

Large predatory marine reptiles from the Albian-Cenomanian of Annopol, Poland

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ABSTRACT - During the Early-Late Cretaceous transition, marine ecosystems in Eurasia hosted a diverse set of large predatory reptiles that occupied various niches. However, most of our current knowledge of these animals is restricted to a small number of bonebed-like deposits. Little is known of the geographical and temporal extent of such associations. The middle Albian - middle Cenomanian phosphorite-bearing succession exposed at Annopol, Poland produces numerous ichthyosaurian and plesiosaurian fossils. These are mostly isolated skeletal elements (e.g. teeth, vertebrae), but disarticulated partial skeletons and an articulated, subvertically embedded ichthyosaur skull are also available. The following taxa are identified: *'Platypterygius'* sp., cf. Ophthalmosaurinae, Ichthyosauria indet, *Polyptychodon interruptus*, Pliosauridae indet., Elasmosauridae indet. and Plesiosauria indet. The large-sized ichthyosaur *'Platypterygius'* and the pliosaurid *Polyptychodon interruptus* predominate within the upper Albian - middle Cenomanian deposits. The Annopol record, combined with data from England, France and western Russia, suggests that *'Platypterygius'* and *Polyptychodon interruptus* formed a long-term, stable ecological sympatry in marine ecosystems of the European archipelago, at least during the Albian - middle Cenomanian. In addition, the marine reptile assemblage from Annopol is distinct from other Eurasian ecosystems in containing also elasmosaurids in its Albian portion.

KEYWORDS: Plesiosauria, Ichthyosauria, sympatry, mid-Cretaceous, Poland.

1. INTRODUCTION

The sympatric association of large-sized apex predators belonging to the same taxon or not, irrespective of taxonomic level (i.e. species, family, order), appears to be common both in ancient and present-day marine ecosystems. It occurs notably in modern killer whales (Foote *et al.* 2011), Late Jurassic metriorhynchid crocodyliforms (Young *et al.* 2012), latest Cretaceous mosasauroids and selachians (Vincent *et al.* 2013) and Miocene sharks and raptorial sperm whales (Lambert *et al.* 2010). However, the geographical extent of these associations appears reduced, and they are usually poorly known in the fossil record.

A similar association of large predators has been reported among marine reptile assemblages from a limited number of upper Albian - upper Cenomanian (mid-Cretaceous) successions in western Europe, with the co-occurrence of the platypterygiine ophthalmosaurid *'Platypterygius'* and the pliosaurid *Polyptychodon interruptus*. This association has been documented mainly from the Cambridge Greensand Member (i.e. the base of the Lower Chalk Formation) in eastern England (Owen, 1840, 1851-1864; Seeley, 1869, 1876; Fischer *et al.* 2014c) and in the Paris Basin (Sauvage, 1882; Buffetaut, Tomasson & Tong, 2003; Buffetaut *et al.* 2005; Fischer, 2012). Eichwald (1853, 1865-1868) and Kiprijanoff (1881, 1882, 1883a, b) described similar associations in the Kursk region in Russia. With the exception of the French specimens, these remains usually occur in reworked bonebed-like deposits that originated during the early phases of the great mid-Cretaceous transgression. The time represented by such bone-rich deposits is comparatively limited geologically speaking, at least for the English greensands (Cookson & Hughes, 1964; Hopson, 2005; Woods *et al.* 2008; Hopson, Wilkinson & Woods, 2008; Owen, 2012).

Here we report the presence of the *'Platypterygius'*-*Polyptychodon interruptus* association from the mid-Cretaceous condensed succession exposed in Annopol, Poland, also documenting the remainder of the ichthyosaur/plesiosaur fauna from this area. This locality has yielded a vast number of vertebrate and

invertebrate fossils, providing a representative picture of the evolution of a marine ecosystem within the European archipelago at the transition between the Early and Late Cretaceous.

In addition to ichthyosaur and plesiosaur remains, those of bony fish, lamniform sharks, chimaeroids (Radwanski, 1968; Marcinowski & Radwanski, 1983; Popov & Machalski, 2014), marine protostegid turtles and pterosaurs (ornithocheiroids and/or azhdarchoids; see Machalski & Martill, 2013) are known from the Annapol succession. Prior to the present study, remains of ichthyosaurs and plesiosaurs had merely been listed but not described by Samsonowicz (1925, 1934), Ciesliński & Milakovic (1962), Marcinowski & Radwanski (1983) and Machalski, Komorowski & Harasimiuk (2009).

2. INSTITUTIONAL ABBREVIATIONS

CAMSM: Sedgwick Museum of Earth Sciences, Cambridge University, Cambridge, UK; IRSNB: Royal Belgian Institute of Natural Sciences, Brussels, Belgium; MWGUW: Geological Museum of the Geology Department, University of Warsaw, Poland; NHMUK: Natural History Museum, London, UK; RGHP: Réserve Naturelle Géologique de Haute-Provence, Digne-les-Bains, France; SSU: Saratov State University, Saratov, Saratov Oblast', Russian Federation; ZPAL: Institute of Paleobiology, Polish Academy of Sciences, Warszawa, Poland.

3. GEOLOGICAL SETTING

The ichthyosaur and plesiosaur remains studied come from the middle Albian - middle Cenomanian part of the mid-Cretaceous (uppermost lower Albian - lower Turonian) condensed, phosphorite-bearing succession as exposed along the limbs of the Annapol anticline, central Poland (Fig. 1a-c). This area is located on the east bank of the Wisła (Vistula) River, central Poland (Fig. 1b) and forms the easternmost part of the so-called Mesozoic Border of the Holy Cross Mountains (Marcinowski & Radwanski, 1983).

The lithology of the mid-Cretaceous succession in the Annapol area was described by Samsonowicz (1925, 1934), Pozaryski (1947), Ciesliński (1959) and Walaszczyk (unpub. M. Sc. thesis, University of Warszawa, 1984; 1987). The biostratigraphy is based mainly on ammonites, inoceramid bivalves and foraminifera (Samsonowicz, 1925, 1934; Pozaryski, 1947; Ciesliński, 1959, 1987; Marcinowski, 1980; Marcinowski & Radwanski, 1983, 1989; Peryt, 1983; Marcinowski & Walaszczyk, 1985; Marcinowski & Wiedmann, 1985, 1990; Walaszczyk, 1992). The upper Albian ammonite biostratigraphy has recently been revised by Machalski & Kennedy (2013).

The Annapol sedimentary succession may be subdivided into a series of thin transgressive units, based on the model of Juignet (1980) and Hancock (1990). These units are capped by layers of reworked phosphatic nodules and clasts (often with biostratigraphically mixed ammonite assemblages), hardgrounds and burrowed omission and/or erosional surfaces (Fig. 1a), reflecting a series of regression maxima. The general bathymetrical trend for the Annapol area is an upwards deepening, as based on foraminiferal spectra and facies development (Walaszczyk, 1987). The latter is characterized by a transition from terrigenous to pelagic carbonate sedimentation, reflecting the progress of the mid-Cretaceous transgression (Ciesliński, 1976; Marcinowski & Radwanski, 1983). At the bottom, the uppermost lower Albian sands and sandstones (unit 2 in Fig. 1a) contain *Ophiomorpha* burrows which point to deposition in extremely shallow-marine, marginal environments (compare Radwanski, Wysocka & Gorka, 2012). There are pure chalks with flints at the top of succession, corresponding to a relatively deep-marine, offshore setting (unit 9 in Fig. 1a). The Annapol succession is extremely thin in comparison to coeval strata in adjacent areas due to its position on a submarine high (Ciesliński, 1976).

The fossils studied originate from the interval that ranges from the upper part of the Albian unit 2 to the top of the Cenomanian unit 5 (Fig. 1a). Most of this material is from unit 3, which is the most fossiliferous interval of the succession. This unit was deposited during a transgressive pulse in the late Albian ('Vraconnian', *sensu* Amédéo, 2002) in a relatively shallow-marine, highly productive, phosphogenic environment (Walaszczyk, 1987). Unit 3 and the top of the underlying unit 2 were collectively referred to as 'the Phosphorite Bed' by previous authors (e.g. Marcinowski & Wiedmann, 1985; Marcinowski & Radwanski, 1983, 1989). Those authors subdivided the Phosphorite Bed into two parts: a lower and an upper part (Marcinowski & Radwanski, 1983, fig. 6). More detailed accounts on fossil content, sedimentary history and stratigraphy of particular bone-bearing levels of the Annapol succession are given in Popov & Machalski (2014) and Machalski & Kennedy (2013).

4. MATERIAL AND METHODS

The present study is based on two collections. One of these was amassed before World War II by Jan Samsonowicz, a geologist who discovered the Annopol anticline in 1923. This collection is housed at the Geological Museum of the Geology Department, University of Warsaw (prefix MWGUW). Another collection was acquired from the Annopol area between 2008 and 2012 by a team led by one of us (MM); additionally, a large and well-preserved pliosaur tooth found in 2014 by the same team has been included in this study. All this material is housed at the Institute of Paleobiology, Polish Academy of Sciences (abbreviated ZPAL), Warsaw, Poland.

4.a. MWGUW collection

This collection is composed exclusively of isolated skeletal elements (teeth and vertebrae, 30 in total) with no precise provenance data within the Annopol area and succession. There are nine teeth and two vertebrae of ichthyosaurs and ten teeth and nine vertebrae of pliosaurs. In view of their preservation, all these fossils must have come from the Albian part of the succession (top of unit 2 and unit 3; see Fig. 1a). Based on the nature of the matrix, which is sandy in the former and marly in the latter, only specimens with adhering matrix in the MWGUW collection can be more precisely allocated either to the upper part of unit 2 or to unit 3 (Właszczyk, 1987). As a consequence, middle or early late Albian and a late late Albian age, respectively, is assigned to these specimens. As far as the specimens without adhering matrix in the MWGUW material are concerned, their most probable source is unit 3 (especially its lower part) based on *in situ* observations at the succession exposed underground (see the following section). Consequently, they are most probably of late late Albian age.

4.b. ZPAL collection

The ZPAL collection is dominated by ichthyosaur remains, including 20 isolated teeth and vertebrae, five disarticulated partial skeletons and a single skull. In addition, six plesiosaur teeth are available. In view of the presence of disarticulated partial skeletons, the number of catalogued elements is larger than that of individuals.

The remains studied come from two localities situated along the northern limb of the Annopol anticline (Fig. 1b): (1) the abandoned underground phosphorite mine *Jan 1*; and (2) a series of temporary trenches dug near the old sandstone quarry south of the village of Kopiec, referred to as the locality of Kopiec.

The specimens from the mine have been collected *in situ* in the interval from the bottom of unit 3 to the top of unit 5 (Fig. 1a); they are therefore late late Albian - middle Cenomanian in age. They represent a full spectrum of preservational states, from isolated elements (teeth and vertebrae), through rare disarticulated partial skeletons to a single articulated (albeit fragmentary) skull. In the Albian part of the underground succession, the majority of the well-preserved isolated elements without adhering phosphatic matrix were recovered from the lowermost part of unit 3.

In contrast, the material collected at Kopiec is represented exclusively by isolated elements. It stems from screen washing of Albian sands, forming the matrix of the Phosphorite Bed of previous authors (top of unit 2 and unit 3). Two screens with an 8 mm mesh size and a motorized pump were used. Material for washing was amassed by using an excavator and, for technical reasons, this could be subdivided into two portions. The lower portion corresponded approximately to the highest 20 cm of unit 2 and the lowest 20 cm of unit 3 (Fig. 1a). The upper portion corresponded to the remainder of unit 3. However, some mixing of sediment (and fossils) from these (and adjacent) horizons was unavoidable during field operations.

For reasons outlined above, the stratigraphic position of the reptile remains recovered by means of screen washing at Kopiec could not be precisely determined. Specimens with adhering phosphatic matrix form a notable exception, analogous to the MWGUW collection. An assignment either to the phosphorite bed at the top of unit 2 or to the phosphorite bed in the upper part of unit 3 is possible, based on differences in phosphatic matrix. Specimens without adhering matrix are thought to have come mostly from unit 3, in particular from its lower part, based on the abovementioned relationships observed in the mine. These loose elements are therefore most probably of late late Albian age.

Figure 1. (a) The mid-Cretaceous succession of the Annapol anticline (modified after Machalski & Kennedy, 2013). 1: quartzose sandstone and quartz sands truncated by a burrowed surface; 2: quartz sands with glauconite, rare sandstone nodules near the top, and phosphate horizon at the top; 3: quartz sands and marly sands with glauconite and phosphate horizon in the upper part; 4: quartzose marls with glauconite and phosphates; 5: glauconitic marls infilling earlier burrows, truncated by a hardground; 6: strongly glauconitic marls with burrowed omission surface at the top; 7: limestone with rare phosphatic nodules and glauconite, capped by a hardground; 8: marls with burrowed surface at the top; 9: marls with black flint; b: burrows, **(b)** Sketch map of the Annapol anticline with localities studied and location within Poland (HCM: Holy Cross Mountains), **(c)** A section exposed underground at the Annapol mine.

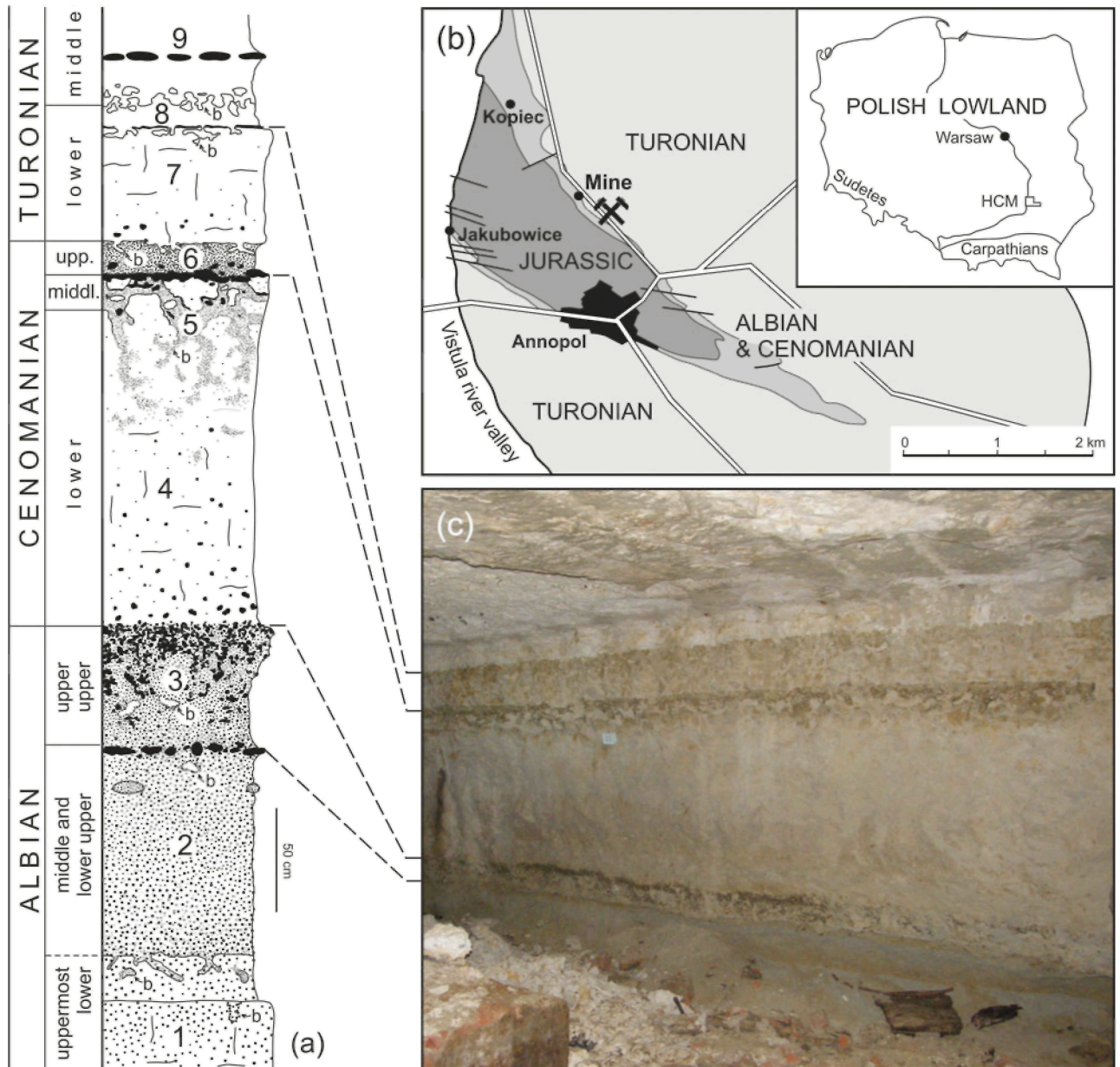
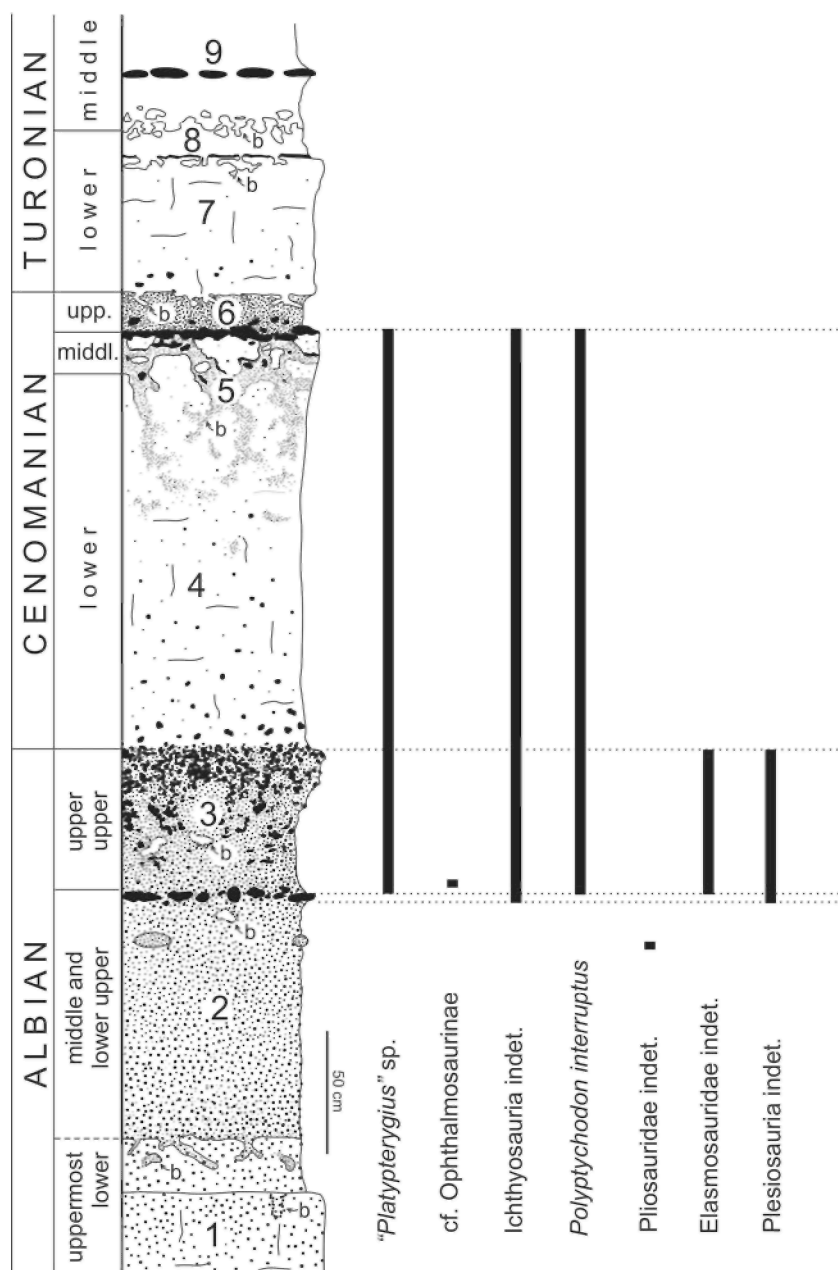


Figure 2. Stratigraphic ranges of the ichthyosaurs and plesiosaurs in the Annapol succession. See Figure 1 caption for characteristics of particular units.



5. TAPHONOMY

Seven ichthyosaur and plesiosaur taxa have been identified in the Annapol sedimentary succession (Fig. 2). These are *'Platypterygius'* sp., cf. Ophthalmosaurinae, Ichthyosauria indet, *Polyptychodon interruptus*, Pliosauridae indet., Elasmosauridae indet. and Plesiosauria indet. All these taxa are present in the Albian horizon corresponding to the Phosphorite Bed of previous authors (see Section 3) which ranges from the top of unit 2 to the top of unit 3 in the present subdivision. Only three taxa - *'Platypterygius'* sp., Ichthyosauria indet. and *Polyptychodon interruptus* - are recorded higher up section, ranging to the middle Cenomanian unit 5 (Fig. 2). The material studied can be assigned to three taphonomic categories: (1) isolated skeletal elements; (2) disarticulated partial skeletons; and (3) articulated partial skeletons.

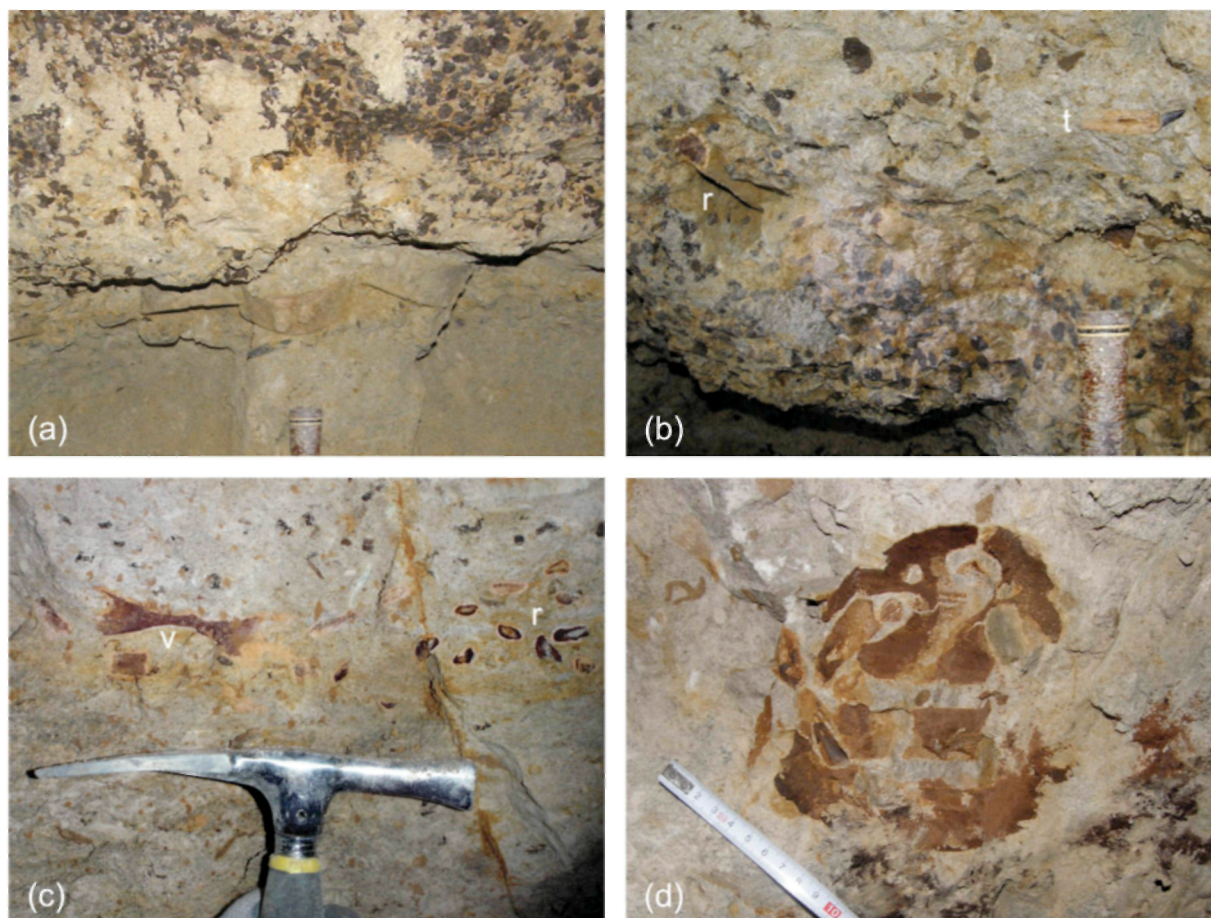
The isolated skeletal elements are mainly vertebrae, tooth crowns and complete teeth (with preserved roots), 52 specimens in total. Most of these are well preserved and show no or few signs of mechanical abrasion. The isolated tooth crowns may represent specimens shed by the living animals during feeding. In contrast, complete teeth and isolated vertebrae may stem either from drifting carcasses or from those which had already landed on

the sea floor, subsequently ripped apart by scavengers.

Disarticulated partial skeletons are represented by remains of five ichthyosaur individuals: (1) skull elements of an ichthyosaur referred to as cf. *Ophthalmosaurinae* (ZPAL V38/51-57, 59); (2) an accumulation of vertebrae of an indeterminate ichthyosaur (ZPAL V 38/433, 875, 888, 889), both from the lower part of unit 3; (3) jaw elements and well-preserved teeth of '*Platypterygius*' sp. (jaw bones ZPAL V38/4, 65-69, 72-81, 83-88; complete teeth ZPAL V38/1-3, 23-26, 28) from the upper part of unit 3; (4) jaw elements and teeth of '*Platypterygius*' sp. (V38/38, 44-47; teeth ZPAL V38/18, 42, 43; Fig. 3b) and (5) vertebrae and ribs of an indeterminate ichthyosaur (ZPAL V38/49 and 100-106; Fig. 3c), both from the lower part of unit 4. Only one of these concentrations (3) occurs in a reworked horizon (phosphatic bed in the upper part of unit 3). This taphonomic category most probably involves parts of carcasses dismembered on the sea floor by scavengers (as suggested by bite marks seen on some bones and teeth), subsequently locally reworked in the case of the individual characterized above.

Articulated partial skeletons are represented by a single fragmentary skull of an indeterminate ichthyosaur of early Cenomanian age. It is embedded in the limestone matrix of unit 4, and still visible in cross-section in the roof of an underground gallery (Fig. 3d). A fragmentary rostrum of the same individual is preserved in close proximity. The orientation of the skull suggests vertical landing of an ichthyosaur on the sea floor and embedding of its head in subvertical position within a firm mud. It can be speculated that the touchdown was relatively 'hard', which resulted in fracture of the rostrum. The remainder of the body was probably removed by scavengers and sea currents, possibly even prior to the lithification and eventual erosional truncation of the deposit containing these. Similar cases from Jurassic strata have been described by Martill (1993), Wetzel & Reisdorf (2007) and Wahl (2009).

Figure 3. Multielement finds of (a, c, d) *Ichthyosauria* indet. and (b) '*Platypterygius*' sp., based on field photographs at the Anapol mine, (a) Group of vertebrae, including ZPAL V.38/888 (see Fig. 5f), lower part of unit 3, late late Albian. (b) Ribs (r) and teeth (t) of '*Platypterygius*' sp., lower part of unit 4, early Cenomanian. (c) Ribs (r) and vertebrae (v) of *Ichthyosauria* indet., lower part of unit 4, early Cenomanian, photo A. Komorowski. (d) Skull in transverse cross-section, *Ichthyosauria* indet., top of unit 4, early Cenomanian.



6. SYSTEMATIC PALAEOLOGY

Order ICHTHYOSAURIA de Blainville, 1835
Family OPTHALMOSAURIDAE Baur, 1887
Subfamily PLATYPTERYGIINAE Arkhangel'sky, 2001 (*sensu* Fischer *et al.* 2012)
Genus: *Platypterygius* von Huene, 1922

Note. Recent analyses suggest that the platypterygiine *Platypterygius* is not monophyletic, representing a variety of platypterygiine taxa. The name has been applied by default for Cretaceous ichthyosaur material for decades (Fischer, 2012; Fischer *et al.* 2014a). An unambiguous definition of *Platypterygius* is still required (Fischer *et al.* 2014a); in the meantime, we use the name '*Platypterygius*' for large-sized platypterygiine specimens from the mid-Cretaceous of Eurasia.

'*Platypterygius*' sp.
Figures 4a-d, 5d

Referred material. Disarticulated partial skeletons: (1) fragmentary jaw bones ZPAL V38/4, 65-69, 72-81, 83-88 and complete teeth ZPAL V38/1-3, 23-26, 28. 2; and (2) fragmentary ribs ZPAL V38/38, 44-47 associated with teeth ZPAL V38/18, 42, 43. Isolated teeth: ZPAL V 38/427, 440, 690, 856 and MWGUW ZI/60/12-18.

Horizon and age. Units 3-5, late late Albian - middle Cenomanian.

Description and comparison. Teeth constitute the sole diagnostic element in the present material. The crown is conical, robust and straight (Figs 4a-c, 5d). Numerous apicobasal striations texture the black enamel and most apices are broken and polished, indicating moderate to intense tooth wear (e.g. Fischer *et al.* 2011a). Larger-sized teeth have a rugose enamel, a ridged acellular cementum ring and deep apicobasal grooves texturing the root, a combination commonly found in some species referred to as '*Platypterygius*' (e.g. Kiprijanoff, 1881; Bardet, 1989; Bardet, Wellenhofer & Herm, 1994; Maxwell, Caldwell & Lamoureux, 2011; Fischer, 2012). The root is square in cross-section, as in platypterygiine ophthalmosaurids (Fischer *et al.* 2012) and bulbous, as in some species referred to as *Platypterygius* (Fischer, 2012; Fischer *et al.* 2014c). More posterior teeth are smaller and have a 'ratched-shaped' apex. Their enamel is devoid of apicobasal striation.

Subfamily OPTHALMOSAURINAE Baur, 1887
(*sensu* Fischer *et al.* 2012)
cf. OPTHALMOSAURINAE
Figure 5e-1

Referred material. A disarticulated partial skeleton: skull elements ZPAL V38/51-57, 59, including ZPAL V38/51 (quadrate), ZPAL V38/52 (tooth), ZPAL V38/53 (angular), ZPAL V38/55 (surangular) and ZPALV38/57Gugal).

Horizon and age. Lower part of unit 3, late late Albian.

Description and comparison. The posterior half of the jugal (ZPAL V38/57) is preserved (Fig. 4g). The anterior ramus is slender but its cross-section is rounded. Posteriorly, the jugal forms a thin lamella; posteroventrally, this lamella possesses a facet, presumably for the anterior ramus of the quadratojugal, terminating anteriorly by a prominent and right-angled hook, a feature previously recorded solely in the ophthalmosaurine *Ophthalmosaurus icenicus* (see A. M. Kirton, unpub. Ph.D. thesis, University of Newcastle upon Tyne, UK, 1983).

The quadrate (ZPAL V38/51) (Fig. 4h, i) is ear shaped as in most ophthalmosaurids, but the articular condyle appears robust and markedly convex ventrally, unlike many ophthalmosaurids where the condyle is divided by a deep and wide groove (*Ophthalmosaurus icenicus*, A. M. Kirton, unpub. Ph.D. thesis, University of Newcastle upon Tyne, UK, 1983; *Sveltonectes insolitus*, VF personal observation of holotype IRSNB R269; *Acamptonectes densus*, see Fischer *et al.* 2012; *Platypterygius australis*, see Kear, 2005). Quite similar condyles are also found in some specimens of '*Platypterygius*' from the Cambridge Greensand Member (Grey Chalk Formation) as well (VF. personal observation of CAMSM and NHMUK material). A high bony ridge borders the stapedia facet of the quadrate ventrally.

Figure 4. Ophthalmosaurid ichthyosaurs. (a-d) Associated teeth of 'Platypterygius' sp. (a, b) ZPAL V38/2. (c, d) ZPAL V.38/3, upper part of unit 3, late late Albian, mine, (e-l) Associated elements of a partial skull of cf. Ophthalmosaurinae, lower part of unit 3, late late Albian, mine, (e, f) Posterior tooth ZPAL V.38/52 in (e) labial and (f) anterior or posterior views, (g) Partial left jugal ZPAL V.38/57 in lateral view; note the posterior process on the posteroventral margin of the jugal. (h, i) Partial right quadrate ZPAL V.38/51 in (h) lateral and (i) anterior views, (j) Left surangular ZPAL V38/55 in lateral view, (k, l) Left angular ZPAL V38/53 in (k) dorsal and (l) lateral views.

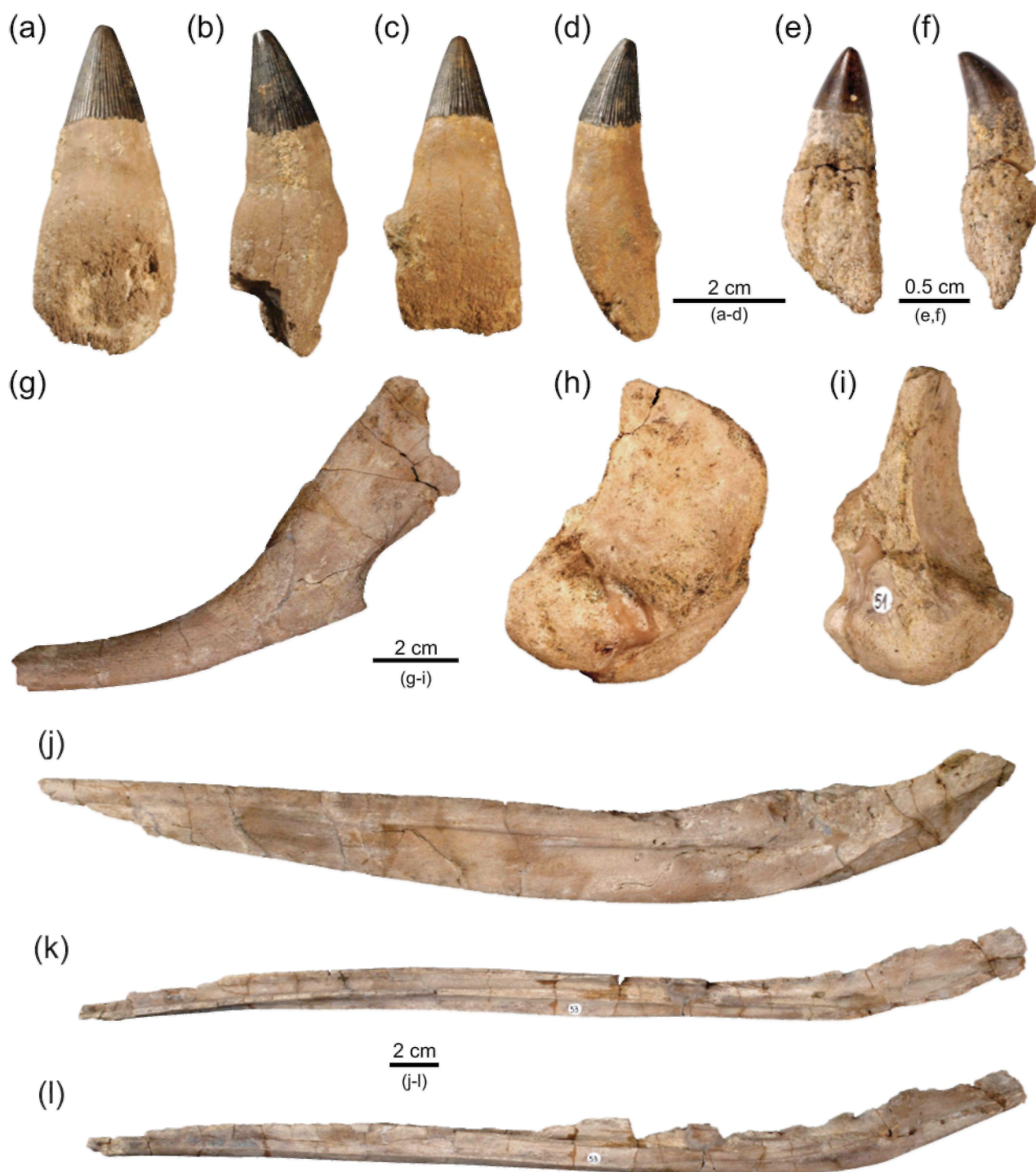
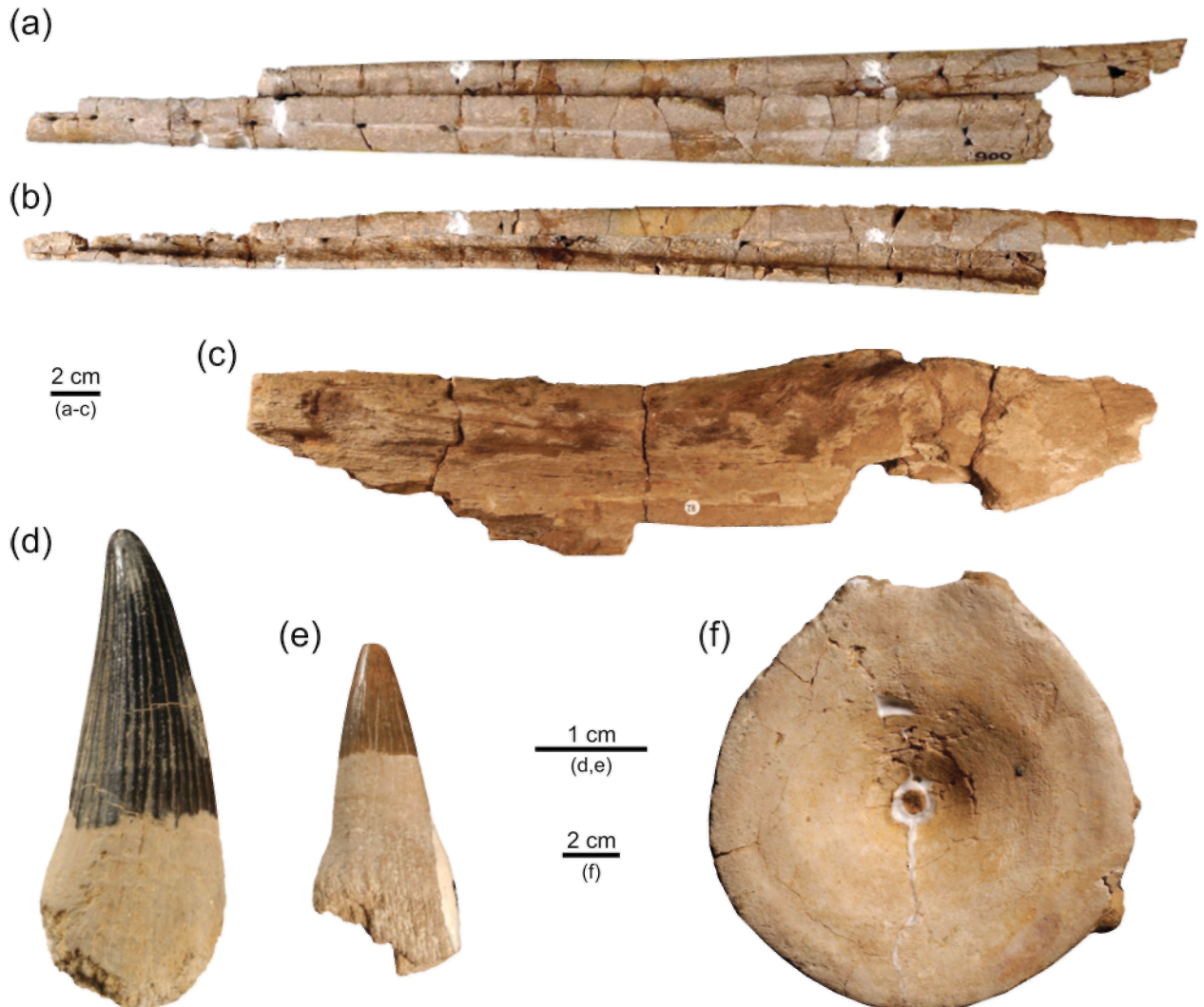


Figure 5. (a-c, e-f) Ichthyosauria indet. and (e) ophthalmosaurid ichthyosaurs, (a, b) Rostrum fragment ZPAL V.38/900, lower part of unit 3, late late Albian, Kopiec, in (a) lateral and (b) 'dental/apical' views, (c) Large right surangular ZPAL V38/82, upper part of unit 3, late late Albian, mine, in medial view, (d) Isolated tooth ZPAL V.38/427 of 'Platypterygius' sp., lower part of unit 3, late late Albian, mine, (e) Isolated tooth MWG UW ZI/60/12 which shares affinities with *Sisteronia seeleyi*, probably unit 3, late late Albian, locality unknown, (f) Large anterior caudal centrum ZPAL V.38/888, lower part of unit 3, late late Albian, mine.



The surangular (ZPAL V38/55) (Fig. 4j) is slender and slightly arched; together with the angular, this suggests a slender and curved mandible, distinct from that of contemporaneous '*Platypterygius*' from Eurasia with straight and robust jaws (e.g. Bardet, Wellnhofer & Herm, 1994; Fischer, 2012; Fischer *et al.* 2014c). The coronoid process is low and barely protrudes from the body of the surangular. A fossa surangularis is present, unlike in *Sveltonectes insolitus* (Fischer *et al.* 20116).

The angular (ZPAL V 3 8/53) is well preserved and nearly complete (Fig. 4k, 1), measuring 33 cm in length. It is flattened markedly differing from that of *Pervushovisaurus*, in which it is thick and semi-circular in cross-section (Fischer *et al.* 2014a). In dorsal view, the angular is slightly sinusoidal, denoting the widening of the mandible behind the symphysis. The angular forms a typical '3'-shaped dorsal surface whose lateral groove houses the ventral edge of the surangular. The ventral edge is rounded and smooth.

A small posterior tooth (ZPAL V38/52) (Fig. 4e, f), the crown enamel of which is totally devoid of apicobasal ridges. Among ophthalmosaurids, small tooth size with enamel texture reduction is usually found in ophthalmosaurines (e.g. Fischer *et al.* 20146). The root is rounded in cross-section; this cannot be used as a criterion for taxonomic assignment since the root of posterior teeth usually lacks clade-specific synapomorphies

in ophthalmosaurids (Fischer *et al.* 2014c). *Identification.* Among ophthalmosaurids, the combination of slender jaws and small teeth lacking enamel texture is found in, yet does not strictly characterize, ophthalmosaurines (Fischer *et al.* 20146; Roberts *et al.* 2014). The jugal also possesses a feature that was previously restricted to *Ophthalmosaurus icenicus*, further suggesting ophthalmosaurine affinities. However, the paucity of the present remains does not allow unambiguous placement of this specimen. We therefore refer to it as cf. Ophthalmosaurinae.

ICHTHYOSAURIA indet.

Figure 5a-c, e-f

Referred material. Disarticulated partial skeletons: (1) vertebrae ZPAL V38/433, 875, 888, 889; and (2) fragmentary ribs ZPAL V 38/49, 100-106. Articulated partial skeleton: skull fragment still in quarry face of underground gallery with detached jaw fragment ZPAL V38/50. Isolated jaw fragments: ZPAL V38/35, 82 (surangular), 449, 900 (rostrum). Isolated centra: ZPAL V38/193, and MWGUW ZI/60/11, 28.

Horizon and age: Top of unit 2, middle or early late Albian; unit 3, late late Albian; unit 4, early Cenomanian.

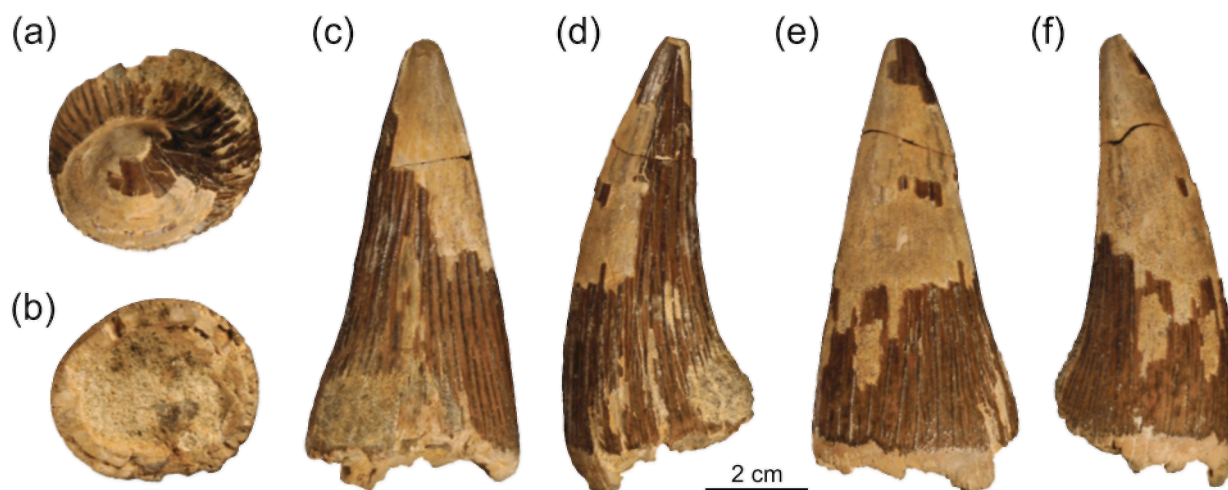
Description and comparison. A partial surangular (ZPAL V38/82, see Fig. 5c) lacks distinguishing features, except for its large size. A large coronoid process is present unlike in the surangular of cf. Ophthalmosaurinae (ZPAL V38/55). The surangular lacks a M.a.m.e process, unlike in some ophthalmosaurine taxa (*O. icenicus*, *A. densus*; see Fischer *et al.* 2012). Contrary to *Sveltonectes insolitus* (Fischer *et al.* 2011b), the fossa surangularis appears well developed.

Of the single rostrum (ZPAL V38/900, see Fig. 5a, b) it cannot be determined whether this originated from a premaxilla or a dentary. Anteriorly, the labial and lingual walls of the groove form numerous shallow dorsoventral ridges separating functional teeth as in many ichthyosaurs (e.g. Kiprijanoff, 1881; Godefroit, 1993; Bardet *et al.* 1997; Fischer *et al.* 2011a). This rostrum appears much more slender than that of contemporaneous taxa referred to as 'Platypterygius' (see Owen, 1851-1864; Fischer, 2012; Fischer *et al.* 2014c) and likely belongs to a discrete taxon with a distinct ecological niche.

The following can be stated of the dentition (MWGUW ZI/60/12, see Fig. 5e, f). While belonging to a platypterygiine ophthalmosaurid as indicated by the quadrangular cross-section of the root, the crown of the present tooth appears much more slender and the enamel possesses delicate apicobasal ridges and invariably has a distinct orange colour. This tooth morphotype bears many similarities to that of *Sisteronia seeleyi*, which was described from coeval strata in England and France (Fischer *et al.* 2014c). Because it is poorly represented in the Anopol assemblage, we regard these specimens as Ichthyosauria indet. until more diagnostic remains are recovered.

The centra (ZPAL V38/193, ZPAL V 38/433, 875, 888, 889) are large (diameter 12 cm), forming the posterior dorsal-anterior caudal region. The centra are weakly amphicelous and their rib facets are prominent (Fig. 5f). These centra are thick, with a *HIL* ratio of c. 2.

Figure 6. *Polyptychodon interruptus*, isolated tooth ZPAL V.38/2034, probably unit 3, late late Albian, Kopic, in (a) occlusal, (b) basal, (c) anterior, (e) posterior, and (d, f) labial/lingual views.



Order PLESIOSAURIA de Blainville, 1835 Family PLIOSAURIDAE Seeley, 1874
Genus: *Polyptychodon* Owen, 1841
Polyptychodon interruptus Owen, 1841
Figures 6, 7a-c

Referred material. Isolated incomplete teeth: MWGUW ZI/60/1, 19-27, ZPAL V38/170, 443, 689, 893, 894, 2034.

Horizon and age. Unit 3, late late Albian; unit 4, early Cenomanian; unit 5, middle Cenomanian.

Description and comparison. All teeth are large and stout, with a rounded basal cross-section (Fig. 6). They bear strong and coarse ridges all along the crown except, when complete, for the last upper quarter where only a very few of them reach the apex. These crowns are only slightly posteriorly recurved and bear carinae that are visible only near the apex (see ZPAL 893; Fig. 7a). Most of these teeth exhibit a broken apex (Fig. 7b, c) with a rounded and polished top surface, probably indicating a preference for large fleshy and bony prey such as large fish and marine reptiles (see Massare, 1987).

Such teeth are fairly common in Albian strata across Europe (N.B., personal observation) and are most often referred to *Polyptychodon* Owen, 1841. As already noted (i.e. Albright, Gillette & Titus, 2007; Schumacher, 2008), the status of this genus remains highly problematic as it is based on very few and poorly diagnostic material, mainly isolated teeth. Indeed Owen defined two species from the Albian greensands of England in his 'Odontography' (Owen, 1841), namely *P. continuus* and *P. Interruptus*. These were differentiated on the basis of tooth enamel ornament, the former having ridges reaching the apex and the latter lacking these. Another species, *Polyptychodon hudsoni*, was described much later based on a skull fragment of Turonian age from Texas (Welles & Slaughter, 1963). *Polyptychodon continuus* was regarded as a *nomen vanum* by Welles (1962).

Identification. Although displaying general pliosaurid morphology and pending a detailed review of this genus, the teeth from Annopol are here referred to *Polyptychodon interruptus*, the only pliosaurid taxon currently recognized in the Albian succession of the European Archipelago (e.g. Bardet & Godefroit 1995; Buffet et al. 2005). The general characteristics of the teeth (size, robustness, intense apical wear) of *Polyptychodon* strongly indicate that this genus was an apex predator in the Annopol trophic web and in coeval European environments.

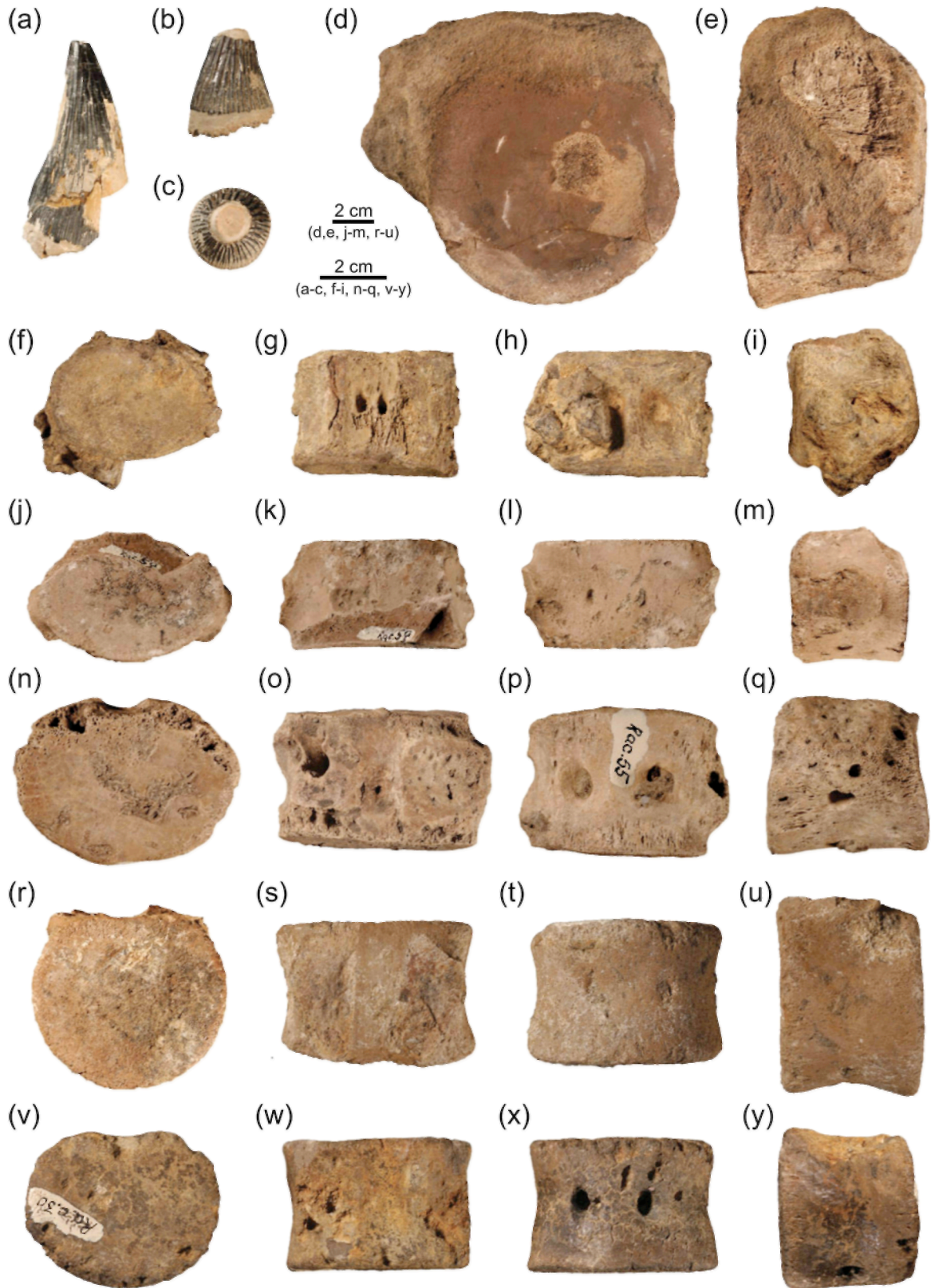
PLIOSAURIDAE indet.
Figure 7d, e

Referred material. MWGUW ZI/60/2.

Horizon and age. Top of unit 2 (based on adhering matrix), middle or early late Albian.

Description. A large (diameter > 10 cm), amphicoelous centrum (Fig. 7d, e), which, although poorly preserved is typically shorter than tall and wide, so that it may belong to an indeterminate pliosaurid. It could correspond to *Polyptychodon*, but this cannot be demonstrated beyond doubt as this centrum is not diagnostic and there is also an absence of homologous material for comparison; as noted above, *Polyptychodon* is mainly known from isolated teeth.

Figure 7. Plesiosaurs. (a-c) Teeth of *Polyptychodon interruptus*. (a) ZPAL V.38/893, unit 5, middle Cenomanian, mine, tooth in labial/lingual view, (b, c) MWGUW ZI/60/1, probably unit 3, late late Albian, locality unknown, tooth in (b) labial/lingual and (c) occlusal views, (d-y) Isolated centra, localities unknown, (d, e) Pliosauridae indet, MWGUW ZI/60/2, unit 2 (matrix), middle or early late Albian, cervical centrum, (f-q) Elasmosauridae indet. (f-i) MWGUW ZI/60/3, unit 3 (matrix), late late Albian, posterior cervical centrum, (j-m) MWGUW ZI/60/5, unit 3, late late Albian (matrix), posterior cervical centrum, probably more posterior than MWGUW ZI/60/3. (n-q) MWGUW ZI/60/4, probably unit 3, late late Albian, pectoral centrum, (r-y) Plesiosauria indet. (r-u) MWGUW ZI/60/7, unit 2 (matrix), middle or early late Albian, median dorsal centrum, (v-y) MWGUW ZI/60/6, probably unit 3, late late Albian, posterior dorsal centrum, (d, f, j, n, r, v) Articular view; (g, k, o, s, w) dorsal view; (h, i, p, t, x) ventral view; (e, i, m, q, u, y) lateral view.



Superfamily PLESIOSAUROIDEA Gray, 1825 (*sensu* Welles, 1943)
Family ELASMOSAURIDAE Cope, 1869
ELASMOSAURIDAE indet.
Figure 7f-q

Referred material. MWGUW ZI/60/3-5.

Horizon and age: Unit 3 (based on adhering matrix), late late Albian.

Description. Several centra from the posterior part of the neck and probably from the pectoral region of the vertebral column are preserved. None of these retain the neural arch, indicating that they belong to immature specimens.

MWGUW ZI/60/3 (Fig. 7f-i) is a posterior cervical centrum, which is much wider than high and long. The articular surfaces are amphicoelous, ovoid in shape without any ventral notch, and with roughly rounded margins. The dorsal surface bears two large median foramina, close set on the floor of the neural canal. The articular surfaces for the neural arch are long and narrow. The ventral surface bears two large foramina subcentralia separated by a rounded median ridge. The lateral surfaces are poorly preserved. On one of them a longitudinal crest might be preserved, just dorsal of remains of bone that could belong to the corresponding rib.

MWGUW ZI/60/5 (Fig. 7j-m) is another posterior cervical centrum, probably more posterior than MWGUW ZI/60/3, because of the rib facet that is situated higher on the lateral surface of the centrum. It is much wider than high and long and its weight is of note. The articular surfaces are amphicoelous, ovoid in shape without a ventral notch and bear sharp margins. The dorsal surface is poorly preserved. The ventral surface is convex from side to side and bears two foramina subcentralia that are wider apart than in MWGUW ZI/60/3, another feature pointing to a more posterior position of this vertebra in the cervical series. The rib processes are situated medially on the lateral surfaces; they are rounded and large, occupying half of the lateral surface.

MWGUW ZI/60/4 (Fig. 7n-q) is probably a pectoral centrum. It is much wider than high and long. The articular surfaces are amphicoelous, ovoid without a ventral notch and with well-defined margins. The dorsal surface bears large articular facets for the neural arch that also extend in part, along the upper part of the lateral surface, indicating that this centrum was probably the last pectoral vertebra. The ventral surface is convex from side to side and bears two very large foramina subcentralia, separated from each other. The lateral surfaces as a whole are pierced by large foramina from place to place and its texture appears highly spongiouse.

Identification. In view of their general morphology and proportions, in particular the ovoid shape of the articular surface, which is much wider than high and long, these centra could belong, among Plesiosauroidea, to an indeterminate elasmosaurid. These centra lack the ventral notch that gives the articular surface a typical 'binocular' shape. The ventral notch is common in Late Cretaceous elasmosaurids (e.g. Bardet, Godefroit & Sciau, 1999; O'Keefe, 2001; Druckenmiller & Russell, 2008), but is commonly absent from Early Cretaceous representatives of this group (Sachs & Kear, 2014).

PLESIOSAURIA indet.
Figure 7r-y

Referred material. Isolated dorsal centra: MWGUW ZI/60/6-10.

Horizon and age. Top of unit 2, based on adhering matrix, middle or early late Albian; unit 3, based on adhering matrix, late late Albian.

Description. Of all specimens only the centra are preserved, indicating that the neural arches were not fused and that specimens were immature.

All exhibit more or less the same characteristics, being all slightly wider than high and long and bearing either rounded or slightly oval articular surfaces which are slightly concave to amphicoelous and with, when preserved, well-defined margins. The lateral and ventral surfaces are regularly convex and smooth from side to side and bear foramina subventralia more or less upwardly facing, depending on their respective position in the dorsal series. These foramina are generally large. The dorsal surface bears large, hourglass-shaped articular surfaces for the neural arch. The floor of the neural canal is flat and narrow and devoid of foramina.

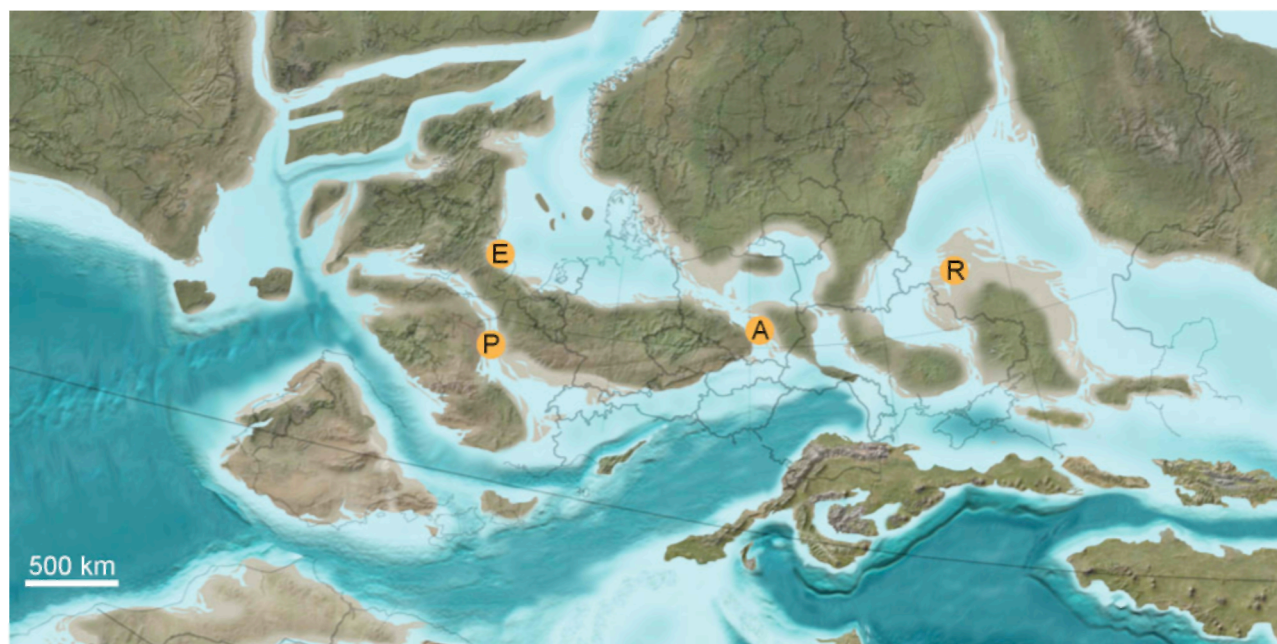
The more elliptical centra (MWGUW ZI/60/6; see Fig. 7v-y) could correspond to posterior dorsals located near the sacral series whereas the largest, such as MWGUW ZI/60/7 (Fig. 7r-u), are probably median dorsals.

Identification. Plesiosaur dorsal vertebrae are very poorly diagnostic as a whole and even the classic dichotomy between Plesiosauroidea and Pliosauroida cannot be determined on the basis of the present material. The dorsal vertebrae from Annopol can therefore only be referred to indeterminate members of the clade Plesiosauria, although it is clear that in this lot some specimens might correspond either to the pliosaurid *Polyptychodon* or to an elasmosaurid taxon; this cannot be demonstrated however because of the lack of associated material.

7. DISCUSSION

Localities with mid-Cretaceous strata across Europe and Russia have yielded abundant vertebrate remains, mainly of late Albian age (Fig. 8). Arguably, the best known are the Gault and Upper Greensand formations, in addition to the Cambridge Greensand Member of the Grey Chalk Formation of east and southeast England which contain thousands of isolated remains of marine reptiles (e.g. Seeley, 1869; Fischer *et al.* 2014c). The predators at the top of the food chains in these marine ecosystems were the ophthalmosaurid ichthyosaur '*Platypterygius*' sp. and the pliosaurid plesiosaur *Polyptychodon interruptus* (Owen, 1860; Seeley, 1876; Bardet, 1992; Bardet & Godefroit, 1995), but also yield diversified assemblages of distinct, smaller ichthyosaurs (Bardet, 1992; Fischer *et al.* 2014c) and plesiosaurs (Owen, 1851-1864; Seeley, 1869; Lydekker, 1889). However, these are in need of a modern revision.

Figure 8. Palaeobiogeography of the Albian marine reptile faunas of Eurasia. I: ichthyosaurian; P: plesiosaurian;



	SW England E	Paris Basin P	Annopol A	W Russia* R
Top predators	' <i>Platypterygius</i> ' sp. (I) <i>Polypychodon interruptus</i> (P)	' <i>Platypterygius</i> ' sp. (I) <i>Polypychodon interruptus</i> (P)	' <i>Platypterygius</i> ' sp. (I) <i>Polypychodon interruptus</i> (P)	' <i>Platypterygius</i> ' sp. (I) <i>Polypychodon interruptus</i> (P)
Smaller predators	<i>Sisteronia seeleyi</i> (I) Ophthalmosaurinae indet. (I) Plesiosauria indet.	Ichthyosauria indet. Plesiosauria indet.	Ichthyosauria indet. cf. Ophthalmosaurinae (I) Elasmosauridae (P)	Ichthyosauria indet. Polycotylidae (P)

*preliminary results of an ongoing study. Palaeogeographical map courtesy of R. Blakey.

The Paris Basin yielded some late Albian marine reptiles. Again, this assemblage is characterized by the co-occurrence of *Polyptychodon interruptus* (see Sauvage, 1878; Buffetaut *et al.* 2005; numerous unregistered specimens in local museums in this region, N.B., personal observation) and frequent large platypterygiine ophthalmosaurids referred to as *Platypterygius* (Barrois, 1875; Sauvage, 1882; Buffetaut, 1977; Debris, 1977, 1978; Buffetaut *et al.* 2005; Colleté, 2010; Fischer, 2012; Y.F., personal observation from several local museums in this region). Similar ichthyosaur remains have also been recorded from southern France (Pouech, 1881; Bardet *et al.* 1991; Fischer *et al.* 2014c) and Italy (Capellini, 1890; Sirotti & Papazzoni, 2002), but many of them have yet to be properly described (V Fischer, unpub. PhD thesis, Université de Liège, 2013).

Coeval assemblages have also been described from the Kursk area, western Russia (Eichwald 1853, 1865-1868; Kiprijanoff, 1881, 1882, 1883a, b; Rozhdestvenskiy, 1973), including a new locality, currently under investigation. A preliminary assessment of this material suggests the presence of '*Platypterygius*' sp. and *Polyptychodon interruptus*, as well as new small ichthyosaurs and abundant polycotyloid remains, the latter feature distinguishing this locality from other European localities.

Although most marine reptilian specimens have been found as isolated bones at Annapol, of note is the co-occurrence from the late late Albian to the middle Cenomanian of both '*Platypterygius*' and *Polyptychodon*, as well as of elasmosaurids and a smaller ichthyosaur taxon (Figs 2, 8). As a result, the Annapol marine reptile assemblage appears to have been distinct from other coeval Eurasian ecosystems in containing elasmosaurids, but was similar in being dominated by the ubiquitous apex predators '*Platypterygius*' and *Polyptychodon* and in also containing at least one smaller ichthyosaur which occupied an ecological niche different from that of '*Platypterygius*'.

To summarize, these two top predators formed a stable ecological sympatry along the entire NW margin of the Tethys Ocean during the Albian - middle Cenomanian. Conversely, each of these ecosystems (SW England Paris Basin, Annapol in Poland western Russia) differed in marine reptilian taxa that occupied lower trophic levels: elasmosaurids in Annapol, polycotyloid plesiosaurs in western Russia and abundant ophthalmosaurid ichthyosaurs in SW England (Fig. 8). These differences perhaps reflect regional differences in ecosystems and/or profound modifications occurring at a very fine temporal scale.

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REFERENCES

- ALBRIGHT, B. L., GILLETTE, D. D. & TITUS, A. L. 2007. Plesiosaurs from the Upper Cretaceous (Cenomanian-Turonian) Tropic Shale of southern Utah, Part 1: New records of the plesiosaur *Brachauchenius lucasi*. *Journal of Vertebrate Paleontology* 27, 31-40.
- AMEDRO, F. 2002. Plaidoyer pour un étage Vraconnien entre l'Albien sensu stricto et le Cénomanién (système Crétacé). *Académie Royale de Belgique, Classe des Sciences* 4, 1-128.
- ARKHANGELSKY, M. S. 2001. On a new ichthyosaur of the genus *Otschevia* from the Volgian Stage of the Volga region near Ulyanovsk. *Paleontological Journal* 35, 629-34.
- BARDET, N. 1989. Un crâne d'Ichthyopterygia dans le Cénomanién du Boulonnais. *Mémoires de la Société Académique du Boulonnais* 6, 1-31.
- BARDET, N. 1992. Stratigraphie evidence for the extinction of the ichthyosaurs. *Terra Nova* 4, 649-56.
- BARDET, N., DUFFAUD, S., MARTIN, M., MAZIN, J.-M., PEREDA SUBERBIOLA, X. & VIDIER, J.-P. 1997. Découverte de l'ichthyosaure *Ophthalmosaurus* dans le Tithonien (Jurassique supérieur) du Boulonnais, Nord de la France. *Neues Jahrbuch für Geologie*

und *Paläontologie Abhandlungen* **205**, 339-54.

BARDET, N. & GODEFROIT, P. 1995. *Plesiosaurus houzeaui* Dollo, 1909 from the Upper Campanian of Cipro (Belgium) and a review of the Upper Cretaceous plesiosaurs from Europe. *Bulletin de l'Institut Royal des Sciences Naturelles de Belgique, Sciences de la Terre* **65**, 179-86.

BARDET, N., GODEFROIT, P. & SCIAU, J. 1999. A new elasmo-saurid plesiosaur from the Lower Jurassic of southern France. *Palaeontology* **42**, 927-52.

BARDET, N., MAZIN, J.-M., AZÉMA, C., BÉGOUEN, V & MASURE, E. 1991. "L'ichthyosaure de Bedeille" (Ar-iège, France): examen palynologique de la gangue et mise au point stratigraphique. *Bulletin de la Société Géologique de France* **162**, 897-903.

BARDET, N., WELLNHOFER, P. & HERM, D. 1994. Discovery of ichthyosaur remains (Reptilia) in the upper Cenomanian of Bavaria. *Mitteilungen aus der Bay-erischen Staatssammlung für Paläontologie und Histor-ische Géologie* **34**, 213-20.

BARROIS, C. 1875. Les reptiles du terrain crétacé du nord-est du Bassin de Paris. *Bulletin Scientifique, Historique et Littéraire du Nord* **6**, 11.

BAUR, G. 1887. On the morphology and origin of the Ichthy-opterygia. *American Naturalist* **21**, 837-40.

BLAINVILLE, H. M. D., de. 1835. Description de quelques espèces de reptiles de la Californie, précédée de l'analyse d'un système général d'érpetologie et d'amphibiologie. *Nouvelles Annales du Muséum d'Histoire Naturelle Paris* **4**, 233-96.

BUFFETAUT, E. 1977. Remarques préliminaires sur l'ichthyosaure de Saint-Jouin (76). *Bulletin de la Société Géologique de Normandie et Amis du Muséum du Havre* **64**, 17-9.

BUFFETAUT, E., COLLETE, C., DUBUS, B. & PETIT, J.-L. 2005. The "sauropod" from the Albian of Mesnil-Saint-Père (Aube, France): a pliosaur, not a dinosaur. *Notebooks on Geology, Letter* 2005/01, 5.

BUFFETAUT, E., TOMASSON, R. & TONG, H. 2003. Restes fossiles de grands reptiles jurassiques et crétacés dans l'Aube (France). *Bulletin d'Information des Géologues du Bassin de Paris* **40**, 33-43.

CAPELLINI, G. 1890. *Ichthyosaurus campylodon* e tronchi di cicadee nelle argille scagliose dell'Emilia. *Memorie della Reale Accademia delle Scienze di Bologna* **4**(10), 431-50.

CIESLINSKI, S. 1959. The Albian and Cenomanian in the northern periphery of the Holy Cross Mountains (stratigraphy based on cephalopods). *Prace Instytutu Geologicznego* **28**, 1-95 (in Polish, English summary).

CIESLINSKI, S. 1976. Development of the Danish-Polish furrow in the Góry Świętokrzyskie region in the Albian, Cenomanian and Lower Turonian. *Biuletyn Instytutu Geologicznego* **295**, 249-71 (in Polish, English summary).

CIESLINSKI, S. 1987. Albian and Cenomanian inoceramids in Poland and their stratigraphic significance. *Biuletyn Instytutu Geologicznego* **354**, 11-62 (in Polish, English summary).

CIESLINSKI, S. & MIŁAKOVIC, B. 1962. Kręgowce i flora kredowa z obrzeżenia Gór Świętokrzyskich. *Biuletyn Instytutu Geologicznego* **174**, 245-66.

COLLETÉ, C. 2010. *Stratotype Albien*. Paris: Muséum National d'Histoire Naturelle, Méze: Biotope, Orléans: BRGM, 332 pp.

COOKSON, I. C. & HUGHES, N. F. 1964. Microplankton from the Cambridge Greensand (mid-Cretaceous). *Palaeontology* **7**, 37-59.

COPE, E. D. 1869. Synopsis of the Extinct Batrachia and Reptilia of North America, Part I. *Transactions American Philadelphia Society New Series* **14**, 1-235.

DEBRIS, J.-P. 1977. Découverte d'un crâne d'ichthyosaurien dans l'Albien de Saint-Jouin. *Bulletin de la Société Géologique de Normandie et Amis du Muséum du Havre* **64**, 13-6.

DEBRIS, J.-P. 1978. Données nouvelles sur les Ichthyosauriens de l'Albien de Saint Jouin (76). *Bulletin Trimestriel de la Société Géologique de Normandie et Amis du Muséum du Havre* **65**, 25-8.

DRUCKENMILLER, P. S. & RUSSELL, A. P. 2008. A phylogeny of Plesiosauria (Sauropterygia) and its bearing on the systematic status of *Leptocleidus* Andrews 1922. *Zoo-taxa* **1863**, 1-120.

EICHWALD, K. E. 1853. Einige paläontologische Bemerkungen über den Eisensand von Kursk. *Bulletin de la Société Impériale des Naturalistes de Moscou* **2**, 209-31.

EICHWALD, K. E. 1865-1868. *Lethaea Rossica ou Paléontologie de la Russie*. **Second** Volume. Période Moyenne. Stuttgart: E. Schweizerbart (E. Koch), 1304 pp.

FISCHER, V. 2012. New data on the ichthyosaur *Platypterygius hercynicus* and its implications for the validity of the genus. *Acta Palaeontologica Polonica* **57**, 123-34.

FISCHER, V., ARKHANGELSK M. S., NAISH, D., STENSHIN, I. M., USPENSKY, G. N. & GODEFROIT, P. 2014a. *Simbirskiasaurus* and *Pervushoviasaurus* reassessed: implications for the taxonomy and cranial osteology of Cretaceous platypterygiine ichthyosaurs. *Zoological Journal of the Linnean Society* **111**, 822-41.

FISCHER, V., ARKHANGELSKY, M. S., USPENSKY, G. N., STENSHIN, I. M. & GODEFROIT, P. 2014b. A new Lower Cretaceous ichthyosaur from Russia reveals skull shape conservatism within Ophthalmosaurinae. *Geological Magazine* **151**, 60-70.

FISCHER, V., BARDET, N., GUIOMAR, M. & GODEFROIT, P. 2014c. High diversity in Cretaceous ichthyosaurs from Europe prior to their extinction. *PLoS ONE* **9**, e84709.

FISCHER, V., CLÉMENT, A., GUIOMAR, M. & GODEFROIT, P. 2011a. The first definite record of a Valanginian ichthyosaur and its implication for the evolution of post-Liassic Ichthyosauria. *Cretaceous Research* **32**, 155-63.

FISCHER, V., MAISCH, M. W., NAISH, D., LISTON, X., KOSMA, R., JOGER, U., KRÜGER, F. X., PARDO-PÉREZ, X., TAINSH, X. & APPLEBY, R. M. 2012. New ophthalmosaurid ichthyosaurs from the Early Cretaceous of Europe demonstrate extensive ichthyosaur survival across the Jurassic-Cretaceous boundary. *PLoS ONE* **7**, e29234.

- FISCHER, V, MASURE, E., ARKHANGELSKY, M. S. & GODEFROIT, P. 2011b. A new Barremian (Early Cretaceous) ichthyosaur from western Russia. *Journal of Vertebrate Paleontology* 31, 1010-25.
- FOOTE, A. D., MORIN, P. A., DURBAN, J. W., WILLERSLEV, E., ORLANDO, L. & GILBERT, M. T. P. 2011. Out of the Pacific and back again: insights into the matrilineal history of Pacific killer whale ecotypes. *PLoS ONE* 6, e24980.
- GODEFROIT, P. 1993. Les grands ichthyosaures sinémuriens d'Arlon. *Bulletin de l'Institut Royal des Sciences Naturelles de Belgique, Sciences de la Terre* 63, 25-71.
- GRAY, X E. 1825. A synopsis of the genera of reptiles and Amphibia, with a description of some new species. *Annals of Philosophy* 26, 193-217.
- HANCOCK, X M. 1990. Sea-level changes in the British region during the Late Cretaceous. *Proceedings of the Geologists' Association* 100 (for 1989), 565-94.
- HOPSON, P. M. 2005. A stratigraphical framework for the Upper Cretaceous Chalk of England and Scotland with statements on the Chalk of Northern Ireland and the UK Offshore Sector. *British Geological Survey Research Reports* RR/05/01, 1-102.
- HOPSON, P. M., WILKINSON, I. P. & WOODS, M. A. 2008. A stratigraphical framework for the Lower Cretaceous of England. *British Geological Survey Research Reports* RR/08/03, 1-87.
- HUENE, F VON. 1922. *Die Ichthyosaurier des Lias und ihre Zusammenhänge*. Berlin: Verlag von Gebrüder Borntraeger, 114 pp.
- JUIGNET, P. 1980. Transgressions-régressions, variations eustatiques et influences tectoniques de l'Aptien au Maastrichtien dans le Bassin de Paris occidental et sur la Bordure du Massif Armoricain. *Cretaceous Research* 1, 341-57.
- KEAR, B. P. 2005. Cranial morphology of *Platypterygius longmani* Wade, 1990 (Reptilia: Ichthyosauria) from the Lower Cretaceous of Australia. *Zoological Journal of the Linnean Society* 145, 583-622.
- KIPRIJANOFF, W 1881. Studien über die fossilen Reptilien Russlands. Teil 1. Gattung *Ichthyosaurus* König aus dem severischen Sandstein oder Osteolith der Kreide-Gruppe. *Mémoires de l'Académie Impériale des Sciences de St.-Petersbourg* 28(7), 1-103.
- KIPRIJANOFF, W. 1882. Studien über die fossilen Reptilien Russlands. Teil 2. Gattung *Plesiosaurus* Conybeare aus dem severischen Sandstein oder Osteolith der Kreide-gruppe. *Mémoires de l'Académie Impériale des Sciences de St.-Petersbourg* 30(7), 1-55.
- KIPRIJANOFF, W. 1883a. Studien über die fossilen Reptilien Russlands. Teil 3. Gruppe *Thaumatosauria* n. aus der Kreide-Formation und dem Moskauer Jura. *Mémoires de l'Académie impériale des Sciences de St.-Petersbourg* 31(7), 1-57.
- KIPRIJANOFF, W. 1883b. Studien über die fossilen Reptilien Russlands. Teil 4. Ordnung *Crocodylia* Opper. Inde-terminirte fossile Reptilien. *Mémoires de l'Académie impériale des Sciences de St.-Petersbourg* 31(7), 1-29.
- LAMBERT, O., BIANUCCI, G., POST, K, DE MUIZON, C, SALAS-GISMONDI, R., URBINA, M. & REUMER, J. 2010. The giant bite of a new raptorial sperm whale from the Miocene epoch of Peru. *Nature* 466, 105-8.
- LYDEKKER, R. 1889. Catalogue of the fossil Reptilia and Amphibia in British Museum (Natural History). Part II. Containing the orders Ichthyopterygia and Sauropterygia. London: Printed by Orders of the Trustees of the British Museum, 307 pp.
- MACHALSKI, M. & KENNEDY, W. J. 2013. Oyster-bioimmured ammonites from the Upper Albian of Annapol, Poland: stratigraphic and palaeobiogeographic implications. *Acta Geologica Polonica* 63, 545-54.
- MACHALSKI, M., KOMOROWSKI, A. & HARASIMIUK, M. 2009. New chances in the quest for Cretaceous marine vertebrates in abandoned phosphate mine at Annapol on Vistula River. *Przegląd Geologiczny* 57, 638-41 (in Polish, English summary).
- MACHALSKI, M. & MARTILL, D. M. 2013. First pterosaur remains from the Cretaceous of Poland. *Annales Societatis Geologorum Poloniae* 83, 99-104.
- MARCINOWSKI, R. 1980. Cenomanian ammonites from German Democratic Republic, Poland, and the Soviet Union. *Acta Geologica Polonica* 30, 215-325.
- MARCINOWSKI, R. & RADWANSKI, A. 1983. The mid-Cretaceous transgression onto the Central Polish Uplands (marginal part of the Central European Basin). *Zitteliana* 10, 65-96.
- MARCINOWSKI, R. & RADWANSKI, A. 1989. Stratigraphic approach to the mid-Cretaceous transgressive sequence of the Central Polish Uplands. *Cretaceous Research* 10, 153-72.
- MARCINOWSKI, R. & WALASZCZYK, I. 1985. Middle Cretaceous deposits and biostratigraphy of the Annapol section, Central Polish Uplands. *Osterreichische Akademie der Wissenschaften, Schriftenreihe der Erd-wissenschaftlichen Kommissionen* 7, 21-41.
- MARCINOWSKI, R. & WIEDMANN, J. 1985. The Albian ammonite fauna of Poland and its palaeogeographical significance. *Acta Geologica Polonica* 35, 199-219.
- MARCINOWSKI, R. & WIEDMANN, J. 1990. The Albian ammonites of Poland. *Palaeontologia Polonica* 50, 1-94.
- MARTILL, D.M. 1993. Soupy substrates: a medium for the exceptional preservation of ichthyosaurs of the Posidonia Shale (Lower Jurassic) of Germany. *Kaupia - Darmstädter Beiträge zur Naturgeschichte* 2, 11-91.
- MASSARE, J. A. 1987. Tooth morphology and prey preference of Mesozoic marine reptiles. *Journal of Vertebrate Paleontology* 7, 121-37.
- MAXWELL, E. E., CALDWELL, M. W. & LAMOUREUX, D. O. 2011. Tooth histology in the Cretaceous ichthyosaur *Platypterygius australis*, and its significance for the conservation and divergence of mineralized tooth tissues in amniotes. *Journal of Morphology* 212, 129-35.
- O'KEEFE, F. R. 2001. A cladistic analysis and taxonomic revision of the Plesiosauria (Reptilia: Sauropterygia). *Acta Zoologica Fennica* 213, 1-63.
- OWEN, R. 1840. Report on British fossil reptiles. *Reports of the British Association for the Advancement of Science* 9, 43-126.

- OWEN, R. 1841. *Odontography; or, a Treatise on the Comparative Anatomy of the Teeth; their Physiological Relations, Mode of Development, and Microscopic Structure, in the Vertebrate Animals*. London: Hippolyte Baillière, 655 pp.
- OWEN, R. 1851-1864. *A Monograph on the Fossil Reptilia of the Cretaceous Formations*. London: The Palaeontographical Society, pp. 118.
- OWEN, R. 1860. Note on some remains of *Polyptychodon* from Dorking. *Quarterly Journal of the Geological Society London* 16, 262-3.
- OWEN, H. G. 2012. The Gault Group (Early Cretaceous, Albian), in East Kent, S.E. England; its lithology and ammonite biozonation. *Proceedings of the Geologists' Association* 123, 742-65.
- PERYT, D. 1983. Planktonic foraminiferal zonation of Mid-Cretaceous of the Annopol Anticline (Central Poland). *Zitteliana* 10, 575-83.
- POPOV, E. V & MACHALSKI, M. 2014. Late Albian chimaeroid fishes (Holocephali, Chimaeroidei) from Annopol, Poland. *Cretaceous Research* 47, 1-18.
- POUECH, J. J. 1881. Note sur un fragment de mâchoire d'un grand Saurien trouvé à Bedeille (Ariège). *Bulletin de la Société Géologique de France* 10(3), 79—87.
- POZARYSKI, W. 1947. A phosphate deposit of the northeastern margin of the Holy Cross Mountains. *Biuletyn Państwowego Instytutu Geologicznego* 27, 1-56 (in Polish, English summary).
- RADWANSKI, A. 1968. *Ischyodus thurmanni* Pictet & Campiche and other chimaeroid fishes from the Albian-Cenomanian of the Holy Cross Mountains (Poland). *Acta Palaeontologica Polonica* 13, 315-22 (in Polish, English summary).
- RADWANSKI, A., WYSOCKA & A. G6rka, M. 2012. Miocene burrows of the ghost crab *Ocypode* and their environmental significance (Mykolaiv Sands, Fore-Carpathian Basin, Ukraine). *Acta Geologica Polonica* 62, 217-29.
- ROBERTS, A. X, DRUCKENMILLER, P. S., SÆTRE, G.-P. & HURUM, J. H. 2014. A new Upper Jurassic ophthalmo-saurid ichthyosaur from the Slottsmoya Member, Agard-hfjellet Formation of central Spitsbergen. *PLoS ONE* 9, e103152.
- RozHDESTVENSKIY, A. K. 1973. The study of Cretaceous reptiles in Russia. *Paleontological Journal* 2, 206-14.
- SACHS, S. & KEAR, B. P. 2014. Postcranium of the paradigm elasmosaurid plesiosaurian *Libonectes morgani* (Welles, 1949). *Geological Magazine*, published online 20 November 2014. doi:10.1017/S0016756814000636.
- SAMSONOWICZ, J. 1925. Esquisse géologique des environs de Rachôw sur la Vistule et les transgressions de l'Albien et du Cénomanien dans les sillons nord-européen. *Spra-wozdanja Państwowego Instytutu Geologicznego* 3, 45-118 (in Polish, French summary).
- SAMSONOWICZ, J. 1934. Explication de la feuille Opatów (zone 45, colonne 33). Service géologique de Pologne, Carte Géologique Générale de la Pologne au 100.000-e. Warszawa: Państwowy Instytut Geologiczny, 97 pp.
- SAUVAGE, H. E. 1878. Prodrome des Plésiosauriens et des Elasmosauriens des formations Jurassiques supérieures de Boulogne-sur-Mer. *Annales des Sciences Naturelles, Zoologie* 13, 1-38.
- SAUVAGE, H. E. 1882. Recherches sur les reptiles trouvées dans le Gault de l'Est du Bassin de Paris. *Mémoires de la Société Géologique de France* 2(3), 21-4.
- SCHUMACHER, B. A. 2008. On the skull of apliosaur (Plesio-sauria; Pliosauridae) from the Upper Cretaceous (Early Turonian) of the North American Western Interior. *Transactions of the Kansas Academy of Science* 111, 203-18.
- SEELEY, H. G. 1869. *Index of the Fossil Remains of Aves, Ornithosauria and Reptilia, from the Secondary System of Strata arranged in the Woodward Museum of the University of Cambridge*. Cambridge: Deighton, Bell and Co., 143 pp.
- SEELEY, H. G. 1874. Note on some of the generic modifications of the plesiosaurian pectoral arch. *Quarterly Journal of the Geological Society London* 30, 436-49.
- SEELEY, H. G. 1876. On an associated series of cervical and dorsal vertebrae of *Polyptychodon*, from the Cambridge Upper Greensand, in the Woodwardian Museum of the University of Cambridge. *Quarterly Journal of the Geological Society London* 32, 433-6.
- SIROTTI, A. & PAPAZZONI, C. 2002. On the Cretaceous ichthyosaur remains from the Northern Apennines (Italy). *Bol-lettino della Società Paleontologica Italiana* 41, 237-48.
- VINCENT, P., BARDET, N, HOUSSAYE, A., AMAGHZAZ, M. & MESLOUH, S. 2013. New plesiosaur specimens from the Maastrichtian Phosphates of Morocco and their implications for the ecology of the latest Cretaceous marine apex predators. *Gondwana Research* 24, 796-805.
- WAHL, W. 2009. Taphonomy of a nose-dive: bone and tooth displacement and mineral accretion in an ichthyosaur skull. *Paludicola* 7, 107-16.
- WALASZCZYK, I. 1987. Mid-Cretaceous events at the marginal part of the Central European Basin (Annopol-on-Vistula section, Central Poland). *Acta Geologica Polonica* 37, 61-74.
- WALASZCZYK, I. 1992. Turonian through Santonian deposits of the Central Polish Upland; their facies development, inoceramid paleontology and stratigraphy. *Acta Geologica Polonica* 42, 1-122.
- WELLES, S. P. 1943. Elasmosaurid plesiosaurs with description of new material from California and Colorado. *Memoirs of the University of California* 13, 125-254.
- WELLES, S. P. 1962. A new species of elasmosaur from the Aptian of Colombia and a review of the Cretaceous plesiosaurs. *University of California Publications in Geological Sciences* 44, 1-96.
- WELLES, S. P. & SLAUGHTER, B. H. 1963. The first record of the Plesiosaurian genus *Polyptychodon* (Pliosauridae) from the New World. *Journal of Paleontology* 37, 131-3.
- WETZEL, A. & REISDORF, A. G. 2007. Ichnofabrics elucidate the accumulation history of a condensed interval containing a vertically emplaced ichthyosaur skull. In: *Ichnology at the Crossroads: A Multidimensional Approach to the Science of Organism-Substrate*

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WOODS, M. A., WILKINSON, G. K., BOOTH, K. A., FARRANT, A. R., HOPSON, P. M. & NEWELL, A. J. 2008. A reappraisal of the stratigraphy and depositional development of the Upper Greensand (Late Albian) of the Devizes district, southern England. *Proceedings of the Geologists' Association* 119, 229-44.

YOUNG, M. T., BRUSATTE, S. L., DE ANDRADE, M. B., DESOJO, J. B., BEATTY, B. L., STEEL, L., FERNANDEZ, M. S., SAKAMOTO, M., RUIZ-OMENACA, J. I. & SCHOCH, R. R. 2012. The cranial osteology and feeding ecology of the metriorhynchidrocodylomorph genera *Dakosaurus* and *Plesiosuchus* from the Late Jurassic of Europe. *PLoS ONE* 7, e44985.