Tarascosaurus* (Le Loeuff and Buffetaut, 1991, Le Loeuff, 1992), but no definitive evidence currently supports this attribution (Isasmendi et al., 2021, 2022; Malafaia et al., 2025).

The middle and upper Campanian abelisaurid record in France and Spain is comparatively richer. In Provence, indeterminate abelisaurid remains have been reported from the Trets-La Boucharde site (Fig. 9, loc. 7), Velaux-Bastide Neuve (Fig. 9, loc. 8), and the Fox-Amphoux area (Fig. 9, loc. 9) (Tortosa et al., 2014). The Trets-La Boucharde specimen from the “Begudian fluvio-lacustrine sandstones” was previously described as an abelisauroid (Allain and Pereda-Suberbiola, 2003), and it is currently regarded as an indeterminate abelisaurid by various authors (Carrano and Sampson, 2008; Tortosa et al., 2014). The abelisaurid remains reported by Tortosa et al. (2014) from Velaux-Bastide Neuve (“Begudian sandstones”; Garcia et al., 2010; Cincotta et al., 2015) and the Fox-Amphoux area (middle–late Campanian to early Maastrichtian? “Grès à reptiles”) comprise a set of isolated teeth that are yet to be published.

*Arcovenator escotae* was erected based on a set of cranial, axial and appendicular elements recovered from the upper Campanian Pourrières-Jas Neuf Sud locality (Fig. 9, loc. 10) of the lower “Argiles rutilantes” Fm (Tortosa et al., 2014). This is the most complete European abelisaurid currently known and has been classified as a majungasaurine (Tortosa et al., 2014). In addition to the holotype and several referred specimens reported by Tortosa et al. (2014), other skeletal elements from Upper



944 Cretaceous deposits of France and the Iberian Peninsula have also been assigned to  
 945 *Arcovenator* or closely related forms (Tortosa et al., 2014; Pérez-García et al., 2016;  
 946 Isasmendi et al., 2022; Malafaia et al., 2025). This is the case of the late Campanian to  
 947 early Maastrichtian Pourcieux specimen from the Les Tuillières site (“Rognacian”  
 948 coarse sandstone) (Fig. 9, loc. 15) in Provence. This specimen was first regarded as  
 949 Abelisauridae (Buffetaut et al., 1998, Rauhut, 2003), and Abelisauridae indet. or  
 950 Carcharodontosauridae (Carrano and Sampson, 2008), before Tortosa et al. (2014)  
 951 referred it to ?*Arcovenator* sp. The coeval “Grès à reptiles” red beds of Cruzy in  
 952 Languedoc (Fig. 9, loc. 14) has yielded several teeth as well as cranial and postcranial  
 953 fragmentary remains, which may be attributable to medium- to large-sized abelisaurids  
 954 (Buffetaut et al., 1999; Buffetaut, 2005; Ósi and Buffetaut, 2011; Tortosa et al., 2014),  
 955 possibly related to *Arcovenator* (Tortosa et al., 2014). In the Iberian Peninsula, isolated  
 956 teeth from Laño (County of Treviño), Armuña (Segovia), Chera 2 (Valencia), and  
 957 Poyos (Guadalajara) were assigned to *Arcovenator* or closely related forms (Pérez-  
 958 García et al., 2016; Isasmendi et al., 2022; Malafaia et al., 2025; this work). The largest  
 959 set of isolated teeth assigned to *Arcovenator* sp. come from the upper Campanian  
 960 Sedano Fm of Laño (Fig. 9, loc. 11) (Isasmendi et al., 2022), which has also yielded  
 961 additional postcranial theropod remains. It must be noted that a dorsal neural arch  
 962 previously assigned to *Rhabdodon* (Pereda-Suberbiola and Sanz, 1999) and the caudal  
 963 vertebra herein described can be referred to cf. *Arcovenator*. The coeval Vegas de  
 964 Matute Fm of Armuña (Fig. 9, loc. 12) and Sierra Perenchiza Fm of Chera 2 (Fig. 9, loc.  
 965 13) have also yielded several isolated teeth referred to cf. *Arcovenator* and *Arcovenator*  
 966 sp., respectively (Pérez-García et al., 2016; Company et al., 2005; this work). It is worth  
 967 noting that the isolated tooth was subject of varying taxonomic interpretations, having  
 968 been referred to Theropoda indet. (Company, 2005), ?Neoceratosauria indet. (Company

969 et al., 2009), and cf. *Arcovenator* (Isasmendi et al., 2022). Additionally, the pedal  
 970 ungual phalanges recovered from the Armuña site were initially assigned to Theropoda  
 971 indet. (Pérez-García et al., 2016) and later reinterpreted as Abelisauridae indet.  
 972 (Isasmendi et al., 2024). Two dental morphotypes from the late Campanian–early  
 973 Maastrichtian Poyos site (Villalba de la Sierra Fm; Gil et al., 2004; Ortega and Pérez-  
 974 García, 2009) (Fig. 9, loc. 18) were described and referred to an abelisaurid closely  
 975 related to *Arcovenator* by Malafaia et al. (2025). This site has additionally yielded non-  
 976 dental theropod remains initially referred to Abelisauroida indet. (Ortega et al., 2019;  
 977 Pérez-García et al., 2019) and later to Abelisauridae indet. (Malafaia et al., 2025),  
 978 pending further detailed studies. Some theropod remain from the Lo Hueco site (Fig. 9,  
 979 loc. 17) were regarded as Theropoda indet. whereas others were referred to  
 980 Abelisauridae indet. (Ortega et al., 2015, 2022).  
 981 The abelisaurid fossil record from the Maastrichtian is comparatively less abundant than  
 982 that from the Campanian (Isasmendi et al., 2024). The early Maastrichtian Montrebei  
 983 site (La Posa Formation; 'Grey Garumnian' of the Tremp Group; Fondevilla et al., 2019)  
 984 of the Tremp Syncline (Fig. 9, loc. 20) in Lleida (Iberian Peninsula) yielded an isolated  
 985 tooth initially assigned to Theropoda indet. (Torices et al., 2015), but its subsequent  
 986 restudy has permitted its referral to Abelisauridae indet. In the western section of the  
 987 Tremp Syncline, abelisaurid teeth were also reported from the late Maastrichtian Blasi  
 988 1, 2B and 3 (Fig. 9, loc. 21) and the latest Maastrichtian 172-i/04/e (Fig. 9, loc. 22)  
 989 sites, located at the top of the Arén Sandstone and the lower part of the Tremp Group  
 990 ("Grey Garumnian") (Canudo et al., 2016; Puértolas-Pascual et al., 2018; Isasmendi et  
 991 al., 2024). Several teeth from the Blasi sites, along with the specimen from 172-i/04/e,  
 992 were previously referred to Theropoda indet. by Torices et al. (2015) and Puértolas-  
 993 Pascual et al. (2018) and later assigned to three different Abelisauridae indet.

994 morphotypes by Isasmendi et al. (2024). At the Cassagnau 1 locality (Fig. 9, loc. 16),  
995 situated within the late Maastrichtian Auzas Marls Fm in Haute-Garonne, an isolated  
996 tooth, originally referred to Theropoda indet. by Laurent (2003), was later reinterpreted  
997 as belonging to Abelisauridae by Csiki-Sava et al. (2015). In addition to isolated teeth,  
998 the Maastrichtian French deposits have yielded several indeterminate abelisaurid  
999 postcranial remains. These are the limb bones (femur and tibia) initially assigned to  
1000 Neoceratosauria by Valentin et al. (2012), recovered at the Vitrolles-La Plaine site (Fig.  
1001 9, loc. 19) from the “Rognacian clays and mottled marls” deposits in Bouches-du-  
1002 Rhône. This site has been dated as possibly late Maastrichtian (Valentin et al., 2012) or  
1003 also as late Campanian–early Maastrichtian (Fondevilla et al., 2019).

1004 Taken together, the abelisaurid fossil record across Ibero-Armorica reveals a complex  
1005 and temporally extensive presence of this clade, spanning from the Albian to the latest  
1006 Maastrichtian, with a roughly 10-million-year gap during the Turonian–Santonian  
1007 without any abelisaurid fossil record. Despite their relatively fragmentary nature, these  
1008 remains, ranging from isolated teeth to partial skeletons, provide compelling evidence  
1009 for the persistence and diversification of abelisaurids within the European archipelago  
1010 throughout the Late Cretaceous. The distribution of taxa such as *Genusaurus*,  
1011 *Caletodraco*, *Arcovenator*, and additional indeterminate forms suggests multiple  
1012 evolutionary lineages, potentially some of them stemming from an Albian stock and  
1013 others likely representing subsequent dispersal events, undergoing insular  
1014 diversification through the Late Cretaceous. Moreover, the taxonomic revisions of  
1015 previously misassigned material emphasize the need for ongoing reevaluation of  
1016 fragmentary theropod remains. Such efforts are crucial to refine our understanding of  
1017 abelisaurid paleobiogeography and systematics in Europe.

1018

## 1019 CONCLUSIONS

1020 The systematic, morphometric and cladistic studies carried out with the tooth sample  
1021 from late Campanian Chera 2 (Valencia), early Maastrichtian Montrebei (Lleida) and  
1022 Campanian–Maastrichtian Viso (Beira Litoral) localities from the Iberian Peninsula  
1023 have allowed to reassign these elements to abelisaurids. Although incomplete, the  
1024 Montrebei (DPM-MON-T10 1) and Viso (MG 73) specimens differ from the dentition  
1025 of the already erected members of Abelisauridae. On the other hand, the crown-shape,  
1026 disposition of the carinae and denticle shape exhibited by CH 168 tooth from Chera 2  
1027 are similar to those found in *Arcovenator* teeth. Hence, the Montrebei and Viso teeth are  
1028 now considered to belong to Abelisauridae indet., while the Chera 2 specimen is  
1029 attributed to *Arcovenator* sp. In addition, the postcranial remains (axial elements)  
1030 recovered from the Laño site (County of Treviño) are here regarded to as an abelisaurid,  
1031 mainly based on the presence of a centrodiapophyseal lamina on the preserved  
1032 transverse process in the caudal vertebra (MCNA 17433). In addition, the shape and  
1033 orientation of the transverse processes of the dorsal and caudal vertebrae suggest  
1034 majungasaurine affinities. Therefore, these features, together with the presence of  
1035 *Arcovenator* teeth in the site, allow us to assign these elements to cf. *Arcovenator*. The  
1036 revision of the Laño postcranial remains has allowed to refute earlier interpretations that  
1037 assigned the anterior dorsal neural arch (MCNA 8366) to *Rhabdodon*. Furthermore, the  
1038 Viso isolated tooth, now assigned to Abelisauridae indet., is the first unequivocal  
1039 abelisaurid remain from Portugal.

1040 The fossil record indicates that abelisaurids were among the most prevalent theropod  
1041 groups in the Late Cretaceous faunas across the Ibero-Armorican landmasses. Indeed,  
1042 the reclassification of mid- to large-sized isolated teeth from Ibero-Armorica as  
1043 belonging to abelisaurids, rather than to carcharodontosaurids or closely related forms,

indicates that carcharodontosaurians were likely extinct in Ibero-Armorica by the Cenomanian, and that abelisaurids had already taken over as the dominant large theropods in those ecosystems. The abelisaurid fossil record from Ibero-Armorica reveals a complex and temporally extensive presence of this clade, spanning from the Albian to the latest Maastrichtian, with a significant gap of approximately 10 million years during the Turonian–Santonian, in which no abelisaurid fossils have been found yet. Despite the generally fragmentary nature of these remains, ranging from isolated teeth to partial skeletons, they provide strong evidence for the persistence and diversification of abelisaurids across the European archipelago. The distribution of taxa such as the noasaurid or possibly furileusaurian *Genusaurus*, the putative furileusaurian *Caletodraco*, the majungasaurine *Arcovenator*, and other indeterminate forms suggests the presence of multiple evolutionary lineages, some potentially originating from an Albian stock, while others might have resulted from several dispersal events, undergoing insular diversification throughout the Late Cretaceous. Additionally, the taxonomic revisions of previously misclassified material underscore the need to continuously reevaluate fragmentary theropod remains, thereby refining our understanding of abelisaurid paleobiogeography and systematics in Europe.

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## REFERENCES

- Allain, R., & Taquet, P. (2000). A new genus of Dromaeosauridae (Dinosauria, Theropoda) from the Upper Cretaceous of France. *Journal of Vertebrate Paleontology*, 20(2), 404–407. [https://doi.org/10.1671/0272-4634\(2000\)020\[0404:ANGODD\]2.0.CO;2](https://doi.org/10.1671/0272-4634(2000)020[0404:ANGODD]2.0.CO;2)
- Allain, R., & Pereda-Suberbiola, X. (2003). Dinosaurs of France. *Comptes Rendus Palevol*, 2(1), 27–44. [https://doi.org/10.1016/S1631-0683\(03\)00002-2](https://doi.org/10.1016/S1631-0683(03)00002-2)
- Allain, R., Xaisanavong, T., Richir, P., & Khentavong, B. (2012). The first definitive Asian spinosaurid (Dinosauria: Theropoda) from the early cretaceous of Laos. *Naturwissenschaften*, 99(5), 369–377. <https://doi.org/10.1007/s00114-012-0911-7>

- 1094 Alonso, A., Meléndez, N., & Mas, J. R. (1991). Sedimentación lacustre durante el  
1095 Cretácico en la Cordillera Ibérica, España. *Acta geológica hispánica*, 26, 35–54.
- 1096 Amudeo-Plaza, J., Soto-Acuna, S., Ugalde, R., Martínez, P., & Rubilar-Rogers, D.  
1097 (2023). Reassessment of theropod material from Pichasca, Northern Chile:  
1098 Presence of Abelisauridae (Theropoda: Ceratosauria) from the Quebrada La  
1099 Totorá Beds (Albian-Turonian). *Journal of South American Earth Sciences*, 129,  
1100 104494. <https://doi.org/10.1016/j.jsames.2023.104494>
- 1101 Antunes, M. T., & Sigogneau-Russell, D. (1991). Nouvelles données sur les dinosaures  
1102 du Crétacé supérieur du Portugal. *Comptes Rendus de l'Académie des Sciences*,  
1103 *Paris, II*, 313, 113–119.
- 1104 Antunes, M. T., & Sigogneau-Russell, D. (1992). La faune de petits dinosaures du  
1105 Crétacé terminal portugais. *Comunicações dos Serviços Geológicos de Portugal*,  
1106 78(1), 49–62.
- 1107 Ardèvol, L., Klimowitz, J., Malagón, J., & Nagtegaal, P. J. C. (2000). Depositional  
1108 Sequence Response to Foreland Deformation in the Upper Cretaceous of the  
1109 Southern Pyrenees, Spain. *AAPG Bulletin*, 84(4), 566–587.  
1110 <https://doi.org/10.1306/C9EBCE55-1735-11D7-8645000102C1865D>
- 1111 Astibia, H., Buffetaut, E., Buscalioni, A. D., Cappetta, H., Corral, C., Estes, R., García-  
1112 Garmilla, F., Jaeger, J. J., Jiménez-Fuentes, E., Le Loeuff, J., Mazin, J. M., Orue-  
1113 Etxebarria, X., Pereda-Suberbiola, J., Powell, J. E., Rage, J. C., Rodríguez-  
1114 Lázaro, J., Sanz, J. L., & Tong, H. (1990). The fossil vertebrates from Laño  
1115 (Basque Country, Spain); new evidence on the composition and affinities of the  
1116 Late Cretaceous continental faunas of Europe. *Terra Nova*, 2(5), 460–466.
- 1117 Astibia, H., Murelaga, X., Pereda-Suberbiola, X., Elorza, J. J., & Gómez-Alday, J. J.  
1118 (1999). Taphonomy and palaeoecology of the Upper Cretaceous continental

1119 vertebrate-bearing beds of the Laño quarry (Iberian Peninsula). *Estudios del*  
1120 *Museo de Ciencias naturales de Alava*, 14(Número especial 1), 43–104.

1121 Baiano, M. A., Coria, R. A., Canale, J. I., & Gianechini, F. A. (2021). New abelisaurid  
1122 material from the Anacleto Formation (Campanian, Upper Cretaceous) of  
1123 Patagonia, Argentina, shed light on the diagnosis of the Abelisauridae  
1124 (Theropoda, Ceratosauria). *Journal of South American Earth Sciences*, 110,  
1125 103402. <https://doi.org/10.1016/j.jsames.2021.103402>

1126 Baiano, M. A., Coria, R., Chiappe, L. M., Zurriaguz, V., & Coria, L. (2023). Osteology  
1127 of the axial skeleton of *Aucasaurus garridoi*: phylogenetic and paleobiological  
1128 inferences. *PeerJ*, 11, e16236. <https://doi.org/10.7717/peerj.16236>

1129 Barbosa, B. P., Soares, A. F., Rocha, R. B., Manuppella, G., & Henriques, M. H.  
1130 (2008). Geological map of Portugal at a scale of 1:50.000: explanatory news of  
1131 the sheet 19-A (Cantanhede). Departamento de Geologia, Instituto Nacional de  
1132 Engenharia, Tecnologia e Inovação, Lisboa.

1133 Benson, R. B. (2010). A description of *Megalosaurus bucklandii* (Dinosauria:  
1134 Theropoda) from the Bathonian of the UK and the relationships of Middle  
1135 Jurassic theropods. *Zoological Journal of the Linnean Society*, 158, 882–935.  
1136 <https://doi.org/10.1111/j.1096-3642.2009.00569.x>

1137 Berreteaga, A. (2008). Estudio estratigráfico, sedimentológico y paleontológico de los  
1138 yacimientos con fósiles de vertebrados del Cretácico final de la Región Vasco-  
1139 Cantábrica (Tesis Doctoral, Facultad de Ciencia y Tecnología, Universidad del  
1140 País Vasco, Leioa).

1141 Bonaparte, J. F. (1991). The gondwanian theropod families Abelisauridae and  
1142 Noasauridae. *Historical Biology*, 5(1), 1–25.  
1143 <https://doi.org/10.1080/10292389109380385>



- 1144 Bonaparte, J. F. (1996). Cretaceous tetrapods of Argentina. *Münchner*  
 1145 *Geowissenschaftliche Abhandlungen*, 30(7), 73–130.
- 1146 Bonaparte, J. F., & Novas, F. E. (1985). *Abelisaurus comahuensis* n. g., n. sp.,  
 1147 Carnosauria del Cretacico tardia de Patagonia. *Ameghiniana*, 22, 259–265.
- 1148 Brochu, C. A. (2003). Osteology of *Tyrannosaurus rex*: insights from a nearly complete  
 1149 skeleton and high-resolution computed tomographic analysis of the skull. *Journal*  
 1150 *of Vertebrate Paleontology*, 22(4 suppl.), 1–138.  
 1151 <https://doi.org/10.1080/02724634.2003.10010947>
- 1152 Buffetaut, E. (2005). Late Cretaceous vertebrates from the Saint-Chinian area (southern  
 1153 France): a review of previous research and an update on recent finds. *Acta*  
 1154 *Palaeontologica Romaniae*, 5(3), 39–48.
- 1155 Buffetaut, E. (2024). The discontinuous fossil record of European Abelisauridae and its  
 1156 significance for abelisaurid evolution and palaeobiogeography. *Evolução*, 3, 19–  
 1157 21.
- 1158 Buffetaut, E. (2025). Furileusaurian osteological characters in *Genusaurus sisteronis*  
 1159 Accarie et al., 1995, an abelisaurid dinosaur from the Albian (Lower Cretaceous)  
 1160 of south-eastern France. *Carnets natures*, 12, 79–88.
- 1161 Buffetaut, E., Mechin, P., & Mechin-Salessy, A. (1988). Un dinosaure théropode  
 1162 d'affinités gondwaniennes dans le Crétacé supérieur de Provence. *Comptes*  
 1163 *Rendus de l'Académie des Sciences II*, 306, 153–158.
- 1164 Buffetaut, E., Le Leuff, J., Tong, H., Duffaud, S., Cavin, L., Garcia, G., & Ward, D.  
 1165 (1999). Un nouveau gisement de vertébrés du Crétacé supérieur à Cruzy (Hérault,  
 1166 Sud de la France). *Comptes Rendus de l'Académie des Sciences II*, 328, 203–208.  
 1167 [https://doi.org/10.1016/S1251-8050\(99\)80097-4](https://doi.org/10.1016/S1251-8050(99)80097-4)

1168 Buffetaut, E., Tong, H., Girard, J., Hoyer, B., & Párraga, J. (2024). *Caletodraco*  
 1169 *cottardi*: A New Furileusaurian Abelisaurid (Dinosauria: Theropoda) from the  
 1170 Cenomanian Chalk of Normandy (North-Western France). *Fossil Studies*, 2(3),  
 1171 177–195. <https://doi.org/10.3390/fossils2030009>

1172 Calvo, J. O., Rubilar-Rogers, D., & Moreno, K. (2004). A new abelisauridae  
 1173 (Dinosauria: Theropoda) from northwest Patagonia. *Ameghiniana*, 41(4), 555–  
 1174 563.

1175 Canale, J. I., Scanferla, C. A., Agnolin, F. L., & Novas, F. E. (2009). New carnivorous  
 1176 dinosaur from the Late Cretaceous of NW Patagonia and the evolution of  
 1177 abelisaurid theropods. *Naturwissenschaften*, 96, 409–414.  
 1178 <https://doi.org/10.1007/s00114-008-0487-4>

1179 Candeiro, C. R. A., & Martinelli, A. G. (2005). Abelisauroida and  
 1180 Carchodontosauridae (Theropoda, Dinosauria) in the Cretaceous of South  
 1181 America. Paleogeographical and geocronological implications. *Sociedade &*  
 1182 *Natureza, Uberlândia*, 17, 5–19.

1183 Canudo, J. I., Oms, O., Vila, B., Galobart, À., Fondevilla, V., Puértolas-Pascual, E.,  
 1184 Sellés, A. G., Cruzado-Caballero, P., Dinarès-Turell, J., Vicens, E., Castanera, D.,  
 1185 Company, J., Burrell, L., Estrada, R., Marmi, J., & Blanco, A. (2016). The upper  
 1186 Maastrichtian dinosaur fossil record from the southern Pyrenees and its  
 1187 contribution to the topic of the Cretaceous-Palaeogene mass extinction event.  
 1188 *Cretaceous Research*, 57, 540–551. <https://doi.org/10.1016/j.cretres.2015.06.013>

1189 Carrano, M. T., & Sampson, S. D. (2008). The phylogeny of ceratosauria (Dinosauria:  
 1190 Theropoda). *Journal of Systematic Palaeontology*, 6(2), 183–236.  
 1191 <https://doi.org/10.1017/S1477201907002246>

1192 Carrano, M. T., Sampson, S. D., & Forster, C. A. (2002). The osteology of  
 1193 *Masiakasaurus knopfleri*, a small abelisauroid (Dinosauria: Theropoda) from the  
 1194 Late Cretaceous of Madagascar. *Journal of Vertebrate Paleontology*, 22(3), 510–  
 1195 534. [https://doi.org/10.1671/0272-4634\(2002\)022\[0510:TOOMKA\]2.0.CO;2](https://doi.org/10.1671/0272-4634(2002)022[0510:TOOMKA]2.0.CO;2)

1196 Carrano, M. T., Benson, R. B., & Sampson, S. D. (2012). The phylogeny of tetanurae  
 1197 (Dinosauria: Theropoda). *Journal of Systematic Palaeontology*, 10(2), 211–300.  
 1198 <https://doi.org/10.1080/14772019.2011.630927>

1199 Cerroni, M. A., Canale, J. I., Novas, F. E., & Paulina-Carabajal, A. (2022). An  
 1200 exceptional neurovascular system in abelisaurid theropod skull: new evidence  
 1201 from *Skorpiovenator bustingorryi*. *Journal of Anatomy*, 240(4), 612–626.  
 1202 <https://doi.org/10.1111/joa.13258>

1203 Cincotta, A., Yans, Y., Godefroit, P., Garcia, G., Dejax, J., Benammi, M., Amico, S., &  
 1204 Valentin, X. (2015). Interated paleoenvironmental reconstruction and taphonomy  
 1205 of a unique Upper Cretaceous vertebrate-bearing locality (Velaux, southeastern  
 1206 France). *PLOS ONE*, 10(8), e0134231.  
 1207 <https://doi.org/10.1371/journal.pone.0134231>

1208 Company, J. (2005). Vertebrados continentales del Cretácico superior (Campaniense-  
 1209 Maastrichtiense) de Valencia (Tesis Doctoral, Facultad de Ciencias Biológicas,  
 1210 Universitat de València, Valencia).

1211 Company, J., & Szentesi, Z. (2012). Amphibians from the Late Cretaceous Sierra  
 1212 Perenchiza Formation of the Chera Basin, Valencia Province, Spain. *Cretaceous*  
 1213 *Research*, 37, 240–245. <https://doi.org/10.1016/j.cretres.2012.04.003>

1214 Company, J., Feist, M., Peyrot, D., Barrón, E., Robles, F., Pereda-Suberbiola, X., &  
 1215 Ruiz-Omeñaca, J. I. (2005). Stratigraphic position and palaeoenvironmental traits

1216 of the Late Cretaceous vertebrate-bearing sites of Chera (Valencia, Spain), based  
 1217 on micropalaeontological data. *Kaupia*, 14, 76.

1218 Company, J., Torices, A., Pereda-Suberbiola, X., & Ruiz-Omeñaca, J. (2009). Theropod  
 1219 teeth from the Late Cretaceous of Chera (Valencia, eastern Spain). *Journal of*  
 1220 *Vertebrate Paleontology*, 29, 81A.

1221 Coria, R. A., Chiappe, L. M., & Dingus, L. (2002). A new close relative of *Carnotaurus*  
 1222 *sastrei* Bonaparte 1985 (Theropoda: Abelisauridae) from the Late Cretaceous of  
 1223 Patagonia. *Journal of Vertebrate Paleontology*, 22(2), 460–465.  
 1224 [https://doi.org/10.1671/0272-4634\(2002\)022\[0460:ANCROC\]2.0.CO;2](https://doi.org/10.1671/0272-4634(2002)022[0460:ANCROC]2.0.CO;2)

1225 Corral, J. C., Pueyo, E. L., Berreteaga, A., Rodríguez-Pintó, A., Sánchez, E., & Pereda-  
 1226 Suberbiola, X. (2016). Magnetostratigraphy and lithostratigraphy of the Laño  
 1227 vertebrate-site: Implications in the uppermost Cretaceous chronostratigraphy of  
 1228 the Basque-Cantabrian Region. *Cretaceous Research*, 57, 473–489.  
 1229 <https://doi.org/10.1016/j.cretres.2015.07.015>

1230 Csiki-Sava, Z., Buffetaut, E., Ósi, A., Pereda-Suberbiola, X., & Brusatte, S. L. (2015).  
 1231 Island life in the Cretaceous-faunal composition, biogeography, evolution, and  
 1232 extinction of land-living vertebrates on the Late Cretaceous European archipelago.  
 1233 *ZooKeys*, 469, 1–161. <https://doi.org/10.3897/zookeys.469.8439>

1234 Currie, P. J., & Zhao, X. J. (1993). A new carnosaur (Dinosauria, Theropoda) from the  
 1235 Jurassic of Xinjiang, People's Republic of China. *Canadian Journal of Earth*  
 1236 *Sciences*, 30(10), 2037–2081. <https://doi.org/10.1139/e93-179>

1237 Currie, P. J., Rigby Jr., J. K., & Sloan, R. E. (1990). Theropod teeth from the Judith  
 1238 River Formation of southern Alberta, Canada. In K. Carpenter, & P. J. Currie,  
 1239 (Eds.). *Dinosaur Systematics: perspectives and approaches* 8 (pp. 107–125).  
 1240 Cambridge University Press.

- 1241 Díez-Canseco, D., Arz, J. A., Benito, M. I., Díaz-Molina, M., & Arenillas, I. (2014).  
 1242 Tidal influence in redbeds: A palaeoenvironmental and biochronostratigraphic  
 1243 reconstruction of the Lower Tremp Formation (South-Central Pyrenees, Spain)  
 1244 around the Cretaceous/Paleogene boundary. *Sedimentary Geology*, 312, 31–49.  
 1245 <https://doi.org/10.1016/j.sedgeo.2014.06.008>
- 1246 Eichenseer, H. (1988). Facies Geology of Late Maastrichtian to Early Eocene Coastal  
 1247 and Shallow Marine Sediments (Tremp-Graus Basin, Northeastern Spain) (Tesis  
 1248 Doctoral, Institut und Museum für Geologie und Paläontologie, Universität  
 1249 Tübingen, Tübingen).
- 1250 Evers, S. W., Rauhut, O. W. M., Milner, A. C., McFeeters, B., & Allain, R. (2015). A  
 1251 reappraisal of the morphology and systematic position of the theropod dinosaur  
 1252 *Sigilmassasaurus* from the “middle” Cretaceous of Morocco. *PeerJ*, 3, e1323.  
 1253 <https://doi.org/10.7717/peerj.1323>
- 1254 Filippi, L. S., Méndez, A. H., Gianechini, F. A., Valieri, R. D. J., & Garrido, A. C.  
 1255 (2018). Osteology of *Viavenator exxoni* (Abelisauridae; Furileosauria) from the  
 1256 Bajo de la Carpa Formation, NW Patagonia, Argentina. *Cretaceous Research*, 83,  
 1257 95–119. <https://doi.org/10.1016/j.cretres.2017.07.019>
- 1258 Floquet, M., Alonso, A., & Melendez, A. (1982). Cameros-Castilla. El Cretácico  
 1259 Superior. In A. García (Ed.). *El Cretácico de España*. Editorial Complutense.
- 1260 Fondevilla, V., Dinarès-Turell, J., & Oms, O. (2016). The chronostratigraphic  
 1261 framework of the South-Pyrenean Maastrichtian succession reappraised:  
 1262 Implications for basin development and end Cretaceous dinosaur faunal turnover.  
 1263 *Sedimentary Geology*, 337, 55–68. <https://doi.org/10.1016/j.sedgeo.2016.03.006>
- 1264 Fondevilla, V., Riera, V., Vila, B., Dinarès-Turell, J., Vicens, E., Gaete, R., Oms, O., &  
 1265 Galobart, À. (2019). Chronostratigraphic synthesis of the latest Cretaceous

1266 dinosaur turnover in south-western Europe. *Earth-Science Reviews*, 191, 168–189.  
 1267 <https://doi.org/10.1016/j.earscirev.2019.01.007>

1268 Galton, P. M. (1996). Notes on Dinosauria from the Upper Cretaceous of Portugal.  
 1269 *Neues Jahrbuch für Geologie und Paläontologie Monatshefte*, 2, 83–90.

1270 Garcia, G., Duffaud, S., Feist, M., Marandat, B., Tambareau, Y., Villatte, J., & Sigé, B.  
 1271 (2000). La Neuve, gisement à plantes, invertébrés et vertébrés du Bégudien  
 1272 (Sénonien supérieur continental) du bassin d’Aix-en-Provence. *Geodiversitas*,  
 1273 22(3), 325–348.

1274 Garcia, G., Amico, S., Fournier, F., Thouand, E., & Valentin, X. (2010). A new  
 1275 titanosaur genus (Dinosauria, Sauropoda) from the Late Cretaceous of southern  
 1276 France and its paleobiogeographic implications. *Bulletin de la Société Géologique*  
 1277 *de France*, 181, 269–277. <https://doi.org/10.2113/gssgfbull.181.3.269>

1278 Gianechini, F. A., Apesteguía, S., Landini, W., Finotti, F., Valieri, R. J., & Zandonai, F.  
 1279 (2015). New abelisaurid remains from the Anacleto Formation (Upper  
 1280 Cretaceous), Patagonia, Argentina. *Cretaceous Research*, 54, 1–16.  
 1281 <https://doi.org/10.1016/j.cretres.2014.11.009>

1282 Gil, J., Carenas, B., Segura, M., García-Hidalgo, J. F., & García, A. (2004). Revisión y  
 1283 correlación de las unidades listostratigráficas del Cretácico Superior de la región  
 1284 central y oriental de España. *Revista de la Sociedad Geológica de España*, 17,  
 1285 249–266.

1286 Goloboff, P. A., & Morales, M. E. (2023). TNT version 1.6, with a graphical interface  
 1287 for MacOS and Linux, including new routines in parallel. *Cladistics*, 39, 144–153.  
 1288 <https://doi.org/10.1111/cla.12524>.

1289 Gómez, J. J., Sandoval, J., Aguado, R., O’Dogerthy, L., & Osete, M. L. (2019). The  
 1290 Alpine Cycle in Eastern Iberia: Microplate Units and Geodynamic Stages. In C.

1291 Quesada, & J. T. Oliveira (Eds.). *The Geology of Iberia: A Geodynamic*  
 1292 *Approach, volume 3: The Alpine Cycle. Regional Geology Reviews* (pp. 15–28).  
 1293 Springer.

1294 Gómez-Alday, J. J. (1999). Stratigraphy and depositional environments of the Upper  
 1295 Cretaceous of the Laño quarry. Evidence of diapiric activity. *Estudios del Museo*  
 1296 *de Ciencias Naturales de Álava*, 14(Número especial 1), 29–35.

1297 Hammer, Ø., Harper, D. A. T., & Ryan, P. D. (2001). PAST: paleontological statistics  
 1298 software package for education and data analysis. *Palaeontologia Electronica*,  
 1299 4(1), 1–9.

1300 Hendrickx, C., & Mateus, O. (2014). Abelisauridae (Dinosauria: Theropoda) from the  
 1301 Late Jurassic of Portugal and dentition-based phylogeny as a contribution for the  
 1302 identification of isolated theropod teeth. *Zootaxa*, 3759, 1–74.

1303 Hendrickx, C., Hartman, S. A., & Mateus, O. (2015a). An overview of non-avian  
 1304 theropod discoveries and classification. *PalArch's Journal of Vertebrate*  
 1305 *Palaeontology*, 12(1), 1–73.

1306 Hendrickx, C., Mateus, O., & Araújo, R. (2015b). A proposed terminology of theropod  
 1307 teeth (Dinosauria, Saurischia). *Journal of Vertebrate Paleontology*, 35, e982797.  
 1308 <https://doi.org/10.1080/02724634.2015.982797>

1309 Hendrickx, C., Mateus, O., Araújo, R., & Choiniere, J. (2019). The distribution of  
 1310 dental features in non-avian theropod dinosaurs: Taxonomic potential, degree of  
 1311 homoplasy, and major evolutionary trends. *Palaeontologia Electronica*, 22(3), 74.  
 1312 <https://doi.org/10.26879/820>

1313 Hendrickx, C., Tschopp, E., & d. Ezcurra, M. (2020). Taxonomic identification of  
 1314 isolated theropod teeth: the case of the shed tooth crown associated with

1315        *Aerosteon* (Theropoda: Megaraptora) and the dentition of Abelisauridae.

1316        *Cretaceous Research*, 108, 104312. <https://doi.org/10.1016/j.cretres.2019.104312>

1317 Iori, F. V., Araújo-Júnior, H. I. de, Tavares, S. A. S., Silva Marinho, T. da, & Martinelli,

1318        A. G. (2021). New theropod dinosaur from the Late Cretaceous of Brazil

1319        improves abelisaurid diversity. *Journal of South American Earth Sciences*, 112,

1320        103551. <https://doi.org/10.1016/j.jsames.2021.103551>

1321 Isasmendi, E., Torices, A., Canudo, J. I., & Pereda-Suberbiola, X. (2021). Abelisaurid

1322        dinosaurs from the Upper Cretaceous Laño site (Iberian Peninsula). *Ciências da*

1323        *Terra Procedia*, 1, 38–41. <https://doi.org/10.21695/cterraproc.v1i0.401>

1324 Isasmendi, E., Torices, A., Canudo, J. I., Currie, P. J., & Pereda-Suberbiola, X. (2022).

1325        Upper Cretaceous European theropod palaeobiodiversity, palaeobiogeography and

1326        the intra-Maastrichtian faunal turnover: new contributions from the Iberian fossil

1327        site of Laño. *Papers in Palaeontology*, 8(1), e1419.

1328        <https://doi.org/10.1002/spp2.1419>

1329 Isasmendi, E., Pérez-Pueyo, M., Moreno-Azanza, M., Alonso, A., Puértolas-Pascual, E.,

1330        Bádenas, B., & Canudo, J. I. (2024). Theropod teeth palaeodiversity from the

1331        uppermost Cretaceous of the South Pyrenean Basin (NE Iberia) and the intra-

1332        Maastrichtian faunal turnover. *Cretaceous Research*, 162, 105952.

1333        <https://doi.org/10.1016/j.cretres.2024.105952>

1334 Lapparent, A. F., & Zbyszewski, G. (1957). The dinosaurs from Portugal. *Memoire*

1335        *Services Géologiques du Portugal*, 2(nouvelle série), 1–131.

1336 Laurent, Y. (2003). Les faunes de vertébrés continentaux du Maastrichtien supérieur

1337        d'Europe: systématique et biodiversité. *Strata, série 2*, 41, 1–81.



1338 Laurent, Y., Bilotte, M., & Le Loeuff, J. (2002). Late Maastrichtian continental  
1339 vertebrates from southwestern France: correlation with marine fauna.  
1340 *Palaeogeography, Palaeoclimatology, Palaeoecology*, 187(1–2), 121–135.

1341 Lavocat, R. (1955). Sur une portion de mandibule de Théropode provenant du Crétacé  
1342 supérieur de Madagascar. *Bulletin du Muséum National d'Histoire Naturelle de*  
1343 *Paris*, 27(2), 256–259.

1344 Le Loeuff, J., & Buffetaut, E. (1991). *Tarascosaurus salluvicus* nov. gen., nov. sp.,  
1345 dinosaure théropode du Crétacé supérieur du Sud de la France. *Geobios*, 24(5),  
1346 585–594.

1347 Le Loeuff, J. (1992). *Les vertébrés continentaux du Crétacé supérieur d'Europe:*  
1348 *paléoécologie, biostratigraphie et paléobiogéographie* (Tesis Doctoral, Université  
1349 Pierre & Marie Curie, Paris, Paris).

1350 Longrich, N. R., Pereda-Suberbiola, X., Jalil, N. E., Khaldoune, F., & Jourani, E.  
1351 (2017). An abelisaurid from the latest Cretaceous (late Maastrichtian) of Morocco,  
1352 North Africa. *Cretaceous Research*, 76, 40–52.  
1353 <https://doi.org/10.1016/j.cretres.2017.03.021>

1354 Maddison, W. P., & Maddison, D. R. (2011). *Mesquite: a modular system for*  
1355 *evolutionary analysis*. Retrieved May 23, 2012, from <http://mesquiteproject.org>.

1356 Madsen, J. H. Jr (1976). *Allosaurus fragilis*: a revised osteology. *Utah Geological*  
1357 *Survey Bulletin*, 109, 1–163.

1358 Madsen, J. H. Jr, & Welles, S. P. (2000). *Ceratosaurus* (Dinosauria, Theropoda), a  
1359 revised osteology. *Utah Geological Survey, Miscellaneous Publication*, 2, 1–80.

1360 Malafaia, E., Mocho, P., Escaso, F. & Ortega, F. (2020). A new carcharodontosaurian  
1361 theropod from the Lusitanian Basin: evidence of allosauroid sympatry in the

1362 European Late Jurassic. *Journal of Vertebrate Paleontology*, 40(1), e1768106.  
 1363 <https://doi.org/10.1080/02724634.2020.1768106>

1364 Malafaia, E., Escaso, F., Coria, R. A., & Ortega, F. (2023). An Eudromaeosaurian  
 1365 Theropod from Lo Hueco (Upper Cretaceous. Central Spain). *Diversity*, 15(2),  
 1366 141. <https://doi.org/10.3390/d15020141>

1367 Malafaia, E., Mocho, P., Escaso, F., Narvaéz, I., & Ortega, F. (2024). Taxonomic and  
 1368 stratigraphic update of the material historically attributed to *Megalosaurus* from  
 1369 Portugal. *Palaeontologia Polonica*, 69(2), 127–171.  
 1370 <https://doi.org/10.4202/app.01113.2023>

1371 Malafaia, E., Escaso, F., Coria, R. A., Pérez-García, A., & Ortega, F. (2025). Theropod  
 1372 teeth from the Upper Cretaceous of central Spain: Assessing the  
 1373 paleobiogeographic history of European abelisaurids. *Cretaceous Research*, 168,  
 1374 106072. <https://doi.org/10.1016/j.cretres.2024.106072>

1375 Marmi, J., Blanco, A., Fondevilla, V., Dalla Vecchia, F. M., Sellés, A. G., Vicente, A.,  
 1376 Martín-Closas, C., Oms, O., & Galobart, A. (2016). The Molí del Baró-1 site, a  
 1377 diverse fossil assemblage from the uppermost Maastrichtian of the southern  
 1378 Pyrenees (north-eastern Iberia). *Cretaceous Research*, 57, 519–539.  
 1379 <https://doi.org/10.1016/j.cretres.2015.06.016>

1380 Marsh, O. C. (1881). Principal characters of American Jurassic dinosaurs. Part V.  
 1381 *American Journal of Science*, 21(Series 3), 417–423.

1382 Marsh, O. C. (1884). Principal characters of the American Jurassic dinosaurs. Part VIII.  
 1383 The order Theropoda. *American Journal of Science*, 27, 329–340.

1384 Martín-Chivelet, J., Floquet, M., García-Senz, J., Callapez, P. M., Lopez-Mir, B.,  
 1385 Muñoz, J. A., Barroso-Barcenilla, F., Segura, M., Ferreira Soares, A., Morgado  
 1386 Dinis, P., Fonseca Marques, J., & Arbues, P. (2019). Late Cretaceous post-rift to

1387 convergence in Iberia. In C. Quesada, & J. T. Oliveira (Eds.). *The geology of*  
1388 *Iberia: A geodynamic approach, Vol. 3, The Alpine Cycle*. Springer.

1389 Méndez, A. H. (2014). The caudal vertebral series in abelisaurid dinosaurs. *Acta*  
1390 *Palaeontologica Polonica*, 59(1), 99–107. <https://doi.org/10.4202/app.2012.0095>

1391 Méndez, A. H., Gianechini, F. A., Paulina-Carabajal, A., Filippi, L. S., Juárez-Valieri,  
1392 R. D., Cerda, I. A., & Garrido, A. C. (2022). New furileusaurian remains from La  
1393 Invernada (northern Patagonia, Argentina): A site of unusual abelisaurids  
1394 abundance. *Cretaceous Research*, 129, 104989.  
1395 <https://doi.org/10.1016/j.cretres.2021.10498>

1396 Meso, J. G., Hendrickx, C., Baiano, M. A., Canale, J. I., Salgado, L., & Díaz Martínez,  
1397 I. (2021). Isolated theropod teeth associated with a sauropod skeleton from the  
1398 Late Cretaceous Allen Formation of Río Negro, Patagonia, Argentina. *Acta*  
1399 *Palaeontologica Polonica*, 66(2), 409–423.  
1400 <https://doi.org/10.4202/app.00847.2020>

1401 Mey, P. H. W., Nagtegaal, P. J. C., Roberti, K. J., & Hartevelt, J. J. A. (1968).  
1402 Lithostratigraphic subdivision of post-Hercynian deposits in the south-central  
1403 Pyrenees, Spain. *Leidse Geologische Mededelingen*, 41, 221–228.

1404 Mutti, E., & Sgavetti, M. (1987). Sequence stratigraphy of the Upper Cretaceous Aren  
1405 strata in the Aren-Orcau region, south-central Pyrenees, Spain: Distinction  
1406 between eustatically and tectonically controlled depositional sequences. *Annali*  
1407 *dell'Università di Ferrara*, 1, 1–22.

1408 Nagtegaal, P. J. C., Van Vliet, A., & Brouwer, J. (1983). Syntectonic coastal offlap and  
1409 concurrent turbidite deposition: The Upper Cretaceous Aren sandstone in the  
1410 South-Central Pyrenees, Spain. *Sedimentary Geology*, 34, 185–218. [https://](https://doi.org/10.1016/0037-0738(83)90086-6)  
1411 [doi.org/10.1016/0037-0738\(83\)90086-6](https://doi.org/10.1016/0037-0738(83)90086-6).

1412 Novas, F. E. (1991). Relaciones filogeneticas de los dinosaurios teropodos  
 1413 ceratosaurios. *Ameghiniana*, 28(3–4), 410.  
 1414 Novas, F. E., Chatterjee, S., Rudra, D. K., & Datta, P. M. (2010). *Rahiolisaurus*  
 1415 *gujaratensis*, n. gen. n. sp., a new abelisaurid theropod from the Late Cretaceous  
 1416 of India. In S. Bandyopadhyay (Ed.). *New Aspects of Mesozoic Biodiversity* (pp.  
 1417 45–62). Springer.  
 1418 Novas, F. E., Agnolín, F. L., Ezcurra, M. D., Porfiri, J., & Canale, J. I. (2013).  
 1419 Evolution of the carnivorous dinosaurs during the Cretaceous: the evidence from  
 1420 Patagonia. *Cretaceous Research*, 45, 174–215.  
 1421 <https://doi.org/10.1016/j.cretres.2013.04.001>  
 1422 O'Connor, P. M. (2007). The postcranial axial skeleton of *Majungasaurus*  
 1423 *crenatissimus* (Theropoda: Abelisauridae) from the Late Cretaceous of  
 1424 Madagascar. *Journal of Vertebrate Paleontology*, 27(S2), 127–163.  
 1425 [https://doi.org/10.1671/0272-4634\(2007\)27\[127:TPASOM\]2.0.CO;2](https://doi.org/10.1671/0272-4634(2007)27[127:TPASOM]2.0.CO;2)  
 1426 Ortega, F., & Pérez-García, A. (2009). cf. *Lirainosaurus* sp. (Dinosauria: Titanosauria)  
 1427 en el Cretácico Superior de Sacedón (Guadalajara). *Geogaceta*, 46, 87–90.  
 1428 Ortega, F., Bardet, N., Barroso-Barcenilla, F., Callapez, P. M., Cambra-Moo, O.,  
 1429 Daviero-Gomez, V., Díez Díaz, V., Domingo, L., Elvira, A., Escaso, F., García-  
 1430 Oliva, M., Gomez, B., Houssaye, A., Knoll, F., Marcos-Fernandez, F., Martín, M.,  
 1431 Mocho, P., Narvaez, I., Perez-García, A., Peyrot, D., Segura, M., Serrano, H.,  
 1432 Torices, A., Vidal, D., & Sanz, J. L. (2015). The biota of the Upper Cretaceous  
 1433 site of Lo Hueco (Cuenca, Spain). *Journal of Iberian Geology*, 41, 83–99.  
 1434 [https://doi.org/10.5209/rev\\_JIGE.2015.v41.n1.48657](https://doi.org/10.5209/rev_JIGE.2015.v41.n1.48657)  
 1435 Ortega, F., Escaso, F., Mocho, P., Narváez, I., & Pérez-García, A. (2019). Eggs and  
 1436 bones: a preliminary comparison between the Upper Cretaceous faunas of the

1437 Poyos, Portilla and Lo Hueco sites (Villalba de la Sierra Formation. Central  
 1438 Spain). *X Congreso Latinoamericano de Paleontología* (p. 106). El Salvador.  
 1439 Ortega, F., Malafaia, E., Escaso, F., & Coria, R. A. (2022). New material of theropods  
 1440 (Abelisauroidea?) from Lo Hueco (Late Cretaceous. Cuenca, Central Spain).  
 1441 *PalaeoVertebrata, special vol. 1*, 147. <https://doi.org/10.18563/pv.eavp2022>  
 1442 Ósi, A., & Buffetaut, E. (2011). Additional non-avian theropod and bird remains from  
 1443 the early Late Cretaceous (Santonian) of Hungary and a review of the European  
 1444 abelisauroid record. *Annales de Paléontologie*, 97(1–2), 35–49.  
 1445 <https://doi.org/10.1016/j.annpal.2011.07.001>  
 1446 Ósi, A., Apesteguía, S., & Kowalewski, M. (2010). Non-avian theropod dinosaurs from  
 1447 the early Late Cretaceous of Central Europe. *Cretaceous Research*, 31(3), 304–  
 1448 320. <https://doi.org/10.1016/j.cretres.2010.01.001>  
 1449 Pereda-Suberbiola, X., & Sanz, J. L. (1999). The ornithomimid dinosaur *Rhabdodon* from  
 1450 the Upper Cretaceous of Lano (Iberian peninsula). *Museo de Ciencias naturales*  
 1451 *de Alava*, 14(Número especial 1), 257–272.  
 1452 Pereda-Suberbiola, X., Astibia, H., Murelaga, X., Elorza, J. J., & Gómez-Alday, J. J.  
 1453 (2000). Taphonomy of the Late Cretaceous dinosaur-bearing beds of the Laño  
 1454 Quarry (Iberian Peninsula). *Palaeogeography, Palaeoclimatology,*  
 1455 *Palaeoecology*, 157(3–4), 247–275. [https://doi.org/10.1016/S0031-](https://doi.org/10.1016/S0031-0182(99)00169-8)  
 1456 [0182\(99\)00169-8](https://doi.org/10.1016/S0031-0182(99)00169-8)  
 1457 Pereda-Suberbiola, X., Corral, J. C., Astibia, H., Badiola, A., Bardet, N., Berreteaga, A.,  
 1458 Buffetaut, E., Buscalioni, A. D., Cappetta, H., Cavin, L., Díez Díaz, V.,  
 1459 Gheerbrant, E., Murelaga, X., Ortega, F., Pérez-García, A., Poyato-Ariza, F.,  
 1460 Rage, J. C., Sanz, J. L., & Torices, A. (2015). Late cretaceous continental and  
 1461 marine vertebrate assemblages from the Laño quarry (Basque-Cantabrian Region,

1462 Iberian Peninsula): an update. *Journal of Iberian Geology*, 41, 101–124.  
 1463 [https://doi.org/10.5209/rev\\_JIGE.2015.v41.n1.48658](https://doi.org/10.5209/rev_JIGE.2015.v41.n1.48658)

1464 Pérez-García, A., Ortega, F., Bolet, A., Escaso, F., Houssaye, A., Martínez-Salanova, J.,  
 1465 de Miguel Chaves, C., Mocho, P., Narvaez, I., Segura, M., Torices, A., Vidal, D.,  
 1466 & Sanz, J. L. (2016). A review of the upper Campanian vertebrate site of Armuña  
 1467 (Segovia Province, Spain). *Cretaceous Research*, 57, 591–623.  
 1468 <https://doi.org/10.1016/j.cretres.2015.08.008>

1469 Pérez-García, A., Gascó, F., & Ortega, F. (2019). The Upper Cretaceous Poyos site: a  
 1470 large dinosaur nesting area in Central Spain. *X Congreso Latinoamericano de*  
 1471 *Paleontología* (p. 108). El Salvador.

1472 Pérez-García, A., Bardet, N., Fregenal-Martínez, M. A., Martín-Jiménez, M., Mocho,  
 1473 P., Narváez, I., Torices, A., Vullo, R., & Ortega, F. (2020). Cenomanian  
 1474 vertebrates from Algora (central Spain): New data on the establishment of the  
 1475 European Upper Cretaceous continental faunas. *Cretaceous Research*, 115,  
 1476 104566. <https://doi.org/10.1016/j.cretres.2020.104566>

1477 Pérez-Pueyo, M., Cruzado-Caballero, P., Moreno-Azanza, M., Vila, B., Castanera, D.,  
 1478 Gasca, J. M., Puértolas-Pascual, E., Bádenas, B., & Canudo, J. I. (2021). The  
 1479 Tetrapod Fossil Record from the Uppermost Maastrichtian of the Ibero-  
 1480 Armorican Island: An Integrative Review Based on the Outcrops of the Western  
 1481 Tremp Syncline (Aragón, Huesca Province, NE Spain). *Geosciences*, 11(4), 1–  
 1482 162. <https://doi.org/10.3390/geosciences11040162>

1483 Platt, N. H. (1989). Lacustrine carbonates and pedogenesis: sedimentology and origin of  
 1484 palustrine deposits from the Early Cretaceous Rupelo Formation, W Cameros  
 1485 Basin, N Spain. *Sedimentology*, 36(4), 665–684. [https://doi.org/10.1111/j.1365-](https://doi.org/10.1111/j.1365-3091.1989.tb02092.x)  
 1486 [3091.1989.tb02092.x](https://doi.org/10.1111/j.1365-3091.1989.tb02092.x)

- 1487 Platt, N. H., & Wright, V. P. (1992). Palustrine carbonates and the Florida Everglades;  
 1488 towards an exposure index for the fresh-water environment?. *Journal of*  
 1489 *Sedimentary Research*, 62(6), 1058–1071. [https://doi.org/10.1306/D4267A4B-](https://doi.org/10.1306/D4267A4B-2B26-11D7-8648000102C1865D)  
 1490 2B26-11D7-8648000102C1865D
- 1491 Peyrot, D., Barron, E., Pereda-Suberbiola, X., & Company, J. (2020). Vegetational  
 1492 composition of the Upper Cretaceous vertebrate site of Chera (Valencia, Spain)  
 1493 and its significance in mosaic vegetation from southwestern Europe. *Cretaceous*  
 1494 *Research*, 106, 104254. <https://doi.org/10.1016/j.cretres.2019.104254>
- 1495 Pol, D., & Rauhut, O. W. M. (2012). A Middle Jurassic abelisaurid from Patagonia and  
 1496 the early diversification of theropod dinosaurs. *Proceedings of the Royal Society*  
 1497 *B*, 279(1741), 3170–3175. <https://doi.org/10.1098/rspb.2012.0660>
- 1498 Puértolas-Pascual, E., Arenillas, I., Arz, J. A., Calvín, P., Ezquerro, L., García-Vicente,  
 1499 C., Pérez-Pueyo, M., Sánchez-Moreno, E. M., Villalaín, J. I., & Canudo, J. I.  
 1500 (2018). Chronostratigraphy and new vertebrate sites from the upper Maastrichtian  
 1501 of Huesca (Spain), and their relation with the K/Pg boundary. *Cretaceous*  
 1502 *Research*, 89, 36–59. <https://doi.org/10.1016/j.cretres.2018.02.016>
- 1503 Rauhut, O. W. M. (2003). The interrelationships and evolution of basal theropod  
 1504 dinosaurs. *Special Papers in Palaeontology*, 69, 1–213.
- 1505 Richter, U., Mudroch, A., & Buckley, L. G. (2013). Isolated theropod teeth from the  
 1506 Kem Kem beds (early Cenomanian) near Taouz, Morocco. *Paläontologische*  
 1507 *Zeitschrift*, 87, 291–309. <https://doi.org/10.1007/s12542-012-0153-1>
- 1508 Rosell, J., Linares, R., & Llompart, C. (2001). El “Garumniense” prepirenaico. *Revista*  
 1509 *de la Sociedad Geológica de España*, 14, 47–56.

1510 Ruiz-Omeñaca, J. I., Vullo, R., Bernárdez, E., & Buscaloni, Á. D. (2009). El primer  
 1511 resto directo de terópodo del Cenomaniense de la Península Ibérica: el diente de  
 1512 Limanes (Oviedo, Asturias). *Geogaceta*, 47(3-4), 29–32.

1513 Sales, M. A., Lacerda, M. B., Horn, B. L., de Oliveira, I. A., & Schultz, C. L. (2016).  
 1514 The “ $\chi$ ” of the matter: testing the relationship between paleoenvironments and  
 1515 three theropod clades. *PLOS ONE*, 11(2), e0147031.  
 1516 <https://doi.org/10.1371/journal.pone.0147031>

1517 Santos Brilhante, N., de França, T. C., Castro, F., Sanches da Costa, L., Currie, P. J.,  
 1518 Kugland de Azevedo, S. A., & Delcourt, R. (2022). A dromaeosaurid-like claw  
 1519 from the Upper Cretaceous of southern France. *Historical Biology*, 34(11), 2195–  
 1520 2204. <https://doi.org/10.1080/08912963.2021.2007243>

1521 Sauvage, H. E. (1897–1898). Fossil vertebrates from Portugal. Contributions to the  
 1522 study of fish and reptiles from the Jurassic and Cretacic. *Direction des Travaux*  
 1523 *Géologiques du Portugal*, 29, 1–46.

1524 Serrano-Martínez, A., Ortega, F., Sciscio, L., Tent-Manclús, J. E., Fierro Bandera, I., &  
 1525 Knoll, F. (2015). New theropod remains from the Tiouraren Formation (?Middle  
 1526 Jurassic, Niger) and their bearing on the dental evolution in basal tetanurans.  
 1527 *Proceedings of the Geologists' Association*, 126(1), 107–118.  
 1528 <https://doi.org/10.4202/app.00101.2014>.

1529 Smith, J. B. (2007). Dental morphology and variation in *Majungasaurus crenatissimus*  
 1530 (Theropoda: Abelisauridae) from the Late Cretaceous of Madagascar. *Society of*  
 1531 *Vertebrate Paleontology Memoir*, 8, 103–126.

1532 Smith, J. B., & Dodson, P. (2003). A proposal for a standard terminology of anatomical  
 1533 notation and orientation in fossil vertebrate dentitions. *Journal of Vertebrate*



1534 *paleontology*, 23(1), 1–12. <https://doi.org/10.1671/0272->  
1535 4634(2003)23[1:APFAST]2.0.CO;2

1536 Smith, J. B., Vann, D. R., & Dodson, P. (2005). Dental morphology and variation in  
1537 theropod dinosaurs: implications for the taxonomic identification of isolated teeth.  
1538 *The Anatomical Record Part A: Discoveries in Molecular, Cellular, and*  
1539 *Evolutionary Biology: An Official Publication of the American Association of*  
1540 *Anatomists*, 285(2), 699–736. <https://doi.org/10.1002/ar.a.20206>

1541 Soto, M., Delcourt, R., Langer, M. C., & Perea, D. (2022). The first record of  
1542 Abelisauridae (Theropoda: Ceratosauria) from Uruguay (Late Jurassic,  
1543 Tacuarembó Formation). *Historical Biology*, 35(12), 2362–2371.  
1544 <https://doi.org/10.1080/08912963.2022.2140425>

1545 Torices Hernández, A. (2002). Los dinosaurios terópodos del Cretácico Superior de la  
1546 Cuenca de Tremp (Pirineos Sur-Centrales, Lleida). *Coloquios de Paleontología*,  
1547 53, 139–146.

1548 Torices, A., Barroso-Barcenilla, F., Cambra-Moo, O., Pérez-García, A., & Segura, M.  
1549 (2012). Palaeontological and palaeobiogeographical implications of the new  
1550 Cenomanian vertebrate site of Algora, Guadalajara, Spain. *Cretaceous Research*,  
1551 37, 231–239. <https://doi.org/10.1016/j.cretres.2012.04.004>

1552 Torices, A., Currie, P. J., Canudo, J. I., & Pereda-Suberbiola, X. (2015). Theropod  
1553 dinosaurs from the upper cretaceous of the south pyrenees basin of Spain. *Acta*  
1554 *Palaeontologica Polonica*, 60(3), 611–626. <https://doi.org/10.4202/app.2012.0121>

1555 Tortosa, T., Buffetaut, E., Vialle, N., Dutour, Y., Turini, E., & Cheylan, G. (2014). A  
1556 new abelisaurid dinosaur from the Late Cretaceous of southern France:  
1557 Palaeobiogeographical implications. *Annales de Paleontologie*, 100(1), 63–86.  
1558 <https://doi.org/10.1016/j.annpal.2013.10.003>

- 1559 Valentin, X., Godefroit, P., Tabuce, R., Vianey-Liaud, M., Wu, W., & García, G.,  
 1560 (2012). First late Maastrichtian (latest Cretaceous) vertebrate assemblage from  
 1561 Provence (Vitrolles-la-Plaine, southern France). In P. Godefroit, (Ed.). *Bernissart*  
 1562 *Dinosaurs and Early Cretaceous Terrestrial Ecosystems* (pp. 582–597). Indiana  
 1563 University Press.
- 1564 Vicente, A., Villalba Breva, S., Ferràndez-Cañadell, C., & Martín-Closas, C. (2014).  
 1565 Revision of the Maastrichtian-Palaeocene charophyte biostratigraphy of the  
 1566 Fontllonga reference section (southern Pyrenees, Catalonia, Spain). *Geologica*  
 1567 *Acta*, 14(4), 349–362. <https://doi.org/10.1344/GeologicaActa2016.14.4.2>
- 1568 Vicente, A., Martín-Closas, C., Arz, J. A., & Oms, O. (2015). Maastrichtian-basal  
 1569 Paleocene charophyte biozonation and its calibration to the Global Polarity Time  
 1570 Scale in the southern Pyrenees (Catalonia, Spain). *Cretaceous Research*, 52(A),  
 1571 268–285. <https://doi.org/10.1016/j.cretres.2014.10.004>
- 1572 Vullo, R., Neraudeau, D., & Lenglet, T. (2007). Dinosaur teeth from the Cenomanian of  
 1573 Charentes, western France: evidence for a mixed Laurasian-Gondwanan  
 1574 assemblage. *Journal of Vertebrate Paleontology*, 27(4), 931–943.  
 1575 [https://doi.org/10.1671/0272-4634\(2007\)27\[931:DTFTCO\]2.0.CO;2](https://doi.org/10.1671/0272-4634(2007)27[931:DTFTCO]2.0.CO;2)
- 1576 Wilson, J. A., Sereno, P. C., Srivastava, S., Bhatt, D. K., Khosla, A., & Sahni, A.  
 1577 (2003). A new abelisaurid (Dinosauria Theropoda) from the Lameta Formation  
 1578 (Cretaceous, Maastrichtian) of India. *Contributions from the Museum of*  
 1579 *Paleontology. University of Michigan*, 31(1), 1–42.
- 1580
- 1581 **Appendices**
- 1582 **Supplementary material 1.** Measurements of the studied teeth, datasets, and DA  
 1583 results.

1584 **Supplementary material 2.** Nexus file of the dental matrix.

1585 **Supplementary material 3.** Abelisauridae and possible Abelisauridae occurrences and  
1586 locality list for the Upper Cretaceous of the Ibero-Armorican Domain.

1587 **Figure captions**

1588 **Figure 1.** Geologic and chronostratigraphic setting of the uppermost Cretaceous fossil  
1589 sites where the studied remains were recovered. **1**, map of the Iberian Mesozoic basins  
1590 showing the locations of the Chera 2, Laño, Montrebei, and Viso fossil sites (modified  
1591 from Gómez et al., 2019); and **2**, chronostratigraphic position of the Chera 2, Laño,  
1592 Montrebei and Viso fossil sites.

1593 **Figure 2.** Abelisauridae indet. teeth from Montrebei and Viso. Morphotype 1 (**DPM-**  
1594 **MON-T10 1**) **1**, mesial denticles; crown in **2**, mesial; **3**, lingual; **4**, distal; and **5**, labial  
1595 views; **6**, close up of enamel texture; and **7**, basal cross section in basal view.

1596 Morphotype 2 (**MG 73**) tooth in **8**, mesial; **9**, lingual; **10**, distal; **11**, labial; and **12**, basal  
1597 views (modified from Malafaia et al., 2024). Abbreviations: **cv**, cervix; **dt**, denticle; **mc**,  
1598 mesial carina; **dc**, distal carina; **dl**, dentine layer; **iet**, irregular enamel texture; **pc**, pulp  
1599 cavity; **tu**; transverse undulation. Scale bar equals 1 cm, except for denticles and texture  
1600 (5 mm).

1601 **Figure 3.** cf. *Arcovenator* axial elements from Laño. Anterior dorsal vertebra (**MCNA**  
1602 8366) in **1**, posterior; **2**, lateral; **3**, anterior; and **4**; dorsal views. Caudal vertebra  
1603 (**MCNA 17433**) in **5**, ventral; **6**, anterior; **7**, lateral; and **8**, posterior views.

1604 Abbreviations: **cdl**, centrodiapophyseal lamina; **fch**, facet for chevron; **ns**, neural spine;  
1605 **poz**, postzygapophysis; **tp**, transverse process; **vg**, ventral groove. Scale bar equals 5  
1606 cm.

1607 **Figure 4.** *Arcovenator* sp. tooth from Chera 2 (**CH 168**). **1**, mesial denticles; crown in  
1608 **2**, mesial; **3**, lingual; **4**, distal; **5**, labial; and **6**, basal views. **7**, close up of enamel

1609 texture; and **8**, distal denticles. Abbreviations: **dt**, denticle; **mc**, mesial carina; **dc**, distal  
1610 carina; **dl**, dentine layer; **iet**, irregular enamel texture; **pc**, pulp cavity. Scale bar equals 1  
1611 cm, except for denticles and texture (5 mm).

1612 **Figure 5.** Graphical results of the LDAs (LDA 1 and 3) carried out at the clade-level. **1**,  
1613 LDA performed using database 1 at clade-level. Axis 1 accounts for 47.67 % of the  
1614 variance (Eigenvalue = 2.6735), and Axis 2 accounts for 24.67 % of the variation  
1615 (Eigenvalue = 1.3703). **2**, LDA performed using database 2 at the clade-level. Axis 1  
1616 accounts for 55.62 % of the variance (Eigenvalue = 2.6056), and Axis 2 for 30.06 % of  
1617 the variation (Eigenvalue = 1.4081). Silhouettes courtesy of Scott Hartman.

1618 **Figure 6.** Graphical results of the LDA performed with the Ibero-Armorican abelisaurid  
1619 teeth (database 3). Axis 1 accounts for 49.29 % of the variance (Eigenvalue = 2.8359),  
1620 and Axis 2 for 26.72 % of the variation (Eigenvalue = 1.5371).

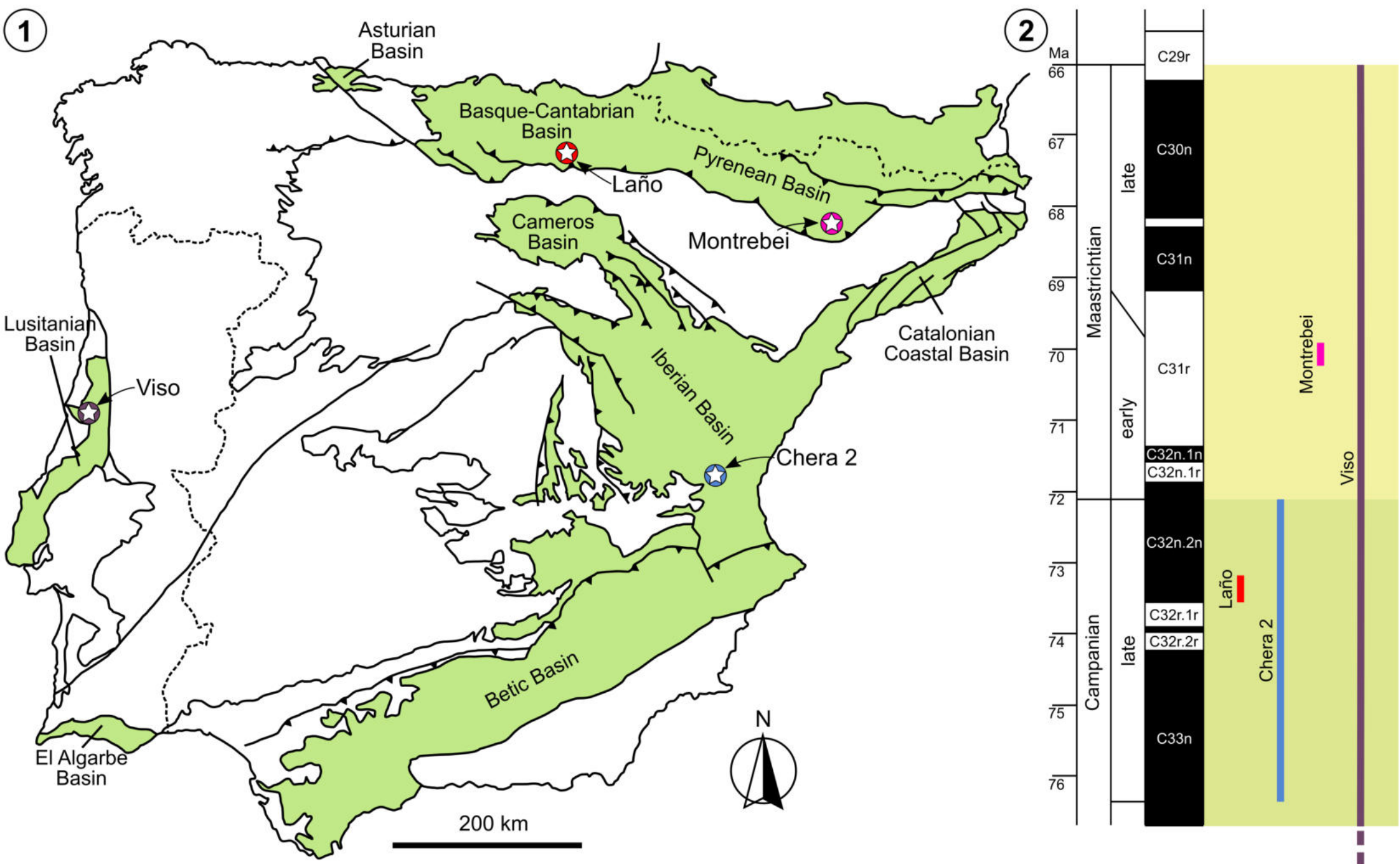
1621 **Figure 7.** Results of the phylogenetic analyses performed for specimens **MG** 73 and  
1622 **CH** 168 based on the dentition-based data matrix (constrained search) of Hendrickx et  
1623 al. (2020). **1**, simplified single MPT (CI = 0.203; RI = 0.542) obtained from the  
1624 phylogenetic analysis, showing the position of specimen **MG** 73 from Viso (highlighted  
1625 in bold). **2**, simplified consensus tree (CI = 0.199; RI = 0.452) obtained from five MPTs  
1626 recovered in the phylogenetic analysis, showing the position of specimen **CH** 168 from  
1627 Chera 2 (highlighted in bold).

1628 **Figure 8.** Simplified strict consensus tree (CI = 0.1809; RI = 0.376) obtained from  
1629 seven MPTs recovered in the phylogenetic analysis based on the dentition-based data  
1630 matrix (constrained search) of Hendrickx et al. (2020), showing the position of **DPM-**  
1631 **MONT-T1** 1 from Montrebei (highlighted in bold).

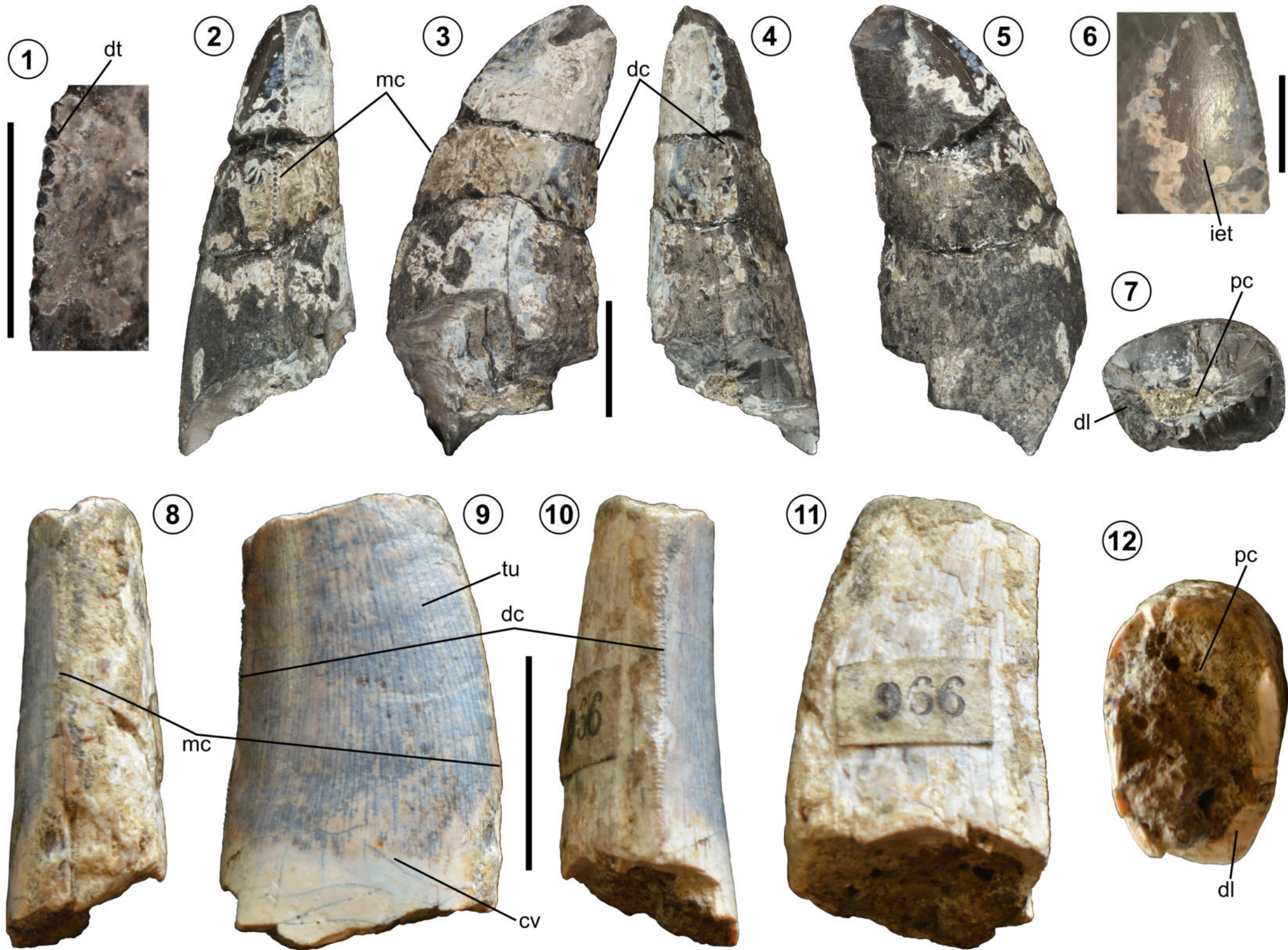
1632 **Figure 9.** Paleogeographic and temporal distribution of the currently known Ibero-  
1633 Armorican abelisaurids and large-sized theropod remains that likely belong to

1634   Abelisauridae. **1**, map of the Ibero-Armorican Mesozoic basins (modified from  
1635   Buffetaut et al., 2024; Malafaia et al., 2025) showing the location where Late  
1636   Cretaceous Ibero-Armorican abelisaurid or large-sized theropod remains have been  
1637   recovered. **2**, chronological distribution of the Late Cretaceous abelisaurids or large-  
1638   sized theropods that may be identified as Abelisauridae (modified from Tortosa et al.,  
1639   2014). Abelisaurid- or large-size theropod-bearing sites or areas: 1, Saint-Jouin-  
1640   Bruneval; 2, “La Buzinie”; 3, Limanes; 4, Algora; 5, Viso; 6, Lambeau du Beausset; 7,  
1641   Trets-La Boucharde; 8, Velaux-La Bastide Neuve; 9, Fox-Amphoux; 10, Pourrières-Jas  
1642   Neuf Sud; 11, Laño; 12, Armuña; 13, Chera 2; 14, Cruzy; 15, Pourcieux-Les Tuillières;  
1643   16, Cassagnau 1; 17, Lo Hueco; 18, Poyos; 19, Vitrolles-La Plaine; 20, Montrebei; 21,  
1644   Blasi; 22, 172-i/04/e.

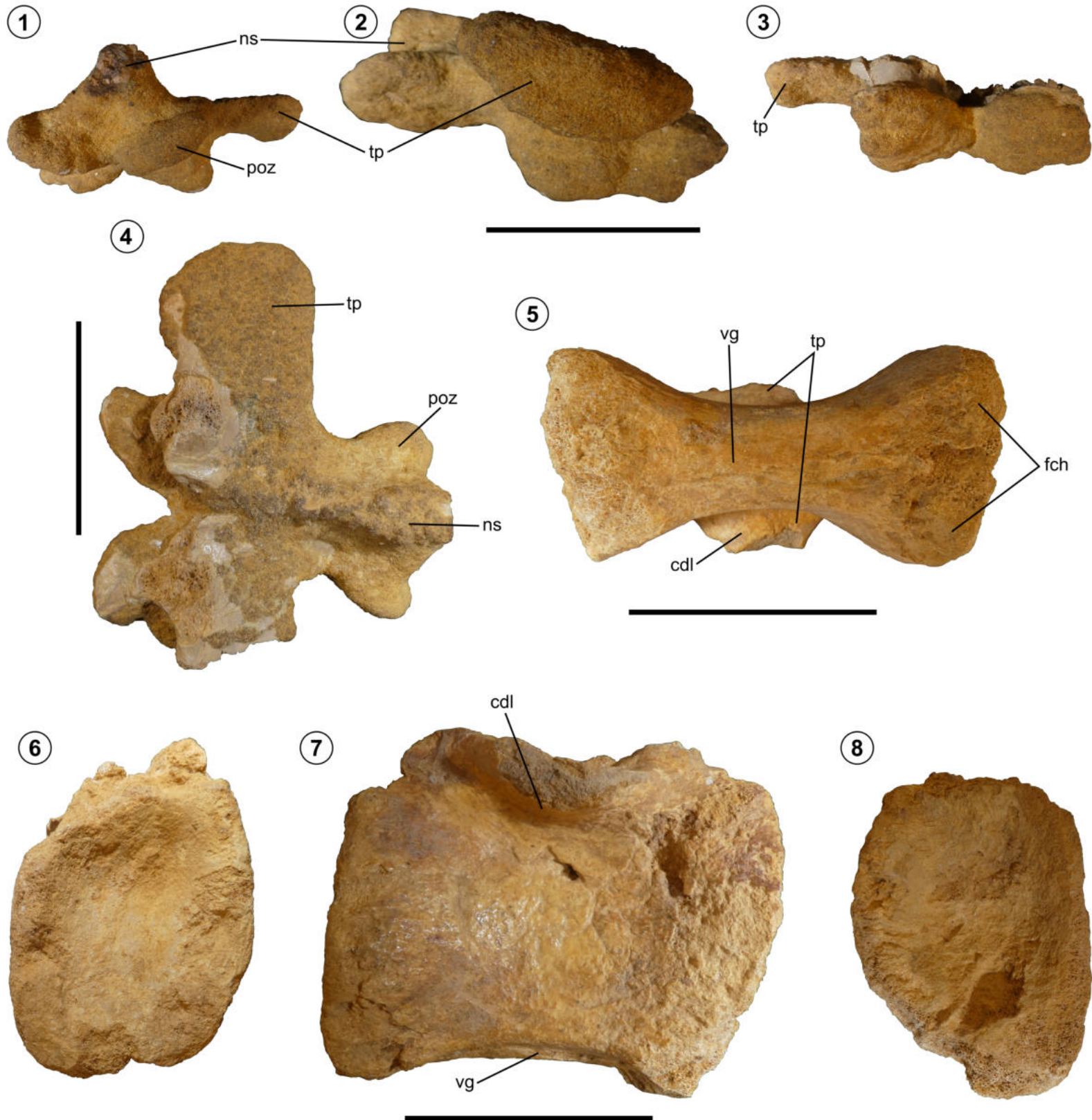
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①

dt



②

mc



③

dc



④

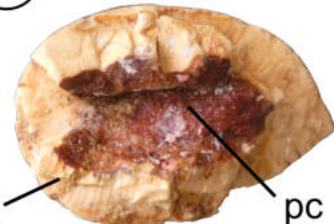


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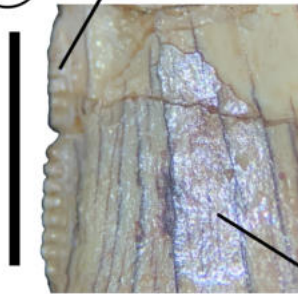
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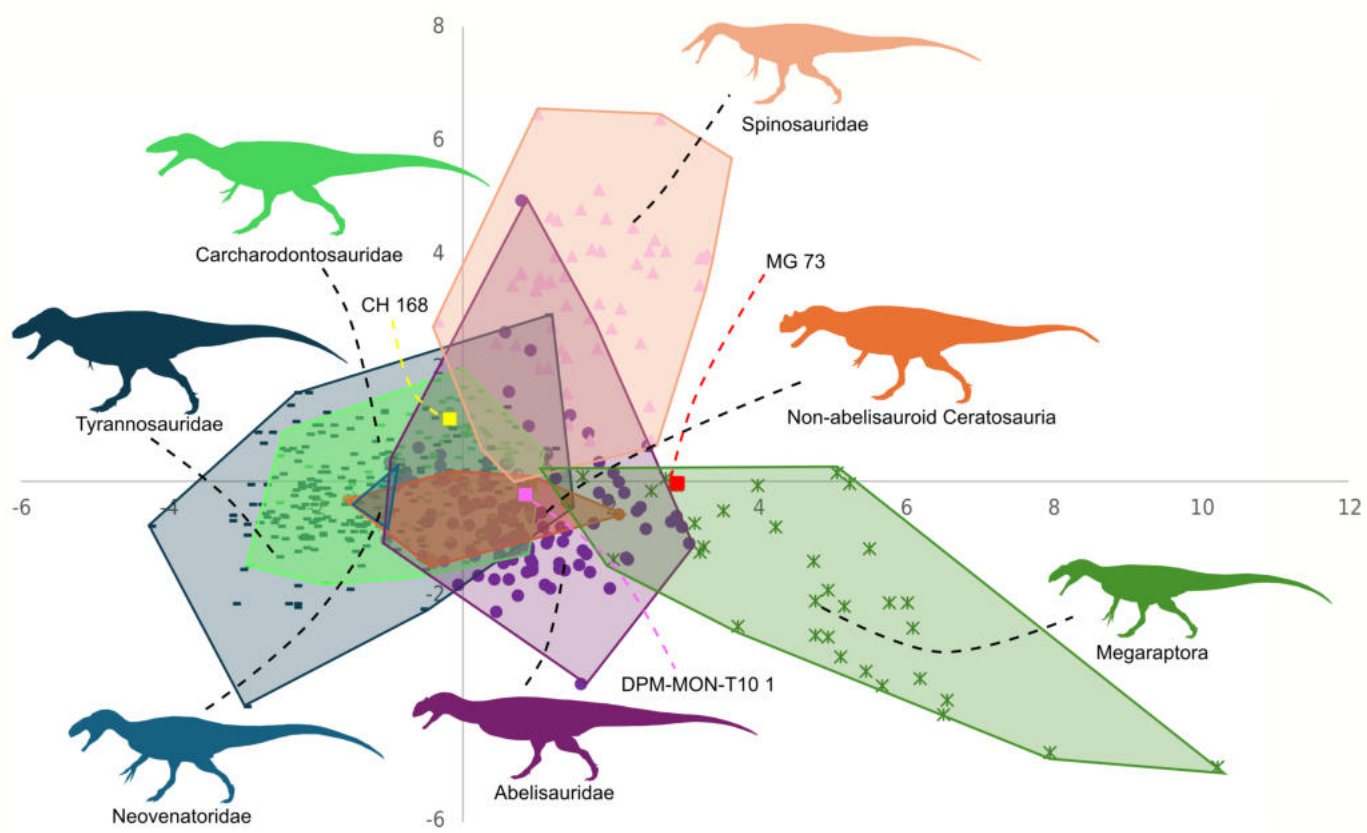
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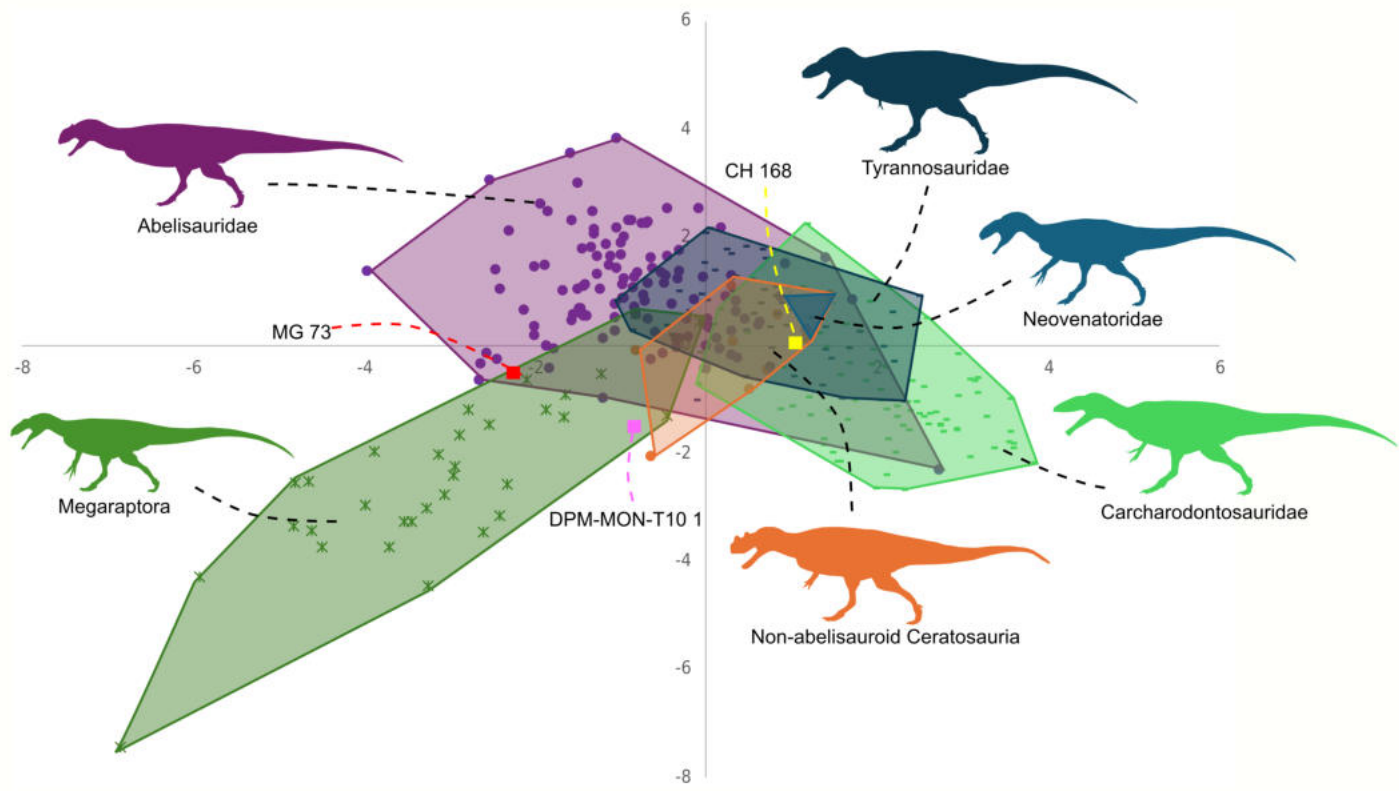
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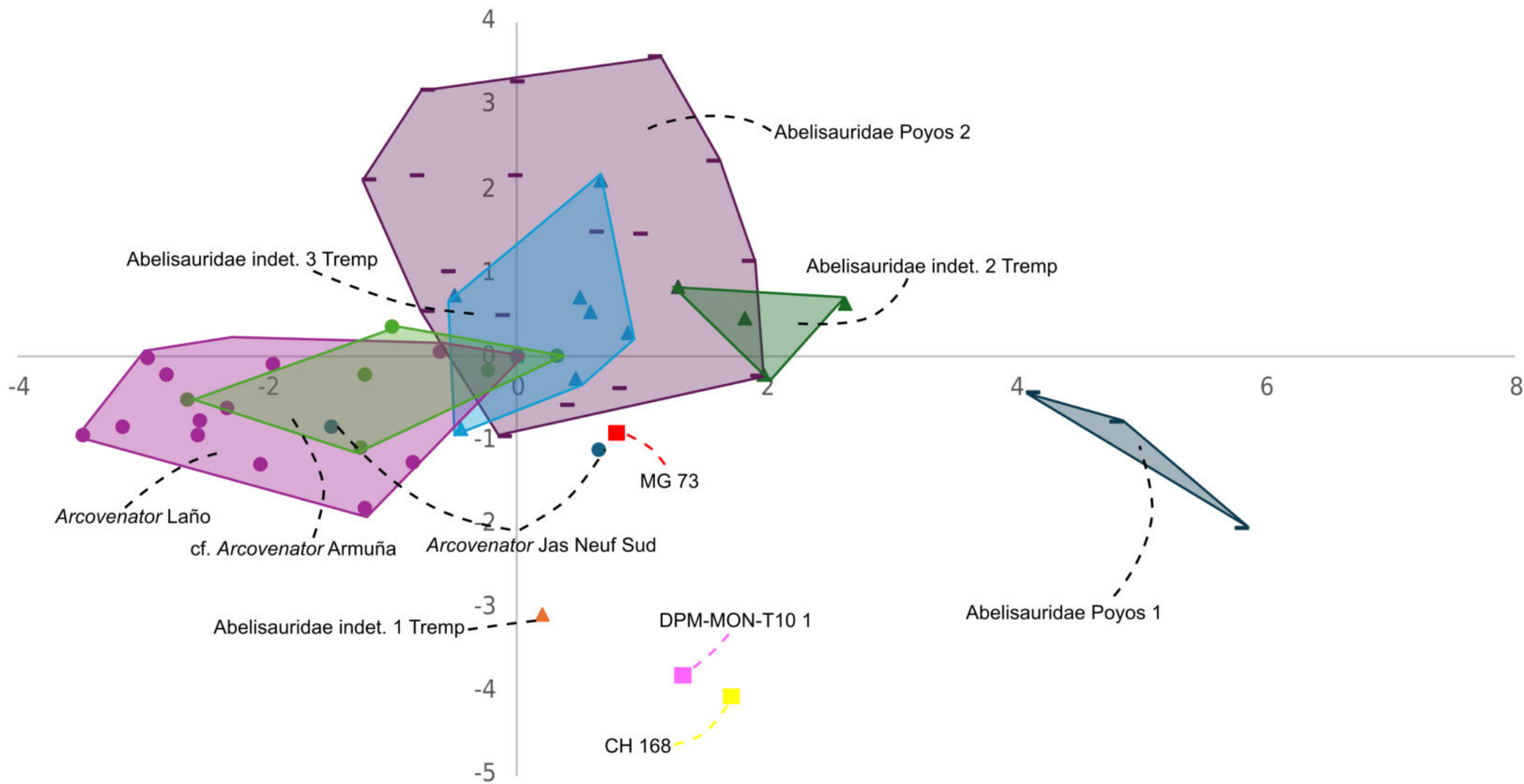


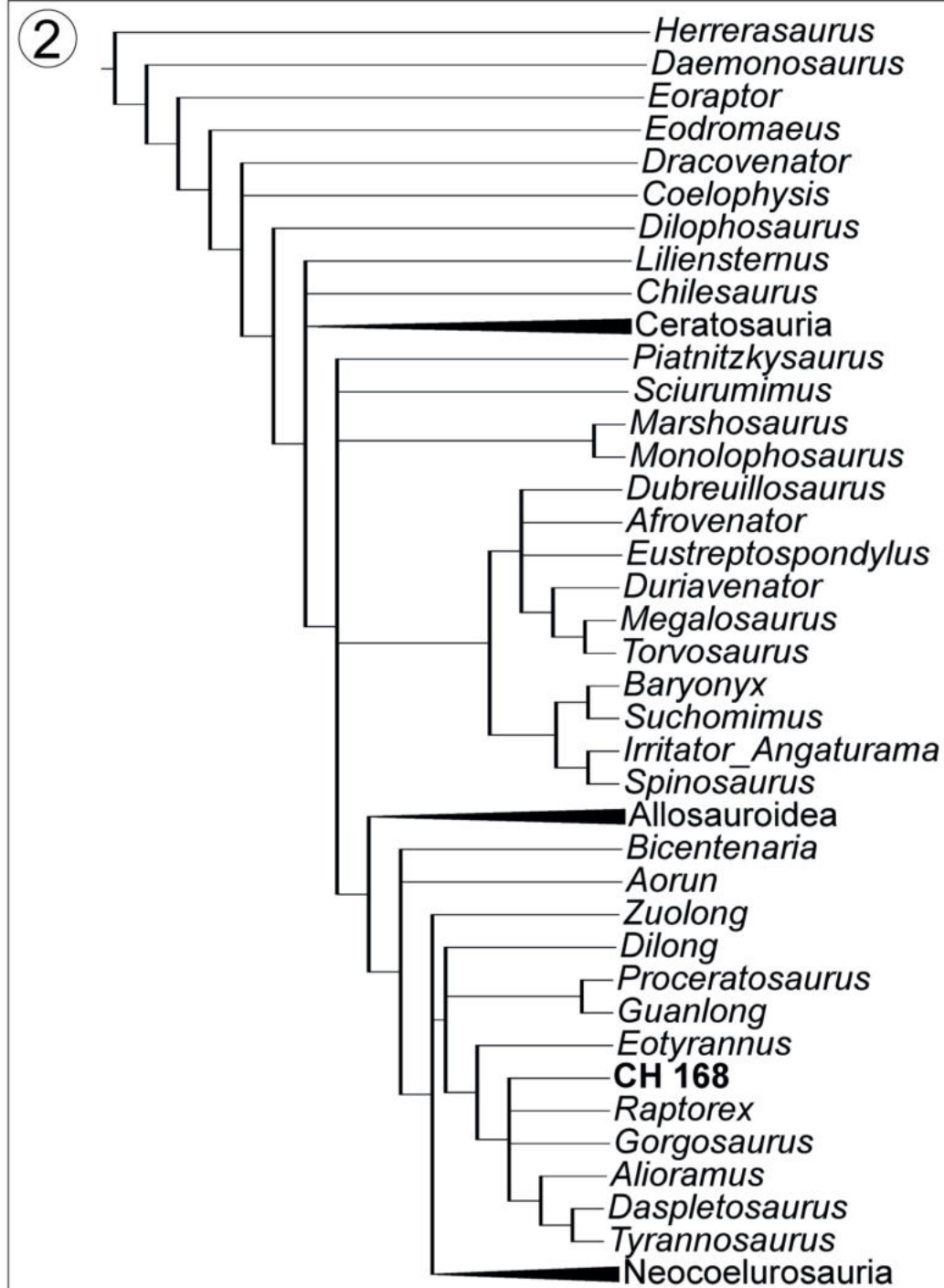
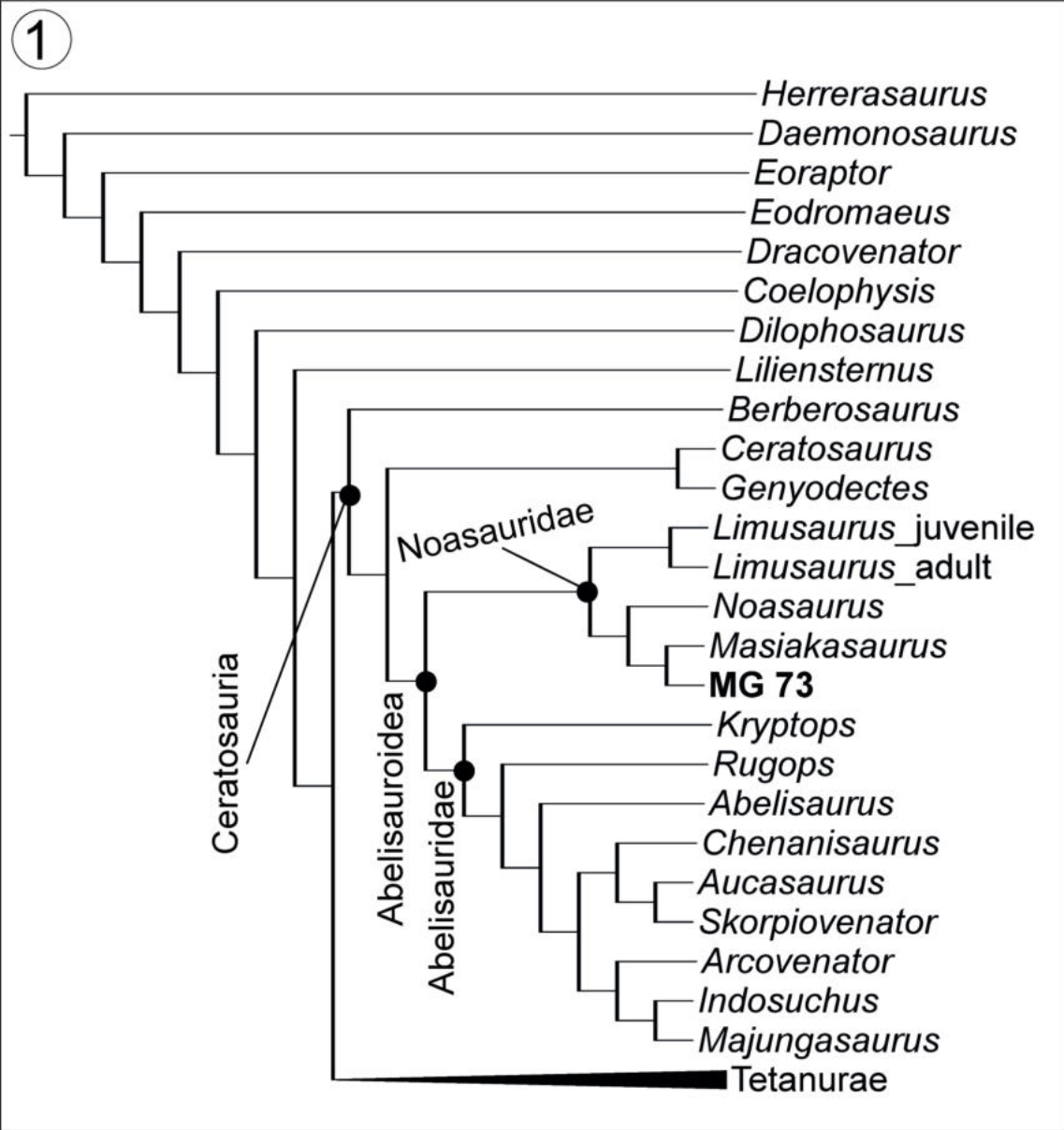
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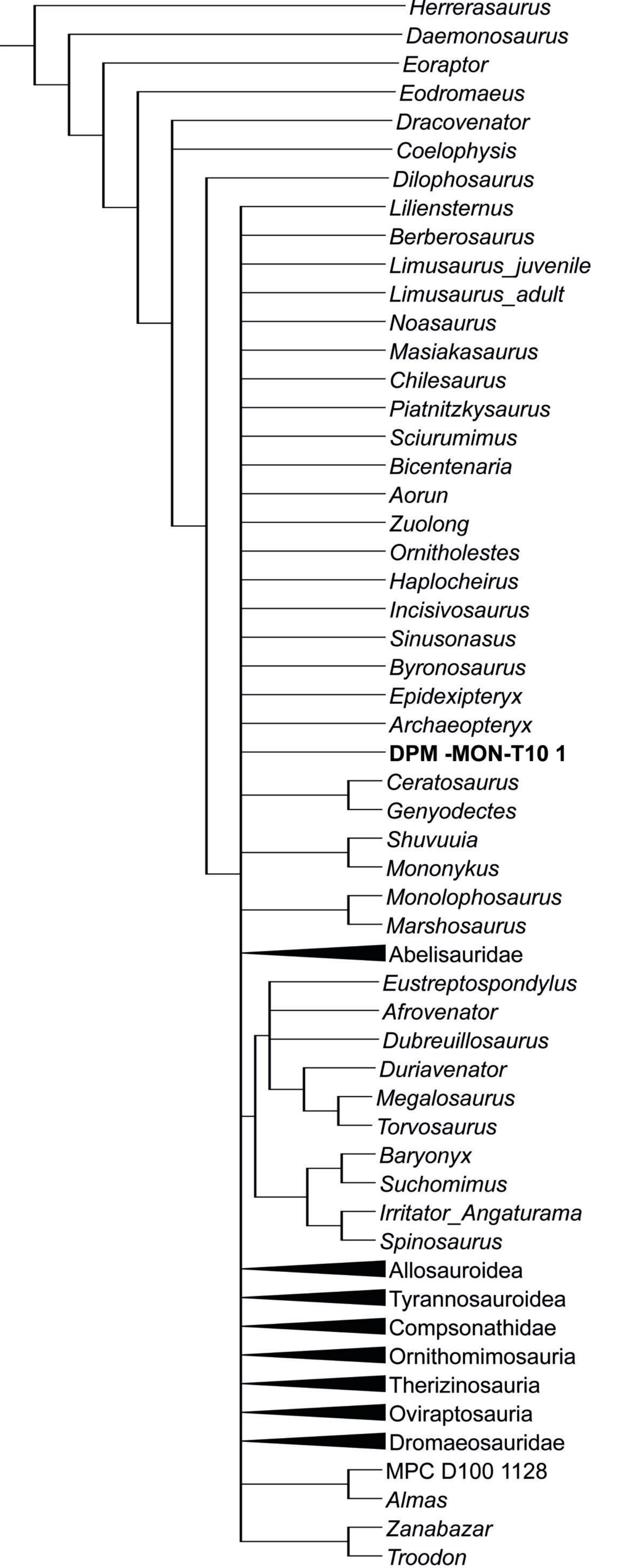
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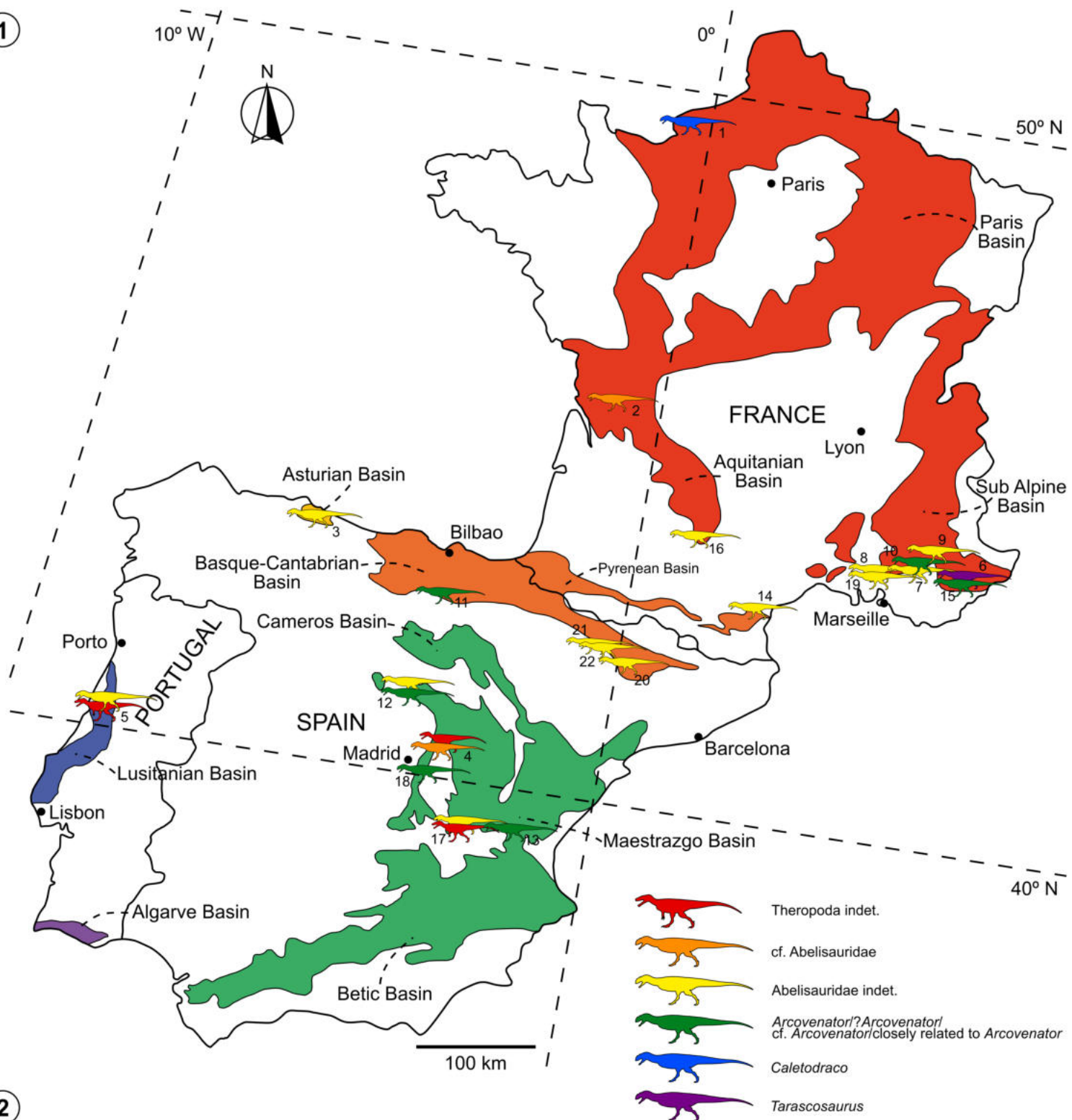




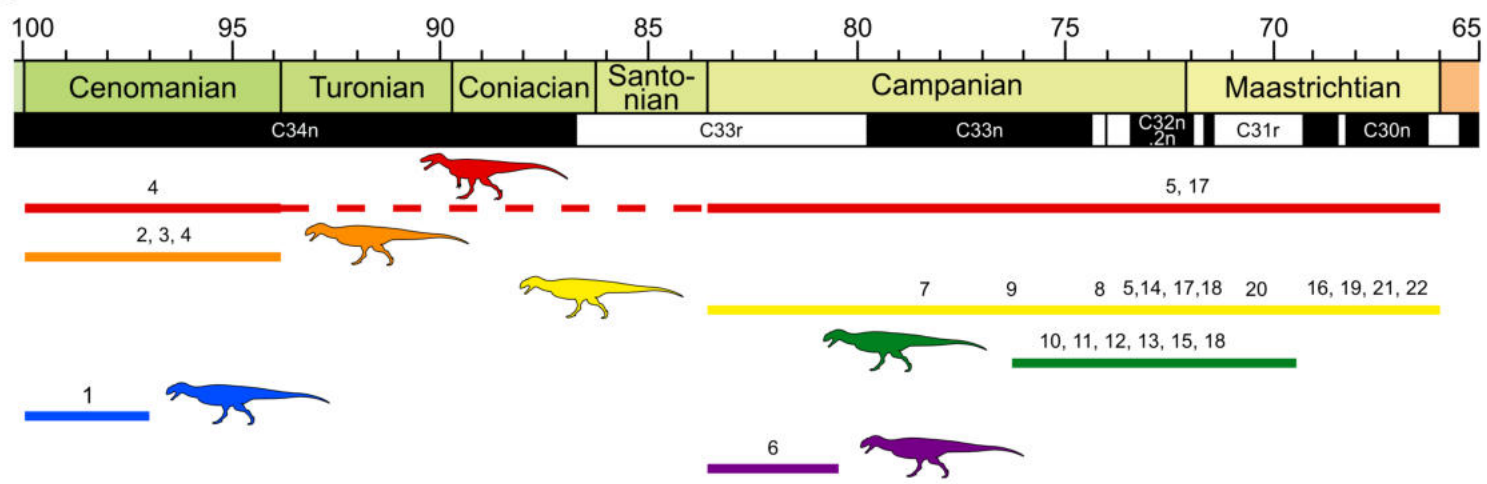




1



2



**TABLE 1. Results of the discriminant analyses performed with database 1, 2 and 3. Databases in Supplementary material 1.**

|                                                                  | LDA 1 (clade)         |                              | LDA 2 (taxa)            |                         | LDA 3 (clade)   |                 | LDA 4 (taxa)          |                       | LDA 5 (Ibero-armorica)          |                                                                                |
|------------------------------------------------------------------|-----------------------|------------------------------|-------------------------|-------------------------|-----------------|-----------------|-----------------------|-----------------------|---------------------------------|--------------------------------------------------------------------------------|
|                                                                  | Classification        | Jackknifed                   | Classification          | Jackknifed              | Classification  | Jackknifed      | Classification        | Jackknifed            | Classification                  | Jackknifed                                                                     |
| <b>M<br/>G<br/>73</b>                                            |                       | Abelisauridae                | <i>Carnotaurus</i>      | <i>Carnotaurus</i>      |                 | Megaprotora     | <i>Carnotaurus</i>    | <i>Carnotaurus</i>    | <i>Arcovenator</i> Jas Neuf Sud | <i>Arcovenator</i> Jas Neuf Sud                                                |
| <b>C<br/>H<br/>16<br/>8</b>                                      |                       | Non-abelisaurid Ceratosauria |                         | <i>Gorgosaurus</i>      | Tyrannosauridae | Tyrannosauridae |                       | <i>Gorgosaurus</i>    |                                 | Abelisauridae indet. closely related to <i>Arcovenator</i> Poyos. Morphotype 1 |
| <b>D<br/>P<br/>M<br/>-<br/>M<br/>O<br/>N-<br/>T1<br/>0<br/>1</b> | Carcharodontosauridae | Carcharodontosauridae        | <i>Acrocanthosaurus</i> | <i>Acrocanthosaurus</i> |                 | Megaprotora     | <i>Aucanthesaurus</i> | <i>Aucanthesaurus</i> |                                 | Abelisauridae indet. 1 Western Tropic Syncline                                 |