

1 **Middle Triassic sharks from the Catalan Coastal Ranges (NE Spain)**
2 **and faunal colonization patterns during the westward transgression of**
3 **Tethys**

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16
17 **Abstract.** Palaeogeographic changes that occurred during the Middle Triassic in the
18 westernmost Tethyan domain were governed by a westward marine transgression of the
19 Tethys Ocean. The transgression flooded wide areas of the eastern part of Iberia,
20 forming new epicontinental shallow-marine environments, which were subsequently
21 colonized by diverse faunas, including chondrichthyans. The transgression is recorded
22 by two successive transgressive–regressive cycles: (1) middle–late Anisian and (2) late
23 Anisian–early Carnian. Here, we describe the chondrichthyan fauna recovered from
24 several Middle Triassic stratigraphic sections (Pelsonian-Longobardian) located at the
25 Catalan Coastal Basin (western-most Tethys). The assemblage consists of isolated teeth
26 of the species *Hybodus plicatilis*, *Omanoselache bucheri*, *O. contrarius* and

27 *Pseudodalatias henarejensis*. Our data complement a series of recent studies on
28 chondrichthyan faunas from Middle-Late Triassic marine basins of the Iberian
29 Peninsula, allowing us to evaluate patterns of faunal colonization. Sharks that reached
30 East Iberia during the first transgressive pulse (Anisian) show strong affinities with the
31 Sephardic bioprovince (related with the Neotethys Ocean) whereas a later increase in
32 chondrichthyan diversity (during the Ladinian transgressive pulse) is due to the arrival
33 of taxa with affinities to European faunas (Palaeotethys). This contrasts with recently
34 evaluated colonization patterns of invertebrates and conodonts, where Middle–Upper
35 Anisian assemblages are composed of taxa with affinities to the Alpine/Germanic
36 bioprovince whereas faunas from the Sephardic bioprovince only reached the shallow
37 waters of the Iberian peninsula during the second (Ladinian) transgressive pulse. The
38 differences between chondrichthyans vs. invertebrate and conodonts colonisation
39 patterns are explained here on the basis of differing dispersal strategies. Our finding
40 suggests the existence, during the Anisian, of connections between Palaeo- and
41 NeoTethys through the western Cimmerian microcontinent, which served as pathways
42 for the entrance of chondrichthyans from the Sephardic bioprovince.

43 *Keywords:* dispersal strategies; palaeocurrents; Anisian; Ladinian; coastal
44 chondrichthyans.

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48 **1. Introduction**

49 The Triassic is a key period to understand the evolutionary history of
50 chondrichthyans, with the decline or extinction of several major Palaeozoic groups
51 (such as xenacanthiforms, symmoriiforms, ctenacanthiformes and non-chimaeroid
52 holocephalans), the increasing dominance of hybodontiforms, and the emergence of the

53 modern sharks (elasmobranchii). The Triassic was also an important period for the
54 Iberian Peninsula in terms of palaeogeography. During these times, the western Tethys
55 domain was characterised by important palaeogeographic changes and the subsequent
56 advance of the broad westward marine transgression of the Tethys Sea that reached East
57 Iberia during the Middle Triassic (Anisian) (Escudero-Mozo et al., 2015). This
58 transgression, for the first time during the Mesozoic, flooded wide areas of the eastern
59 part of Iberia, connecting some internal basins (i.e. Iberian and Catalan), which until
60 this moment had been independent (e.g. Arche et al., 2004; Galán-Abellán, et al., 2013;
61 Escudero-Mozo et al., 2015), causing the demise of continental sedimentation and the
62 consequent deposition of marine carbonates and mixed coastal facies (Muschelkalk).

63 In East Iberia, this general transgression is recorded by two transgressive-
64 regressive cycles that show important palaeogeographic and fossil content changes
65 (Escudero-Mozo et al., 2015). The first one (middle–late Anisian) took place from the
66 NE to the Catalan Coastal Basin reaching the Iberian Basin through a narrow corridor
67 bounded by palaeohighs. The second transgressive-regressive cycle (late Anisian–early
68 Carnian) was characterised by an extensive transgression that covered a wide area of E
69 Iberia including some of the previous topographic highs. After that, during the early
70 Carnian, this area is characterised by the development of an extensive siliciclastic-
71 evaporitic setting of the Keuper. These transgressive events resulted in the evolution of
72 a diversity of epicontinental shallow coastal environments and associated colonization
73 by marine faunas, including chondrichthyans. Accordingly, Middle Triassic faunas from
74 the Iberian Peninsula reflect the transformation from continental settings to a wide
75 variety of coastal and marine depositional environments owing to the Muschelkalk
76 transgression, which occurred in the middle Anisian (Escudero Mozo et al., 2015).

77 In this paper, we document for the first time the presence of chondrichthyans in
78 the Middle Triassic rocks of the Catalan Coastal ranges (CCR). This paper is
79 complementary to Pla et al.'s (2013) and Manzanares et al.'s (2018) studies on Middle–
80 Late Triassic chondrichthyans from other areas of the Iberian Peninsula, i.e. the Iberian
81 (IR) and Betic ranges (BR), respectively. Together, these three papers give the first
82 general overview of the chondrichthyan communities that inhabited the shallow
83 epicontinental seas of the westernmost Tethys during the Middle to Late Triassic. This
84 allows us to evaluate here the patterns of colonization of the new shallow epicontinental
85 seas of the western Tethys followed by coastal sharks and to compare it with
86 invertebrate faunas. Findings require some adjustments of previous palaeogeographic
87 reconstructions of the westernmost Tethyan realm for the Middle Triassic.

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89 **2. Geographical and geological settings**

90 The Catalanian Coastal range is located in the north-eastern Iberian Peninsula
91 (Fig. 1A) and includes two mountain chains that run parallel to the Mediterranean Coast
92 in a NE–SW direction, from the Gerona region towards the Ebro delta, separated by the
93 Prelitoral Depression (Sopeña and De Vicente, 2004; Escudero-Mozo et al., 2015).

94 The Catalanian Basin constituted an intracratonic basin that was mainly developed
95 during Mesozoic times when it experienced some extensional periods with several syn-
96 rift and post-rift phases (Vargas et al., 2009; De Vicente et al., 2009). It is dominated by
97 longitudinal NE–SW faults and to a lesser extent by a conjugated NW–SE fault system
98 (Vegas, 1975); the latter divided this basin into three domains with different subsidence
99 (from SW to NE): Priorat-Baix Ebre, Prades, and Gaiá-Montseny (Marzo, 1980; Calvet
100 and Ramón, 1987; Ramón and Calvet, 1987; Gaspar-Escribano et al., 2004). During the
101 Cenozoic, Alpine compressional tectonics gave rise to the present-day Iberian range and

102 Catalan Coastal range.

103 In the Catalonian Basin, the succession shows the three typical subdivisions of
104 the Germanic Triassic, namely Buntsandstein, Muschelkalk and Keuper facies. In the
105 Catalan Coastal range, the Muschelkalk facies is represented by two carbonate units
106 (“Lower” and “Upper” Muschelkalk) that record the deposition of shallow to outer
107 carbonate ramps, separated by a mixed evaporite-siliciclastic unit (“Middle
108 Muschelkalk”). This facies, combined with the uppermost part of the Buntsandstein (or
109 Röt) facies and the lower part of the Keuper facies, records two major transgressive–
110 regressive cycles, respectively of middle–late Anisian and late Anisian–early Carnian
111 age (Escudero-Mozo et al., 2014, 2015), each one related to the development of a wide
112 carbonate ramp. The studied sections are constituted from materials of the “Lower
113 Muschelkalk” (L’Ametlla section) from the Gaia–Montseny domain, and “Upper
114 Muschelkalk” (Rasquera-Benifallet and Tivissa sections) from the Priorat-Baix-Ebre
115 (Fig.1B).

116 In the Gaia–Montseny domain, the Lower Muschelkalk shows an average
117 thickness of 120 m, showing a sharp lower contact with lutite, carbonate and evaporite
118 units (Röt facies) and a transitional upper contact with the Middle Muschelkalk. The
119 unit was divided by Calvet and Ramón (1987) and Calvet et al. (1990) into four
120 members, which correspond to El Brul (shallow subtidal to supratidal deposits), Olesa
121 (lagoonal bioclastic carbonates), Vilella Baixa (open marine burrowed deposits) and
122 Coldejou (white peritidal dolomite facies). Based on ammonoids, foraminifera, pollen
123 and brachiopods, the Lower Muschelkalk unit is dated as middle–upper Anisian (Calvet
124 and Marzo, 1994; Goy, 1995; López-Gómez et al., 1998; Márquez-Aliaga et al., 2000).
125 Chondrichthyan remains are concentrated in different levels of the Olesa unit.

126 The L'Ametlla section is located close to the village of L'Ametlla del Vallès in
127 the province of Barcelona, Catalonia, 35 km northeast of Barcelona City (coordinates at
128 the base; 41° 43' 18" N and 2° 15' 26" W; Fig. 1A). The section and its fossil record
129 have been studied previously by Márquez-Aliaga et al. (2000), Plasencia (2009, and
130 references therein).

131 In the Baix-Ebre Priorat domain, the Upper Muschelkalk carbonates show a
132 gradual to sharp lower contact with the "Middle" Muschelkalk and a gradual upper
133 contact with the Keuper, and show an average thickness of 140 m. In this area, the
134 carbonate unit was divided by Calvet et al. (1987) into several members from base to
135 top that are: 1) Rojals Member (shallow subtidal to supratidal oolitic limestone and
136 dolostone); 2) Benifallet Member (low energy subtidal bioturbated limestone and
137 dolostone); 3) Rasquera Member (outer to middle ramp limestone, dolostone and shales
138 with *Daonella*); 4) Tivissa Member (outer to inner ramp carbonates); and 5) Capafons
139 Member (peritidal dolomites, marls and shales). Based on the ammonoids yielded by
140 the Rasquera and Tivissa members and the sequence stratigraphic analysis, the Upper
141 Muschelkalk has been dated as late Illyrian–late Longobardian (Goy, 1995; Escudero-
142 Mozo et al., 2015). Conodonts from the same samples confirm a Ladinian age
143 (Plasencia, 2009).

144 Both the Rasquera–Benifallet and Tivissa sections expose sediments of the five
145 members as defined by Calvet et al. (1987). These sections are located in the province
146 of Tarragona, Catalonia (Fig. 1A). The Rasquera–Benifallet section is along the road of
147 Rasquera to Benifallet villages (coordinates at the base 40° 59' 38" N and 0° 32' 22"
148 W), and the Tivissa section is close to Tivissa village (coordinates at the base 41° 02'
149 33" N and 0° 44' 48" W). Samples that have yielded chondrichthyan remains came from

150 both the Rasquera and Tivissa members (Fig. 1B). Conodonts from the same samples
151 confirm a Ladinian age (Plasencia, 2009).

152

153 **3. Methodology**

154 The specimens studied here were obtained by acid etching in 5–10% acetic acid,
155 of more than 60 Kg of limestones (in samples of around 1.5 kg each) from the
156 Rasquera–Benifatell, Tivissa and Ametlla sections. After dilution residues were sieved
157 with sieves meshes of 2, 0.125 and 0.063 mm respectively, which created three fractions
158 refer as the “total sample”. Figure 1B shows the stratigraphic position of all samples
159 that have yielded chondrichthyan remains. Fossils were picked under a binocular
160 microscope and photographed with an HITACHII S-4800 Electronic Microscope hosted
161 at the Microscopy Services of the University of Valencia. The archived specimens are
162 mainly isolated teeth and scales, housed in the Museum of Natural History of the
163 University of Valencia (former Geological Museum of Valencia University), Spain
164 (MGUV-36111 to MGUV-36113; and MGUV-36117 to MGUV-36131).

165

166 **4. Results**

167 The chondrichthyan assemblage found in the Middle Triassic of the Catalan
168 Basin includes four species belonging to three different genera; i.e. *Pseudodalatias*
169 *henarejensis*, *Omanoselache bucheri*, *O. contrarius* and *Hybodus plicatilis* (Fig. 2). A
170 detailed Systematic Palaeontology section is provided as Supplementary Material.

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172

173 **5. Discussion**

174

175 5.1. Middle Triassic sharks from the Catalan Basin in the context of the Iberian
176 Peninsula

177 This association is similar to, but less diverse than, the chondrichthyan faunas
178 recently described by Pla et al. (2013) and Manzanares et al. (2018) from other Middle-
179 Late Triassic basins of the Iberian Peninsula (see also Botella et al., 2009; Ferrón et al.,
180 2014; Manzanares et al., 2017). In fact, *P. henarejensis*, *O. bucheri*, *O. contrarius* and
181 *H. plicatilis* are a typical selection of Middle Triassic forms from Iberia being also
182 present in the Ladinian of both the Iberian and Betic Ranges (Fig. 3 B).

183 In general, the Middle Triassic shark communities from the epicontinental seas
184 of the westernmost margin of the Tethys ocean (i.e. east of the actual Iberian Peninsula)
185 were largely dominated, in terms of diversity of species and abundance, by durophagous
186 sharks with crushing dentitions (i.e. *O. bucheri*, *O. contrarius*, *Lissodus* aff. *L. lepagei*,
187 *Lonchidion derenzii*), well adapted for grabbing and crushing prey with a hardened
188 exoskeleton (e.g. crustaceans and mollusks). The teeth in general are small, suggesting
189 that these sharks were also of small size, possibly in turn subsidiary prey of some of the
190 larger aquatic reptiles, e.g. pachypleurosaurs, nothosaurs or ichthyosaurians, that are
191 common in the same areas (Fortuny et al., 2011, Márquez-Aliaga et al., 2019), being
192 these (most probably) the apex predator of these shallow marine environments.

193 The stratigraphic distribution of taxa in East Iberia (i.e. Catalanian, Iberian and
194 Betic basins) is summarized in Figure 3 A. Figure 3B provided synthetic information on
195 the specific abundance. Teeth of *O. bucheri*, *O. contrarius* and *H. plicatilis* from
196 samples of the Olesa Unit in section L'Ametlla represent the first report of Anisian
197 (Pelsonian–lower Illyrian) sharks of the Iberian Peninsula showing that chondrichthyans
198 reached the epicontinental shallow waters of NE Iberia during the first (middle–late
199 Anisian) Triassic transgression. The stratigraphic record of all these three species

200 extends up into the Carnian (Late Triassic) in the Iberian Peninsula, being also the most
201 representative and abundant taxa.

202 On the other hand, an important increase of the chondrichthyan diversity in the
203 Triassic Iberian basins occurs in the Longobardian, during the second transgressive–
204 regressive cycle, with the appearance of several taxa, such as *Palaeobates*
205 *angustissimus*, *Lissodus* aff. *L. lepagei*, *Rhomaleodus*, or *Pseudodalatias henarejensis*
206 (Fig. 3A).

207

208 5.2 Biogeographical affinities of chondrichthyan vs. invertebrate faunas from East 209 Iberia during the Triassic westward transgressions of the Tethys Sea

210 Recent evaluations on the evolution of invertebrate faunas from the Middle
211 Triassic carbonate platforms in eastern Iberia reflect a drastic palaeobiogeographic
212 change during the Upper Anisian. Thus, fossil associations (mainly bivalves,
213 foraminifers, brachiopods and conodonts) from the Landete Formation (Iberian Range)
214 and lower Muschelkalk (Catalan Coastal Range), both dated as Middle–Upper Anisian
215 (Pelsonian–early Illyrian), are almost exclusively comprised by taxa with affinities to the
216 Alpine/Germanic bioprovince, related with the Palaeotethys Ocean. This has been
217 interpreted as the Iberian and Catalonian Basins acting as a palaeogeographic gulf
218 opening northwards, only connected with the Palaeotethys. The presence of
219 geographical barriers should have prevented the entrance of faunas from the Sephardy
220 bioprovince, related with the Neotethys (Escudero-Mozo et al. 2015; see fig. 19). In
221 contrast, fossil associations of invertebrates and conodonts from the upper Anisian to
222 upper Ladinian (late Illyrian–Longobardian; i.e. Cañete Fm., Iberian Range and upper
223 Muschelkalk, Catalan Coastal Range) show a mixed origin, where, along with endemic
224 species, typical species of both the Alpine/Germanic and Sephardic bioprovinces

225 appear. This palaeobiogeographical heterogeneity of faunas has been linked to the
226 second and most severe, transgressive–regressive cycle (late Anisian–early Carnian),
227 which allowed for the colonization of shallow waters of the Iberian Peninsula by faunas
228 coming from both Tethys provinces: the Germanic/Alpine to the north and the
229 Sephardic to the south.

230 Noticeably, the stratigraphic distribution of Middle–Upper Triassic
231 chondrichthyans in the Iberian Peninsula differs from this pattern of colonization. Thus,
232 taxa that reached the new shallow-water environments of NE Iberia during the middle
233 Anisian transgression (*i.e.* *Hybodus plicatilis*, *O. bucheri* and *O. contrarius*) do not show
234 the clear Alpine/Germanic affinities reflected in invertebrate and conodont faunas. Only
235 *Hybodus plicatilis* has been reported from other localities in Europe (see supplementary
236 information.) Poland (Niedzwiedzki, 2008); Luxembourg (Delsate and Duffin, 1999);
237 Bulgaria (Stefanov, 1966); Germany and UK (Korneisel et al., 2015). However, this
238 taxon might be a cosmopolitan species from the Tethys realm as teeth assigned to
239 *Hybodus* sp. aff. *Hybodus plicatilis* have been reported from the Middle Triassic of
240 Saudi Arabia (Vickers-Rich et al., 1999). In European localities, *H. plicatilis* occurs in
241 association with other “*Hybodus*”, *Acrodus*, *Polyacrodus*, *Lissodus* and some
242 neoselachian species (Niedzwiedzki, 2008; Delsate and Duffin, 1999; Korneisel et al.,
243 2015), which are absent in the Anisian of the Iberian Peninsula. In the same sense, the
244 genus *Omanoselache* has never been reported from the other Triassic localities in
245 Europe apart from the Iberian Peninsula occurrences. *O. contrarius* occurs in the
246 Middle Triassic of Canada (Johns et al., 1997) and in the Middle–Late Triassic of
247 Gualing, China (Chen et al., 2007), whereas *O. bucheri* is known from the Middle
248 Triassic of Nevada, U.S.A. (Rieppel et al., 1996; Cuny et al., 2001). Pla et al. (2013)
249 suggested that this species might have reached the Iberian Peninsula through the

250 Sephardic province (i.e., the southern margin of the Tethys Ocean). The recent
251 discoveries of several new *Omanoselache* species from the Middle Permian (Koot et al.,
252 2013) and Lower Triassic (Koot et al., 2015) of Oman confirms the Sephardic affinities
253 of the genus. Then again, the increase of the chondrichthyan diversity occurred during
254 the second transgressive–regressive cycle (Longobardian, Fig 3A) is due to the
255 appearance of several taxa with clear affinities with Alpine/Germanic faunas, such as
256 *Palaeobates angustissimus*, *Lissodus*, *Rhomaleodus*, or *Pseudodaltias* (Fig. 3).

257

258 *5.3 Dispersal strategies, oceanic palaeocurrents and readjustment of palaeogeographic*
259 *reconstructions; an overview.*

260 Oceanic currents therefore act as distinctive unidirectional corridors for passive
261 dispersion of organisms with a microscopic pelagic life-stage (Zinsmeister and
262 Emerson, 1979; Jablonski and Lutz, 1983). Consequently, palaeontologists have for a
263 long time recognized that distributions of some marine fossils are useful in
264 reconstructing ancient marine current patterns. As Jablonski and Lutz (1983) pointed
265 out, low (or high) faunal similarity between regions does not necessarily imply large (or
266 small) geographic disjunctions, but rather it can reflect the prevailing current system.

267 In this sense, dispersion of bivalves, ammonites and foraminifers was largely
268 determined by the dominant systems of marine currents. Most marine invertebrates
269 include a microscopic, free-living, pelagic life history stage—the larval form. Larvae
270 feed in the plankton (except for lecithotrophs) exhibiting long pelagic durations and
271 high dispersal capacity. Distribution of these taxa is largely determined by ocean
272 currents transporting larvae between distant patches of suitable habitat for adults, often
273 over great distances (Scheltema, 1986; Jablonski and Lutz, 1983; Pechenik, 1999;
274 Kinlan et al., 2005; Trakhtenbrot et al., 2005; Trembl et al., 2008; Wangensteen et al.,

275 2017). On the other hand, ammonites underwent a direct development lacking a distinct
276 larval stage (as other, extant cephalopods: nautilus, cuttlefishes, squids and octopuses)
277 consisting of four ontogenetic stages (embryonic/egg, neanic, juvenile, and adult; see
278 e.g. Westermann, 1996; Klug et al 2015; Zell and Stinnesbeck, 2016; Tajika *et al*,2018).
279 Nevertheless, most authors consider that ammonite hatchlings – the neanic stage – were
280 planktic, with ocean currents enhancing their dispersion. In fact, the patchy yet
281 widespread palaeogeographic distribution of numerous ammonite species suggests that
282 extensive planktic dispersal with ocean currents played an important role in their
283 dispersion patterns (Zell and Stinnesbeck, 2016). The same is valid for foraminifers.
284 Thus, for benthic attached species, such as those fossils found in NE Iberia (L. Marquez
285 pers. comm. (2018), but see also the list of species in Escudero-Mozo et al., 2015), the
286 most common ways of dispersion were, based on living forms (1) release of gametes,
287 zygotes, or of embryonic agamonts/gamonts to the water column, and (2) adaption to a
288 meroplanktonic juvenile stage with passive spread by currents. Moreover, shallow-
289 water taxa can also be transported long distances by floating objects, larger animals
290 and/or by suspended particles (Myers, 1936, Murray, 2006, Nomura et al, 2010, Alve
291 1999).

292 In contrast, chondrichthyans lack any planktic stage but undergo a direct
293 development after internal fertilization. They possess large eggs with a high amount of
294 yolk and produce ovoviviparous or oviparous juveniles (there are some viviparous
295 species). In any case, the distribution of the chondrichthyan species largely depends on
296 the dispersal abilities of adults and much less on marine currents. In this sense, it is well
297 known that the vagility of sharks is directly proportional to body size (e.g. Musick et al.,
298 2004). As a whole, small-bodied coastal sharks present a very low vagility and short
299 forage distances, with oceanic distances, deep channels or strong regional currents

300 acting as dispersal barriers (Musick et al., 2004; Munroe et al., 2015; Whitney et al.,
301 2012). Thus, dispersion of coastal species usually requires continuous appropriate
302 habitats across regionally small geographical scales.

303

304 *5.4. Palaeogeography and dominant marine currents at the westernmost Tethys*

305 To consider the different dispersal capability of organisms is important in order
306 to evaluate the timing and nature of palaeobiogeographical barriers. Taxa with planktic
307 stages can provide considerable information for reconstructing marine palaeocurrent
308 systems while taxa lacking these stages (or with low-dispersal larvae) can be used to
309 infer continuity among continental shelves, both groups being affected differentially by
310 palaeobiogeographic barriers (see e.g. Shuto, 1974; Jablonski and Lutz, 1983).

311 Following these general rules, the existing differences in the colonizing patterns of
312 invertebrate vs chondrichthyan faunas in the Triassic shallow waters of E Iberia can be
313 reasonably well explained. However, it requires some adjustments of previous
314 palaeogeographic reconstructions of the westernmost Tethyan realm (see Escudero-
315 Mozo et al. 2015; their figure 19 and references therein).

316 We suggest that the colonization by bivalves, ammonites and foraminifers, during both
317 the late Pelsonian–early Illyrian and the late Illyrian–Longobardian marine
318 transgressions, was largely determined (or enhanced) by the prevailing marine currents
319 in the westernmost Tethys and by the transgression of the sea itself (Fig. 4).

320 Thus, in late Pelsonian–early Illyrian times, marine currents from the
321 Palaeotethys enhanced the arrival of planktic organisms to the new Catalan and Iberian
322 marine basins. At this time, the Cimmerian microcontinent, which controlled the
323 regional palaeogeography in the westernmost Tethys (Yin and Song, 2013; Escudero-
324 Mozo et al., 2015), would have acted as a barrier for marine currents (if any) of the

325 incipient Neotethys. However, the occurrence of chondrichthyans from the Sephardic
326 domain indicates the formation of narrow, probably temporary connections (acting as
327 alternative routes for the entrance of sharks) between Palaeo- and Neotethys during the
328 northward displacement of the Cimmerian microcontinent (Fig. 4). During the next
329 greater and more generalized sea level rise (Illyrian–Longobardian transgression), the
330 Tethys sea flooded vast areas of the westernmost Cimmerian microcontinent, such as
331 Alboran, Majorca, Minorca, Sardinia, Corsica, Tuscany, Apulia, Mani (Escudero-Mozo,
332 2015: figure 19). Subsequently, the Palaeotethys and Neotethys became broadly
333 connected, allowing for the entrance of planktonic organisms (i.e. bivalve larvae, neatic
334 ammonites and foraminifers, among others) into E Iberia, which were transported by
335 marine currents from the Neotethys (Fig. 4).

336 Notably, during both colonization events, conodonts follow the same dispersion
337 patterns as invertebrates and not those of chondrichthyans. Conodonts are considered by
338 some authors a basal group of vertebrates (e.g. Donoghue et al., 2000) although their
339 placement in this group is seriously questioned by other authors (e.g. Turner et al.,
340 2010). Little is known about their reproductive strategies (with the finding of putative
341 conodont eggs, von Bitter and Pohl, 2009), although it has been suggested, based on
342 their crown growth and function (see Martínez-Pérez et al, 2014), that conodonts could
343 had developed a larval stage (Shirley et al., 2018). Data here discussed are compatible
344 with conodonts having a passive dispersive planktic stage. Planktic larval stages (or
345 eggs) are present in several osteichthyan (bony fishes; common in teleosteans) but
346 unknown in other fishes. A larval stage is present in agnathan lampreys (but not in
347 mixines) and consists of a large sedentary burrowed larva in freshwater streams (e.g.
348 Kelly and King, 2001), and could to be present in other extinct Palaeozoic agnathan
349 fishes (Ferrón et al. 2018). The cosmopolitanism and high dispersal of Palaeozoic

350 conodonts might also agree with a planktic dispersion drifted by ocean currents;
351 however, further and more extensive studies on this topic are necessary to support this
352 idea.

353 It should further be noted that, despite that our interpretation are well supported
354 by data from Pla et al.'s (2013), Manzanares et al. (2018) and here, the number of
355 specimens and taxa in the Iberian Peninsula is relatively low. Moreover, triassic shark
356 faunas are still poorly known in general. Therefore, new findings could change (or
357 refine) this interpretation in the future.

358

359 **6. Conclusions**

360 This work completes a series of studies on the chondrichthyan faunas from
361 Middle–Late Triassic of Iberia which has changed the previously inaccurate perception
362 that chondrichthyans were rare, or completely absent, in the Iberian Peninsula. These
363 studies provide now a detailed overview of the evolution of chondrichthyan
364 communities in the new shallow-water environments developed in E Iberia
365 (westernmost Tethys) during the Triassic westward transgression of the Tethys Sea.
366 Chondrichthyan communities were largely dominated by small durophagous coastal
367 sharks with crushing dentitions well adapted for preying on crustaceans and mollusks.
368 Shark that reach NE Iberia during the first transgressive pulse (middle–late Amirian) of
369 the Tethys Sea show clear affinities with taxa from the Sephardic bioprovince, related to
370 the Neotethys Ocean (i.e. *Omanoselache*). The notable increase in chondrichthyan
371 diversity that occurs in E Iberia during the second transgressive pulse (Ladinian) is due
372 to the arrival of new taxa with clear affinities with European faunas from the
373 Palaeotethys Ocean (i.e. *Paleobates angustissimus*, *Lissodus*, *Hybodius*, *Rhomaleodus*,
374 or *Pseudodalatias*). This colonization pattern is somewhat inverse to those recently

375 evaluated for other fossil groups. Thus, Anisian bivalves, foraminifers, brachiopods and
376 conodont faunas from E Iberia are almost exclusively composed of taxa from the
377 Alpine/Germanic bioprovince. In contrast invertebrate and conodonts from the
378 Sephardic bioprovince reach the shallow waters of the Iberian Peninsula for the first time
379 during the second (Ladinian) transgressive pulse.

380 The existing differences between the colonization patterns of sharks and other
381 groups are explained on the basis of their dissimilar dispersion strategies: dispersion of
382 bivalves, ammonites and foraminifers, is largely determined by oceanic currents
383 whereas dispersion of coastal sharks depends on adult vagility (very low) and requires
384 continuous appropriate habitats. Based on this, we propose some adjustments of
385 previous palaeogeographic reconstructions including: (1) the existence of narrow,
386 probably short-term, shallow-water corridors between Palaeo- and Neo-Tethys during
387 the Anisian; (2) the Cimmerian microcontinent would have acted as a barrier for the
388 oceanic currents in the incipient Neotethys, while currents from the Palaeotethys
389 transported planktic organisms to the new Catalan and Iberian Marine Basins, and (3)
390 during the next sea-level rise (Ladinian), the Tethys sea overlapped vast areas of the
391 westernmost Cimmerian microcontinent allowing marine currents from the Neotethys to
392 extend toward the Palaeotethys reaching E Iberian epicontinental seas.

393

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402

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634

635 **Figure captions**

636 **Fig. 1. Geographical and geological settings** A) Geographical setting of
637 Catalonian Coastal Range (CCR) with location of sections studied here, modified from
638 Escudero-Mozo (2015); B) Stratigraphic columns of the sections studied. Only levels
639 where chondrichthyans remains were recovered are marked in the columns; for
640 complete columns see Plasencia (2009).

641

642 **Fig. 2. Chondrichthyan teeth from the Middle Triassic of the Catalan Coastal**
643 **Ranges, Spain.** A-D) *Hybodus plicatilis*; A–B) labial and lingual view, MGUV-36111;
644 C–D) basal and occlusal view, MGUV-36112; E–J) *Omanoselache bucheri* E) labial
645 view, MGUV-36121; F) labial view, MGUV-36122; G–H) labial and lingual view,
646 MGUV-36123; I–J) labial and lingual view, MGUV-36127; K–P) *Omanoselache*
647 *contrarius*; K–L) labial and lingual view, MGUV-36127; M–N) labial and lingual view,
648 MGUV-36120; O–P) MGUV-36118. Q–R) *Pseudodalatias henarejensis* in labial and
649 lingual view, respectively, MGUV-36113. All scale bars equal 100 μm .

650

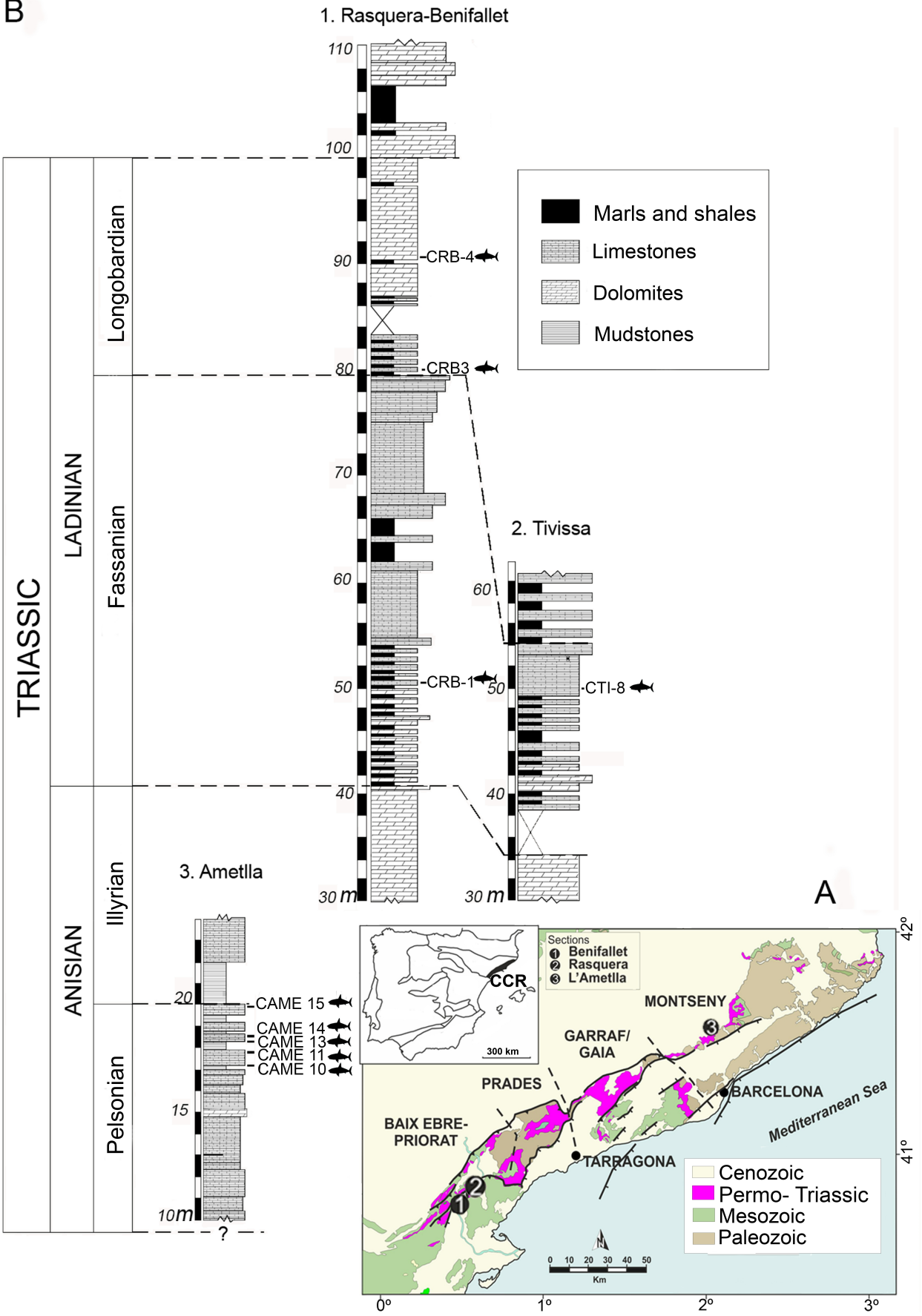
651 **Fig. 3. Stratigraphic distribution and abundance of chondrichthyans in Middle-Late**
652 **Triassic of the Iberian Peninsula** A) Stratigraphic distribution of taxa. *H. plicatilis*, *O.*
653 *bucheri*, *O. contrarius* and *P. henarejensis* are the typical fauna that can be found in any
654 Middle Triassic basin of the Iberian Peninsula. *Lissodus* aff. *L. lepagei* appears only in
655 the Iberian Ranges and Betic Ranges, whereas *Hybodus bugarensis* and *Palaeobates*
656 *angustissimus* have been only found in the Iberian Ranges. *Lonchidion derenzii* and cf.
657 *Rhomaleodus budurovi* are only present at the Betic Ranges. B) Histograms showing the
658 number of chondrichthyan teeth recovered (for each species) from in the three basins of
659 the Iberian Peninsula considered in this study (the Iberian Range, the Betic Range and
660 the Catalan-Coastal Range).

661

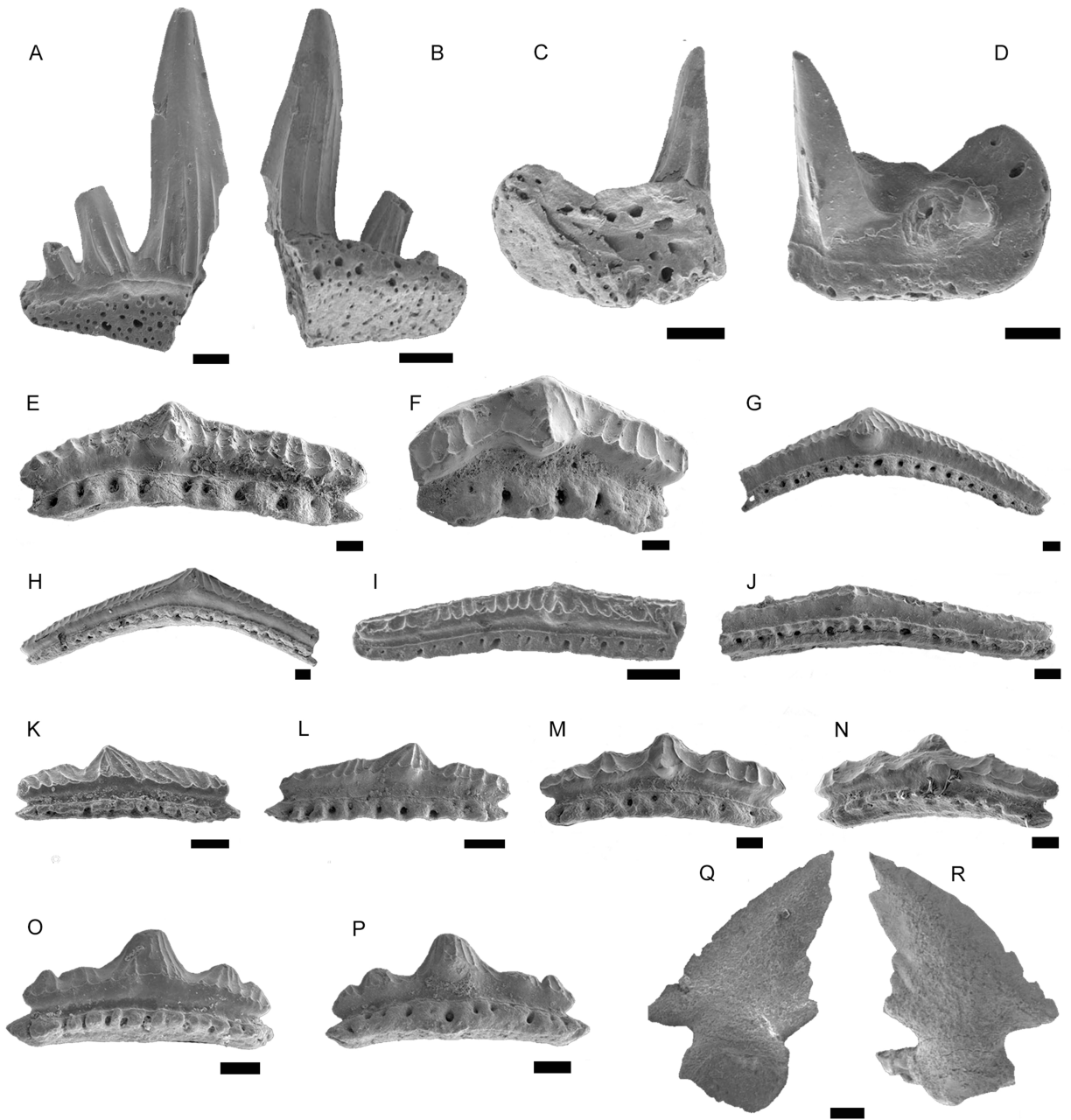
662 **Fig. 4. Faunal colonization patterns of East Iberia during the Anisian and**
663 **Ladinian westward transgression of the Tethys ocean.** A) Global palaeogeographic
664 map, Early Triassic (Induan), with schematic surface-current circulation. B–C) detailed
665 palaeogeographic reconstructions of the westernmost Tethyan realm for the Anisian (B)
666 and Ladinian (C) with interpreted colonization routes for chondrichthyans and

667 invertebrates. Palaeogeographic maps modified from Scotese (1994) (A), Stampfli and
668 Borel (2002) and Escudero-Mozo et al. (2015) (B); oceanic currents interpreted
669 according with Yan and Zhao (2001); Crasquin et al. (2010) and Martindale et al.
670 (2019). The black star marks the position of the Catalan Coastal Range, the black
671 circle marks the Betic Range, the black diamond represents the Iberian Range.

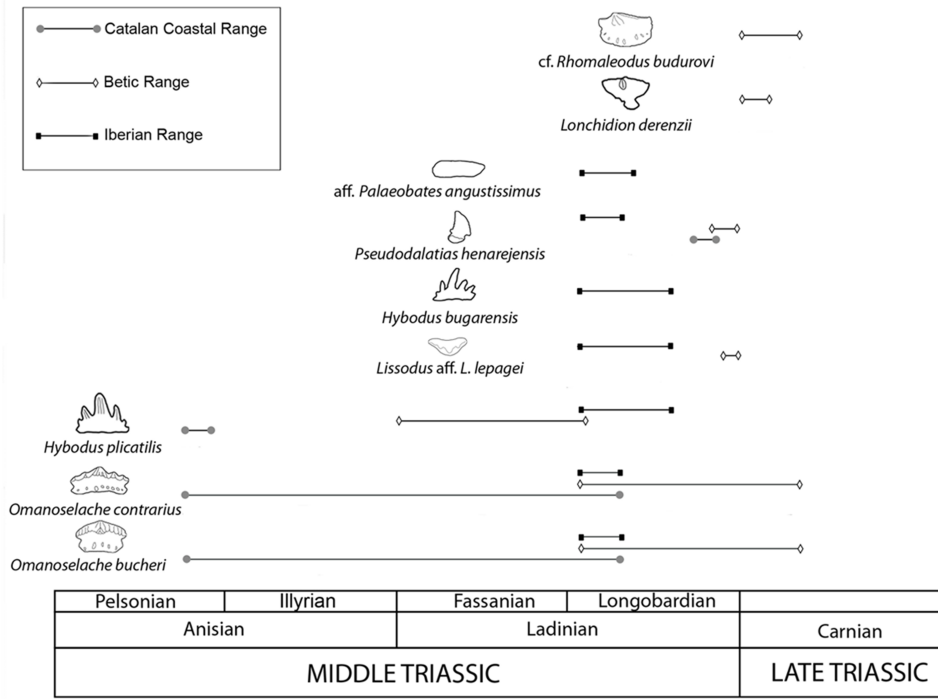
B



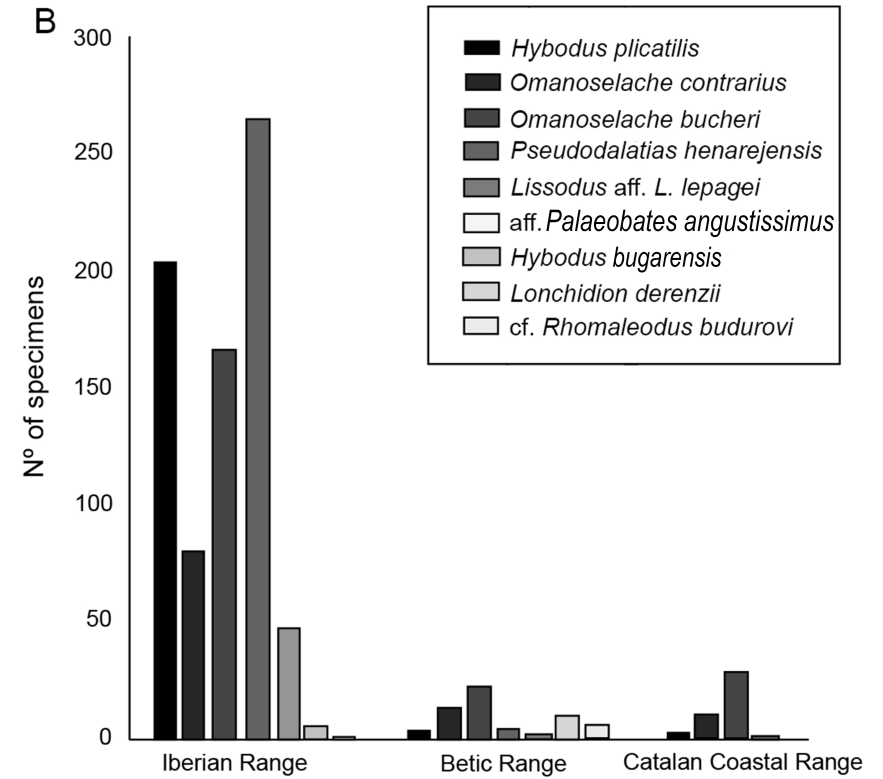
A



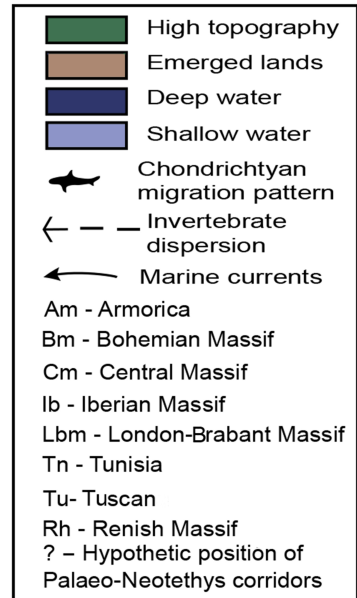
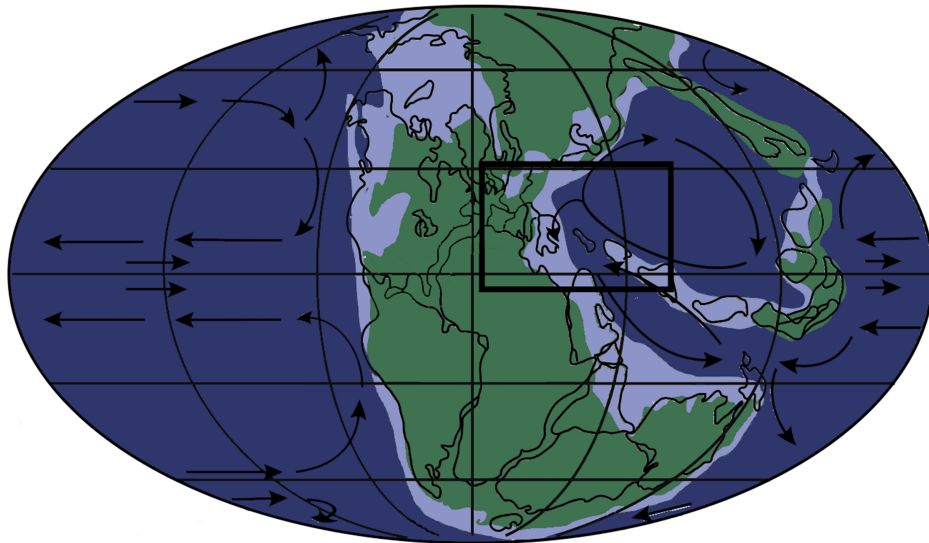
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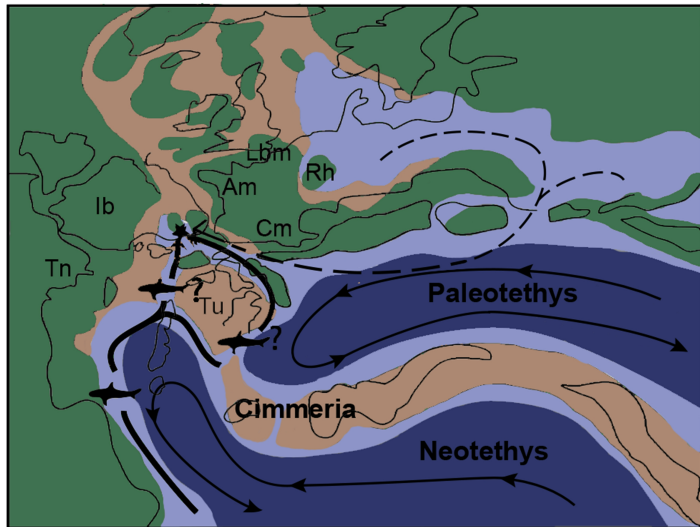
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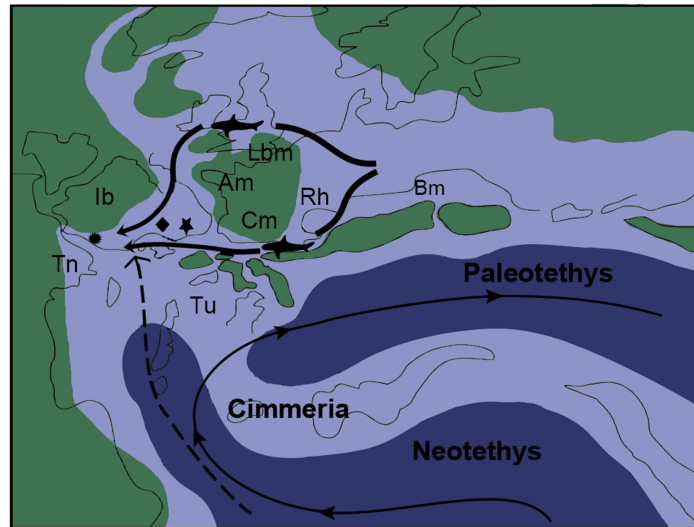
A Early Triassic (250 Ma)



B Anisian (240 Ma)



C Ladinian (230 Ma)



We confirm that there are no known conflicts of interest associated with this publication and there has been no significant financial support for this work that could have influenced its outcome. The manuscript has been read and approved by all named authors and that there are no other people who satisfied the criteria for authorship who are not listed.

We further confirm that the order of authors listed in the manuscript has been approved by all of us. We confirm that we have given due consideration to the protection of intellectual property associated with this work and that there are no impediments to publication, including the timing of publication, with respect to intellectual property. In so doing we confirm that we have follow the regulations of our institutions concerning intellectual property.

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Supplementary material

Systematic Palaeontology

Class CHONDRICHTHYES Huxley, 1880

Subclass ELASMOBRANCHII Bonaparte, 1838

Order EUSELACHII Hay, 1902

Superfamily HYBODONTOIDEA Owen, 1846

Family HYBODONTIDAE Owen, 1846

Genus *HYBODUS* Agassiz, 1837

Type species—*Hybodus reticulatus* Agassiz, 1837

Occurrence— Lower Jurassic, Lias of Lyme Regis, England (Woodward, 1916).

HYBODUS PLICATILIS Agassiz, 1843

Figure 2, A–D

Material—Two incomplete teeth from the Rasquera–Benifallet section (CAME-14 level: MGUV-36111/-36112).

Description—Most of our material is broken, with only almost complete two multicuspid teeth. They show a central cusp flanked by up to two pairs of cusplets, all of them lingually inclined (Fig. 2 A–D). Ornamentation ridges run from the tooth shoulder towards the tip of the cusp and cusplets. In the main cusp, the ridges do not reach the apical part; they cannot be seen in the lateral cusplets due to their damage (Fig. 2A-B). The vascularization is of the anaulacorhize type (Fig. 2 A-B).

Remarks —Several recent authors agree that the genus *Hybodus* is a polyphyletic group and that it does not correspond to a natural group (Rees, 1999; Underwood and Rees, 2002; Rees and Underwood, 2008; Ginter et al., 2010; Cappetta, 2012, Leuzinger et al., 2017), a statement with which we fully agree. A large number of isolated teeth described in

the literature have been assigned to *Hybodus*, although a revision of this material is required to verify, which of them fit completely the original diagnosis of the genus. However, this is beyond the scope of the present work; nevertheless, our material shows characteristics of the type species *Hybodus reticulatus*: well-defined and well-ornamented conical cusps; lateral cusplets decreasing in height, frequently inclined lingually; and a fairly shallow base (see Maisey, 1987; Duffin, 1993; Ginter et al., 2010). Thus, we choose to maintain the name of the genus *Hybodus* until a revision of the genus is done.

Occurrence—Muschelkalk, Middle Triassic of Schweningen, Germany (Agassiz, 1843); Muschelkalk, Middle Triassic of Monte Giorgio, Switzerland (von Meyer, 1849; Rieppel, 1981); Muschelkalk, Middle Triassic of Luxemburg (Delsate, 1992, 1993); Middle Triassic, Iberian Ranges, Spain (Pla et al., 2013), Middle–Late Triassic of the Betic Ranges, Spain (Manzanares et al., 2018); Middle Triassic, Catalan Coastal Ranges, Spain.

Order *INCERTAE SEDIS*

Family HOMALODONTIDAE Mutter, De Blanger and Neuman, 2008

Genus *OMANOSELACHE* Koot, Cuny, Tintori and Twitchett, 2013

Type species—*Omanoselache hendersoni* Koot, Cuny, Tintori and Twitchett, 2013

Occurrence— Middle Permian, Wordian from the Khuff Formation, Oman (Koot et al., 2013)

OMANOSELACHE BUCHERI (Cuny, Rieppel and Sander, 2001)

Figure 2, E–J

Material—Eight complete teeth and more than 20 fragments from the Rasquera–Benifallet section (CRB-1 level: MGUV-36121/-36122/-36126/-361627/-36128; and CRB-3 levels MGUV-MGUV-361216/36128; MGUV/-36123/-36130/-36131), l’Ametlla section

(CAME-12: MGUV-36124; and CAME-13 levels; MGUV-36125) and the Tivissa section (CT-8 level: MGUV-36129).

Description—Teeth are labiolingually compressed and elongated mesiodistally. All display a well-developed occlusal crest and a pyramidal main cusp (Fig. 2 E-H). Undulated vertical ornamentation ridges originate on the occlusal crest and appear on the labial and lingual sides (Fig. 2 E-F, I). A rounded lingual peg, characteristic of this species, is located under the main cusp (Fig. 2 E-G, I). Vascularisation is anaulacorhize with a row of well-developed foramina opening on the labial and lingual sides (Fig. 2 E, G).

Remarks—*Omanoselache bucheri* teeth are the most widespread, abundant and well-preserved chondrichthyan remains from the Iberian Peninsula (Pla et al., 2013; Manzanares et al., 2017). Pla et al. (2013), described three different morphotypes related to the different position they occupied in the jaws: symphyseal, mesial, or latero-distal teeth. Most of our material can be classified as their mesial teeth. These teeth exhibit one main cusp and are arched, showing a ‘boomerang’ appearance (Fig. 2 G-H). A few show a blunt and reduced main cusp and an overall more rectilinear appearance (Fig. 2 I-J), which are the characteristics of latero-distal teeth. The presence of a lingual peg, but no labial one, and the rounded edges (Fig. 2E-F, G, I) permit us to differentiate them from *Omanoselache contrarius* (Johns, Barnes and Orchard, 1997).

Occurrence—Middle Triassic, North-West of Nevada (Rieppel et al., 1996; Cuny et al., 2001); Middle Triassic, Iberian Ranges, Spain (Pla et al., 2013); Middle–Late Triassic from the Betic Ranges of Spain (Manzanares et al., 2018); Middle Triassic, Catalan Coastal Ranges, Spain.

OMANOSELACHE CONTRARIUS (Johns, Barnes and Orchard, 1997)

Figure 2 K-P

Material—Four complete teeth and several fragments from the Rasquera–Benifallet section (CRB-1 level: MGUV-36117; and CRB-3 levels: MGUV-3619/-36120) and l’Ametlla section (CAME-12 level: MGUV-36118).

Description—Teeth elongated mesiodistally and compressed labiolingually. All of them show one main cusp flanked by up to two lateral cusplets (Fig. 2 K-P). Ornamentation consists of vertical ridges that originate from the apex of the cusps and extend to the tooth shoulder. The teeth present two pegs (labial and lingual) under the main cusp (Fig. 2 M-N). As in *O. bucheri*, the vascularization is anaulacorhize (Fig. 2 K-P).

Remarks—All the teeth recovered of this species present evident wear signs on the cusps.

Occurrence—Middle Triassic of British Columbia, Canada (Johns et al., 1997); Middle–Late Triassic of Yang Liu Jing, China (Chen et al., 2007); Middle Triassic, Iberian Ranges, Spain (Pla et al., 2013); Middle–Late Triassic, Betic Ranges, Spain (Manzanares et al., 2018); Middle Triassic, Catalan Coastal Ranges, Spain.

Order *INCERTAE SEDIS*

Family PSEUDODALATIIDAE Reif, 1978

Genus *PSEUDODALATIAS* Reif, 1978

Type species—*Pseudodalatias bartonensis* (Sykes, 1971)

Occurrence — Upper Triassic, Rhaetian of Barnstone, England (Skyles, 1971)

PSEUDODALATIAS HENAREJENSIS Botella, Plasencia, Márquez-Aliaga, Cuny and Dorka, 2009

Figure 2 Q-R

Material—One tooth from the Tivissa section (CT-8 level: MGUV-36113).

Description—The tooth is almost complete, but poorly preserved with the base missing (Fig. 2Q-R). It shows the characteristic features of *Pseudodalatias henarejensis*: a serrated-triangular morphology, with three (damaged) denticles on the edges and often lacking the base (Botella et al., 2009). The cusp is also damaged and the whole tooth surface is eroded.

Remarks—*Pseudodalatias henarejensis* is an endemic species from the Iberian Peninsula, which shows a dignathic heterodonty, as described in Pla et al. (2013). The tooth found in Tivissa belongs to the lower jaw.

Occurrence —Middle Triassic from the Iberian Chain of Spain (Botella et al., 2009; Pla et al., 2013); Middle–Late Triassic of the Betic Ranges (Manzanares et al., 2018); Middle Triassic of the Catalan Coastal Ranges, Spain.

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