

The relationships of *Cuspicephalus scarfi* Martill and Etches, 2013 and *Normannognathus wellnhoferi* Buffetaut *et al.*, 1998 to other monofenestratan pterosaurs

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Abstract

The evolution of pterodactyloid pterosaurs occurred in a 'modular' fashion with 'pterodactyloid'-type crania and cervical vertebrae evolving in pterodactyloid sister taxa – early monofenestratan pterosaurs – before later postcervical modifications marked the development of the true pterodactyloid condition. This means of evolution creates problems for distinguishing isolated pterodactyloid crania from those of non-pterodactyloid monofenestratans, and has led to uncertainty over the affinities of two Late Jurassic European pterosaurs known only from skulls, *Cuspicephalus scarfi* Martill and Etches, 2013 and *Normannognathus wellnhoferi* Buffetaut *et al.*, 1998. Some aspects of their cranial anatomy suggest affinities to early pterodactyloids – specifically the Germanodactylidae – while others indicate a relationship with a group of non-pterodactyloid monofenestratans, the Wukongopteridae. Here, we characterise the skulls of Jurassic monofenestratans to provide greater insight into the identity of these pterosaurs. We find a suite of characters indicating that *Cuspicephalus* is a wukongopterid, notable for being a particularly large and long snouted member of the group, as well as the youngest, and the first European record of this clade. The affinities of *Normannognathus* are less clear however. We consider its previous allocation to the Germanodactylidae doubtful, and note some similarities it shares with ctenochasmatoid pterodactyloids, but the only known specimen is probably too fragmentary for confident referral to any specific clade within Monofenestrata.

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Introduction

The origin of the Pterodactyloidea is currently a hot topic in pterosaur research. Whereas a clear morphological divide once separated pterodactyloids from other pterosaurs, new discoveries have revealed major stages of their early evolution. These include the purported earliest known member of the Pterodactyloidea (Andres *et al.*, 2014) and 'transitional' taxa bridging pterodactyloid-like anatomy to earlier pterosaurs (Lü *et al.*, 2010; Tischlinger and Frey, 2014). Perhaps the most significant of these 'transitional' species is the Callovian-Oxfordian Tiaojishan Formation species *Darwinopterus modularis* Lü *et al.*, 2010, a small pterosaur with anatomy 'intermediate' between that of pterodactyloids and their historically recognised sister group, the Rhamphorhynchidae (Lü *et al.*, 2010). The anatomy of *Darwinopterus* is noted for its 'modular' nature, combining 'pterodactyloid-grade' head and neck anatomy with non-pterodactyloid postcervical features (Lü *et al.*, 2010). *Darwinopterus* and the Pterodactyloidea appear to form a monophyletic clade, the Monofenestrata, named after the combined nasal and antorbital opening common to all members of this group (Lü *et al.*, 2010). Another recent discovery, a privately-owned complete skeleton from the latest Kimmeridgian Painten Formation of Germany, apparently represents a grade of monofenestratan between *Darwinopterus* and the

Pterodactyloidea (Tischlinger and Frey, 2014). As in *Darwinopterus*, its skull and neck possesses typical ‘pterodactyloid’ features while its postcranial skeleton, despite being relatively pterodactyloid-like, retains clear hallmarks of an earlier pterosaur bauplan. This specimen has not been named because of its lack of public accession and, following Tischlinger and Frey (2014), is hereafter referred to as the ‘Painten Pro-pterodactyloid’.

Since *Darwinopterus* was described, a number of similar taxa have been identified from the Tiaojishan Formation. These include two other *Darwinopterus* species, *D. robustodens* Lü *et al.*, 2011a and *D. linglongtaensis* Wang *et al.*, 2010; as well as *Kunpengopterus sinensis* Wang *et al.*, 2010, *Wukongopterus lii* Wang *et al.*, 2009 and *Changchengopterus pani* Lü, 2009. These taxa are considered to form a clade, the Wukongopteridae (Wang *et al.*, 2010), diagnosed by their combination of relatively derived pterodactyloid-like skulls and

cervical vertebrae with more plesiomorphic, non-pterodactyloid-like postcervical anatomy (Wang *et al.*, 2010; Hone, 2012; Andres *et al.*, 2014). The purported early istiodactylid *Archaeoistiodactylus linglongtaensis* Lü and Fucha, 2010 is also probably a wukongopterid (Martill and Etches, 2010; Witton, 2013) or a close relative of this group (Sullivan *et al.*, 2014). It is highly likely that the Tiaojishan wukongopterids are oversplit (Lü *et al.*, 2012; Witton, 2013).

The characterisation of non-pterodactyloid monofenestratans has solely used relatively complete skeletons unambiguously demonstrating their distinctive, ‘modular’ anatomy. Problems arise when applying these means of identifying wukongopterids to more fragmentary Jurassic pterosaur material, however. Non-pterodactyloid monofenestratan skeletons are distinctive, but their individual ‘modules’ are not strongly apomorphic, instead showing plesiomorphic anatomies which are very simi-

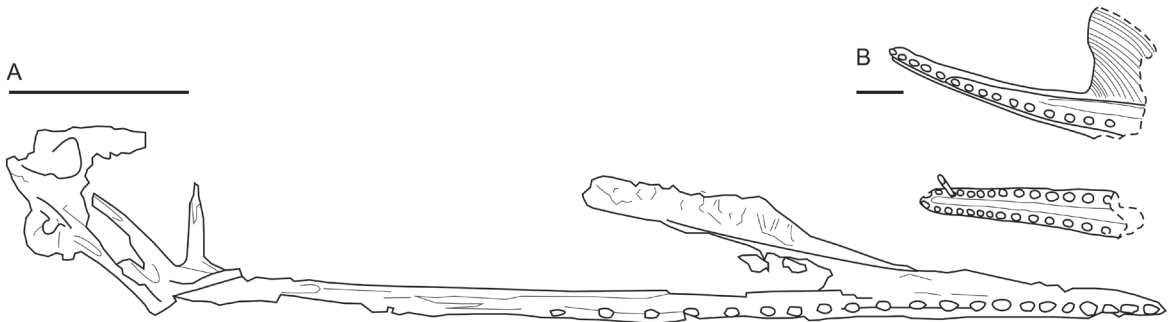


Fig. 1. A, MJML K1918, holotype skull of the long-snouted pterosaur *Cuspicephalus scarfi* Martill and Etches, 2013; B, MGCL 59'583, holotype of *Normannognathus wellnhoferi* Buffetaut *et al.*, 1998. Scale bars represent 50 mm (A) and 10 mm (B).

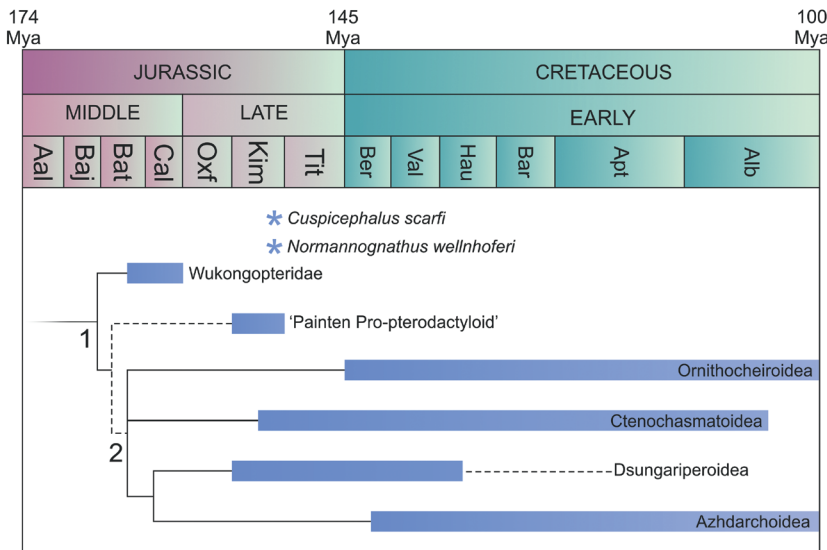


Fig. 2. Simplified stratigraphic distribution of Monofenestratan groups at the Jurassic/Cretaceous boundary, and the stratigraphic position of the two species discussed herein, *Normannognathus wellnhoferi* and *Cuspicephalus scarfi*. Phylogeny largely based on Lü *et al.* (2010, 2012), but the position of the ‘Painten Pro-pterodactyloid’ is inferred from recent work by Tischlinger and Frey (2014). 1, Monofenestrata; 2, Pterodactyloidea. Abbreviations of geologic ages: Aal, Aalenian; Alb, Albian; Apt, Aptian; Bar, Barremian; Baj, Bajocian; Bat, Bathonian; Ber, Berriasian; Cal, Callovian; Hau, Hauterivian; Kim, Kimmeridgian; Oxf, Oxfordian; Tit, Tithonian; Val, Valanginian.

lar to those of other Jurassic pterosaur clades. This raises questions over how precisely incomplete monofenestratan fossils can be classified if evidence of a combined pterodactyloid/non-pterodactyloid bauplan is absent: do they represent wukongopterids, pterodactyloids, or something else entirely? Lü *et al.* (2010) validated this concern when performing separate cladistic analyses of the cranial and cervical, and postcervical anatomy of *Darwinopterus modularis*. The head and neck ‘modules’ were found to nest deeply within the Pterodactyloidea while the postcervical module plotted as the sister taxon to the Rhamphorhynchidae (Lü *et al.*, 2010). This problem has also been borne out in other studies where classifying isolated monofenestratan crania has proved challenging (Martill and Etches, 2013; also see below). The current diagnosis of Wukongopteridae, suggested by Wang *et al.* (2010), is of little help here because it is reliant on characteristics of relatively complete specimens. Most wukongopterid cranial characters provided by Wang *et al.* (2010) are plesiomorphic for the Monofenestrata (‘confluent naris and antorbital fenestra; maxillary ramus of the jugal long, anteriorly projected and splint-like; free lateral nasal process’), and thus are of little significance without associated ‘non-pterodactyloid’-like postcrania. Other suggested cranial characters may be of questionable application to the group (‘quadrate inclined backwards for about 120°; see Martill and Etches [2013] for contrasting measurements). Wukongopterid postcranial characters, relating to neck and forelimb bone length ratios, may be synapomorphic (Wang *et al.*, 2010; Sullivan *et al.*, 2014, but require associations of several bones for their application. With the majority of the Jurassic pterosaur record comprising isolated and fragmented material (*e.g.* Unwin, 1996; Barrett *et al.*, 2008), there is a clear need for greater characterisation of early monofenestratan anatomy to ensure an accurate understanding of Jurassic pterosaur diversity.

Europe’s possible non-pterodactyloid monofenestratans

Three fragmentary pterosaur specimens have been identified as possible non-pterodactyloid monofenestratans, and all are from Europe (Steel, 2010; Andres *et al.*, 2011a, b; Martill and Etches, 2013; Witton, 2013; Tischlinger and Frey, 2014). Each is represented only by cranial remains. They include NHMUK R 464, a fragmentary skull from the Bathonian Taynton Limestone Formation (the ‘Stonesfield Slate’) of Oxfordshire (Steel *et al.*, 2010; Andres *et al.*, 2011a, b). A full appraisal of NHMUK R 464 has yet to be published but is currently underway

(Andres pers. comm.), and we accordingly await publication of this before discussing this specimen further.

MJML K1918, the holotype of the long-snouted pterosaur *Cuspicephalus scarfi* Martill and Etches, 2013 from the Kimmeridgian Kimmeridge Clay of Dorset, UK (Fig. 1A) represents a second possible non-pterodactyloid monofenestratan. Known from a nearly complete skull, Martill and Etches (2013) noted some similarities in cranial and dental features between MJML K1918 and the wukongopterid *Darwinopterus*, but concluded that ‘a close relationship cannot be proved’ (Martill and Etches, 2013: p. 285). This was in part because the specimen also bears several similarities to the pterodactyloid *Germanodactylus* (Martill and Etches, 2013: p. 291), suggesting possible affinities to a more derived monofenestratan clade. The significance of *Cuspicephalus* potentially being related to *Germanodactylus* is confused by the taxonomic controversies surrounding the latter. *Germanodactylus* has been argued as belonging to two different pterodactyloid lineages, Archaeopterodactyloidea (*sensu* Kellner, 2003) or Dsungaripteroidea (*sensu* Unwin, 2003), and the congeneric status, and placement within Pterodactyloidea, of the two recognised *Germanodactylus* species (*G. cristatus* Plieninger, 1901 and *G. rhamphastinus* Wellnhofer, 1970) are also disputed (*e.g.* Maisch *et al.*, 2004; Vidovic and Martill 2014). Irrespective of the outcome of these controversies, that *Cuspicephalus* has been likened to both pterodactyloids and basal monofenestratans validates the problems outlined above concerning to classifying fragmentary monofenestratans.

Considerations of a third potential non-pterodactyloid monofenestratan, MGCL 59’583, are similar to those of *Cuspicephalus*. MGCL 59’583 represents the holotype jaw tips of *Normannognathus wellnhoferi* Buffetaut *et al.*, 1998, from the Upper Kimmeridgian Argiles d’Ecqueville, Normandy, France (Fig. 1B). This taxon has traditionally been likened to the pterodactyloid *Germanodactylus* (Buffetaut *et al.*, 1998; Unwin, 2005) and sometimes considered part of the Germanodactylidae (Buffetaut *et al.*, 1998; Unwin and Heinrich, 1999; Andres and Myers, 2013), a possibly paraphyletic (Unwin and Heinrich, 1999; Maisch *et al.*, 2004) taxon comprising *Normannognathus*, *Tendaguripterus recki* Unwin and Heinrich, 1999, and *Germanodactylus*. However, Martill and Etches (2013) cast doubt on this identification, stating ‘[in] the light of the discovery of non-pterodactyloid monofenestratans, the holotype and only specimen of *Normannognathus* (MGCL 59’583) can no longer be placed in Germanodactylidae with confidence, and should be regarded as Monofenestrata indet.’ (p. 292).

Martill and Etches (2013) made no further elaboration on this point, but clear similarities between the crest morphology, jaw tips and dental alveoli of MGCL 59'583 and those of non-pterodactyloid monofenestrans are good cause for considering the affinities of *Normannognathus* open to question.

If *Cuspicephalus* and *Normannognathus* have affinities with germanodactylids, these pterosaurs represent some of the oldest pterodactyloid material known and certainly the oldest pterodactyloid crania (Fig. 2). If they represent wukongopterids or another form of non-pterodactyloid monofenestrans, they provide important new data on this poorly understood portion of pterosaur evolution. Because evidence presented for both possible identifications is either equivocal (*Cuspicephalus*) or in need of review (*Normannognathus*), we have attempted to resolve the phylogenetic placement of these poorly known taxa via detailed characterisation of non-pterodactyloid monofenestrans skulls. From this, we propose means to distinguish fragmentary skull mate-

rial of early monofenestrans pterosaurs from those of early pterodactyloids, and apply our findings to *Cuspicephalus* and *Normannognathus*.

Institutional abbreviations

BSP, Bayerische Staatssammlung für Paläontologie und Geologie, Munich, Germany; HGM, Henan Geological Museum, China; GPIT, Paläontologische Forschungs, Lehrund Schausammlung, Institut für Geowissenschaften, Universität Tübingen, Tübingen, Germany; IVPP; Institute of Vertebrate Palaeontology and Palaeoanthropology, Beijing, China; MGCL, Musée Géologique Cantonal de Lausanne, France; MJML, Museum of Jurassic Marine Life (The Etches Collection), Kimmeridge, Dorset, UK; NHMUK, Natural History Museum, London, UK; NMING, National Museum of Ireland, Dublin, YH, Yizhou Museum, Yixian, Liaoning Province, China; ZMNH, Zhejiang Museum of Natural History, Hanzhou, Zhejiang Province, China.

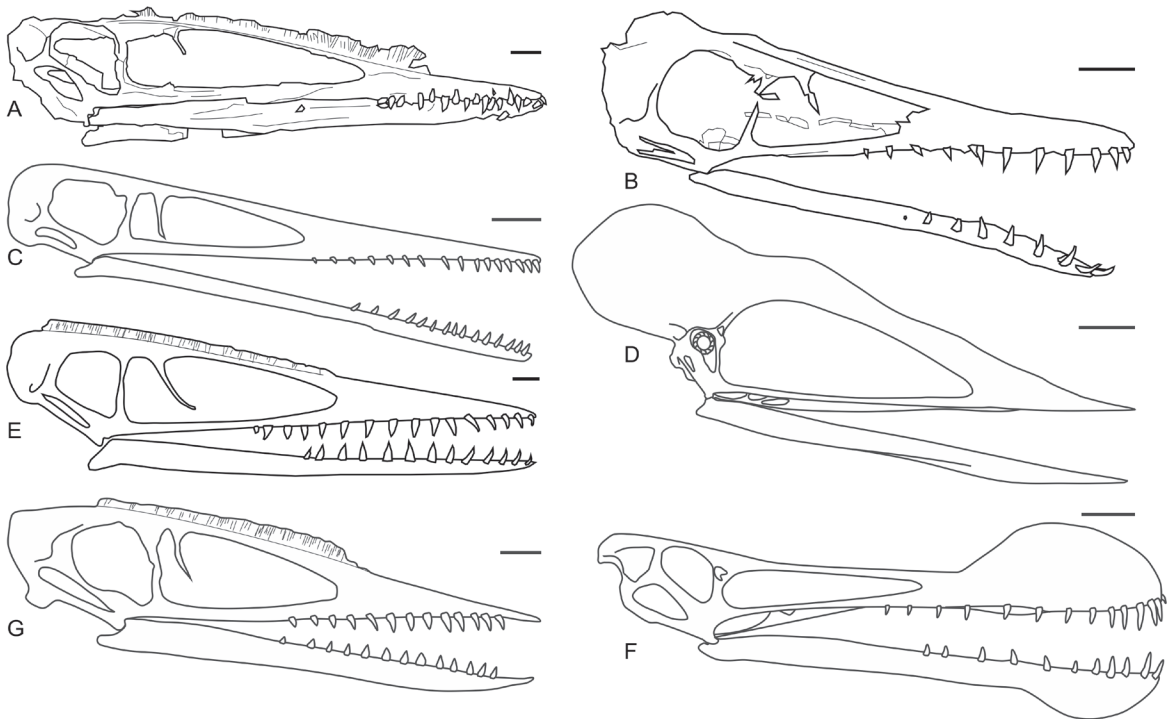


Fig. 3. Monofenestrans skulls. A, the wukongopterid *Darwinopterus robustodens*; B, likely pterodactyloid sister-taxon the 'Painten Pro-pterodactyloid'; C, ctenochasmatoid *Pterodactylus antiquus*; D, azhdarchoid *Tupuxuara leonardii*; E, early dsungaripteroid *Germanodactylus rhamphastinus*; F, ornithocheiroid *Ornithocheirus mesembrinus*; G, early dsungaripteroid *Germanodactylus cristatus*. Scale bars represent 10 mm, except for D and F, which represent 100 mm. A, after Lü *et al.*, 2011a; B, after Tischlinger and Frey, 2014; C and E, after Wellnhofer, 1970; D and F, after Witton, 2013.

Material and methods

Systematic declaration

Many aspects of pterosaur systematics remain contested. Key arguments include the relationships of major taxa, the diagnoses, names and content of many clades, methods to distinguish species and genera, and the number of valid species (e.g. Kellner, 2003, 2010; Unwin, 2003; Lü *et al.*, 2010, 2011b; Andres and Meyers, 2013; Naish *et al.*, 2013). The result is poor consensus on many components of pterosaur phylogeny, competing nomenclatural schemes with contradicting meanings for many established groups (e.g. compare Unwin, 2003; Kellner, 2003; Andres and Meyers, 2013) and multiple names for essentially identical clades (e.g. Novialoidea Kellner, 2003 *vs.* Breviquartossa Unwin, 2003). We are not the first authors to note these issues and the problems they create for modern pterosaur researchers (Naish *et al.*, 2013; Geist *et al.*, 2014): even basic discussion of pterosaur systematics now requires regular citation of the specific taxonomic scheme being followed (e.g. as in Andres and Meyers, 2013; Naish *et al.*, 2013) or frequent mentions of conflicting interpretations (e.g. discussions in Witton, 2013). Pending resolution of these issues, and to ease readability, we follow other authors working in fields with controversial classifications by clearly stating the taxonomic scheme followed by the present article. Unless otherwise mentioned, we follow the nomenclature and taxonomy of Lü *et al.* (2010, 2012). Wang *et al.* (2010) and Andres *et al.* (2014) offer alternative schemes to those used here.

Comparative anatomy

Measurements and observation of the anatomy of *Cuspicephalus scarfi* MJML K1918 were made from the holotype specimen and data in Martill and Etches (2013), while measurements of *Normannognathus wellnhoferi* MGCL 59'583 were obtained from a high-quality cast of the holotype and Buffetaut *et al.* (1998). Data on Tiaojishan wukongopterids and the 'Painten Pro-pterodactylid' were obtained from literature (Lü *et al.*, 2010, 2011a, b; Wang *et al.*, 2009, 2010; Tischlinger and Frey, 2014). Data on *Germanodactylus* specimens and other Jurassic pterodactyls were obtained from specimens, referred material (casts and original specimens), literature (e.g. Wellnhofer, 1970) and photographs.

Results

Cranial characteristics of the Wukongopteridae

Although undeniably pterodactyl-like, the skulls of wukongopterids lack synapomorphies of most major pterodactyl clades (Fig. 3; Kellner, 2003; Unwin, 2003; Andres and Ji, 2008). For example, they lack the rounded and reclined posterior skull regions of ctenochasmatooids, the depressed orbits and edentulous jaws of azhdarchooids, and the unusual rostral anatomy and dentition of ornithocheiroids (Kellner, 2003; Unwin, 2003; Andres and Ji, 2008). The wukongopterid skull most closely resembles those of early dsungaripteroids in overall skull proportions and shape, and particularly that of *Germanodactylus rhamphastinus* (Fig. 3E).

A number of similarities are present between wukongopterids and *Germanodactylus*. Both have striated cranial crests extending to the posterior region of the prenarial rostrum, generally with a gently sloping anterior margin. The crest of *G. rhamphastinus* differs here in terminating almost directly above the anterior margin of the nasoantorbital fenestra. This may reflect a genuine anatomical difference, but we are aware of the problems presented by poor preservation and historically erroneous preparation of pterosaur crests (Bennett, 2013a), as well as the current low number of *G. rhamphastinus* specimens (Bennett, 2006). The discovery of more specimens will clarify this morphology.

The anterior crests of wukongopterids and germanodactylids are relatively low compared to some pterosaurs. Each is lower than the height of the underlying prenarial rostrum, but the same crest portions of *Cycnorhamphus* and *Dsungaripterus* are as tall, if not taller, than the underlying rostrum (Young, 1973; Bennett, 2013b). The rostral indices (Martill and Naish, 2006) of sub-adult or older wukongopterids and *Germanodactylus* all plot within 3.04–5.07, with each group showing a wide range of values within this range (Martill and Etches, 2013). The preorbital skull length is rather short in *G. cristatus* (70% of jaw tip to squamosal length), but the value for *G. rhamphastinus* (78%) is very similar to those measured for wukongopterids (72–78%). Wukongopterid and *Germanodactylus* tooth spacing is relatively uniform, with a slight increase in spacing posteriorly. Wukongopterid and *Germanodactylus* inter-alveoli spacing is generally larger than corresponding alveolus length, although this is less apparent for *G. cristatus* than for *G. rhamphastinus*. Teeth occur under the anterior half of the nasoantorbital fenestra along a relatively straight

ventral skull margin in both wukongopterids and *Germanodactylus*.

In other respects wukongopterid crania differ markedly from skulls of both *Germanodactylus* species. Some distinctions concern characters which have previously suggested a close relationship between the Dsungaripteridae and *Germanodactylus* (Unwin, 2003), such as the expanded exoccipital processes of *Germanodactylus cristatus* (Unwin, 2003). This feature, unknown for any wukongopterid, is rarely discussed for *Germanodactylus* because it is poorly preserved in the holotype slab, BSP 1892 IV 1. However, a large exoccipital can clearly be seen in NMING:F15005, the counter-slab of the *G. cristatus* holotype (Fig. 4,

also see Hone, 2010). It seems that expanded exoccipital processes are common to all dsungaripteroids (Unwin, 2003), although their presence in *G. rhamphastinus* cannot be evaluated because appropriate skull regions are not clearly preserved in any specimens we are aware of.

The orbits of wukongopterids are piriform, as they are in *Germanodactylus* and several other pterodactyloid lineages (see Lü et al., 2006 for a discussion of this character among the Pterodactyloidea). The anterodorsal region of their orbits are convex, this portion of their orbits being partially occupied by lacrimal and nasal bones. Orbits with similarly convex anterodorsal margins evolved repeatedly in pterosaurs, being present in rham-



Fig. 4. Skull of NMING:F15005, *Germanodactylus cristatus*, holotype counter-slab, showing well-preserved expanded exoccipital process. This feature is vaguely discernible on the holotype, but much clearer on this slab. Scale bar represents 50 mm. Photograph courtesy of David Hone.

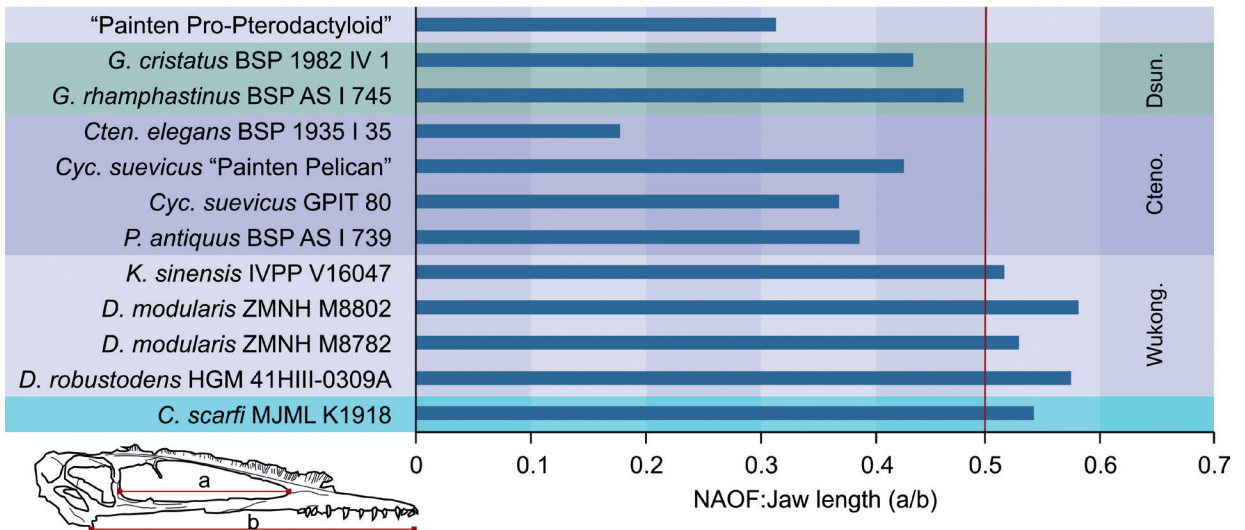


Fig. 5. Nasoantorbital fenestra length against jaw length in Late Jurassic monofenestratans. Cteno., Ctenochasmatoidea; Dsun., Dsungaripteridae; Wukong., Wukongopteridae.

phorhynchids, ctenochasmatooids, ornithocheirids and thalassodromids (e.g. Wellnhofer, 1970, 1975, 1987; Witton, 2013). They do not occur in either *Germanodactylus* species however. The nasal processes of many Jurassic monofenestrans are rather long (e.g. *Germanodactylus*, *Cycnorhamphus*, *Pterodactylus*; see Wellnhofer, 1970; Bennett, 2013b), but are relatively short in wukongopterids. Conversely, the anterior jugal process is suggested to be longer in some wukongopterids (e.g. *D. linglongtaensis*; *Wukongopterus*) than in *Germanodactylus* (e.g. Wang *et al.*, 2009, 2010; Martill and Etches, 2013). Long anterior jugal processes may not be common to all wukongopterids however, being reportedly truncated in at least the holotype of *D. modularis* (Lü *et al.*, 2010). We propose that this feature is variable within wukongopterids, or alternatively, that some interpretations of their jugal margins are erroneous. Cracks and marks interpreted as anterior margins of ‘short’ and ‘long’ jugal processes can be seen on many wukongopterid specimens, suggesting further investigation may be warranted to confirm their anterior jugal limits.

The nasoantorbital fenestra is longer with respect to jaw length in the Wukongopteridae than in other Jurassic monofenestrans (Fig. 5). Nasoantorbital openings are generally less than 40% of jaw length in ctenochasmatooids and the ‘Painten pro-pterodactylid’, 43–48% of the jaw length in *Germanodactylus*, but 51–58% of the jaw length in wukongopterids. Such a value places wukongopterid nasoantorbital openings amongst the longest of any monofenestratan, second only to istiodactylids and azhdarchoids (Kellner, 2003; Unwin, 2003; Witton, 2012). The dorsal margin of the wukongopterid preorbital rostrum, excluding the cranial crest, is concave, which contrasts with the straight dorsal margin of *Germanodactylus* rostra.

The expanded dentition of *Germanodactylus* is not mirrored in any wukongopterid. This is even the case in *Darwinopterus robustodens*, a species named for its relatively robust teeth (Lü *et al.*, 2011a). Wukongopterids have a relatively slender, sharp, and pointed dentition generally similar to that of pterodactyloids like *Pterodactylus* and *Haopterus* (Wellnhofer, 1970; Wang and Lü, 2001). Wukongopterid teeth are also mostly evenly sized, whereas those of *Germanodactylus* expand markedly from the jaw tip with the broadest teeth at the midpoint of the toothrow (Wellnhofer, 1970). The larger teeth of wukongopterids are, like virtually all toothed pterosaurs, found towards the front of the jaw. Wukongopterid toothrows extend to the end of the jaws as they do in *G. rhamphastinus* and most non-dsun-

garipteroid monofenestrans. *Germanodactylus cristatus*, however, has edentulous jaw tips.

The first premaxillary tooth pair arrangement in wukongopterids is distinctive. These alveoli are situated on the anteroventral margin of the jaw, a condition which sees the anterior teeth over-biting the lower jaw (Wang *et al.*, 2009, 2010). This is somewhat reminiscent of the anterior dentition of ornithocheirids (e.g. Wellnhofer, 1987) and some ctenochasmatooids (e.g. Howse and Milner, 1995), but the dentition and tooth arrangement of these pterosaurs is clearly demarked from wukongopterids and is very likely convergent. Wang *et al.* (2009, 2010) considered over-biting first premaxillary tooth pairs autapomorphic for *Wukongopterus lii*, but other wukongopterids demonstrate an identical condition (e.g. Lü *et al.*, 2010; Wang *et al.*, 2010). This feature is likely characteristic of the Wukongopteridae rather than a single wukongopterid genus.

Our assessment suggests that wukongopterid skulls can be distinguished from other Jurassic monofenestrans by not only lacking the well-documented cranial synapomorphies of pterodactylid clades, but also through a unique combination of characters:

1. Striated bony crest lower than the underlying pre-narial rostrum, with sloping anterior margin
2. Anterior crest terminates in the posterior region of the pre-narial rostrum, closer to the anterior border of the nasoantorbital fenestra than the jaw tip
3. Reclined, but not sub-horizontal, occipital regions
4. Piriform orbit
5. Convex anterodorsal orbital margin
6. Short nasal process
7. Unexpanded exoccipital processes
8. Concave dorsal skull surface
9. Straight ventral skull surface
10. Nasoantorbital fenestra over 50% of jaw length
11. Small, equally sized alveoli
12. First alveolus pair located on anterior face of jaw, with mandible over-bitten by first premaxillary tooth pair
13. Regular tooth spacing
14. Intervalveolar spacing generally greater than tooth length
15. Dentition extends under anterior half of the nasoantorbital region
16. Relatively slender, sharply pointed conical teeth

The particularly long nasoantorbital fenestra appears to be the most characteristic feature of wukongopterid skulls even though, as noted above, large nasoantorbital fenestrae are not unique to the Wukongopteridae within Monofenestrata. Among Jurassic pterosaurs

however, wukongopterid nasoantorbital openings are the longest of any group, and only distantly related, derived Cretaceous pterodactyloids demonstrate a similar condition. This feature therefore provides a useful synapomorphy for Wukongopteridae.

Cranial characteristics of the 'Painten pro-pterodactyloid'

Remarkably, the skull of the 'Painten pro-pterodactyloid' (Fig. 3B) bears several features considered characteristic of derived ctenochasmatooids, including a near-circular orbit, almost horizontal occipital region, and a rounded posterior skull (Kellner, 2003; Unwin, 2003; Tischlinger and Frey, 2014). The possession of these features in a taxon clearly demarked from the Pterodactyloidea by its postcrania is quite striking, and complicates our understanding of early pterodactyloid evolution. More pertinent to this study, they also allow for easy distinction of the crania of the 'Painten pro-pterodactyloid' from monofenestratan groups with mostly plesiomorphic skull anatomy, the wukongopterids and germanodactylids.

Within Ctenochasmatoidea, the skull of the 'Painten pro-pterodactyloid' is most similar to that of *Pterodactylus* (Fig. 3C). This occurs through its short nasoantorbital fenestra (Fig. 5), straight ventral skull margin, convex anterodorsal orbital margin, long nasal process, relatively short, pointed teeth and (as seen in juvenile *Pterodactylus*) concave dorsal skull margin. They are primarily differentiated by the very wide spacing and reduced number of teeth of the Painten specimen, as well as its procumbent anterior mandibular dentition (Tischlinger and Frey, 2014). The anterior premaxillary teeth, by contrast, are not procumbent. The regular alveolar spacing of the 'Painten pro-pterodactyloid' is a further distinguishing feature, contrasting with the posteriorly-increasing alveolus spacing of *Pterodactylus* and other Late Jurassic monofenestratans. The heavier construction of the premaxillary rostrum and proportionally shorter, taller skull are also characteristic for the Painten specimen, but these features vary with ontogeny (e.g. Bennett, 1995, 2006, 2013a) and their taxonomic significance is questionable. Likewise, the absence of a striated crest in the Painten specimen might be a useful identifying feature given the propensity of crests in ctenochasmatooids (e.g. Wellnhofer, 1970; Dong, 1980; Bennett, 2013a), but could also reflect ontogenetic or individual variation (Lü et al. 2011b). Studies into the ontogenetic status of the 'Painten pro-pterodactyloid' will hopefully provide some insights into the diagnostic utility of these characters (see Bennett, 1993).

Thus the 'Painten pro-pterodactyloid' can be distinguished from other Jurassic monofenestratans by a combination of 13 character states:

1. Near-horizontal occipital region
2. Rounded posterior skull
3. Sub-circular orbit
4. Convex anterodorsal orbital margin
5. Concave dorsal skull surface
6. Straight ventral skull surface
7. Robust premaxillary rostrum
8. Inter-alveolar spacing much greater than alveolus lengths
9. Consistent alveolus spacing
10. Dentition extends under anterior half of the nasoantorbital region
11. No anteriorly-facing premaxillary teeth at jaw tip
12. Relatively slender, sharply pointed conical teeth
13. Procumbent first and second pairs of mandibular teeth

Most of these characters are not diagnostic in isolation. However, some features of the dentition seem apomorphic. The combination of procumbent anterior mandibular teeth with vertical anterior premaxillary teeth is unique, as is the arrangement of the mandibular dentition, where only the anteriormost two tooth pairs are procumbent, while the remaining teeth are vertical.

Discussion

Cuspicephalus scarfi

Twelve of the 16 wukongopterid features listed above can be evaluated on the holotype of *Cuspicephalus scarfi*. Virtually all of them meet the conditions seen in wukongopterid skulls. These include a low striated crest terminating above the posterior region of the premaxillary rostrum; reclined posterior skull face; piriform orbit; a nasoantorbital fenestra exceeding 50% of the jaw length (at least 54%; Fig 5); small, relatively uniformly-sized alveoli, and a toothrow terminating under the anterior end of the nasoantorbital fenestra. The dorsal margin of the rostrum is not entirely preserved, but it can be reconstructed as gently concave with fair confidence (Martill and Etches, 2013), and the ventral skull margin is straight. The exoccipital processes are unexpanded: they look relatively large on MJML K1918, but this is largely an artefact of distortion around the occipital region, and they are not as prominent as those of *Germanodactylus* or dsungaripterids. Only one feature of MJML K1918 is inconsistent with a wukongopterid

identification: the anterior alveoli of MJML K1918 are separated by less than one alveolus-length, although the spacing of the posterior alveoli is more typical of the wukongopterid condition. The status of the anterodorsal region of the orbit, the presence of anteriorly-facing premaxillary alveoli and the length of the nasal process cannot be confidently determined for MJML K1918. Martill and Etches (2013) indicated that the nasal process may be represented by a small, faint trace in MJML K1918, but its length cannot be accurately assessed. Nevertheless, we conclude that 11 of the 12 observable or inferable characters of *Cuspicephalus* resemble a wukongopterid-like skull configuration and only one, alveolus spacing, shows a slightly different state.

In contrast, *Cuspicephalus* does not possess characters clearly indicative of close relationships to other monofenestratan taxa, including the ‘Painten Pro-pterodactyloid’ and Germanodactylidae. MJML K1918 can be evaluated for nine characters provided here for the ‘Painten Pro-pterodactyloid’ skull but is congruous with only three (concave dorsal rostrum; straight ventral skull; dentition under nasoantorbital fenestra). Similarly, *Cuspicephalus* differs from *Germanodactylus* in lacking a straight dorsal rostral margin and expanded exoccipital processes. Based on alveolus size, it also possessed more gracile teeth which increased in size anteriorly, not medially as in *Germanodactylus*. Martill and Etches (2013) argued that the shortness of the anterior jugal process suggested affinities with *Germanodactylus* but, as noted above, long anterior jugal processes may not be ubiquitous across the Wukongopteridae.

The *Cuspicephalus* skull meets nearly all character conditions of wukongopterid pterosaurs, but lacks many defining characteristics of other Jurassic pterosaurs, supporting the suggested close relationship with *Darwinopterus* (Martill and Etches, 2013) and indicating placement elsewhere among the Monofenestrata is unlikely. Furthermore, its possession of a feature unseen in Jurassic pterosaurs outside of wukongopterids – the especially elongate nasoantorbital fenestra – strongly suggests placement within Wukongopteridae. The differences it has with other wukongopterids – a proportionally long skull, higher tooth count and more condensed alveolus spacing at the jaw tip (Martill and Etches, 2013) – are not problematic for this identification: variable rostrum length and dental counts are typical within pterosaur clades. Indeed, greater rostrum length and tooth counts may be expected for a pterosaur of larger absolute size than its close relatives (see below). We thus consider a wukongopterid placement most likely for *Cuspicephalus*: the significance of this is discussed below.

Normannognathus wellnhoferi

In being represented by less material than *Cuspicephalus*, *Normannognathus* presents a much greater challenge for identification. *Normannognathus* can only be fully evaluated for nine of the features identified in our list of wukongopterid characteristics, with five positive comparisons. It can be compared with six characteristics of the ‘Painten Pro-pterodactyloid’, but none compare favourably.

Normannognathus bears a striated crest extending beyond the nasoantorbital region and terminating with an overturned leading edge. This crest is proportionally very tall – perhaps the tallest of any pterosaur when compared to the underlying rostrum height. We agree with Buffetaut *et al.* (1998) that the crest shape of *Normannognathus* is more reminiscent of *Dsungaripterus* than any other pterosaur, and it certainly differs from the condition in *Germanodactylus*, wukongopterids and the crestless ‘Painten Pro-pterodactyloid’ specimen. Only the ctenochasmatoid *Huanhepterus quingyangensis* offers a crest of similar height when compared to underlying rostral proportions (Dong, 1982). The proximity of the *Normannognathus* anterior crest margin to the nasoantorbital fenestra cannot be evaluated, nor is it clear from the dorsal and ventral rostral margins how the skull shape continued beyond the broken jaw tips. We see no reason to infer a short, *Germanodactylus*-like skull for *Normannognathus* any more than a much longer, lower skull akin to that of the ctenochasmatoids *Feilongus* (Wang *et al.*, 2005) or *Huanhepterus*. MGCL 59’583 possesses a low, concave dorsal rostral margin but also a convex ventral surface, together forming an upturned jaw tip. The upturned jaw of MGCL 59’583 is genuine, but the specimen is slightly distorted and obliquely preserved, so this feature is not as pronounced as it may first appear (Buffetaut *et al.*, 1998). Upturned jaws are known from *Pteranodon*, ctenochasmatoid, dsungaripterid and istiodactylid pterodactyloids (*e.g.* Young, 1973; Bennett, 1996, 2001; Wang *et al.*, 2005; Witton, 2012), but are not apparent in either wukongopterids or the ‘Painten Pro-pterodactyloid’. The rostrum of *Normannognathus* is rather more slender than that of the ‘Painten Pro-pterodactyloid’.

Some aspects of the MGCL 59’583 alveoli match the wukongopterid condition, being of approximately uniform size with the anterior pair situated on the anteroventral surface of the premaxilla. The spacing of the alveoli is generally tighter than those of wukongopterids however, surpassing even the spacing frequency of *Cuspicephalus*. A solitary tooth is preserved in the mandible

Table 1. Skull and wing bone lengths (mm) of complete wukongopterids from the Tiaojishan Formation, used in wingspan estimation of MJML K1918. McIV, metacarpal IV.

Taxon	Specimen	Skull	Hum	Ul	McIV	Wing finger phalanges				Wingspan
						I	II	III	IV	
<i>Darwinopterus linlongtaensis</i>	IVPP V16049	119.2	40.4	58	23.6	46.2	50.7	53.2	53.6	651.4
<i>Kunpengopterus sinensis</i>	IVPP V16047	106.9	36.2	59.2	23	54.2	58	59.2	48.8	677.2
<i>Darwinopterus robustodens</i>	HGM 41HIII-0309A	175	50	80	30	65	75	75	67	884
<i>Darwinopterus modularis</i>	YH-2000	140	44	64	26	48	55	59	53	698

of MGCL 59'583 but does not taper to a sharp point like the teeth of wukongopterids. Rather, it is relatively long and slender, more akin to the teeth of ctenochasmatis and certainly very different to the robust dentition seen in *Germanodactylus* and *Tendaguripterus*. This tooth projects anterolaterally from the alveolus in a ctenochasmatis manner, although additional discoveries are probably required to verify this as the *in vivo* condition and not post-mortem displacement. The alveolar spacing, positions and apparent dental orientations contrast markedly with the distinctive dentition of the 'Painten Pro-pterodactyloid'.

Further features of interest in MGCL 59'583 are the midline grooves on the occlusal surfaces of the upper jaw and mandibular symphysis. The symphyseal trough is wider and deeper than the groove of the upper jaw, which is relatively shallow and only clearly seen at the jaw tip. Among toothed pterosaurs, midline jaw grooves are well documented in ornithocheirids and lonchodectids (e.g. Unwin, 2001), but are poorly known in other

groups. For Jurassic monofenestrans at least, this reflects the frequent lateral crushing of pterosaur skulls. Nevertheless, the germanodactylid *Tendaguripterus* lacks a mandibular groove (Unwin and Heinrich, 1999), as do dsungaripterids (e.g. Young *et al.*, 1973). The Jurassic ctenochasmatis *Gnathosaurus macrurus* bears a symphyseal trough (Howse and Milner, 1995). The status of jaw grooves in wukongopterids and the 'Painten Pro-pterodactyloid' remains unknown.

Normannognathus possesses a mosaic of monofenestrans features with conflicting phylogenetic signals, preventing confident referral to any specific monofenestrans clade, including the Germanodactylidae. Buffetaut *et al.* (1998) referred *Normannognathus* to this group because of its crest morphology, similar tooth distribution to *G. rhamphastinus* and the possible close relationship of *Germanodactylus* to *Dsungaripterus*, which *Normannognathus* resembles by its upturned jaw and crest size. However, these features are now recognised as widely distributed across the Monofenestrans, and no

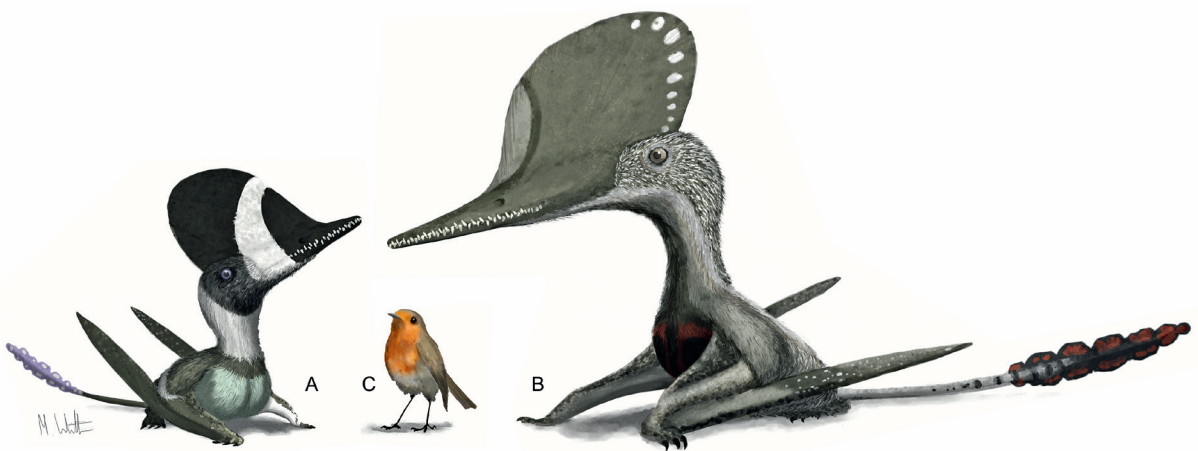


Fig. 6. Life reconstructions of wukongopterid pterosaurs showing size range across the group with extant avian for scale. A, *Darwinopterus robustodens*, the largest Tiaojishan Formation wukongopterid with a 884 mm wingspan; B, *Cuspicephalus scarfi*, with a projected wingspan of 1.2 m; C, European robin, *Erithacus rubecula* (wingspan c., 200 mm).

longer exclusively indicate a close relationship with *Germanodactylus*. Indeed, we do not find any features on MGCL 59'583 which unambiguously indicate germanodactylid affinities and, to the contrary, find much of its anatomy inconsistent with this placement.

What *Normannognathus* represents remains difficult to ascertain, however. It remains a valid genus due to its unique assemblage of characters, but placing it within an existing pterosaur group is problematic. It differs from wukongopterids as much as it resembles them, bears no features of the 'Painten Pro-pterodactylid' or clear synapomorphies of any major pterodactylid clade. The constituent components of *Normannognathus* anatomy – steeply terminating, relatively tall striated crests; slightly upturned, shallow rostra with slender, anteriorly directed teeth and symphyseal troughs – occur in several ctenochasmatoidea including *Huanhepterus* (Dong, 1982); *Cycnorhamphus suevicus* (Bennett, 2013a), *Aurorazhdarcho micronyx* (Bennett, 2013b), *Feilongus youngi* (Wang *et al.*, 2005) and *Gnathosaurus macrurus* (Howse and Milner, 1995). *Normannognathus* may therefore have affinities with the Ctenochasmatoidea, but more complete remains are required to make a confident assessment of its systematic position. At present it may be best considered Monofenestrata *incertae sedis*.

Significance of Cuspicephalus scarfi interpreted as a wukongopterid

The suggestion that wukongopterid skulls can be distinguished from those of other early monofenestratans without associated postcranial material bodes well for further research into this group. However, their low number of cranial synapomorphies proves limiting when considering particularly fragmentary monofenestratan crania, as evidenced by difficulties resolving the affinities of *Normannognathus*. We hope the comparisons made here will be useful in future assessments of Jurassic monofenestratan material, such as the fragmentary crania from the Purbeck Limestone, Kimmeridge Clay and Morrison formations.

The occurrence of a wukongopterid in Kimmeridgian deposits of the southern UK expands the stratigraphic and geographic range of this group significantly. Presently, all wukongopterids are known from the upper Oxfordian or Lower Callovian Tiaojishan Formation of northeast China. *Cuspicephalus scarfi* extends the wukongopterid stratigraphic range into the Kimmeridgian, being the youngest wukongopterid by at least 5 million years, and expands the geographic range of the group to

Europe. This makes wukongopterid palaeobiogeography comparable to that of other Middle and Late Jurassic pterosaur lineages, most of which are distributed across multiple continents or even cosmopolitan in their distribution (see Barrett *et al.*, 2008 and Witton, 2013 for recent overviews of pterosaur palaeobiogeography).

Cuspicephalus further offers new insights into wukongopterid disparity. It is the first wukongopterid to obviously differ from the *Darwinopterus*-like taxa of the Tiaojishan Formation, animals which are so morphologically similar that Lü *et al.* (2011b) suggest they represent a single taxon. The skull of *Cuspicephalus* is proportionally longer and lower than any Tiaojishan form, its teeth more numerous and tightly packed at the jaw tip. This presumably reflects ecological differentiation from the Chinese wukongopterids, perhaps allowing for greater reach during foraging and manipulating relatively small or slippery prey. Moreover, *Cuspicephalus* is also considerably larger than its relatives. At 326 mm long, the skull of *Cuspicephalus* is much longer than those of its fellow wukongopterids and also one of the largest Jurassic pterosaur skulls known. Only the Morrison Formation scaphognathine *Harpactognathus gentryii* is estimated to have a skull of comparable length (280–300 mm; Carpenter *et al.*, 2003). By contrast, the largest Tiaojishan wukongopterid skull (HGM 41HIII-0309A; the osteologically mature holotype of *Darwinopterus robustodens*) is 175 mm long (Fig. 3A), 53% of the skull length of *Cuspicephalus*. A regression of skull length against wingspan of complete Tiaojishan wukongopterids (Table 1) shows that their skull length scales with negative allometry to wingspan ($n = 4$, $r^2 = 0.7997$), predicting a wing spread of 1.2 m for *Cuspicephalus*. This is 35% greater than the 884 mm estimated for *D. robustodens* (Fig. 6) and, while smaller than the predicted 1.8–2.5 m wingspans of the largest Jurassic pterosaurs, such as *Harpactognathus* and *Rhamphorhynchus*, still brings wukongopterids into a new size class of pterosaurs.

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References

- Andres B, Ji Q. 2008. A new pterosaur from the Liaoning Province of China, the phylogeny of the Pterodactyloidea, and the convergence in their cervical vertebrae. *Palaeontology* 51: 453-469.
- Andres B, Myers TS. 2013. Lone Star Pterosaurs. *Earth and Environmental Science Transactions of the Royal Society of Edinburgh* 103: 383-398.
- Andres B, Howard L, Steel L. 2011a. Owen's pterosaurs, old fossils shedding light on new clades. P. 4 in: Forrest R, ed., SVP-CA, 59th Annual Symposium of Vertebrate Palaeontology and Comparative Anatomy, Abstracts of Presentations.
- Andres B, Howard L, Steel L. 2011b. Pterosaurs, modules, and the origin of the Pterodactyloidea. *Journal of Vertebrate Paleontology* 31(A): 62.
- Andres B, Clark J, Xu X. 2014. The earliest pterodactyloid and the origin of the group. *Current Biology* 24: 1011-1016.
- Barrett PM, Butler RJ, Edwards NP, Milner AR. 2008. Pterosaur distribution in space and time: an atlas. *Zitteliana* B28: 61-107.
- Bennett SC. 1993. The ontogeny of *Pteranodon* and other pterosaurs. *Paleobiology* 19: 92-106.
- Bennett SC. 1995. A statistical study of *Rhamphorhynchus* from the Solnhofen Limestone of Germany: year-classes of a single large species. *Journal of Paleontology* 16: 569-580.
- Bennett SC. 1996. Year-classes of pterosaurs from the Solnhofen Limestone of Germany: taxonomic and systematic implications. *Journal of Vertebrate Paleontology* 16: 432-444.
- Bennett SC. 2001. The osteology and functional morphology of the Late Cretaceous pterosaur *Pteranodon*. *Palaeontographica Abteilung A* 260: 1-153.
- Bennett SC. 2006. Juvenile specimens of the pterosaur *Germanodactylus cristatus*, with a review of the genus. *Journal of Vertebrate Paleontology* 26: 872-878.
- Bennett SC. 2013a. New information on body size and cranial display structures of *Pterodactylus antiquus*, with a revision of the genus. *Paläontologische Zeitschrift* 87: 269-289.
- Bennett SC. 2013b. The morphology and taxonomy of the pterosaur *Cynnorhamphus*. *Neues Jahrbuch für Geologie und Paläontologie-Abhandlungen* 267: 23-41.
- Buffetaut E, Lepage J, Lepage G. 1998. A new pterodactyloid from the Kimmeridgian of the Cape de la Héve (Normandy, France). *Geological Magazine* 135: 719-722.
- Carpenter K, Unwin DM, Cloward K, Miles C, Miles C. 2003. A new scaphognathine from the upper Jurassic Morrison Formation of Wyoming, USA. Pp. 45-54 in: Buffetaut E, Mazin JM, eds, *Evolution and Palaeobiology of Pterosaurs*, Geological Society Special Publication 217.
- Dong Z. 1982. A new pterosaur (*Huanhepterus quingyangensis* gen. et sp. nov.) from Ordos, China. *Vertebrata Palasiatica* 20: 115-121.
- Geist NR, Hillenius WJ, Frey E, Jones TD, Elgin RA. 2014. Breathing in a box: Constraints on lung ventilation in giant pterosaurs. *The Anatomical Record* 297: 2233-2253.
- Hone DW. 2010. A short note on modifications to nineteenth century pterosaur specimens held in the National Museum of Ireland–Natural History, Dublin. *Geological Curator* 9: 261-265.
- Hone DW. 2012. Pterosaur research: recent advances and a future revolution. *Acta Geologica Sinica-English Edition* 86: 1366-1376.
- Howse SCB, Milner AR. 1995. The pterodactyloids from the Purbeck Limestone Formation of Dorset. *Bulletin of the Natural History Museum, London* 51: 73-88.
- Kellner AWA. 2003. Pterosaur phylogeny and comments on the evolutionary history of the group. Pp. 105-137 in: Buffetaut E, Mazin JM, eds, *Evolution and Palaeobiology of Pterosaurs*, Geological Society Special Publication 217.
- Kellner AWA. 2010. Comments on the Pteranodontidae (Pterosauria, Pterodactyloidea) with the description of two new species. *Anais da Academia Brasileira de Ciências* 82: 1063-1084.
- Lü JC. 2009. A new non-pterodactyloid pterosaur from Qinglong County, Hebei Province of China. *Acta Geologica Sinica* 83: 189-199.
- Lü JC, Fucha X. 2010. A new pterosaur (Pterosauria) from the Middle Jurassic Tiaojishan Formation of western Liaoning, China. *Global Geology* 13: 113-118.
- Lü JC, Jin X, Unwin DM, Zhao L, Azuma Y, Ji Q. 2006. A new species of *Huaxipterus* (Pterosauria: Pterodactyloidea) from the Lower Cretaceous of Western Liaoning, China with comments on the systematics of tapejarid pterosaurs. *Acta Geologica Sinica* 80: 315-326.
- Lü JC, Unwin DM, Jin X, Liu Y, Ji Q. 2010. Evidence for modular evolution in a long-tailed pterosaur with a pterodactyloid skull. *Proceedings of the Royal Society B* 277: 383-389.
- Lü JC, Xu L, Chang H, Zhang X. 2011a. A new darwinopterid pterosaur from the Middle Jurassic of Western Liaoning, Northeastern China and its ecological implications. *Acta Geologica Sinica* 85: 507-514.
- Lü, JC, Unwin DM, Deeming DC, Jin X, Liu Y, Ji, Q. 2011b. An egg-adult association, gender, and reproduction in pterosaurs. *Science* 331: 321-324.
- Lü JC, Unwin DM, Zhou B, Chunling G, Shen C. 2012. A new rhamphorhynchid (Pterosauria: Rhamphorhynchidae) from the Middle/Upper Jurassic of Qinglong, Hebei Province, China. *Zootaxa* 3158: 1-19.
- Maisch MW, Matzke AT, Sun G. 2004. A new dsungaripteroid pterosaur from the Lower Cretaceous of the southern Junggar Basin, north-west China. *Cretaceous Research* 25: 625-634.
- Martill DM, Naish D. 2006. Cranial crest development in the azhdarchoid pterosaur *Tupuxuara*, with a review of the genus and tapejarid monophyly. *Palaeontology* 49: 925-941.
- Martill DM, Etches S. 2013. A monofenestratan pterosaur from the Kimmeridge Clay Formation (Upper Jurassic, Kimmeridgian) of Dorset, England. *Acta Palaeontologica Polonica* 58: 285-294.
- Naish D, Simpson M, Dyke G. 2013. A new small-bodied azhdarchoid pterosaur from the Lower Cretaceous of England and its implications for pterosaur anatomy, diversity and phylogeny. *PloS one* 8: e58451.
- Plieninger F. 1901. Beiträge zur Kenntnis der Flugsaurier. *Palaeontographica* 48: 65-90.
- Steel L. 2010. The pterosaur collection at the Natural History Museum, London, UK: history, overview, recent curatorial developments and exciting new finds. *Acta Geoscientica Sinica* 31: 59-61.
- Sullivan C, Wang Y, Hone DWE, Wang YQ, Xu X, Zhang F. 2014. The vertebrates of the Jurassic Daohugou Biota of Northeastern China. *Journal of Vertebrate Paleontology* 34: 243-280.
- Tischlinger H, Frey E. 2014. Ein neuer Pterosaurier mit Mosaikmerkmalen basaler und pterodactyloider Pterosaurier aus dem

- Ober-Kimmeridgium von Painen (Oberpfalz, Deutschland) [A new pterosaur with mosaic characters of basal and pterodactyloid Pterosauria from the Upper Kimmeridgian of Painen (Upper Palatinate, Germany)]. *Archaeopteryx* 31: 1-13.
- Unwin DM. 1996. The fossil record of Middle Jurassic Pterosaurs. Pp. 291-304 in: Morales M, ed., *The Continental Jurassic, Museum of Northern Arizona Bulletin* 60.
- Unwin DM. 2001. An overview of the pterosaur assemblage from the Cambridge Greensand (Cretaceous) of Eastern England. *Mitteilungen Museum für Naturkunde Berlin, Geowissenschaftlichen* 4: 189-221.
- Unwin DM. 2003. On the phylogeny and evolutionary history of pterosaurs. Pp. 139-190 in: Buffetaut E, Mazin JM, eds, *Evolution and Palaeobiology of Pterosaurs*, Geological Society Special Publication, 217.
- Unwin DM. 2005. *The Pterosaurs from Deep Time*. Pi Press, New York.
- Unwin DM, Heinrich WD. 1999. On a pterosaur jaw from the Upper Jurassic of Tendaguru (Tanzania). *Mitteilungen aus dem Museum Für Naturkunde in Berlin Geowissenschaftliche Reihe* 2: 121-134.
- Vidovic SU, Martill DM. 2014. *Pterodactylus scolopaciceps* Meyer, 1860 (Pterosauria, Pterodactyloidea) from the Upper Jurassic of Bavaria, Germany: the problem of cryptic pterosaur taxa in early ontogeny. *PLoS one* 9: e110646.
- Wang X, Lü JC. 2001. Discovery of a pterodactylid pterosaur from the Yixian Formation of western Liaoning, China. *Chinese Science Bulletin* 46: 1112-1117.
- Wang X, Kellner AWA, Zhou Z, Campos DA. 2005. Pterosaur diversity and faunal turnover in Cretaceous terrestrial ecosystems in China. *Nature* 437: 875-879.
- Wang X, Kellner AWA, Jiang S, Meng X. 2009. An unusual long-tailed pterosaur with elongated neck from western Liaoning of China. *Anais da Academia Brasileira de Ciências* 81: 793-812.
- Wang X, Kellner AWA, Jiang S, Cheng X, Meng X, Rodrigues T. 2010. New long-tailed pterosaurs (Wukongopteridae) from western Liaoning, China. *Anais da Academia Brasileira de Ciências* 82: 1045-1062.
- Wellnhofer P. 1970. Die Pterodactyloidea (Pterosauria) der Oberjura-Plattenkalke Süddeutschlands. *Bayerische Akademie der Wissenschaften, Mathematisch-Wissenschaftlichen Klasse, Abhandlungen* 141: 1-133.
- Wellnhofer P. 1975. Die Rhamphorhynchoidea (Pterosauria) der Oberjura-Plattenkalke Süddeutschlands. *Palaeontographica A* 148: 1-33, 132-186, 149: 1-30.
- Wellnhofer P. 1987. New crested pterosaurs from the Lower Cretaceous of Brazil. *Mitteilungen der Bayerischen Staatssammlung für Paläontologie und Historische Geologie* 27: 175-186.
- Witton MP. 2012. New insights into the skull of *Isiodactylus latidens* (Ornithocheiroidea, Pterodactyloidea). *PLoS ONE* 7: e33170.
- Witton MP. 2013. *Pterosaurs: Natural History, Evolution, Anatomy*. Princeton University Press.
- Young CC. 1973. Pterosaurian fauna from Wuerho, Sinkiang. Pp. 18-34 in: *Reports of paleontological expedition to Sinkiang II*, Kexue Chubanshe, Nanjing, China.

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