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A new species of *Pleuropholis* (Teleostei: Pleuropholidae) from the Upper Cretaceous Akrabou Formation of southeastern Morocco, with comments on the evolutionary history of the genus

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Credit Author Statement

Authors' contribution. P.M.B. and R.V. conceived the study, determined and assessed morphological traits, discussed the results, wrote the manuscript and made the figures.

ournal Prevension

- 1 A new species of *Pleuropholis* (Teleostei: Pleuropholidae) from the Upper Cretaceous
- 2 Akrabou Formation of southeastern Morocco, with comments on the evolutionary history of
- 3 the genus
- 4
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- 14

15 ABSTRACT

- 16 A pleuropholid fish is described for the first time from the Upper Cretaceous (Cenomanian)
- 17 Akrabou Formation, Agoult locality, southeastern Morocco. The new material, consisting of a
- 18 single complete specimen, is attributed to a new species, *Pleuropholis danielae*, and is diagnosed by
- 19 its slender body shape, the absence of a leptolepid notch in the dentary, a broad maxilla, covering
- 20 almost the entirely lower jaw, and flank scales with an unserrated posterior margin. This species
- 21 represents the last known occurrence of Pleuropholidae, a group so far unknown from the Late
- 22 Cretaceous.
- 23
- 24 Keywords:
- 25 Actinopterygii; Neopterygii; Pleuropholid fish; New taxon; Cenomanian; Africa.
- 26

27 **1. Introduction**

28

29 Pleuropholidae is an extinct family erected by Saint-Seine (1949), with a 30 chronostratigraphic range of Middle Jurassic to Lower Cretaceous (Olive et al., 2022). Known from 31 marine and continental strata in Europe (e.g. Egerton, 1858; Sauvage, 1883; Woodward, 1919; 32 Biese, 1927; Bravi, 1988; Lambers, 1999; Bonde, 2004; Bravi et al., 2014; Schultze and Arratia 33 2015; Poyato-Ariza and Martín-Abad, 2016; Olive et al., 2022), North America (Alvarado-Ortega 34 and Brito, 2016), South America (Brito and Gallo, 2002; Chiappe et al., 1998; Giordiano et al., 2018), Africa (Saint-Seine, 1955) and the Middle East (Janensch, 1925; Basse et al., 1954; El Hajj 35 36 et al., 2021), pleuropholids are currently accepted as basal teleosts (Patterson, 1973, 1977; Pinna, 1996) although their phylogenetic relationships within Teleostei are still unclear (López-Arbarello 37 38 et al., 2008). Today, pleuropholids are represented by five genera (i.e. *Pleuropholis* Egerton, 1858, 39 Parapleuropholis Saint-Seine, 1955, Austropleuropholis Saint-Seine, 1955, Gondwanapleuropholis 40 Brito and Gallo, 2002, and Zurupleuropholis Giordano et al., 2018) and about 24 nominal species 41 comprising relatively small (usually < 10 cm in total length), slender fishes, readily recognized by 42 their small and upturned mouth, the numerous supra and infraorbital bones, and the presence of a single row of deep (i.e. dorsoventrally elongated) flank scales covering almost the whole side of the 43 44 body (Brito and Gallo, 2002; López-Arbarello et al., 2008; Arratia, 2013).

In the present paper we describe a new species of *Pleuropholis* from the Akrabou
Formation, which is clearly distinguishable from all the other known species. In addition to the
inherent importance of this discovery, the presence of this taxon in the Upper Cretaceous of
Morocco increases the distribution and diversity of pleuropholids and represents the youngest
member of this family.

50 The specimen described here comes from the Plattenkalk Konservat-Lagerstätte of Agoult, 51 located at the top of the Gara Sbâa hill, in the Kem Kem region of southeastern Morocco (Cavin et 52 al., 2010: fig. 2; Martill et al., 2011: fig. 1). These transgressive laminated limestones (Gara Sbâa

53	Member) correspond to the base of the Akrabou Formation and assigned to the upper Cenomanian-
54	?lower Turonian (Cavin et al., 2010; Martill et al., 2011). This site has mainly yielded a rich and
55	diverse actinopterygian assemblage (Cavin and Dutheil, 1999; Murray and Wilson, 2009, 2011,
56	2013, 2014; Murray et al., 2013; Taverne and Capasso, 2021); the assemblage also includes
57	ichnofossils, plants, worms, arthropods, echinoderms, elasmobranchs, and an undescribed lizard
58	specimen (Garassino et al., 2008; Guinot et al., 2008; Martill et al., 2011; Engel et al., 2012;
59	Krassilov and Bacchia, 2013; Vullo et al., 2016; Lamsdell et al., 2020). The Agoult fossil
60	assemblage, consisting of both marine and terrestrial organisms, indicates that these laminated
61	carbonates were deposited in a shallow marine coastal environment, most likely a quiet muddy
62	lagoon (Garassino et al., 2008; Cavin et al., 2010; Martill et al., 2011; Ibrahim et al., 2020).
63	
64	2. Material and methods
65	
66	The specimen is cataloged in the collections of the Muséum d'Histoire naturelle de
67	Marrakech, Morocco (MHNM), but is currently housed at the Muséum national d'Histoire naturelle
68	of Paris, France (MNHN), for study, within an agreement between both museums. Photographs
69	were taken using ultraviolet light, following the technique proposed by Tischlinger and Arratia
70	(2013).
71	General anatomical terminology follows Brito (1997). For the anatomy of the caudal region,
72	we used the terminology proposed by Arratia (2008).
73	The publication and the newly described species are registered in Zoobank. The LSID (Life
74	Science Identifier) for the publication is: urn:lsid:zoobank.org:pub:93E130B3-AFBE-480F-BA42-
75	9EA726DDD0E8; the LSID for the new species is: urn:lsid:zoobank.org:act:21F4D5D2-06E9-
76	4846-8504-F43B230DABC6.
77	
78	3. Results

79	
80	3.1. Systematic palaeontology
81	
82	Infraclass Teleostei Müller, 1846 (sensu Arratia, 1999)
83	Order incertae sedis
84	Family Pleuropholidae Saint-Seine, 1949
85	Genus Pleuropholis Egerton, 1858
86	Type species: P. attenuata Egerton, 1858
87	Pleuropholis danielae sp. nov.
88	LSID: urn:lsid:zoobank.org:act:21F4D5D2-06E9-4846-8504-F43B230DABC6.
89	Figs. 1–5
90	Diagnosis. Large-sized pleuropholid differing from all other species of the genus by its slenderer
91	body shape, the absence of a leptolepid notch in the dentary, a broad maxilla, covering almost the
92	entirely lower jaw, and flank scales with an unserrated posterior margin.
93	Holotype. MHNM.KK.393, complete flattened specimen preserved in left lateral view.
94	Etymology. The specific epithet is dedicated to Daniela Vullo, the mother of one of us (R.V.), who
95	passed away in May 2022, for her constant support.
96	Type-locality and horizon. The specimen comes from outcrops of the Gara Sbaa hill, near the
97	locality of Agoult, in the Kem Kem region of southeastern Morocco. Gara Sbaa Member of the
98	Akrabou Formation, Upper Cretaceous (Cenomanian).
99	
100	3.2. Description
101	
102	This is a large-sized <i>Pleuropholis</i> (115 mm in standard length and about 130 mm in total
103	length) and by far one of the largest known pleuropholid fishes. The body is very elongated and

104 slender (standard length/body height = 7.7) (Fig. 1). The deepest part of the body is constant

between the pectoral and the anal fin (body height = 15 mm). The head, including the opercular
series, measures approximately 1/6 of the standard length. The dermal bones and scales are covered
with ganoin.

108 The description of some skull elements is difficult due to poor preservation. Most of the 109 skull bones are recognizable, but in some cases, some elements are damaged. This situation makes 110 it difficult to interpret the shape of several bones, especially in the upper jaw.

111 Only a small fragment of the rostral is preserved and both its shape and suture with the 112 posterior bones are not clearly discernible. The nasal is longer than wide, with its anterior half slightly thinner than the posterior one. The nasal overlaps the frontal posteriorly, and contacts the 113 114 antorbital antero-laterally. The frontal lies just behind the nasal, and is the longest element of the dermal skull, having a length approximately three times that of the nasal. It extends posteriorly to 115 the parietal and dermopterotic. The parietal is a well-developed bone, slightly longer than wide and 116 having approximately half of the length of the frontal. The lateral border of the parietal is almost 117 118 linear and contacts ventrally the dermosterotic. The dermosterotic is a subrectangular bone, placed 119 lateral to the parietal and posterior to the frontal. This bone is directed backwards, narrowing at its 120 most posterior part. The extrascapular is much smaller than the parietal. The occipital sensory canal passes through the mid-length of the extrascapulars. Anteriorly, the extrascapular partly overlaps 121 122 the posterior margins of the parietal and the dermopterotic; posteriorly, it overlaps the anterior 123 margin of the posttemporal, and laterally, it is in contact with the dorsal part of the opercle.

The orbital series is formed by one antorbital, the lacrimal, four infraorbitals, the dermosphenotic, and at least two supraorbitals (Figs. 2, 3). The antorbital is a subtriangular bone; this bone is positioned above the lacrimal. The lacrimal is an elongated bone with the infraorbital canal extending into it. Both antorbital and lacrimal form the anterior margin of the orbit. The first two infraorbitals border the orbit ventrally, followed by two covering the postorbital region of the orbit. The infraorbital canal extends in the middle part of the bones. The posterior infraorbitals border the anterior edge of the vertical limb of the preopercle. Although not very well preserved,

the dermosphenotic is a somewhat diamond-shaped bone that borders the posterior dorsal angle of the orbit. It is in contact with the anterior part of the dorsal edge of infraorbital 4. At least two supraorbitals are preserved, forming the dorsal margin of the orbit.

As a characteristic of the genus, the jaws are short, upturned, and positioned ahead of the orbit. The maxilla is poorly preserved with only few fragments present. However, these fragments point to an extremely broadened bone that covers almost entirely the lower jaws. It has a straight dorsal edge. The supramaxilla is not preserved.

The lower jaw seems to be somewhat triangular; the dentary and the angular bones are exposed laterally (Figs. 2, 3). The dentary is a short stout subtriangular bone that presents a rounded symphysis. The mandibular sensory canal extends along the ventral border of the bone. Contrary to the pattern found in *Pleuropholis decastroi* Bravi, 1988, from the Albian of Italy (cf. Taverne and Capasso, 2019) and *P. germinalis* Olive, Taverne and Brito, 2022 from the Barremian–Aptian of Belgium the new species does not present a leptolepid notch in the median-dorsal part of the dentary. Posteriorly, the dentary sutures with the angular. The jaw looks toothless.

The preopercle is an L-shaped bone in which the horizontal limb is slightly larger than the vertical limb. These limbs are placed perpendicular to each other (Figs. 2, 3). The preopercular canal is close and parallel to the ventral and dorsal borders of the bone. The opercular series consists of the large opercle, a subtriangular subopercle and an elongated interopercle. Only part a relatively narrow branchiostegal ray is preserved.

The posttemporal bone is somewhat oval and located behind the skull; the posterior edge overlies the dorsal end of the supracleithrum. The supracleithrum laterally overlaps the dorsal end of the cleithrum. The cleithrum has an elongated lower arm, less developed than the dorsal one. The pectoral fin is well developed and nine rays are preserved. It is positioned close to the ventral margin of the body, on the lower border between the flank scales and the scales immediately below. The pelvic fin is located at the level of the 15th transverse row of scales. Only four branched rays are preserved. No bones of the pelvic girdle are visible.

Both the dorsal and the anal fins are triangular and located posteriorly in the body, at the level of the 29th and 26th transversal row of scales, respectively (Figs. 1, 4). The dorsal fin presents two short, unsegmented procurrent ray followed by seven principal rays. The first ray is segmented, longer and thicker than the preceding procurrent rays. The longest ray of the dorsal fin is the second principal ray, which is segmented and branched. Elongated fringing fulcra are placed in the anterior margin of the first and the second principal rays.

163 The anal fin is larger than the dorsal fin. Procurrent rays are not preserved in the specimen 164 that has about 13 branched principal rays. The first and the second rays seem to have the same 165 length. The main leading ray is preceded by numerous fringing fulcra. After the third ray, they 166 decrease in size and width.

167 The vertebrae as well as the caudal endoskeleton are completely hidden by the scales and 168 thus not observable in the studied specimen. The caudal fin is almost completely preserved, 169 although some few elements are broken.

The caudal fin has a forked outline. As in other species of the genus (e.g. Egerton 1858: pl.
7, fig. 4; Woodward, 1919: text-fig. 36; Taverne and Capasso, 2019: fig. 8), the upper lobe is
slightly longer than the lower lobe (Figs. 1, 5). There are about fourteen epaxial basal fulcra, a
series of elongated, small, epaxial fringing fulcra, 15 to 16 principal rays, a series of hypaxial
fringing fulcra, segmented procurrent rays, and two hypaxial basal fulcra.

The epaxial basal fulcra are elongated, unsegmented leaf-like elements, partially covering
the next fulcra. The epaxial fringing fulcra are associated with the first and second principal rays.
The first principal ray is shorter than the following segmented-and-branched principal ray. Both,
together form the dorsal leading margin of the fin.

In the lower lobe, the hypaxial procurrent rays are segmented with their terminal segment resembling the fringing fulcra; fringing fulcra extend along the ventral margin of the last principal rays. As in the upper lobe, the last principal ray is shorter than the following segmented-andbranched principal ray (Fig. 5).

183	There are at least 39 lateral rows of thick, ganoid-type scales, from the supracleithrum to the
184	base of the caudal fin (Fig. 5). These scales have different shapes according to the region of the
185	body. The flank scales are deeper than long, progressively decreasing in size posteriorly and
186	becoming diamond-shaped in the caudal region. They present no serrations in the posterior edge.
187	Dorsal to the flank scales there are two rows of small scales. Ventral to the flank scales, there are
188	three rows of small, rectangular scales.
189	
190	4. Discussion
191	
192	Pleuropholis danielae sp. nov. possesses synapomorphies of the family Pleuropholidae,
193	including: the presence of a single row of deepened ganoid scales covering almost the whole side of
194	the body, an edentulous mouth, and a complete orbital series formed by infraorbital and supraorbital
195	bones (cf. Saint-Seine, 1949; Brito and Gallo, 2002; Olive et al., 2022).
196	The presence of an upturned maxilla and a jaw articulation lying anterior to the orbit (cf.
197	Woodward, 1919; Saint-Seine, 1949; Patterson, 1973; Bravi, 1988; Alvarado-Ortega and Brito,
198	2016; Taverne and Capasso, 2019; Olive et al., 2022) places the new taxon in the genus
199	Pleuropholis.
200	One of the biggest problems concerning the genus <i>Pleuropholis</i> is the validity of most of its
201	contained species, especially those from the Jurassic. Therefore, a comprehensive revision of all
202	these taxa would be necessary, especially because many of them lack any autapomorphies to sustain
203	their taxonomic status, or they are based on body proportions from imperfect or distorted specimens
204	(e.g. Patterson, 1973; Alvarado-Ortega and Brito, 2016; Taverne and Capasso, 2019; Olive et al.,
205	2022).
206	Despite these problems, P. danielae sp. nov. clearly differs from the other species of

207 *Pleuropholis* by its considerably more slender body shape. The absence of a leptolepid notch in the

208 dentary distinguishes *P. danielae* n. sp from *P. jamotti* Saint-Seine, 1955 (a species, like *P. lannoyi*

209 from the same locality, that needs taxonomic revision and does not appear to belong to the genus

210 Pleuropholis), P. decastroi Bravi, 1988, and P. germinalis Olive, Taverne and Brito, 2022.

211 Although not well preserved, the maxilla of *P. danielae* sp. nov. seems to be broad, almost entirely

covering the lower jaw, a condition also known in *P. thiollieri* Sauvage, 1883, *P. decastroi* and *P.*

213 *cisnerosorum* Alvarado-Ortega and Brito, 2016. Finally, flank scales with an unserrated posterior

214 margin are known in *P. germinalis, P. longicauda* Egerton, 1858, *P. formosa* Woodward, 1919, *P.*

215 attenuata Egerton, 1858, and P. crassicauda Egerton, 1858.

216 From a palaeobiogeographical point of view, the evolution and distribution of *Pleuropholis* appear to have been centered in the Tethyan realm (Fig. 6). So far, the only unambiguous Middle 217 218 Jurassic occurrence of *Pleuropholis* is from the Bajocian–Bathonian of southern Italy (Bravi et al., 219 2014); this single occurrence, representing the oldest known record of *Pleuropholis*, is indicative of 220 a low taxic diversity at that time and probably reflects a restricted distribution at the beginning of the evolutionary history of the lineage (neoendemism; see Cowman et al., 2017). The genus 221 222 *Pleuropholis* reached its apparent highest diversity during the Late Jurassic–earliest Cretaceous (an 223 observation possibly partly due to a preservational bias; see Cavin, 2010), resulting in several 224 European species described from Cerin and the Meuse department in France (Sauvage, 1883; Saint-Seine, 1949), Bavaria in Germany (Egerton, 1858; Biese, 1927), and the English Purbeck and 225 226 coeval beds in Denmark (Egerton, 1858; Woodward, 1919; Bonde, 2004). Two additional species, 227 P. cisnerosorum from the Kimmeridgian of southern Mexico (Alvarado-Ortega and Brito, 2016) 228 and P. bassei Lehman in Basse et al., 1954 from the Tithonian-Berriasian of Yemen (Basse et al., 1954), indicate westward dispersal to the Hispanic Corridor and eastward dispersal to the Arabian 229 230 platform, respectively; furthermore, pending a revision of *P. jamotti* and *P. lannoyi* Saint-Seine, 231 1955, the genus also occurs in Upper Jurassic freshwater deposits of the Democratic Republic of 232 Congo (Saint-Seine, 1955; Caillaud et al., 2017). During the Early Cretaceous, the genus 233 Pleuropholis appears to have been decreasing in diversity and less widely distributed; it is known 234 from five localities in the Middle East and Western Europe: the Barremian of Lebanon (Janensh,

1924; El Hajj et al., 2021), the Barremian of El Montsec and Las Hoyas in eastern Spain (Wenz, 235 1968; Sanz et al., 1988; Poyato-Ariza and Martín-Abad, 2016), the Barremian–Aptian of Belgium 236 237 (Olive et al., 2022), and the Albian of southern Italy (Bravi, 1988); with the exception of the latter, 238 all these occurrences are from continental localities, indicating that *Pleuropholis* was a euryhaline genus that colonized brackish and freshwater environments at many times between the Late Jurassic 239 240 and the middle Early Cretaceous. The present study reports the first Upper Cretaceous occurrence 241 of Pleuropholis; therefore, P. danielae sp. nov. from the Cenomanian of Agoult, southeastern Morocco, represents the youngest known member of the genus and the last known occurrence of 242 pleuropholid fishes (Fig. 6). The narrowly distributed species P. decastroi (Albian of Italy) and P. 243 244 danielae sp. nov. suggest a range contraction for this formerly widespread genus, with the Mediterranean Tethys as a center of survival (paleoendemism; see Cowman et al., 2017). The new 245 discovery extends the stratigraphic range of this group by about 15 million years and makes 246 *Pleuropholis* one of the most temporally long-lived actinopterygian genera of all time, with a fossil 247 248 record spanning ~75 million years. Based on 1108 time ranges calculated from Cavin et al.'s 249 (2021a) database, Cavin et al. (2021b: fig. 7) found that the average time duration of the ray-finned 250 fish genera having their first occurrence in the Devonian–Palaeocene interval is 13.3 million years all environments combined, with only a dozen genera having a longer range than that of 251 252 Pleuropholis.

253 The discovery of *P. danielae* sp. nov. confirms that several relictual lineages of fishes that 254 had been flourishing during the Late Jurassic persisted until the early Late Cretaceous (a period during which they were probably severely affected by competition with diversifying modern 255 256 teleosts; Guinot and Cavin, 2016), and apparently became extinct at the end of the Cenomanian, 257 possibly in relation to the Cenomanian-Turonian Boundary Event (CTBE). These forms include 258 some holosteans such as the ionoscopiforms Spathiurus (Ionoscopidae) and Petalopteryx 259 (?Ophiopsiellidae) from Lebanon and Agoultichthys (?Ophiopseliidae) from Morocco (Ebert, 2018; 260 El Hossny et al., 2020), the probable ionoscopiform Aphanepygus (?Ophiopsiellidae) from Croatia

261	and Lebanon (Bartram, 1977), and some basal teleosts such as the newly described Pleuropholis
262	(Pleuropholidae) species from Morocco. It is likely that these declining groups, known from very
263	few specimens in Lebanese, Moroccan, and Croatian Cenomanian Konservat-Lagerstätten, were
264	relatively rare in early Late Cretaceous (pre-CTBE) fish faunas and palaeoendemic, i.e.
265	geographically restricted to small areas of the Mediterranean Tethys (Levant Platform, Atlas Gulf
266	of the Saharan Platform, Apulian terrane). This suggests that the apparent absence in Cenomanian
267	strata of other groups of ray-finned fishes with last known occurrences being from Albian deposits
268	(e.g. macrosemiids and callipurbeckiid semionotiforms) could be due to problems with the
269	detection of rare species in the fossil record (rarity bias), especially in undersampled Plattenkalk
270	localities and understudied areas (collecting bias).
271	
272	5. Conclusions
273	
274	The Cenomanian beds of the Akrabou Formation of southeastern Morocco yielded a single
275	specimen of pleuropholid fish, which is recognized as belonging to a new species of <i>Pleuropholis</i> .
276	Interestingly, P. danielae sp. nov. is the first Late Cretaceous Pleuropholidae and thus represents
277	the youngest known member of this family of basal teleosts. This discovery contributes to our
278	understanding of the mid-Cretaceous turnover of marine fish faunas, which is a significant
279	biological event characterized by the apparent extinction of relictual lineages such as pleuropholids
280	and the diversification of modern groups such as neoteleosts.
281	
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283	
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289	
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447 Figure Captions

448

Fig. 1. *Pleuropholis danielae* sp. nov., holotype MHNM.KK.393 photographed under natural light
(A) and UV light (B), in left lateral view. Scale bar equals 10 mm.

451

- 452 Fig. 2. *Pleuropholis danielae* sp. nov., holotype MHNM.KK.393, detail of the head region
- 453 (photograph under UV light) (A) and interpretative line drawing (B). Abbreviations: Ang, angular;
- 454 Ao, antorbital; Cl, cleithrum; De, dentary; Dsp, dermosphenotic; Dpt, dermopterotic; Ex,
- 455 extrascapular; Fr, frontal; Io, infraorbital; Iop, interopercle; La, lacrimal; Mx, maxilla; Na, nasal;
- 456 Op, opercle; Pa, parietal; Pmx, premaxilla; Pop, preopercle; Pt, posttemporal; Ro, rostral; Scl,
- 457 supracleithrum; So, supraorbital; Sop, subopercle. Scale bar equals 10 mm.

458

- Fig. 3. Reconstruction of the skull of *Pleuropholis danielae* sp. nov. Abbreviations: Ang, angular;
 Ao, antorbital; De, dentary; Dsp, dermosphenotic; Dpt, dermopterotic; Ex, extrascapular; Fr,
 frontal; Io, infraorbital; Iop, interopercle; La, lacrimal; Mx, maxilla; Na, nasal; Op, opercle; Pa,
 parietal; Pmx, premaxilla; Pop, preopercle; Pt, posttemporal; Ro, rostral; Smx, supramaxilla; So,
- 463 supraorbital; Sop, subopercle. Scale bar equals 10 mm.

464

- 465 Fig. 4. *Pleuropholis danielae* sp. nov., holotype MHNM.KK.393, detail of the dorsal and anal fins
 466 (photograph under UV light). Scale bar equals 10 mm.
- 467

Fig. 5. *Pleuropholis danielae* sp. nov., holotype MHNM.KK.393, detail of the caudal region
(photograph under UV light) (A) and interpretative line drawing (B). Abbreviations: 1st PR, first
principal ray; ebfu, epaxial basal fulcra; f.f, fringing fulcra; hbfu, hipaxial basal fulcra; l. PR, last
principal ray; v.pr, ventral procurrent rays. Scale bar equals 10 mm.

- 473 Fig. 6. Simplified time-scaled distribution of the genus *Pleuropholis*, showing the various known
- 474 occurrences from the Middle Jurassic to the early Late Cretaceous (see text for details and
- 475 references). Some nominal species (i.e. P. lannoyi from the Kimmeridgian-Tithonian of the
- 476 Democratic Republic of Congo, P. egertoni and P. obtusirostris from the Kimmeridgian of France,
- P. pompecki and P. wagneri from the Kimmeridgian–Tithonian of Germany, and P. crassicauda 477
- 478 and P. longicauda from the Berriasian of England), co-occurring with some of those presented here,
- 479 are not included.

Juinal



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A pleuropholid fish specimen is described from the Cenomanian of Agoult, SE Morocco.

The specimen is assigned to a new species of the genus Pleuropholis, P. danielae.

This is the first known occurrence of Pleuropholidae in the Upper Cretaceous.

Pleuropholid fishes may have become extinct at the Cenomanian–Turonian boundary.

This adds to our understanding of the mid-Cretaceous turnover of marine fish faunas.

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Declaration of interests

 \boxtimes The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

□ The authors declare the following financial interests/personal relationships which may be considered as potential competing interests: