Pterosauria of the Great Oolite Group (Bathonian, Middle Jurassic) of Oxfordshire and Gloucestershire, England

MICHAEL O’SULLIVAN and DAVID M. MARTILL


The current understanding of UK Middle Jurassic pterosaur taxonomy is under-developed, leading to it being previously considered a time of low diversity. This is despite the presence of a productive but under-studied pterosaur-bearing horizon extending over parts of Oxfordshire and Gloucestershire. This unit, informally called the Stonesfield Slate, is part of the Great Oolite Group and it produces the largest number of Middle Jurassic pterosaurs. There are over 200 specimens distributed across museums in the United Kingdom, America, and Australia, almost all of which are accessioned under the genus *Rhamphocephalus* and referred to three species: the type species *Rhamphocephalus prestwichi*, *Rhamphocephalus bucklandi*, and *Rhamphocephalus depressirostris*. This study reviews the British Middle Jurassic Pterosauria assemblage, evaluating both their systematics and taxonomic diversity. The holotype of *Rhamphocephalus*, an isolated skull table, is found to be a misidentified crocodylomorph skull and the genus is considered a nomen dubium. The holotype of *Rhamphocephalus bucklandi* is identified as missing and that of *Rhamphocephalus depressirostris* has characters diagnostic at a family level, not a generic or specific one. Both species are considered dubious. Detailed examination of the entire assemblage shows that rather than being monogenic, the assemblage contains at least five pterosaur taxa, representing three families. This diversity includes the potential earliest occurrences of both Monofenestrata and Pterodactyloidea. A new genus, *Klobiodon rochei* gen. et sp. nov. is described based on a well-preserved mandible. The English Bathonian pterosaur assemblage is shown to be diverse and indicates that, as has been suggested in other studies, the low-diversity signal in the Middle Jurassic is at least partially artificial.

**Key words:** Pterosauria, *Rhamphocephalus*, Jurassic, Bathonian, Taynton Limestone Formation, UK, England, Stonesfield Slate.

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**Introduction**

In recent years there has been a renewed interest in Jurassic pterosaur taxonomic diversity (Butler et al. 2009, 2012, 2013; O’Sullivan 2017; O’Sullivan and Martill 2017; O’Sullivan and Rigby 2017). Historically, the understood pattern of pterosaur evolution has been that pterosaur appear in the Late Triassic and serve almost as background fauna, with no radical morphological shifts for much of the Jurassic. The group then underwent an explosion of diversity in the Late Jurassic and remain highly variable for much of the Cretaceous (Romer 1956; Wellnhofer 1991). This was supported by the high level of pterosaur productivity from Late Jurassic formations in Germany and Asia, e.g., the Solnhofen Limestone Formation which currently contains 11 pterosaur genera while almost all Triassic and Early--Middle Jurassic pterosaur units have only one or two species (Butler et al. 2009; Vidovic and Martill 2014). However, several recent studies (e.g., Butler et al. 2009, 2012, 2013; Tutin and Butler 2017) argue that this is an artificial diversity signal, affected by taphonomic biases. As we move into the Late Jurassic there is an increase in Lagerstätten which produce exceptional numbers of exquisite fossils and it is argued this creates a much higher preservation potential for fragile pterosaur remains, resulting in more species being identified. The smaller number of Lagerstätten in the Triassic and Early–Middle Jurassic is said to result in a much lower yield in pterosaur fossils, whose weaker construction makes them more susceptible to destruction under normal preservational conditions (Butler et al. 2012, 2013). The mentioned studies have used data analyses to test Jurassic pterosaur diversity signals but there has not been a
practical examination done on a pterosaur assemblage due to the low yield of most Late Triassic and Early Jurassic formations. There are, however, exceptions with several formations which have a fossil assemblage comparable to the Late Jurassic Lagerstätten. Of these the most heavily studied is Middle to Late Jurassic Tiaojishan Formation of China (Czerkas and Ji 2002). This produces large numbers of complete and near complete pterosaurs which identifies not only a high number of taxa but several transitional forms (Lü et al. 2010). There is also another, frequently overlooked, source of information found in the United Kingdom. The Bathonian (Middle Jurassic) Great Oolite Group is an oolitic unit exposed in the Oxfordshire and Gloucestershire regions of the UK. It has produced some 215 isolated pterosaur fossils (see SOM 1, Supplementary Online Material available at http://app.pan.pl/SOM/app63-OSullivan_Martill_etal_SOM.pdf). Most of these specimens have been recovered from the Taynton Limestone Formation, colloquially known as the Stonesfield Slate. The material is largely incomplete, isolated and often fragmentary but contains a variety of fossil types ranging from partial mandibles, cervical vertebrae and various appendicular elements (Fig. 1). Despite the large number of specimens, the assemblage contains only one named genus, *Rhamphocephalus* Seeley, 1880, divided into 3 poorly defined species. Since its inception, the genus *Rhamphocephalus* has never been systematically evaluated but its validity has been questioned several times (see below). The high productivity and age of the Great Oolite Group makes it a strong candidate for examining whether Jurassic pterosaur diversity really did “explode” in the Late Jurassic or whether the signal has been artificially inflated by the low yield of older formations. What follows is an historical account of Great Oolite Group pterosaur research and the first detailed examination of the assemblage, testing the validity of established taxa, looking for novel animals and gaining an understanding of what the diversity of the Great Oolite Group pterosaur fauna tells us about the nature of pterosaur diversity in the Jurassic.

**Institutional abbreviations.**—GSM, British Geological Survey, Keyworth, UK; MUM, Manchester University Museum, Manchester, UK; NMW, National Museum of Wales, Cardiff, Wales, UK; NHMUK, Natural History Museum, London, UK; NHMW, Natural History Museum, Vienna, Austria; OUM, Oxford University Museum, Oxford, UK; UUPM, University of Uppsala, Sweden.

**Other abbreviations.**—DPC, deltopectoral crest; MC, metacarpal; WP, wing phalanx.

**Nomenclatural acts.**—The electronic edition of this article conforms to the requirements of the amended International Code of Zoological Nomenclature, and hence the new names contained herein are available under that Code from the electronic edition of this article. This published work and the nomenclatural acts it contains have been registered in ZooBank. The LSID for this publication is: urn:lsid:zoobank.org:act:F79D2658-AF38-4F16-87C2-4BAC9F1021E5. The electronic edition of this work was published in a journal with an ISSN and has been archived. It is available from the following digital repositories: PubMed Central, LOCKSS.

### Historical background

The earliest known record of Bathonian pterosaurs is an account of “fossil bird bones” from the Taynton Limestone Formation of Stonesfield by an anonymous author A.B., appearing in the March edition of the Gentleman’s Magazine of 1757 (AB 1757). This brief mention was made 27 years before the first scientific description of a pterosaur, *Pterodactylus antiquus* Sömmerring, 1812, by Alessandro Collini (1784). Delair and Sarjeant (2002) determined that A.B. was a pseudonym for David Stuart Erskine (later Earl of Buchan), a prolific fossil collector of his day. However, Howlett et al. (2017) argued against this identity, pointing out that the pseudonym was not used by Erskine until 1785 and he would have been only 15 years old at the time of publication. Instead, they suggested that the initials stand for “Artium Baccalaureus” or Bachelor of Arts in Oxford University. A pterosaurian identification can be given to this material based on the work of William Buckland (1836), who (on the suggestion of John Samuel Miller an eminent Bristol based naturalist) later proposed that all supposed avian material from the Taynton Limestone should be considered pterosaur; a point reiterated by several others (e.g., Owen 1859a; Weishampel et al. 2004). Unfortunately, these original specimens were likely subsumed into various 19th century collections and can no longer be traced.

Over the decades that followed several probable Bathonian pterosaur fossils were described (e.g., Parkinson 1811; Ure 1829) but it wasn’t until 1832 that a specimen from the Taynton Limestone Formation was formally acknowledged.
Meyer (1832) erected the taxon Pterodactylus bucklandi for a disarticulated WP3 and WP4 but the specimens were never figured or identified in text. The original description was attributed by Meyer (1832) to the naturalists Spix and Martius, however, there is no record of them describing Stonesfield material. Johann Baptist von Spix (Spix 1817) did publish a description of an associated WPIII and WPIV from the Solnhofen Limestone Formation which he described as a fruit bat, and so it is possible that Meyer (1832) misread this description and mistakenly believed it was from Stonesfield. Regardless, Pterodactylus bucklandi was diagnosed in text on what was perceived as a unique WP ratio, where WPIII was longer than WPIV; and the phalanges being three times the size of the analogous elements in Pterodactylus.

Huxley (1859) discussed several isolated pterosaur elements from the Great Oolite Group, including lower jaws from both Stonesfield and lithologically equivalent strata at Sarsden, also in Oxfordshire (Boneham and Wyatt 1993). Much of this material is now accessioned in the NHMUK, OUM, and GSM collections. The most significant of Huxley’s (1859) mandibles were an unnumbered specimen, hereafter referred to as “Huxley 1859: fig. 2” (Fig. 2A) and GSM 113723 (Fig. 2B). He argued that these jaws represented different species and referred GSM 113723 to the new taxon Rhamphorhynchus depressirostris Huxley, 1859 based on the robust nature of the jaw and shallow anterior rostrum; reduced curvature in the mandible, and 5 teeth in the dentary. The affinities of “Huxley 1859: fig. 2” were less well described with Huxley (1859) figuring the mandible as belonging to either Rhamphorhynchus depressirostris or Pterodactylus bucklandi. In the text, however, Huxley (1859: 666) noted: “… it is impossible to compare the mandibles I have described with the corresponding parts of the species named Pterodactylus bucklandi, inasmuch as none exist; but since a specific name published without a description has no authority, I shall not hesitate to affix the name of Rhamphorhynchus bucklandi to the nearly perfect mandible from Stonesfield, and provisionally, to the other remains in the same area.”

Several other specimens were figured within the text including a posterior mandible, first wing phalanx (WPI), a fourth metacarpal (MCIV), a humerus and 2 scapulocoracoids (Huxley 1859: figs. 4–9). These other remains were referred to Rhamphorhynchus bucklandi Huxley, 1859 without a formal diagnosis.

The pterosaur fauna of the Taynton Limestone Formation was discussed in detail by Phillips (1871). He did not revise the taxonomy but figured some of the fossils, even providing a life-restoration of Rhamphorhynchus bucklandi. Shortly after, Owen (1874) erected three new Pterodactylus species for Stonesfield specimens: Pterodactylus kiddi Owen, 1874, Pterodactylus duncani Owen, 1874, and Pterodactylus aclandi Owen, 1874 for three isolated WPIs (Fig. 3) based on their proportions and size. It is likely that size was used as a diagnostic character based on an assumption that fully volant pterosaurs must have been osteologically mature and therefore significant size differences, as in birds, could be taxonomically diagnostic. It has since been argued that pterosaurs may have been precocial flyers (Unwin 2005) and thus overall size may have only limited taxonomic application, although a recent analysis of a pterosaur nesting site suggested that newly hatched pterosaurs were unable to fly (Wang et al 2014, 2017).

An isolated cranium (OUM J.28266) from the Great Oolite Group was made the holotype of Rhamphorhynchus prestwichi Seeley, 1880 (see Systematic palaeontology). This specimen is exposed on a limestone slab in dorsal view and was assigned to Pterosaurus based on its perceived thin bone walls. Seeley (1880) noted that the arrangement of bones was more crocodilian than pterosaurian and considered this construction diagnostic of the new taxon. Significantly he (Seeley 1880: 30) stated: “I shall be quite prepared to find
that all the ornithosaurians from Stonesfield belong to this or an allied genus which had *Rhamphorhynchus* for its nearest ally."

Five years later, Seeley (1885) briefly noted that the Stonesfield Slate pterosaur assemblage included *Rhamphorhynchus* and a new genus named *Dolichorhamphus* Seeley, 1885 for which there is no description or referred specimens. Later authors (Romer 1956; Kuhn 1967; Wellnhofer 1978) would treat *Dolichorhamphus* as a junior synonym of *Rhamphocephalus*. Lydekker (1888) re-evaluated the Stonesfield assemblage and synonymized *Rhamphorhynchus bucklandi* and *Rhamphorhynchus depressirostris* with *Rhamphocephalus* while maintaining the separate species. He also presented a revised diagnosis for *Rhamphocephalus depressirostris* Lydekker, 1888 that included features from a newly discovered mandible (NHMUK PV OR 47991; Fig. 4) with in-situ teeth. *Rhamphocephalus* was now defined by the anterior teeth being taller than the posterior and a constriction of the cranium near the orbits. Lydekker (1888) also included the fusion of the scapula and coracoid as a character, a now well-known ontogenetic feature. Lydekker (1888) then provided new definitions for both species: *Rhamphocephalus depressirostris* Lydekker, 1888 and *Rhamphocephalus depressirostris* Lydekker, 1888. *Rhamphocephalus bucklandi* he characterized as possessing more than 5 mandibular tooth pairs, its large size and the inferior border of the mandibular ramus being markedly concave. *Rhamphocephalus depressirostris* was diagnosed by having 5 mandibular tooth pairs and a prominent ventral dip to the jaw line. Several other specimens were placed in *Rhamphocephalus* with size determining species assignment. In the process, Lydekker (1888) synonymized *Pterodactylus duncani* and *Pterodactylus kiddi* with *Rhamphocephalus bucklandi*, and *Pterodactylus aclandi* with *Rhamphocephalus depressirostris*.

Other investigations into the Bathonian pterosaur assemblage involved several researchers. Owen (1859b) identified two cervical vertebrae as *Pterodactylus bucklandi* based on their occurrence in the same formation (Andres et al. [2010] noted that these were unusual for non-pterodactyloid cervicals in that they appeared to possess postexapophyses ventrolateral to the posterior condyle). Seeley (1901) attempted to establish the proportions of *Rhamphocephalus* relative to the Toarcian *Campylognathoides Strand*, 1928 from the Lower Jurassic of Germany using isolated appendicular elements. He argued that *Campylognathoides* may be a junior synonym of *Rhamphocephalus*, based on some perceived crossover of the measurements of the wing and hind limb elements. Arthaber (1922) referred to *Rhamphocephalus prestwichi* as *Rhamphorhynchus prestwichi* Arthaber, 1922, perhaps unaware of Lydekker’s work (Lydekker 1888) or simply choosing to ignore it. Between 1926 and 2011, *Rhamphocephalus* received very little attention with the genus attracting only brief mentions or overviews in the literature (e.g., Kuhn 1967; Wellnhofer 1978; Bennett 1996; Sayão 2003; Andres et al. 2010) and receiving limited evaluation. An exception is Unwin (1996) who provided a slightly expanded diagnosis of *Rhamphocephalus*, including postcranial characters such as the sacral vertebral count. He argued that all Stonesfield
material except for the holotype of Rhamphocephalus prestwichi should be synonymized with Rhamphocephalus bucklandi, considering Rhamphocephalus prestwichi highly unusual in that its skull table was massive, texturally coarse and unusually elongate with large temporal openings. He also suggested that Rhamphocephalus prestwichi was not a pterosaur, but subsequent authors continued to treat it as one. Martill (2010) provided a review to the history of pterosaurs in England, including a very brief overview of the Bathonian material currently assigned to the various Rhamphocephalus species, but passed no comment on the validity of Rhamphocephalus. Buffetaut and Jeffery (2012) mentioned the holotype of Rhamphocephalus prestwichi briefly but erroneously referred to it as Rhamphocephalus sedgwicki Buffetaut and Jeffery, 2012.

Geological setting

The Great Oolite Group is a Bathonian (165–166 ma) carbonate dominated stratigraphic unit cropping out widely in central and eastern England, with pterosaur material recovered only from the Oxfordshire and Gloucestershire outcrops (Boneham and Wyatt 1993; Benton and Spencer 1995; Gradstein et al. 2004; Fig. 5). It comprises several oolitic limestone formations with interbedded marly and sandy limestone, silty sandstone, and thin mudstone layers (Arkell 1933; Sellwood et al. 1985; Boneham and Wyatt 1993) which occur throughout the sequence (Arkell 1933; Sellwood et al. 1985). Some of the limestones of the Great Oolite Group split in a “slatey” fashion and thus were commercially mined as a tiling stone between the 17th and early 20th centuries (Plot 1677; Aston 1974; Martill 2010). Mining was concentrated around the village of Stonesfield, Oxfordshire (Fig. 6), where the mines have since been declared the Stonesfield Slate Mines SSSI (Aston 1974).
Most pterosaur specimens from the Great Oolite Group accessioned in British collections are recorded as coming from the Stonesfield Slate (see SOM 1) but as discussed by Boneham and Wyatt (1993), confusion surrounds the source horizons for the tilestones. It has alternately been identified as a sedimentary unit occurring at the top (Hull 1860; Woodward 1894) or bottom (Plot 1677) of the Taynton Limestone Formation; a member within the Sharp’s Hill Formation (Sellwood and McKerrow 1974); or as a unit at the top of the Sharp’s Hill Formation (Arkell 1947; Torrens 1980). Boneham and Wyatt (1993) attempted to identify the source formation of the Stonesfield Slate using cored boreholes drilled from the Stonesfield mines in 1991. They determined that the Stonesfield Slate material was collected from three levels within the Procerites hodsoni Zone (Wyatt 1996) of the Taynton Limestone Formation. The Stonesfield Slate Member was identified as a sequence of interbedded sandstone, siltstone and oolites from the central part of the Taynton Limestone Formation. The Stonesfield Slate Member facies was found to be recurrent throughout the Great Oolite Group in the Stonesfield area and it was concluded that the informal name had been applied to material from other members or even other formations. There is thus doubt as to whether all the material within British collections accessioned under the Stonesfield Slate label is from the Stonesfield Member, from another unit within the Taynton Limestone Formation or even from another formation within the Great Oolite Group. In this study, Middle Bathonian specimens are treated as being from the Taynton Limestone Formation unless explicitly accessioned under another stratigraphic name.

Material and methods

The studied collections house 215 specimens with the majority held at the NHMUK and OUM, 70% of the total assemblage. A complete list of accessioned fossils can be found in the SOM 1.

While most Bathonian pterosaur specimens were freely available for study, two specimens have been excluded from this analysis. NHMUK PV R 464 is a partial pterosaur skull accessioned in the NHMUK. A highly significant fossil, it is currently being studied by Brian Andres. Another specimen, OUM J.01419 is an isolated jaw from the OUM collection described by Buffetaut and Jeffery (2012) as a pterodactyloid that, if correct would make it one of the oldest pterodactyloid pterosaurs. Andres et al. (2014) expressed concern about this identification, suggesting it most likely belonged to a crocodylomorph and consequently it is excluded from this analysis.

This study also does not further evaluate Pterodactylus kiddi, Pterodactylus duncani, and Pterodactylus aclandi. These species were erected for isolated wing phalanges without accompanying diagnoses. For a WPI to be diagnostic of a taxon it must have at least 1 exceptional feature, which these three fossils do not possess, having the typical morphology of Great Oolite Group phalanges (see below; SOM 2). The lack of any distinguishing characteristics leads these three species to be considered nomina dubia and consequently the taxa are not further examined within the text. The phalanges themselves are described as they do have some familial level taxonomic significance.

Systematic palaeontology

See SOM 1 for a complete list of studied specimens. The Bathonian pterosaur assemblage includes numerous high-quality specimens such as a small number of cranial and mandibular specimens, several axial elements and examples of appendicular bones. While a large percentage of the fossils are sufficiently complete to be fully described (examples shown in Fig. 7), reviewing every well-preserved specimen lacking notable diagnostic features is beyond the scope of this paper. Instead, descriptions of the better examples are to be found in SOM 2.

Clade Reptilia Laurenti, 1768
Clade Archosauria Cope, 1869
Clade Pterosauria Kaup, 1834
Non-monofoenestratan pterosaur indet.  

Fig. 8A.
Material.—MUM STR1244b, left MCIV, Stonesfield, Oxfordshire (Bathonian, Middle Jurassic).

Description.—MUM STR1244b (Fig. 8) is a 20 mm long left MCIV exposed in dorsal view with an enlarged large double condyle. It is preserved in anterior view with an expanded ventral condyle. It is short and broad with a length/width ratio of approximately 4:1, falling within the typical range of non-monofenestratan pterosaurs (Wild 1984; Wellnhofer 1991; Martill et al. 2013), and well below the 9:1–20:1 range seen in monofenestratans such as *Darwinopterus* Lü, Unwin, Jin, Liu, and Ji, 2010.

Remarks.—More precise identification of MUM STR1244b is difficult as pterosaur metacarpals in anteroposterior views are only infrequently figured and their diagnostic features less well documented than those exposed in dorsal, ventral, cranial, and caudal views. We restrict our identification to Non-Monofenestrata indet.

Clade Breviquartossa Unwin, 2003

Family Rhamphorhynchidae Seeley, 1870

Remarks.—Rhamphorhynchidae is one of the best-known Jurassic pterosaur groups. First named by Seeley (1870) for its most famous member *Rhamphorhynchus* Meyer, 1846 the family is divided into the subfamilies Rhamphorhynchinae and Scaphognathinae. Rhamphorhynchines possess slender skulls with a well-developed elongate anterior prow, procumbent fangs, tall neural spines and some of the longest wing/body ratios of any non-pterodactyloid pterosaur with wingspans reaching 2 m in some species (Witton 2013; O’Sullivan and Martill 2017). Scaphognathines have thicker skulls, more robust cervical vertebrae, shorter prows, more vertically directed teeth and generally shorter wingspans (Unwin 2005; Witton 2013). Rhamphorhynchidae is the most common pterosaur group in the Bathonian of Britain, making up more than 90% of the assemblage and including at least 3 different species, including a novel
taxon. Excluded from this categorization is the holotype of *Rhamphocephalus prestwichi*, which is instead identified as non-pterosaurian material (see below).

**Genus Klobiodon nov.**

*Type species:* *Klobiodon rochei* gen. et sp. nov.; see below.

*Etymology:* From the Greek *klobion*, small cage and *don*, tooth; in reference to the large anterior laniaries that appear to form a fish grab.

*Diagnosis.*—As for the type and only species (see below).

**Klobiodon rochei** sp. nov.

*Fig. 4.*

1878 *Pterodactylus raptor*; Waterhouse 1878: 34.

1888 *Rhamphocephalus depressirostris*; Lydekker 1888: 36.


2012 *Rhamphocephalus depressirostris* (Lydekker, 1888); Steel 2012: 1347.

*ZooBank LSID:* urn:lsid:zoobank.org:act:F79D2658-AF38-4F16-87C2-4BAC9F1021E5

*Etymology:* After comic book artist Nick Roche. Comic books are a medium where extinct animals are portrayed in an increasingly scientifically accurate manner, and Roche’s work in the late 2000s was one of the earlier examples in this renaissance in palaeoart.

*Holotype:* NHMUK PV OR 47991, a right lower jaw in right lateral view.

*Type locality:* Stonesfield, Oxfordshire, UK.

*Type horizon:* Stonesfield Slate Member, Taynton Limestone Formation, Bathonian, Middle Jurassic.

*Material.*—Holotype and OUM 28410, an isolated mandibular symphysis from the type locality.

*Diagnosis.*—A rhamphorhynchine pterosaur possessing a unique combination of fang-like laniaries and short but robust medial teeth; medial teeth with a vertical height at least 1.3 times the width of the alveolar base; a posterior laniary 1.4–2.4 times the height of the first medial tooth; the longest laniaries are at least 1.5 times the depth of the dentary at its deepest point.

*Description.*—NHMUK PV OR 47991 (Fig. 4, Table 1) is a 140 mm long jaw preserved in left lateral view with a broken mandibular symphysis and posterior ramus. Using *Rhamphorhynchus* (NMHU PV R 37002) and *Dorygnathus* (MBR 1920.16) as proxies (Padian 2008a; Bonde and Leal 2015) NHMUK PV OR 47991 is estimated to be missing approximately 22% of the total jaw, giving it an estimated original length of ~180 mm. The mandible has a consistent depth of 13 mm posterior to the diverging rami at the mandibular symphysis. From the mandibular symphysis to the anterior break, the jaw depth increases to 17 mm. The ramus curves gently ventrally through an arc of approximately 165°. Mandibular specimen NHMUK PV OR 47991 preserves the second and third anterior teeth as well as the first two medial teeth. Two morphotypes are present; elongate recurved anterior laniaries and shorter, straighter medially positioned teeth. A concave margin on the dorsal surface anterior to the first preserved tooth is identified by Lydekker (1888) as the alveolus for the first tooth. The first preserved laniary is 26 mm long with an 8.6 mm diameter and is directed anteriorly 120° relative to the jaw line. It is elongate and strongly recurved throughout its length. The second preserved is 19 mm long with an 8 mm wide alveolus. It is more robust than its predecessor and less curved distally. Its mesial margin is broader and more sharply recurved relative to the first preserved tooth. It is angled at 110° relative to the jaw line. There is a marked size differentiation between the anterior laniaries and the medial teeth. Even assuming the first medial tooth is a replacement tooth due to its small size, such teeth are always at least 60% erupted (Fastnacht 2008). Thus, the third preserved tooth is at least 1.4–2.4 times the height of the first medial tooth, giving the dental profile a stepped appearance (Fig. 3). The first medial tooth is 7.9 mm tall with a 6.5 mm base. It is squat and more triangular than the highly recurved laniaries. Like the posterior-most laniary it has a slightly recurved mesial margin and a straight, less expanded distal margin. The second medial tooth is 10.8 mm tall and with a 7.2 mm base. Despite the size difference, the two medial teeth are similarly shaped. Based on a comparison with the jaw of NHMUK PV R 37002, *Klobiodon* is estimated to have achieved an adult wingspan of ~2 m.

Besides holotype, only one other specimen can confidently be referred to *Klobiodon*. OUM J.28410 (Fig. 9) is an isolated mandibular symphysis with three complete alveoli, 63 mm anteroposteriorly and 13 mm dorsoventrally at its deepest point. It is broken posterior to the third alveolus, around the midpoint of the incomplete fourth alveolus and the alveoli are approximately 8 mm wide mesiodistally. The first alveolus is directed dorsoanteriorly, whereas the second and third alveoli are more dorsally oriented. The anterior symphysis is developed into a large sweeping prow which makes up 28% of the preserved jaw. The large alveoli and the relatively deep jaw are similar to the arrangement of the holotype of *Klobiodon rochei*. However, *Klobiodon rochei* is primarily defined on dental characters whereas OUM J.28410 lacks teeth. Therefore, it can only tentatively be placed in the genus but is sufficiently similar in its overall morphology to be provisionally identified as *Klobiodon cf. rochei*.

*Remarks.*—Basal pterosaur mandibles are highly variable between taxa (Fig. 10) and often taxonomically informative. The Triassic pterosaurs *Austriadactylylus* Dalla Vecchia, Wild, Hopf, and Reitner, 2002 and *Preondactylylus* Wild, 1984 have similar dental morphologies with tightly packed,

**Table 1. Measurements for the teeth of NHMUK PV OR 47991.**

<table>
<thead>
<tr>
<th>Tooth</th>
<th>Crown length (mm)</th>
<th>Alveolar width (mm)</th>
<th>Crown width (mm)</th>
<th>Distance to next tooth (mm)</th>
<th>Tooth/Jaw depth</th>
<th>Tooth length as a ratio</th>
</tr>
</thead>
<tbody>
<tr>
<td>2nd laniary</td>
<td>26</td>
<td>8.6</td>
<td>1.7</td>
<td>9.3</td>
<td>1.7</td>
<td>3.3</td>
</tr>
<tr>
<td>3rd laniary</td>
<td>18.7</td>
<td>8</td>
<td>1.2</td>
<td>7.2</td>
<td>1.2</td>
<td>2.4</td>
</tr>
<tr>
<td>1st medial</td>
<td>7.9</td>
<td>6.5</td>
<td>1.4</td>
<td>10.3</td>
<td>0.6</td>
<td>1</td>
</tr>
<tr>
<td>2nd medial</td>
<td>10.8</td>
<td>7.2</td>
<td>1.6</td>
<td>NA</td>
<td>0.8</td>
<td>1.4</td>
</tr>
</tbody>
</table>
sub-triangular serrated teeth (Dalla Vecchia et al. 2002; Dalla Vecchia 2003). *Eudimorphodon* Zambelli, 1973, *Cariniasaurus* Dalla Vecchia, 2009, and *Caviramus* Fröbisch and Fröbisch, 2006 share similar complex dentitions, each with tightly spaced heterodont teeth and relatively enlarged anterior laniaries (Wild 1984; Stecher 2008; Dalla Vecchia 2009). *Dimorphodon* Buckland, 1829 from the lowermost Jurassic of southern England caniform teeth which are tightly packed in the anterior half of the jaws, becoming more widely spaced with the gaps reducing posteriorly (Buckland 1829; Padian 1984a).

*Campylognathoides* Strand, 1928 from the Toarcian of southern Germany has several anterior enlarged caniform teeth comparable in size to the anterior laniaries seen in several of the more basal pterosaurs (Plieninger 1894). Scaphognathines (sensu Lü et al. 2012) are well known for their jaws whose depth can be as much as 1.4 times the length of the longest dentary tooth crown (Goldfuss 1831; Wellnhofer 1991; Cheng et al. 2012). The tip of the jaw possesses a short prow, less prominent than that of rhamphorhynchines (Wellnhofer 1975; Cheng et al. 2012). The teeth are slightly taller medially, giving the dentition a slightly arched profile (Wellnhofer 1975; Cheng et al. 2012; Bennett 2014) and widely spaced. The curvature and robustness of the teeth in scaphognathines is somewhat variable between taxa and their position in the jaw (Carpenter et al. 2003; Cheng et al. 2012; Bennett 2014).

Rhamphorhynchines (sensu Lü et al. 2012) possess large,
procumbent fang-like laniaries (Wellnhofer 1975, 1978; He et al. 1983; Padian 2008b; Hone et al. 2012; Lü et al. 2012) that can mesh together to form a “fish-grab” (Wellnhofer 1991; Kellner and Tomida 2000; Unwin 2003), a cage-like structure at the anterior rostrum associated which some authors associate with an at least partially piscivorous diet (e.g., Wellnhofer 1991). The mandibular symphysis in rhamphorhynchines develops into a hooked anterior prow that may vary in length ontogenetically (Wellnhofer 1975, 1978). The Toarcian *Dorygnathus* has three enlarged recurved laniaries, which become less procumbent posteriorly. Directly behind the posterior most laniary there is a marked step in tooth height, with the next tooth being 16–33% the height of the last laniary. Posteriorly the successive teeth are of a similar height, are more erect and have a high triangular outline in lateral aspect. The Chinese rhamphorhynchine *Angustinaripterus* He, Yang, and Su, 1983 has teeth of relatively equal height with the anterior third becoming more strongly procumbent. The Late Jurassic rhamphorhynchines *Bellubrunnus* Hone, Tischlinger, Frey, and Röper, 2012 from the Late Jurassic of Solnhofen and *Qinglongopterus* Lü, Unwin, Zhao, Gao, and Shen, 2012 from the Oxfordian of China have a similar dentition to *Rhamphorhynchus* with the teeth being slightly procumbent and fang-like, and the anteromesial teeth being the most anteriorly inclined. The teeth show a similar profile to those of as *Scaphognathus* Goldfuss, 1831 with the “arch” created by the elongate third tooth. The Middle Jurassic scaphognathine *Jianchangnathus robustus* Cheng, Wang, Jiang, and Kellner, 2012 possesses a similar dental arrangement to *Scaphognathus* but with much larger, more anteriorly oriented laniaries, deep jaw and convex prow. Bennett (2014) argued that these features did not generically separate the two taxa and referred *J. robustus* to *Scaphognathus robustus*.

Wukongopteridae (sensu Wang et al. 2009) have numerous slightly recurved, well-spaced teeth (Wang et al. 2009; Lü et al. 2011; Martill and Etches 2012). Ctenochasmatoids (sensu Unwin 2003; also Pterodactylidae sensu Pereda Suberbiola et al. 2012) possess a wide variety of dental morphologies or multiple cusps of *Austriadactylus*, *Preondactylus*, *Eudimorphodon*, *Caviromus*, or *Carniadactylus*.*Dimorphodon* and *Campylognathoides* lack laniaries and have more closely spaced alveoli. Wukongopterid teeth have tighter spacing, straighter jaws and do not possess laniaries. The curvature of the jaw and the size of the teeth show that *Klobiodon* belongs in Rhamphorhynchinae (sensu Lü et al. 2011), the parent clade of Scaphognathinae and Rhamphorhynchinae.

In *Klobiodon* the longest tooth is much larger relative to the depth of the jaw than is common for scaphognathines (Cheng et al. 2012; Bennett 2014). At least five rhamphorhynchine taxa are known with lower jaws preserved and can be compared directly with *Klobiodon*. Of these five taxa, the lower jaws of *Bellubrunnus* and *Qinglongopterus* are crushed dorsoventrally and difficult to compare to *Klobiodon* (Hone et al. 2012; Lü et al. 2012). *Angustinaripterus* lacks complete teeth but the alveoli are of a relatively consistent size (He et al. 1983). This lack of medial/anterior tooth differentiation distinguishes it from *Klobiodon*. *Dorygnathus* and *Rhamphorhynchus* show some size differentiation in the teeth. In *Rhamphorhynchus* this differentiation is subtle with a slight increase in the tooth size towards the middle of the jaw followed by a gradual decrease posteriorly. This gives it a dental profile not dissimilar to *Scaphognathus* but distinct from *Klobiodon*. In contrast, *Dorygnathus* shares several similarities with *Klobiodon*. Like *Klobiodon* the jaws of *Dorygnathus* can possess a gentle curvature throughout their length, developing into a robust and well-developed prow (Padian 2008b; Fig. 10). *Dorygnathus* also possesses a well-developed differentiation between small medial teeth and large somewhat recurved laniaries. These similarities do suggest that *Klobiodon* is more closely allied to *Dorygnathus* than other rhamphorhynchine pterosaurs, however, *Klobiodon* does possess a combination of characters that distinguish it from *Dorygnathus*. While both pterosaurs have small medial teeth, in *Dorygnathus* they are smaller, more closely spaced and less robust than in *Klobiodon* (Padian 2008a). In *Klobiodon* the posterior most laniary is 1.4–2.4 times the size of the first medial tooth. In *Dorygnathus* (SMNS 55886; Padian 2008b), the posterior most laniary is between 2.6 and 3.7 times the size of the first medial tooth. In the Vienna specimen of *Dorygnathus* (NHMW 1911/0001/0023 and is close to the size of the *Klobiodon* holotype [Padian 2008b]), the ratio is 3.3. This difference is a consequence of the medially positioned teeth being much larger and more robust in *Klobiodon* compared to *Dorygnathus*, with a tooth length/ alveolar width ratio in the anterior medial teeth of at least 1.3. In the above *Dorygnathus* specimens, the ratio ranges between 2.6 and 6, depending on the mesiodistal width of the tooth. The laniaries of *Klobiodon* are robust, with broad alveoli while those in *Dorygnathus* are not only more strongly anteriorly inclined but comparatively thinner mesiodistally. In *Klobiodon* the longest tooth in the jaw is at least 1.5 times the dorsoventral depth of the jaw at its deepest point. Contrasting this, in NHMW 1911/0001/0023 the tooth/jaw ratio is around 2.6 and in *Dorygnathus* UUPM R156 the ratio increases to 3. This is due to *Klobiodon*’s laniaries being shorter relative to the depth of the jaw whereas *Dorygnathus* has elongate laniaries set in a comparatively narrower mandible (Padian 2008b). Within Scaphognathinae the most comparable animals are the two species of *Scaphognathus*. *Scaphognathus crassirostris* Goldfuss, 1831 possess smaller less procumbent teeth and
is thus a poor comparison with *Klobiodon*. *Scaphognathus robustus* on the other hand has more comparable laniaries, being larger and more procumbent. This may suggest a possible relationship, but the taxa can be distinguished on several characters. Unlike *Klobiodon*, *S. robustus* lacks the strong size differentiation between the anterior and medial teeth seen in *Klobiodon*, with the most posterior tooth being almost 1/1 in scale to the anterior fangs. In *S. robustus* the longest tooth/jaw depth ratio is 0.8, less than is seen in *Klobiodon* and is due to the greater depth of its jaw.

While the elongated laniaries, convex anterior prow and deep jaw may indicate a relationship between *Klobiodon* and *S. robustus*, the prominent differentiation between the large anterior laniaries and small medial teeth are more like *Dorygnathus* which also possess a convex prow and anteriorly oriented fangs. The similarities with both taxa, combined with the continued lack of broad consensus in the structure of non-monofenestratan phylogeny (e.g., Lü et al. 2012; Andres and Myers 2013; Vidovic and Martill 2014) recommends caution is assigning the new genus to either Scaphognathinae or Rhamphorhynchinae. *Klobiodon rochelii* is therefore conservatively identified here as a member of the Rhamphorhynchidae rather than assigned to either subgroup.

Remarks.—NHMUK PV OR 47991 was first mentioned by Waterhouse (1878) in a vertebrate acquisitions list for the NHMUK, who confirmed the specimen was donated by Robert Marsham. In the list Waterhouse (1878: 34) can be quoted as listing NHMUK PV OR 47991 as “mandible, right ramus, of *Pterodactylus raptor*, of Stonesfield Slate, Stonesfield”. The name *Pterodactylus raptor* was taken from an unpublished manuscript by Richard Owen. No formal description or figure reference has been associated with this name but Ingles and Sawyer’s (1979) compilation of NHMUK illustrations referenced NHMUK PV OR 47991 as being illustrated in Folio 201A of Richard Owen’s unpublished Collection of Drawings under the name *Pterodactylus raptor* (Michael Hanson, personal communication 2017). Lydekker (1880) placed NHMUK PV OR 47991 in *Rhamphocephalus* and identified it as a large example of *Rhamphocephalus depressirostris*. He argued that it possessed two characteristics of the species: five teeth in the dentary and a concave ventral jaw line. It was considered the best example of *Rhamphocephalus depressirostris* in the NHMUK collections and was noteworthy for its relatively large size. Lydekker’s (1880) description of NHMUK PV OR 47991 was written immediately after his new diagnosis of *Rhamphocephalus depressirostris* and this may explain some confusion on the part of Benton and Spencer (1995) who mistakenly identified NHMUK PV OR 47991 as the type specimen for *Rhamphocephalus depressirostris*.

*Stratigraphic and geographic range.—*NHMUK PV OR is most likely from the Stonesfield Slate Member of the Taynton Limestone Member, collected from the Stonesfield region of Oxfordshire.

Genus *Rhamphocephalus* Seeley, 1880 (nomen dubium)

**Remarks.**—The genus *Rhamphocephalus* is identified in this study as a non-diagnostic thalattosuchian crocodylomorph (see Non-pterosaurian section below). This assignment raises questions as to the taxonomic identification of the holotypes of “*Pterodactylus* bucklandi” and “*Rhamphorhynchus* depressirostris”, two isolated jaws described below.

“*Pterodactylus* bucklandi” Meyer, 1832 (nomen dubium)

(= Rhamphorhynchinae indet.)

**Fig. 2A.**

1832 *Pterodactylus bucklandi*; Meyer 1832: 27–30.
1859 *Rhamphorhynchus bucklandi* (Meyer, 1832); Huxley 1859: 658–670.
1859 *Pterodactylus bucklandi* Meyer, 1832; Owen 1859b: 169.
1871 *Rhamphorhynchus bucklandi* (Meyer, 1832); Phillips 1871: 224.
1888 *Rhamphocephalus bucklandi* (Meyer, 1832); Lydekker 1888: 34–363.
1907 *Rhamphocephalus bucklandi* (Meyer, 1832); Seitz 1907: 289–6291.
1978 *Rhamphocephalus bucklandi* (Meyer, 1832); Wellnhofer 1978: 41.
2003 *Rhamphocephalus bucklandi* (Meyer, 1832); Sayão 2003: 335.
2003 *Rhamphocephalus bucklandi* (Meyer, 1832); Unwin 2003: 177.
2008 *Rhamphocephalus bucklandi* (Meyer, 1832); Barrett et al. 2008: 68.
2012 *Rhamphocephalus bucklandi* (Meyer, 1832); Buffetaut and Jeffery 2012: 1.
2012 *Rhamphocephalus bucklandi* (Meyer, 1832); Steel 2012: 1341.
2013 *Rhamphocephalus bucklandi* (Meyer, 1832); Witton 2013: 125.

**Holotype.**—“Huxley 1859: fig. 2”, a lower jaw in left lateral view. The whereabouts of the specimen is unknown.

**Type locality.**—Smith’s Quarry, Sarsden, Oxfordshire, UK.

**Type horizon.**—Great Oolite Group, Bathonian, Middle Jurassic.

**Material.**—NHMUK PV R 28610, 32752, 37765, 38014, 38015, 38016, 38017, 38019, 38020, 40126, 47994, 47999a, 1028, 1029, 1030, 1824, 2637, 6749, 6750 (Steel 2012); OUM J.28275, J.28537, J.28303 (Huxley 1859), isometrically preserved from the Stonesfield region of Oxfordshire with no cross over with the holotype of “*Pterodactylus* bucklandi”.

**Description.**—The holotype (“Huxley 1859: fig. 2”) of *Rhamphocephalus bucklandi* (Fig. 2A) is an isolated jaw with 5 alveoli and a short anterior prow. The holotype was (and remains) lost but as the other specimens figured by Huxley (1859) are accessioned in the OUM collections, it is likely it may have been held there at one point. Based on Huxley’s (1859) figure 2, it is a rhamphorhynchine jaw due to its anterior prow, its being relatively shallow and its low number of teeth. As we cannot locate the specimen, our only reference point being a drawing (which the holotype of *Rhamphocephalus prestwichi* highlights can be a poor representation of the actual specimen) and Huxley’s (1859) only
identifier being it is morphologically distinct from *R. depressirostris*, we cannot recommend a further detailed analysis. As there is no crossover between Huxley’s (1859) figure and any of the appendicular material assigned to the genus by Lydekker (1888), we also identify the name as inapplicable to anything but “Huxley 1859: fig. 2” and transfer all remaining material to either Pterosauria indet. or Rhamphorhynchinae indet. (see SOM 1). This unfortunately puts “*Pterodactylus bucklandi*” in a nebulous position. We can confidently state that the number of teeth, slim jaw and distinct prow would place the taxon within Rhamphorhynchinae. We can also say that the teeth seem less broad and the jaw less dorsoventrally thick than *Klobiodon* but unless the holotype is one day uncovered, we regrettable cannot test the taxon any further.

Remarks.—Placed within Rhamphorhynchinae indet. due to its proved mandibular symphysis and alveoli indicating broad, well-spaced teeth. The holotype has been lost.

“*Rhamphorhynchus depressirostris*” Huxley, 1859 (nomen dubium)

(= ?Scaphognathinae indet.)

Fig. 11.

1888 *Rhamphocephalus depressirostris* (Huxley, 1859); Lydekker 1888: 34–36.
1978 *Rhamphocephalus depressirostris* (Huxley, 1859); Wellnhofer 1978: 41.
1996 *Rhamphocephalus depressirostris* (Huxley, 1859); Unwin 1996: 293.

2003 *Rhamphocephalus depressirostris* (Huxley, 1859); Unwin 2003: 177.
2008 *Rhamphocephalus depressirostris* (Huxley, 1859); Barrett et al. 2008: 69.
2012 *Rhamphocephalus depressirostris* (Huxley, 1859); Buffetaut and Jeffery 2012: 1.
2012 *Rhamphocephalus depressirostris* (Huxley, 1859); Steel 2012: 1341.
2013 *Rhamphocephalus depressirostris* (Huxley, 1859); Witton 2013: 125.

Holotype: GSM 113723, a three-dimensional lower jaw symphysis preserved in left lateral view.

Type locality: Smith’s Quarry (51°54’8.62"N 1°34’44.74″W), Sarsden, Oxfordshire, UK.

Type horizon: Fuller’s Earth Formation, Bathonian, Middle Jurassic.

Material.—Holotype and NHMUK PV R 40126, isolated limb material; Taynton Limestone Formation (Bathonian, Middle Jurassic), Stonesfield, Oxfordshire (Steel 2012).

Description.—GSM 113723 is a three-dimensional and nearly complete lower jaw symphysis with partial rami (Fig. 11). It is 87 mm long and contained within a block of oolitic limestone typical of the Stonesfield Slate Member. The medial left and lateral right rami are both obscured by matrix, as is the dorsal aspect of the symphysis. It has been damaged with the right ramus broken off and reattached. The anterior symphysis is missing, exposing its oval cross-section and thin bone walls. The ventral posterior symphysis is bowed indicating the presence of a prow. In ventral view the symphysis has a relatively deep sulcus at its posterior boundary. There are five alveoli preserved on the rami, with the fifth alveolus of both rami bearing an in-situ tooth. There are no alveoli preserved posterior of these teeth. The alveoli are 5–6 mm mesiodistally and spaced 3–8 mm apart, with the spacing increasing posteriorly. The
anterior alveoli are slightly splayed, giving the dorsal symphysis an undulose margin and hinting that the teeth projected anterolaterally. The preserved teeth are 15 mm long, thin and peg-like, with teeth directed anteriorly at ~70° relative to the jaw line.

The elongate jaw, thin bone walls, the dental arrangement and smooth bone texture confirm GSM 113723 as a pterosaur. The low tooth count and simple elongate teeth distinguish GSM 113723 from the more complex tooth pattern of basal non-rhamphorhynchid pterosaurs. The height and number of the medially placed teeth also serves to separate it from Jurassic monofenestratans, which tend to have numerous relatively low medial teeth (Wellnhofer 1978, 1991; Wang et al. 2010). GSM 113723 is confidently placed in Rhamphorhynchidae due to the narrow mandibular symphysis, reduced tooth count and wide spacing of the teeth. The apparent lack of teeth posterior to the fifth alveolus may ally it with Scaphognathinae as these pterosaurs have no more than 5–6 teeth in the lower jaw (Unwin 2003). Tall, peg-like teeth, wide medial spacing and relatively short prows are also found in scaphognathines (Cheng et al. 2012; Bennett 2014; Zhou 2014), being similar to the jaw of Scaphognathus (= Jianchangnathus) robustus (Bennett 2014), a scaphognathine from the Tiaojishan Formation of China.

GSM 113723 is identified here as ?Scaphognathinae but the question remains: can it be assigned to a genus or species? Neither Huxley’s (1859) or Lydekker’s (1888) definition of Rhamphocephalus depressirostris (robust jaw, shallow anterior rostrum, reduced curvature in the mandible, 5 teeth) are diagnostic by modern taxonomic standards. Unfortunately, the taphonomy of the jaw limits morphological taxonomy. The preserved tooth is clearly distinct from Klobiodon but is still similar to what we see in several Jurassic pterosaur taxa (see above) and lacks the distinct ratios inform the definition of Klobiodon. There is a deep mandibular sulcus which can be seen at the posteroventral prow but this character could be ontological. Overall, morphological definition is limited. We also cannot suggest the erection of a chronotaxon, not only because it runs counter to the taxonomic practices of this research but also because named scaphognathines occur in contemporaneous formations (e.g., Cheng et al. 2012). Therefore, GSM 113723 is not currently considered a strong candidate for a generic or specific name despite representing a distinct mandibular morphotype within the assemblage. Regrettably this puts “Rhamphocephalus” depressirostris in a similar nebulous position as “Pterodactylus” bucklandi, if for different reasons. However, if GSM 113723 can at some point undergo further preparation or perhaps CT-scanning, we may reveal currently obscured diagnostic characters.

Remarks.—Alleged attribution to Scaphognathinae Hooley, 1913 is based on the morphology of the teeth and consistent increase in the alveolar spacing towards the posterior end of the jaw.

Other rhamphorhynchid remains

Other identifiable rhamphorhynchid fossils consists of axial and dental remains collected from the Taynton Limestone Formation of Stonesfield and the Chipping Norton Formation of Chipping Norton, both in Oxfordshire (Bathonian, Middle Jurassic). Full locality information is available in SOM 1.

Scapulocoracoids.—OUM J.28294 (Fig. 12B) is a partial left scapulocoracoid preserved in posterolateral view, heavily worn around the glenoidal region. The scapula is 37 mm long and the coracoid is 23 mm long. The coracoid possesses a low biceps tubercle that extends 2 mm ventrally and 6 mm proximodistally. The glenoid is posterolaterally positioned and limited to the scapula. The angle formed between the scapula and coracoid is at 75°. A second complete scapulocoracoid, OUM J.28295 (Fig. 12A) is near complete and preserved in anteroposterior view. It is missing the proximal ends of both the scapula and the coracoid as well as the surface bone of the glenoid. In the case of the glenoid, this exposes the internal trabeculae. The scapula is 28 mm long and the coracoid being 40 mm long. The angle between them is approximately 75°. The restriction of the glenoid to the scapula identifies them as non-pterodactyloid. The proximal coracoid is slenderer than Campylognathoides or Dimorphodon (Buckland 1829; Padian 2008a). The low biceps tubercle is like Dorygnathus, Darwinopterus, and Campylognathoides (Padian 2008a; Lü et al. 2010) but lower than other basal pterosaurs. The angle of 75° falls into the range shared by several basal pterosaurs (Wild 1984; Padian 2008b; Witton 2013). The glenoid limited to the scapula and approximately 10 mm proximodistally. The lateral scapulocoracoid is less inclined than in Rhamphorhynchus but in a similar position to Dorygnathus and Scaphognathus. The coracoid distinguishes it from Darwinopterus where it is wider dorsoventrally and does not taper proximally (Lü et al. 2010). Based on the angle formed by the scapula and coracoid, along with the thin coracoid morphology, the scapulocoracoids are identified as ?Rhamphorhynchidae indet.

A complete right scapulocoracoid (OUM J.28297) is preserved in medial view (Fig. 13) and is the most complete pterosaur shoulder element in the assemblage, with a 62 mm long scapula ramus and a 60 mm coracoid ramus. The two elements form an angle of ~70°. The glenoid is restricted to the scapula and is ~16 mm long. The acrocoracoid process is rounded and extends 4.5 mm distal of the glenoid. The coracoid bows slightly behind its scapular articulation. The sternocoracoidal joint is 3 mm wide dorsoventrally and is more rounded on its ventral surface. The biceps tubercle is very low, extending 1 mm ventrally and 7–9 mm proximodistally. OUM J.28297 is identified as a non-pterodactyloid by the glenoid restriction and the coracoid not exceeding the scapula in length. However, the scapula is of sub-equal length to the coracoid whereas in most basal pterosaurs, the scapula is between 124–60% the length of the coracoid (Buckland 1829; Wild 1984; Padian 2008a, b). In Sericipterus (Andres
et al. 2010) and Rhamphorhynchus the scapula and coracoid are of sub-equal lengths. Although this suggests a rhamphorhynchine affinity, the same situation occurs in Darwinopterus (Lü et al. 2011). However, Darwinopterus has a larger biceps tubercle than OUM J.28297 while in rhamphorhynchines the size of the biceps tubercles is variable (Padian 2008b; Andres et al. 2010; O'Sullivan and Martill 2015). Based on the ratio of the scapula to the coracoid and the profile of the biceps tubercle, OUM J.28297 is tentatively identified as ?Rhamphorhynchinae indet.

Humeri.—OUM J.23043 (Fig. 14A) is a complete left humerus exposed in dorsal view, 90 mm long proximodistally with a diaphysis 10 mm wide anteroposteriorly at its medial point with a length/width ratio of 9/1. The medial process and distal articulation are preserved primarily as an external mould on the rock surface, but with a clearly distinct outline. The diaphysis develops a gentle anterior bowing distally, angled at approximately 165° and distally has a round anterior articulation which extends 8 mm anterior to the shaft. The medial process has a triangular outline and is 10 mm long anteroposteriorly. The DPC is 13 mm anteroposteriorly and 19 mm proximodistally. The anterior margin is rounded, giving the DPC a short tongue-like shape (Fig. 14A). OUM J.23043 is one of the largest Jurassic pterosaur humeri known, 7% larger than the largest Dorygnathus humerus and 13% that of the largest Rhamphorhynchus (Wellnhofer 1975; Padian 2008b). It can be distinguished from via the morphology of the DPC and medial process. Eudimorphodon (Wild 1984), C aviationus (Stecher 2008) and Campylognathoides (Padian 2008a) have enlarged quadrangular DPC. Wukongopterids have similar morphologies to OUM J.23043 but the DPC is less robust (Lü et al. 2011). The medial process of aurorazhdarchids and basal ctenochasmatoids are distally deflected away from the proximal articulation (Wellnhofer 1978; Vidovic and Martill 2014). OUM J.23043 lacks the medial pinching of the DPC seen in Nesodactylus Colbert, 1969 and some examples of Rhamphorhynchus. The DPC of Scaphognathus
is a like OUM J.23043 with a comparable placement of the medial process. However, the proximal margin of the DPC is straighter and the DPC is less strongly deflected (Bennett 2014). *Sericipterus* has a higher length/width ratio than OUM J.23043 (O’Sullivan et al. 2013) and a more elongate DPC (Andres et al. 2010). The humerus of *Dorygnathus* is like OUM J.23043 (Padian 2008b) but the DPC is generally more elongate. Ultimately OUM J.23043 does not perfectly correlate with any described pterosaur humerus but the DPC, medial process and the gentle curvature of the diaphysis is most like *Scaphognathus* (Cheng et al. 2012; Bennett 2014), and thus it is assigned to ?Scaphognathinae indet. *Scaphognathus* humeri make up at least 6% of the total wingspan (Bennett 2014), suggesting a total wingspan of 1.5 m.

NHMUK PV R 40126b (Fig. 14B) is a small 15 mm long humerus, and 1.5 mm wide medially. It is almost complete, missing only the proximal humeral head, medial process and some of the distal articulation. The DPC is 2.4 mm anteriorly, semi-tongue shaped with a broad sub-rectangular anterior margin and a slight proximodistal pinching towards the extremity of the DPC. NHMUK PV R 40126b is identified as ?Rhamphorhynchinae indet based on the slight elongation and minor pinching of the DPC. At 15 mm long NHMUK PV R 40126b is one of the smallest pterosaur humeri known and based on similarly sized examples of *Rhamphorhynchus* from the Solnhofen Limestone (Wellnhofer 1975), is most likely a juvenile with a 290–340 mm wingspan. NHMUK PV R 40126c (Fig. 14C) is a right humerus preserved in dorsal view. It is 8 mm long proximodistally with a 6 mm wide diaphysis, giving it a length/width ratio of 13:1. It is missing a large section of its humeral head and distal diaphysis. The distal articulation is not preserved. The diaphysis curves anteriorly at 170–175°. NHMUK PV R 40126c does preserve the medial process. The DPC has a rounded anterior margin. The high length/width ratio in combination with the anteriorly bowed shaft of NHMUK PV R 40126c is indicative of it being a possible rhamphorhynchine identification. NHMUK PV R 40126c is identified here as Rhamphorhynchidae, most likely ?Rhamphorhynchinae. The possibility that NHMUK PV R 40126c represents an adult example of NHMUK PV R 40126b and that the straighter shaft in the latter specimen is an ontogenetic feature cannot be ruled out. However, without any intermediaries the two morphologies are considered sufficiently different to rule out an ontogenetic transition. Thus, NHMUK PV R 40126a–c confirms the presence of a second large humeral morphotype that can be assigned to Rhamphorhynchinae rather than Scaphognathinae.
NHMUK PV R 28160a is a left pterosaur humerus preserved in dorsoposterior view (Fig. 14D). It is 14 mm proximodistally with a 1–2 mm medial diaphysis. It is anteriorly bowed through an arc of ~160°. The medial process and DPC both project into and are partially overlain by the matrix. The diaphysis has several “dimples” across the posterior margin corresponding with an immature bone texture as illustrated by Tumarkin-Deratzian et al. (2007). This, combined with its small size, indicates that NHMUK PV R 28160a is a juvenile. The DPC proximally deflected ventral margin. It is similar in overall morphology to OUM J.23043 and is thus also identified as ?Scaphognathitinae indet. It should be noted that Unwin (2015) considered NHMUK PV R 28160a as a possible monofenestratan humerus but did not explain his reasoning in detail.

**Wing phalanges.**—There are several WPIs within the Bathonian pterosaur assemblage (Fig. 2; see SOM 1), although their preservation is variable. The WPIs are of a seemingly uniform morphology, being relatively large and robust elements between 101–142 mm long. Each WPI diaphysis is bowed medially such that the proximal and distal termini are angled at 170–175° relative to the midpoint. The presence or absence of a posterior longitudinal groove is indeterminate in all specimens given their preservation, however, NHMUK PV R 40126 C (Fig. 2B) is not only free of surrounding sediment but broken medially, revealing a triangular cross-section with a relatively thick bone wall and a more rounded anterior surface. Taxonomic identification of these phalanges is problematic given their isolated nature. The consistent robustness and slightly bowed diaphysis suggest they are most likely from a single taxon with distinctive anterior bowing. This is unusual as most pterosaurs have straight proximal phalanges (Wellnhofer 1991; Witton 2013). Unwin (2003) suggests that a bowed phalanx may be a dsungaripteroid apomorphy but notes that a dsungaripteroid presence in the Taynton Limestone is problematic as presently there is no substantive evidence for Dsugaripteroidea in strata older than Kimmeridgian (Fastnacht 2008). Anterior bowing is, however, found in several Dorygnathus specimens (Padian 2008b) and it is therefore not a dsungaripteroid apomorphy. The phalanges of dsungaripteroids are more elongate (Young 1964) than those of the British Bathonian pterosaurs while those of Dorygnathus (e.g., SMNS 56255; Padian 2008b) are shorter and somewhat broader, as in NHMUK PV R 40216. He we identify the phalanges as Rhamphorhynchidae indet. Based on their similarity to the phalanges of Dorygnathus the wingspans of these pterosaurs are estimated to be between 1.4 m and 2 m.

**Mandibles.**—NHMUK PV R 1824 (Fig. 15), is broken transversely and divided into part and counterpart. The jaw is missing most of the anterior symphysis. It is 100 mm long with rami approximately 13 mm deep dorsoventrally. Both rami preserve five alveoli but with no teeth. The alveolar sections of the rami comprise approximately 49% of the total ramus length. The posterior alveoli are dorsally placed with the more anterior alveoli becoming more laterally placed. The alveoli range between 4–5 mm mesiodistally and are set 6–8 mm apart. The taxonomic identity of NHMUK PV R 1824 is difficult to determine. The number of medial teeth posterior to the symphysis, their relatively uniform size and spacing suggest it is a rhamphorhynchid but it is unclear if it is distinct from or synonymous with Klobiodon rochei. While the holotype of Klobiodon rochei has similar anterior alveolar spacing, the rami of NHMUK PV R 1824 appears more gracile than Klobiodon rochei, and the rami lack the gentle curve seen in Klobiodon. The rami remain relatively straight but curve ventrally proximal to the posterior articulation. It is tentatively identified as Rhamphorhynchidae indet.

**Cranial material.**—UMZC T.718 (Fig. 16) is a semi-3D posterior skull, 52 mm anteroposteriorly and 37 mm dorsoventrally. It is catalogued under the genus ‘Rhamphinion
Fig. 16. Partial skull of “Rhamphinion jenkensi” Padian, 1984b (UMZC T.718) from the Taynton Limestone Formation (Bathonian, Middle Jurassic), Stonesfield, Oxfordshire, UK; in left lateral view (matrix digitally removed).

Rhamphinion jenkensi” Padian, 1984b, an Early Jurassic American genus from the Kayenta Formation of north-eastern Arizona. While the holotype of Rhamphinion is from the same region of the skull, there is very little proportional similarity between the two fossils (see Padian 1984: fig. 1 for comparison) and as the identification has no accompanying descriptive text, this identification is considered here to be unsupported. It preserves the quadrate, quadratojugal, jugal, postorbital and squamosopostorbital bar. The quadrate is 32 mm dorsoventrally and 1.6 mm medially in lateral view. It is elongated, strap-like in posterior view and angled posteriorly at 118°. Ventrally it has a well-developed and rounded condyle for articulating with the mandibular glenoid. The condyle is ~3.8 mm wide. Only the postorbital and quadrate processes of the squamosal are preserved but their full extent is obscured by fractures and some fusion of the skull. The postorbital is tri-radiate, 8 mm anteroposteriorly and 11 mm dorsoventrally. The squamosoparietal bar is elongate and sub-rectangular, 11 mm long anteroposteriorly and 4 mm wide dorsoventrally. The quadratojugal is sub-triangular, with slightly elongated jugal and quadrate processes. The quadratojugal is well-developed, 2.3 mm dorsoventrally and 6.7 mm anteroposteriorly, with the jugal being the largest element in the specimen. Including its processes, it is 34 mm anteroposteriorly and ~6 mm dorsoventrally. It is a sub-quadrangular bone, although the maxillary process is more of a flange, being large and sub-rounded where it borders the antorbital fenestra. The postorbital process is the longest of the four but the point of contact with the postorbital is one of the most damaged regions of bone, obscuring its true length. It is estimated to be 8–16 mm long. The postorbital and lacrimal processes form an angle of ~70° around the ventral margin of the orbit. UMZC T.718 preserves three fenestrae in varying degrees of completeness. The superior temporal fenestra is missing its dorsal half. It is bounded by the squamosal and postorbital and is 14 mm anteroposteriorly. The ventral margin is smooth and sub-oval. The anterior border is more vertical than the posterior, and the anteroventral margin is shallower than the posterior. The inferior temporal fenestra is the only complete fenestra in the specimen. It is 28 mm dorsoventrally and 14 mm anteroposteriorly. The ventral margin is smooth and sub-oval. The anterior border is more vertical than the posterior, and the anteroventral margin is shallower than the posterior. Its boundary includes the quadrate, quadratojugal, jugal, postorbital and squamosal. It possesses an irregular piriform morphology outline, with the dorsal half being wider with a relatively straight dorsal boundary in comparison to the ventral half’s thinner, more angular appearance. The orbit is missing its dorsal half but is clearly the largest of the three fenestrae. It is bounded by the jugal ventrally and the postorbital posteriorly. Like the inferior temporal fenestra, the orbit is piriform with the ventral boundary approximately 10 mm wide in comparison to the 24 mm dorsally positioned widest point. UMZC T.718 is identified as a non-monofenestrate based on its low-lying antorbital fenestra, the relatively shallow angle of the quadrate and its tetraradiate jugal. Generic identification is problematic given the limited information: the angle of the quadrate at 118° distinguishes it from Dimorphodon (Buckland 1829), Campylognathoides (Padian 2008a) and Anurognathus (Bennett 2007) but is comparable to several other basal pterosaurs, including Eudimorphodon, Austriadactylus and Dorygnathus (Wild 1984; Padian 2008a, b). The angle formed by the dorsal processes of the jugal is typical of basal pterosaurs more derived than Campylognathoides or Eudimorphodon (Wellnhofer 1975; He et al 1983; Wild 1984; Padian 2008b; Bennett 2014). The broad sub-rounded inferior temporal fenestra is distinct from most basal pterosaurs (Wild 1984; Padian 2008a) but a similar condition is seen in several rhamphorhynchids (He et al. 1983; Padian 2008b; Cheng et al. 2012; Bennett 2014). Based on the angle of the quadrate combined with the ventral orbital and inferior temporal fenestra morphology, UMZC T.718 is identified here as ?Rhamphorhynchidae.

OUM J.28409 (Fig. 17) is an 83 mm long isolated and near-complete left maxilla and a fragment of the posterior premaxilla. Much of the premaxilla and the posterior maxilla are absent. The surface texture appears irregular, but this is a taphonomic artefact as most of the external bone wall has been eroded away anteriorly, revealing the internal trabeculae. The centrally positioned nasal process is angled posteriorly at 70–90°. While the posterior margin of the antorbital fenestra is not present in the maxilla, the preserved section of the premaxilla defines the anterior border of the nares. The nares is approximately 23 mm anteroposteriorly. The antorbital fenestra is at least 27 mm along the same plane. It has five maxillary alveoli. The smallest is 4 mm mesiodistally, with the subsequent alveoli approximately equidimensional at around 7 mm. OUM J.28409 is identified as pterosaur based on a combination of the nares and antorbital fenestra being proximal to each other and the thin bone walls. It is further be identified as a non-monofenestratin as it possesses differentiated nares and antorbital fe-
nstra. The possession of five relatively widely spaced maxillary tooth pairs is a rhamphorhynchid characteristic and OUM J.28409 is identified as Rhamphorhynchidae indet.

**Teeth.**—Numerous isolated teeth (Fig. 18) have been collected from the Taynton Limestone Formation and accessioned within pterosaur collections. However, due to the difficulty in identifying ex-situ teeth with a simple slender conical, gently curved morphology, many of these have been misidentified. Gently recurved teeth with a slightly sigmoidal shape accessioned as pterosaur might more correctly be assigned to Teleosauridae (Massare 1987). Pterosaur teeth can be identified by the restriction of the enamel to the higher parts of the tooth crown while the base of the crown is exposed dentine. The exposed dentine sometimes extends laterally to a median point of the crown (Witton 2013). Six teeth thought to be pterosaurian, lacking carinae and seemingly possessing restricted enamel are illustrated here (Fig. 18). They share a similar morphology of being relatively elongate and distally recurved, making them likely to be rhamphorhynchine.

**Monofenestrata remains**

Monofenestrata is a recently erected group (Lü et al. 2010) including the transitional Wukongopteridae and the derived Pterodactyloidea. Wukongopteridae includes several taxa showing mosaic evolution with animals possessing derived skull morphology but plesiomorphic bodies. The recognition of this group has not only revolutionised our understanding of pterosaur evolution but complicates Middle Jurassic taxonomy. With few uniquely diagnostic features (Witton et al. 2015) found within the group, isolated remains are easily assigned to either basal or pterodactyloid pterosaurs, and consequently, this has been a major contributor to the conservative approach adopted in this study. Pterodactyloidea is a highly diverse group including at least 11 families. Distinguished from basal pterosaurs by features such as metacarpal length, rostral index, dentition, cervical structure, pelvic morphology and numerous other features, the group includes the most well-known pterosaurs (Wellnhofer 1991; Unwin 2005). The Middle Jurassic has gained increasing relevance in their evolution as several discoveries push the group further and further back (Andres et al. 2014). Establishing if any monofenestratans are present in the Bathonian of Britain has considerable significance to Jurassic pterosaur diversity.

**Vertebrae.**—Whereas several examples of Bathonian pterosaur vertebrae are known (see SOM 1), the most diagnostic specimens are those accessioned under NHMUK PV R 40126a. These are two isolated cervical vertebrae referred to here as CVA and CVB (Fig. 19). CVA is an isolated cervical vertebra within a slab of oolitic limestone while CVB is a compressed, but three dimensional isolated cervical free of matrix. Both were described by Owen (1859b) as coming from the “Stonesfield oolite” but no locality data was provided. CVA is identified as a cervical based on its robustness, the lack of strongly developed horizontal transverse processes and its relatively elongate centrum (Howse 1986). It is 16 mm anteroposteriorly and 12 wide mm transversely. Based on its quadrangular appearance it is identified as a cervical 3–7 (Wellnhofer 1991). In comparison to CVB, CVA is the more poorly preserved. As well as being dorsoventrally compressed, it is heavily fractured along its lateral
margins. Both the pre- and postzygapophyses are preserved but are incomplete. The neural spine has been broken, most likely during compaction. CVB is morphologically similar to CVA and considered a cervical 3–7. It is 25 mm anteroposteriorly (excluding the posterior condyle) and 15 mm medi-ally. It has undergone dorsoventral compression, the loss of the neural spine and the right prezygapophysis. CVB has undergone some repair or preparation as an adhesive fills several fractures in the surface. Despite this damage, CVB is well preserved and provides a clear view of the centrum in ventral view. Ventrolateral to the posterior condyle are two enlarged projections that in dorsal view are slightly dished. Andres et al. (2010) provided a brief description of NHMUK PV R 40126c, identifying these structures as postexapophyses. In the anteroventral centrum a prominent hypapophysis extends posteriorly towards the midpoint of the cervical. On either side of the neural arch are oval structures which are most likely lateral pneumatic foramina (Wellnhofer 1991; Andres and Norell 2005). On either side of the vertebra are transverse processes, broken proximal to the centrum, corresponding to the dorsal articulation loci for the cervical ribs. NHMUK PV R 40126e have been discussed previously by Owen (1859b) and Andres et al. (2010). Owen figured CVB and identified it as “Pterodactylus” bucklandi but did not provide a description. Andres et al. (2010) discussed the vertebrae in comparison with the cervicals of the rhamphothychine Sericipterus from the Shishugou Formation in Xinjiang, China. They noted that NHMUK PV R 40126e cervicals were of a similar size to those of Sericipterus Andres et al. (2010), arguing that they were non-pterodactylid with the unusual feature of possessing postexapophyses. They suggested that development of postexapophyses may correlate with increasing size of cervicals.

Whereas CVA is more limited in its taxonomic information due to taphonomy, the three-dimensional preservation of CVB allows for a more detailed comparison. Howse (1986) considered the possession of cervical ribs as a characteristic of basal pterosaurs, but reduced cervical ribs are known from several pterodactyloid clades (Zhou and Schoch 2011), including the highly derived Azhdarchidae (Witton and Naish 2008). Cervical ribs are not preserved in the holotype of the non-pterodactyloid monofenestratan Darwinopterus modularis Lü, Unwin, Jin, Liu, and Ji, 2010, but there appear to be anterolateral projections on several vertebrae which may correspond to cervical rib facets. Therefore, the presence of these facets is an unreliable character for distinguishing between pterodactyloid and non-pterodactyloid vertebrae. The presence of hypapophyses in combination with the postexapophyses corresponds with Howse’s (1986) descriptions of pterodactyloid cervical vertebrae however, as Andres et al. (2010) note, in most respects these vertebrae appear typically non-pterodactyloid. Most likely these vertebrae are from a non-pterodactyloid monofenestratan, identified here as ?Monofenestrata indet., which have garnered considerable attention for possessing a mosaic of features of both pterodactyloids and non-pterodactyloids (Lü et al. 2010), as seen in NHMUK PV R 40126a.

Metacarpals.—NHMUK PV R 28160b (Fig. 8B) is a 41 mm MCIV with a preserved length/width ratio of 8/1. It has a prominent distal condyle above which is a piece interpreted as either a broken sliver of the shaft of MCIV or the rem-
nants of one of the associated metacarpals I–III. Basal pterosaurs have MCIVs with a maximum length/width ratio of approximately 5:1 whereas pterodactyloids have a length/width ratio of 9:1–20:1 (Wild 1984; Wellnhofer 1991; Martill et al. 2013). Although a ratio of 8:1 in NHMUK PV R 28160b is closer to the pterodactyloid condition, the proximal and distal epiphyses are relatively broader and more like those of basal pterosaurs. The non-pterodactyloid monofenestratan *Darwinopterus* has a MCIV with a ratio of ~10:1, with a similar constricted medial shaft along with wider proximal and distal epiphyses (Lü et al. 2010). NHMUK PV R 28160b is therefore identified here as ?Monofenestra indet.

Suborder Pterodactyloidea Pleninger, 1901

?Pterodactyloidea indet.

Fig. 20.

**Material.**—NHMUK PV R 2637 (Fig. 20) from the Taynton Limestone Formation (Bathonian, Middle Jurassic) of Stonesfield, Oxfordshire, is a synsacrum preserved in dorsal view, 21 mm anteroposteriorly and 14 mm laterally. It is poorly preserved, having been crushed flat and heavily worn. There are at least 4 sacral vertebrae with another preserved anterior to these, although it is uncertain if this is a sacral or a dorsal vertebra. The neural spines have broken and while the transverse processes are all present, they are significantly damaged anteriorly. The transverse processes are approximately 2–3 mm anteroposteriorly with the posterior pairs horizontally oriented while the anterior transverse processes are angled posteriorly at ~160°. The last vertebra is 3 mm anteroposteriorly and its broken transverse processes appear to connect with the ilia. They form the anterior margin of a fenestra twice as broad anteroposteriorly as the previous fenestrae. Synsacs are unique to dinosaurs and pterosaurs among Taynton Limestone vertebrates (Unwin 2003; Norman 1988). NHMUK PV R 2637 is identified as pterosaurian based on its smooth and ultra-thin bone walls, and its relatively broad transverse processes. In basal pterosaurs, the sacrum comprises 4 vertebrae (Unwin 2003, 2005). Wukongopterids have at least 5 sacrals (Lü et al. 2011) and the sacrum tends to be broad mediially. In pterodactyloids, there are at least 5 sacral vertebrae (Wellnhofer 1978) and several have a large fenestra around the first sacral vertebra delineating the anterior and posterior synsacrum. The pterodactyloid synsacrum is elongate, thinning posteriorly throughout its length (Wellnhofer 1991; Witton 2013). The broad anterior fenestra and the elongate sacrum suggests that NHMUK PV R 2637 may be either a basal monofenestratan with a modified synsacrum (Lü et al. 2010) or an indeterminate pterodactyloid. If the former it would represent the first evidence of a monofenestratan modifying a module other than the skull and neck. If the latter it rep-

Fig. 20. A synsacrum of ?Pterodactyloidea indet. (NHMUK PV R 2637) from Taynton Limestone Formation (Bathonian, Middle Jurassic), Stonesfield, Oxfordshire, UK; in dorsal view. Photograph (A) and accompanying diagram (B), black regions represent the sacral fenestration.
resents the earliest occurrence of a pterodactyloid extending
the temporal range of pterodactyloids down into the Middle
Jurassic, before the Callovian–Oxfordian Kryptodrakon
progenitor Andres, Clark, and Xu, 2014.

Non-pterosaurian material

The fragmentary nature of the Bathonian pterosaur assem-
blage has led to several specimens being misidentified as
pterosaur, although most are small isolated specimens of
little note. Buffetaut and Jeffreys (2012) described a sup-
posed pterosaur jaw that Andres et al. (2014) later described
as thalattosuchian, an identification with which we concur.
The most significant misidentified non-pterosaurian fossil is
also the most well-known, the holotype of Rhamphocephalus
prestwichi.

Thalattosuchia incertae sedis
Genus Rhamphocephalus Seeley, 1880

Type species: Rhamphocephalus prestwichi Seeley, 1880; unknown
horizon, Great Oolite Group, Kineton Thorns Quarry, Stow-On-Wold,
Gloucestershire.

Included species: Rhamphocephalus prestwichi Seeley, 1880, “Rham-
phocephalus” bucklandi (Meyer, 1832) and “Rhamphocephalus” de-
pressirostris (Huxley, 1859).

Rhamphocephalus prestwichi Seeley, 1880

Fig. 21.

1880 Rhamphocephalus prestwichi; Seeley1880: 27–30.
1888 Rhamphocephalus prestwichi Seeley, 1880; Newton 1888: 534.
1888 Rhamphocephalus prestwichi Seeley, 1880; Lydekker 1888: 33.
1901 Rhamphocephalus prestwichi Seeley, 1880; Seeley 1901: 153–155.
1922 Rhamphorhynchus prestwichi (Seeley, 1880); Arthaber 1922: 9.
1995 Rhamphocephalus prestwichi Seeley, 1880; Benton and Spencer
1995: 149.
2012 Rhamphocephalus sedgwicki (sic); Buffetaut and Jeffery 2012: 1.
Holotype: Rhamphocephalus prestwichi (OUM J.28266) (Fig. 21A)
poorly preserved cranium presenting a dorsal view of the palate.
Type locality: Kineton Thorns Quarry, Gloucestershire, UK.
Type horizon: Bathonian, Middle Jurassic.
Remarks.—Rhamphocephalus is one of the oldest British
“pterosaur” taxa re-interpreted here as a thalattosuchian
based on its crocodylomorph skull morphology. The type
specimen of Rhamphocephalus prestwichi OUM J.28266
(Fig. 21A) from an unrecorded horizon in the Kineton
Thorns Quarry, Stow-On-Wold, Gloucestershire (Fig. 22)
is a poorly preserved cranium presenting a dorsal view of
the palate. It has been a contested taxon for several years
due to the lack of well-supported diagnostic characters (see
Introduction). Lydekker’s (1888) characters occur plesio-
morphically in several other pterosaur genera (Wellhofer
1991) and the skull arrangement is, as noted by Seeley
(1880), more crocodilian than pterosaurian. When a sim-
plicated reconstruction of OUM J.28266 is presented next to
the skull of Rhamphorhynchus (Fig. 21B), the differences
become apparent. Pterosaur skulls possess large, sometimes
medially positioned orbits which are like the larger fenes-
trae of OUM J.28266 but the nares are positioned more
anteriorly along the rostrum. The antorbital fenestrae are
proximal to the orbits but are laterally placed and do not sig-
nificantly invade the dorsal profile of the skull. The rugose
striations exposed on OUM J.28266 are absent in ptero-
saur bone which has a smoother internal texture (Bennett
1993). Given the lack of morphological convergence be-
tween OUM J.28266 and pterosaur skulls, it can be confi-
dently identified as non-pterosaurian. Its cranial morphol-
gy is instead more like choristodires and thalattosuchians.
Figure 23 reconstructs OUM J.28266 in two orientations,
one presenting it as a near complete medial skull and the
other as an isolated squamosoparietal region. Choristodires
usually have extremely broad superior temporal fenestrae
and orbits more anteriorly positioned than OUM J.28266
and are therefore not a likely candidate. Thalattosuchian
crocodylomorphs such as Steneosaurus Geoffroy, 1825 and
Teleosaurus Geoffroy, 1825 more strongly matches OUM
J.28266 in either orientation with their large superior tem-
poral fenestra separated by a squamosoparietal bar with a
broad ventral base (Walkden et al. 1987), small orbits
proximal to the temporal fenestrae and medially position
choanae corresponding with the elevated domes of OUM
J.28266 (Pierce and Benton 2006). Thus OUM J.28266 is
identified as thalattosuchian but the incompleteness of the
fossil limits the taxonomic resolution. With no identifiable
autapomorphies, the genus Rhamphocephalus cannot be
transferred to Thalattosuchia and is therefore regarded as
a nomen dubium.

Stratigraphic and geographic range.—The holotype of was
collected from the Stonesfield Slate of Kineton, near Stow-
on-Wold prior to 1880 (Seeley 1880). Kineton is approxi-
mately 7 miles from Stow-on-the-Wold, in Gloucestershire,
near the River Windrush (Fig. 22). Fossils were collected
from the Kineton Thorns Quarry where exposures of the
Cotswold Slate Formation and the Eyeford Member of the
Fuller’s Earth Formation underlie the Taynton Limestone
Formation (Boneham and Wyatt 1993; Sumbler et al. 2002).
The exact origin of OUM J.28266 is unrecorded, however,
it is older than the Taynton Limestone Formation pterosaurs
(~167 million years old).

Discussion

Prior to this study the most diverse Middle–Upper Jurassic
pterosaur bearing unit was the Bathonian–Oxfordian
Tiaojishan Formation of Liaoning, China. It is a highly lith-
ologically variable unit with 13 named genera (Lü et al.
2010, 2012; Bennett 2014). The assemblage includes groups
present in the Great Oolite Group like rhamphorhynchids
(3 taxa) and monofenestratans (potentially 7 taxa) as well as 2 anurognathids, a group of highly derived small pterosaurs known for their short robust skulls (Bennett 2007). Unlike the British material, the Tiaojishan pterosaur assemblage is dominated by exceptionally preserved articulated skeletons. With at most 5 potential taxa in the Great Oolite Group the Tiaojishan Formation is clearly the most speciﬁcally diverse, but this does not diminish the importance of the British fossils. There is clear continuity between the pterosaur fauna of Europe and Asia, therefore each unit can suggest what taxa are notable by their absence. The presence of anurognathids in the Tiaojishan Formation and the Late Jurassic of Germany and Kazakhstan (Wellnhofer 1991; Witton 2015) suggests the group may have been present in the Bathonian of Britain. Perhaps their absence from the British assemblage can be attributed to their small size reducing their preservation potential.

While not the most diverse Middle Jurassic pterosaur assemblage, the Great Oolite Group presents several important points regarding pterosaur evolution. The monofenestratans of the Tiaojishan Formation are regarded as the earliest occurrence of the group, dating to the Callovian (Witton 2013). The Great Oolite monofenestratan fossils are reliably dated to the Bathonian, ~2 million years older, and thus represent an even earlier appearance for the clade. It is likely that the origins of Monofenestrata are older still with the Early Jurassic Alkaruen Codorniú, et al. 2016 recovered as the sister taxon to the group (Codorniú et al. 2016). Furthermore, the identification of Cuspicephalus Martill and Etches, 2013, as a late surviving wukongopterid (Witton et al. 2015) suggests the basal monofenestratan lineage was well established in Britain throughout the Jurassic. With regards to the potential for pterodactyloids in the assemblage, if NHMUK PV R 2637 can be placed within the group it would place the evolution of Pterodactyloidea several of million years earlier than previously recorded. Presently the oldest pterodactyloid is Liaodactylus Zhou, Gao, Yi, Xue, Li, and Fox, 2017, a ctenochasmatoid from the Oxfordian of the Tiaojishan Formation and Kryptodrakon from the Callovian–Oxfordian boundary in the Shishugou Formation.

![Fig. 22. Maps showing location of the area of the Kineton Thorns Quarry (asterisks) within the Stow-On-Wold region, Gloucestershire, England, UK.](http://www.cotswoldphotolibrary.com)
of Xinjiang, China, dating to 160 and 162 ma respectively. This is around 4–6 million years younger than the Great Oolite Group pterosaurs, making NHMUK PV R 2637 the oldest pterodactyloid and strongly suggesting the group has an early Middle or perhaps even Lower Jurassic origin. Unfortunately, NHMUK PV R 2637’s identification is tenuous but further examination will hopefully answer if this is an important fossil in pterosaur evolution.

Fig. 23. Explanatory drawings of OUM J.28266 along side the skulls of a choristodire and a thallatosuchian. A. The possible arrangement of the *Rhamphoccephalus prestwichi* Seeley, 1880 (OUM J.28266), holotype reconstructed as a complete skull (A₁) and the frontoparietal region (A₂). B. *Simoedosaurus* Gervais, 1877, a choristodire from the Early Palaeocene of Saskatchewan, Canada; in dorsal view (from Sigogneau-Russell and Russell 1978). C. *Pelagosaurus* Bronn, 1841, a thallatosuchian from the Lower Jurassic of Somerset, England; in dorsal view. B, C not to scale.
With regards to the Great Oolite Group and Jurassic pterosaur diversity, it appears to support the findings of previous studies (Butler et al. 2009, 2012, 2013) that the appearance of a sudden radiation in the Late Jurassic (Kimmeridgian–Tithonian) has been somewhat inflated by the poor yield of most old deposits. The unit may possess up to 5 taxa from 3 groups, including the earliest recorded appearance of the monofenestratans. While not directly comparable to the Solnhofen Limestone’s assemblage of 14 species, it is notably higher than other Lower or Middle Jurassic units (e.g., the Whitby Mudstone Formation which has 2 families [O’Sullivan 2017; O’Sullivan and Martill 2017] or the Posidonia Shale Formation which despite being a pterosaur Lagerstätte includes only 2 genera [Padian 2008a, b]). The Bathonian Great Oolite Group strata are separated from the Kimmeridgian “radiation” by more than 10 million years, yet there’s little suggestion that this is a period of low diversity. Further evidence for pterosaurs remaining diverse for much of the Jurassic, as well as the at least partially artificial nature of the Late Jurassic pterosaur radiation, can be found by comparing U.K. formations with the Tiaojishan Formation of Hebel and Liaoning, China (Lü et al. 2010, 2012; Bennett 2014). As discussed above it contains 13 species, at least 9 of occur in Bathonian–Oxfordian aged strata. Clearly this formation had diverse biota comparable with the Great Oolite Group which it maintained right up until the Late Jurassic transition. Yet if we look at the contemporaneous Oxford Clay Formation of Britain, a Callovian–Oxfordian unit, we find a signal identifying a period of very low diversity with only a small number of fossils preserved and no well-supported taxa (O’Sullivan 2017). The Tiaojishan Formation is a Lagerstätte laid down in a terrestrial environment with a diverse water system, resulting in better preservation and a reasonably high population density (Tan et al. 2006) nature of the Tiaojishan Formation whereas the Oxford Clay Formation is laid down in a marine environment where pterosaurs have a much lower preservation potential and a much lower population density (O’Sullivan 2017). Therefore, it seems likely that as suggested by Butler et al. (2009, 2012, 2013) pterosaur radiation at the end of the Jurassic has been artificially inflated by at least preservation biases. That does not mean that there is no evidence for greater pterosaur taxic diversification in the Late Jurassic. While the Tiaojishan Formation contains 13 species from the Bathonian to the Oxfordian, the Solnhofen Limestone Formation contains many species within the Kimmeridgian and Tithonian, a notably shorter time frame. This may still be a result of the Lagerstätten effect, but we should not assume that while pterosaurs were much more diverse in the Middle Jurassic than traditional models may indicate that there wasn’t an increase of some kind in diversity, possibly associated with a transition from a non-pterodactyloid to a pterodactyloid dominated assemblage.

Conclusions

The Middle Jurassic pterosaur assemblage of the UK is not the most diverse assemblage of its time, but it is still important as it highlights the strong phenotypic and specific variance found in the Middle Jurassic. Lacking any truly exceptional fossils, we have identified multiple lineages and morphotypes and the presence of a new genus. With two diverse assemblages on opposite sides of the world, there is now stronger evidence that the Mid Jurassic Pterosaur Gap is an artefact of low yield and poor preservation. Far from their being an evolutionary bottleneck, pterosaurs appear to have been undergoing diversification from at least the Middle and likely the late Lower Jurassic. We would recommend all early to mid-Jurassic pterosaur material would benefit from further examination and that future research take care to recognise the misleading signals of previous diversity studies.

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