

https://doi.org/10.1038/s42003-024-06132-6

# New insights into pterosaur cranial anatomy: X-ray imaging reveals palatal structure and evolutionary trends

Check for updates

He Chen <sup>1,2,3</sup>, Shunxing Jiang<sup>2</sup>, Alexander W. A. Kellner <sup>1</sup> & Xiaolin Wang <sup>2,3</sup> ⊠

Among the least studied portion of the pterosaur skeleton is the palate, which tends to be poorly preserved and commonly only visible from one side (the ventral portion). Even in well-preserved specimens, the bones tend to be fused, with the limits of individual palatal elements obscured. To shed new light on this region, we employed advanced X-ray imaging techniques on the non-pterodactyloid *Kunpengopterus* (Wukongopteridae), and the pterodactyloids *Dsungaripterus* (Dsungaripteridae), *Hongshanopterus* (Istiodactylidae), and *Hamipterus* (Hamipteridae). Our analyses revealed the presence of sutures between palatal bones in *Dsungaripterus* and *Kunpengopterus*, which resulted in different interpretations of the relation between palatine, ectopterygoid, and pterygoid, leading to a new identification of the palatal openings. Furthermore, our study shows six main observations such as the variation of the angle between the palatine rami and the variation in the relative sizes of the palatal openings. We also point out that the presence of a maxillopalatine fenestra (previously identified as postpalatine fenestra), is unique within Diapsida. Although much more work needs to be done, we showed that advanced X-ray imaging techniques open a window for understanding pterosaur cranial anatomy and provide a new perspective for investigating the evolutionary history of these flying reptiles.

Pterosaurs, the first vertebrates to achieve powered flight, have been a subject of fascination for scientists for decades. Identifying individual bones in the skull of extinct clades can be challenging given the lack of living descendants for comparison. This challenge is particularly challenging for pterosaurs, a group of extinct flying reptiles that lived during the Mesozoic Era. However, an accurate interpretation of their anatomy is the fundamental prerequisite for a wide range of studies, including phylogeny, ecology, biology, and functional morphology<sup>1-9</sup>.

As generally known the normal condition of preservation regarding pterosaurs is that of crushed or flattened specimens and three-dimensionally preserved material is exceedingly rare (e.g.,  $^{10,11}$ ). Additionally, cranial bones, including the palate, often exhibit fusion. Consequently, gaining an understanding of detailed skull architecture, especially the conformation of the palate in pterosaurs, has always been difficult. Furthermore, the identification of the limits of several elements generally still lacks consensus  $^{12-14}$ .

Questions regarding the construction of the pterosaur palate have been the subject of debate since the 19th century<sup>15–22</sup>. While several new specimens have come to light over the years<sup>10,23–37</sup>, the lack of clarity regarding the boundaries between different palatal bones has resulted in an ongoing debate<sup>38–45</sup>. Recently, a specimen of the Jurassic pterosaur *Dorygnathus banthensis*, featuring several unfused cranial bones, presented a very different configuration of the palatal portion of the maxilla<sup>13</sup>. This discovery led to a reinterpretation of the palate<sup>13</sup> with broad implications (e.g., refs. 14,38,41). However, controversy persists, particularly concerning the extent and shape of the palatines and whether a single, generalized palatal pattern configuration can be applied to Pterodactyloidea<sup>23,38,40–43</sup>.

Here we review the palate structure in various pterosaur taxa (Table 1), including both pterodactyloids and non-pterodactyloids. We introduce new insights based on CT-scans and CL-scans of *Kunpengopterus* (Wukongopteridae), *Hongshanopterus* (Istiodactylidae), *Hamipterus* (Hamipteridae), and *Dsungaripterus* (Dsungaripteridae). Our findings shed light on

¹School of Ecology, Sun Yat-sen University, Shenzhen 510006, China. ²Key Laboratory of Vertebrate Evolution and Human Origins, Institute of Vertebrate Paleontology and Paleoanthropology, Chinese Academy of Sciences, Beijing 100044, China. ³University of Chinese Academy of Sciences, Beijing 100049, China. ⁴Laboratory of Systematics and Taphonomy of Fossil Vertebrates, Department of Geology and Paleontology, Museu Nacional/UFRJ, Rio de Janeiro 20940-040, Brazil. ⊠e-mail: wangxiaolin@ivpp.ac.cn

Table 1 | Specimens with palatal information

Specimens	Taxon	Institutional abbreviations
SMNS 18969 (50,184 + 50,914 + 51,827)	Dorygnathus banthensis	Staatliches Museum fur Naturkunde, Stuttgart, Germany
CM 11,424	Campylognathoides liasicus	Carnegie Museum Pittsburgh, USA
GSM 3166	Parapsicephalus purdoni	British Geological Survey, Nottingham, UK
IGO-V 208	Cacibupteryx caribensis	Museo Mario Sánchez Roig, Instituto de Geología y Paleontología, La Habana, Cuba
NHM R 2786 (Nr.55)	Rhamphorhynchus muensteri	Natural History Museum, London, UK
CM 11434	Rhamphorhynchus muensteri	Carnegie Museum Pittsburgh, USA
GPIB 1304 (Nr. 109)	Scaphognathus crassirostris	Geologische Institut fur Palaontologie der Universitat Bonn, Germany
NMS G.2021.6. 2	Dearc sgiathanach	National Museums Scotland, Edinburgh, UK
<sup>a</sup> IVPP V 23674	Kunpengopterus sinensis	Institute of Vertebrate Paleontology and Palaeoantropology, Beijing, China
BSP 1936 I 50	Aurorazhdarcho micronyx	Staatliche Naturwissenschaftliche Sammlungen Bayerns/Bayerische Staatssammlung fur Paläontologie und Geologie, Munich, Germany
<sup>a</sup> IVPP V 14852	Hongshanopterus lacustris	Institute of Vertebrate Paleontology and Palaeoantropology, Beijing, China
IVPP V 18943.1, <sup>a</sup> IVPP V 18943.3	Hamipterus tianshanensis	Institute of Vertebrate Paleontology and Palaeoantropology, Beijing, China
RGM 401 880	Anhanguera spielbergi	National Natuurhistorisch Museum/Naturalis, Leiden, The Netherlands
AMNH 25555	Anhanguera santanae	American Museum of Natural History, New York, USA
AMNH 22555	Anhanguera santanae	American Museum of Natural History, New York, USA
SNSB-BSPG 1982   90	Anhanguera santanae	Staatliche Naturwissenschaftliche Sammlungen Bayerns/Bayerische Staatssammlung fur Paläontologie und Geologie, Munich, Germany
SNSB-BSPG 1982   89	Anhanguera araripensis	Staatliche Naturwissenschaftliche Sammlungen Bayerns/Bayerische Staatssammlung fur Paläontologie und Geologie, Munich, Germany
SAO 16494	Anhanguera araripensis	Sammlung Oberli, St. Gallen, Switzerland
MN 4805-V	Anhanguera blittersdorffi	Museu Nacional/Universidade Federal do Rio de Janeiro, Rio de Janeiro, Brazil
BSP 1987 I 46	Tropeognathus mesembrinus	Staatliche Naturwissenschaftliche Sammlungen Bayerns/Bayerische Staatssammlung fur Paläontologie und Geologie, Munich, Germany
PTH 1951.84	Gnathosaurus subulatus	Philosophische-Theologische Hochschule, Eichstätt, Germany
PMOL-AP00031	Liaodactylus primus	Paleontological Museum of Liaoning, Shenyang Normal University, Shenyang, China
<sup>a</sup> IVPP V 4063, 26257, 26258	Dsungaripterus weii	Institute of Vertebrate Paleontology and Palaeoantropology, Beijing, China
MN 4726-V	Caupedactylus ybaka	Museu Nacional/Universidade Federal do Rio de Janeiro, Rio de Janeiro, Brazil
IMCF 1052	Tupuxuara leonardii	Iwaki Coal and Fossil Museum, Iwaki, Japan
MNHN 1908-24 <sup>31</sup>	Pteranodon	Museum National d'Histoire Naturelle, Paleontologie, Paris, France
No.2594 <sup>64</sup>	Pteranodon ingens	Peabody Museum of Yale University, USA
FMNH 25026	Nyctosaurus gracilis	Field Museum of Natural History, Chicago, USA

<sup>&</sup>lt;sup>a</sup>The specimens have been CT-scanned or CL-scanned in this study.

the boundaries of several palatal bones and propose a new configuration for the palate of numerous pterodactyloid clades. Additionally, we reinterpret distinct configurations for the palate of several non-pterodactyloid clades, offering insights into potential evolutionary trends of this region in pterosaurs.

#### Materials and methods

The objective of this study was to examine the morphology and internal structure of palatal bones using X-ray micro-computed tomography and laminography. The specimens were scanned using the following different types of X-ray instruments, depending on their different preservation and size.

X-ray micro-computed tomography (CT): The specimen of Dsungaripterus (IVPP V 4063) was scanned using a GE v|tome|x m dual tube 300/180 kV system in the Key Laboratory of Vertebrate Evolution and Human Origins, Institute of Vertebrate Paleontology and Paleoanthropology (IVPP), Chinese Academy of Sciences (CAS). The specimen was scanned with a beam energy of 240 kV and a flux of 200  $\mu A$  at a resolution of 35.88  $\mu m$  per pixel using a 360° rotation with a step size of 0.18°. Two specimens of Dsungaripterus (IVPP V 26257, IVPP V 26258) and one specimen of Hamipterus (IVPP V 18943.3) were scanned using the 225 kV micro-computerized tomography in IVPP (developed by the Institute of High Energy Physics, CAS). The specimens of Dsungaripterus and

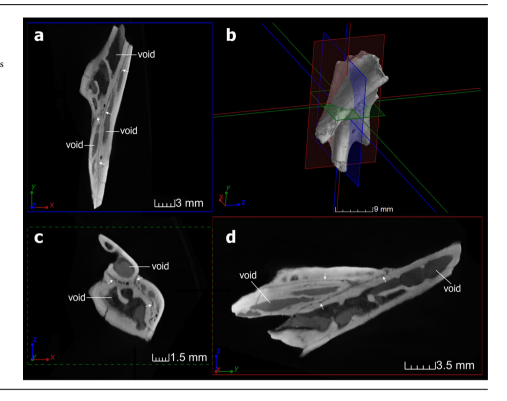
*Hamipterus* were scanned with a beam energy of 130 kV and a flux of 150 μA at a resolution of 34.50 μm per pixel, and 120 kV and a flux of 120 μA at a resolution of 25.09 μm per pixel respectively, using a 360° rotation with a step size of 0.5°. A total of 720 projections were reconstructed in a 2048  $\times$  2048 matrix of 1536 slices using a two-dimensional reconstruction software (IVPP-IHEP) developed by the Institute of High Energy Physics, CAS. Figure 1 shows examples of the scan results of the IVPP V 26257, which are primary data on which the palatal bones were reconstructed.

X-ray micro-computed laminography (CL): The flattened specimens of Kunpengopterus and Hongshanopterus were scanned using the CL scanner in the lab at the IVPP (developed by the Institute of High Energy Physics, CAS for flat specimens). The specimens were scanned with a beam energy of 80 kV and a flux of 60  $\mu A$  at a resolution of 9.02  $\mu m$  per pixel, and 70 kV and a flux of 70  $\mu A$  at a resolution of 33.7  $\mu m$  per pixel, respectively, using a 360° rotation with a step size of 0.5°. A total of 360 image slices with a size of 2048 by 2048 were reconstructed using a modified Feldkamp algorithm developed by the Institute of High Energy Physics, CAS.

The imagery data obtained from CT and CL were using VG Studio Max 3.0 (Volume Graphics, Heidelberg, Germany) to segment, render, and reconstruct the 3D bones (Supplementary Data 1), and make the video 1.

Regarding pterosaur phylogenies, there are several proposals/hypothesis published, with two main phylogenetic schemes (Kellner<sup>2</sup> and Unwin<sup>3</sup>), that have been modified with the addition of new taxa and the employment

Fig. 1 | CT slices of *Dsungaripterus weii* IVPP V 26257. a, c, d CT slices, showing the sutures pointed by arrows and internal structure. b Shows the place where those CT slices are. This specimen's CT-rendered result is shown in Fig. 2c.



of different methodologies (e.g., refs. 4,9,46). Here we have followed mainly Kellner et al.<sup>47</sup>.

#### Reporting summary

Further information on research design is available in the Nature Portfolio Reporting Summary linked to this article.

#### Results and discussion

The utilization of X-ray techniques (including CT-scan and CL-scan) has enabled us to investigate the internal structure of the palate of several pterosaurs, thus revealing the boundaries of several bones despite their strongly fused external surface. The CT-scans of the thick palatal elements in all three *Dsungaripterus* specimens analyzed (e.g., Fig. 1) show irregular edges of the sutures between the bones, resembling the crenelate boundary of postage stamps (e.g., Fig. 1), which differentiate them from cracks or fractures. In the case of the most complete specimen of *Dsungaripterus* (IVPP V 4063), most sutures are clearly identifiable, allowing for the separation of the bones (Fig. 2). However, the CT-scans of the specimen of *Hamipterus* (IVPP V 18943.3) (Fig. 3a, b) studied here did not reveal any sutures, suggesting that all palatal elements are fused.

In the case of the compressed specimens of *Hongshanopterus* and *Kunpengopterus*, we use CL-scan to identify any potential preserved suture. Regarding *Hongshanopterus* (Fig. 3d, e), due to the preservation that shows several fractures, no sutures could be confidently identified. In the case of *Kunpengopterus* (Fig. 3f, g), there are fewer fractures, and the boundary between the ectopterygoid and pterygoid could be seen (Fig. 3f). The remaining palatal elements in this specimen appeared to be fused. Despite the fact that no clear sutures were found in the specimens from *Hamipterus* and *Hongshanopterus* studied here (Figs. 3a, b, 4a, b), and only one suture was observed in *Kunpengopterus* (Fig. 3f, g), CT-scan and CL-scan proved to be very informative since they made it possible to observe the dorsal side of the palate.

# Palatal reconstruction of *Dsungaripterus weii* (Dsungaripteridae)

The CT-scans of IVPP V 4063 provide a relatively clear picture of the anatomy structure of the maxilla, palatine, pterygoid, and ectopterygoid,

which differs from previous work on Dsungaripterus<sup>43</sup>. Except for the contact with the vomer, the boundary of the posteroventral side of the maxilla is well established (Fig. 2a, b, e). The jugal process of the maxilla contacts the jugal (Fig. 2f) ventrally, forming the lateral edge of the palate, and extends posteriorly to the ectopterygoid (Video 1 and Fig. 2b). Compared to the maxilla of *Dorygnathus banthensis*<sup>13</sup> (Fig. 5d), the maxilla of Dsungaripterus presents a process on the ventromedial side directed posteriorly that is here referred to as the palatine process of the maxilla (Video 1 and Fig. 2e). This new process contacts the anterior ramus of the palatine dorsomedially (Video 1 and Fig. 2b, e). With this new interpretation, the most anterior fenestra positioned lateral to the choanae is bordered by the maxilla anteriorly and the palatine posteriorly, and therefore renamed here as the maxillopalatine fenestra. This opening has been called by most authors as the postpalatine fenestra (e.g., refs. 12,31,38) or suborbital fenestra<sup>13</sup>. CT-scans did not reveal any suture at the grooves preserved on the ventral side of the maxilla that has been previously interpreted as the boundaries of the maxilla and the palatine in several pterodactyloid taxa (e.g., refs. 28,38,43,48).

The 3D reconstruction of the palatine in Dsungaripterus shows two dorsoventrally flat palatine rami that form a distinctive y-shape (Video 1 and Fig. 2b-d, g). According to our new interpretation, the palatine contacts all other palatal bones except for the ventral portion of the premaxilla (no information is presently available for the vomer) and is involved in almost all palatal openings, except for the interpterygoid fenestra. The anterior ramus of the palatine ventromedially contacts the palatine process of the maxilla, and both separate the maxillopalatine fenestra medially from the choana. The lateral ramus of the palatine contacts the lateral edge of the palate, separating the maxillopalatine fenestra from the postpalatine fenestra referred to by others as the infraobital vacuity (e.g., refs. 18,36), the pterygoid-ectopterygoid fenestra<sup>13,14</sup>, or the secondary subtemporal fenestra<sup>37,40,42</sup>. The lateral ramus of the palatine is elongated and overlays the anterior ramus on the dorsal side at the posterior region (Video 1 and Fig. 2b-d, g). Medially, this lateral ramus becomes thinner, and contacts the dorsal surface of the pterygoid. At the posterior end, it contacts the opposing palatine, and both form the posterior margin of the choanae. There is a small posterior process that contacts the pterygoid dorsally and the ectopterygoid laterally and contacts the subtemporal fenestra (Video 1 and Fig. 2b-d, g). A

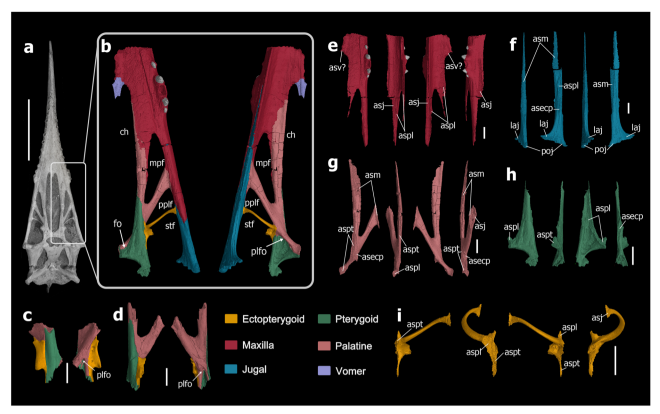


Fig. 2 | The three specimens of *Dsungaripterus weii* and their CT-rendered results. a The skull IVPP V 4063 in ventral view. b CT-rendered the left part of the palate of IVPP V 4063 in ventral and dorsal views. c CT-rendered model of IVPP V 26258, showing the contact region of the right side of the palatine, pterygoid, and ectopterygoid in ventral and dorsal views. d CT-rendered model of IVPP V 26257 showing the contact region of the left side of the palatine, pterygoid, and ectopterygoid, in ventral and dorsal views. e The posterior part of the maxilla in ventral, medial, dorsal, and lateral views. f The maxilla process of the jugal in ventral, medial, dorsal, and lateral views. g The palatine in ventral, medial, dorsal, and lateral views. g The palatine in ventral, medial, dorsal, and lateral views.

views. **h** The pterygoid with unclear posterior boundary in ventral, medial, dorsal, and lateral views. **i** the ectopterygoid in ventral, medial, dorsal, and lateral views. Abbreviations: asecp articular surface of ectopterygoid, asj articular surface of jugal, asm articular surface of maxilla, aspl articular surface of palatine, aspt articular surface of pterygoid, asv articular surface of vomer, ch choana, fo foramen, laj lacrimal process of jugal, mpf maxillopalatine fenestra, plfo palatine foramen, pplf postpalatine fenestra, poj postorbital process of jugal, stf subtemporal fenestra. Scale bars:  $\mathbf{a} = 100 \text{ mm}$ :  $\mathbf{c} - \mathbf{i} = 10 \text{ mm}$ .

small foramen perforates the lateral rami of the palatine at the contact surface with the pterygoid and is here called the palatine foramen (Video 1 and Fig. 2b-d). Since this foramen is ventrally covered by the pterygoid, it can only be observed from the posterodorsal view.

The pterygoid is a plate-like bone with two processes (Video 1 and Fig. 2b, h). The anterior process gradually tapers anteriorly, and it ventrally contacts the palatine anterior ramus. The medial process of the pterygoid covers ventrally the posterior part of the palatine (Video 1 and Fig. 2b, h), and medially contacts its counterpart, separating the choanae and the interpterygoid fenestra. Posterior to the choanae, the pterygoid foramen is observed between the palatine and the pterygoid (Fig. 2b). The articulation surface of the pterygoid with the ectopterygoid shows a rough texture (Video 1 and Fig. 2b, h). In the present specimen (IVPP 4063), the pterygoid fuses with the quadrate and probably also with the basisphenoid. The medial process of the pterygoid could have been anteriorly in contact with the vomer, as the slender vomer expands posteriorly to the posterior margin of the choanae<sup>43</sup>.

Previously, the ectopterygoid was described as the "lateral process of the pterygoid" This bone can be divided into two parts (Video 1 and Fig. 2b, i). The medial part is curved dorsoventrally, flattened, and has the dorsal surface perforated by several foramina (Video 1 and Fig. 2b, i). Medially, it contacts the palatine and the pterygoid. The lateral portion of the ectopterygoid is positioned anterolaterally relative to the medial part, and it is a slender, tubular, and parabolic arc-shaped element, that laterally contacts the jugal (Fig. 2f). The ectopterygoid separates the postpalatine fenestra from the subtemporal fenestra (Video 1 and Fig. 2b).

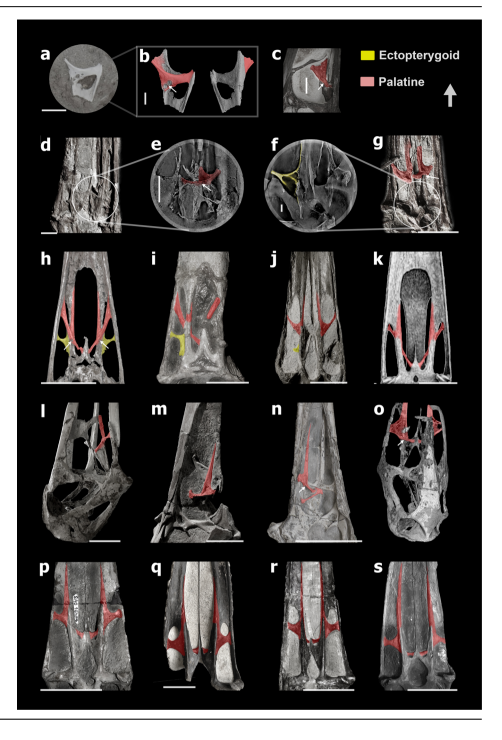
# The palate of *Kunpengopterus sinensis* (Wukongopteridae)

The specimen of *Kunpengopterus sinensis* (IVPP 23674) also preserves a similar "y" shape palatine and also shows a palatine foramen which, as in *Dsungaripterus*, can only be seen from the dorsal side (Fig. 3g). Based on the CL-scans, the bone previously described as the "lateral process of the pterygoid" is the ectopterygoid that shows a clear boundary with the pterygoid (Fig. 3f). There are three palatal openings lateral to the choana, similar to that observed in *Dsungaripterus*. Furthermore, the palatine and ectopterygoid are also similar in both taxa, suggesting that both have developed a similar palatal bone pattern. (Fig. 6b). The main difference is the presence of a small pterygoid foramen in *Dsungaripterus*, while *Kunpengopterus* exhibits a pterygoid fenestra that is more similar to the one observed in *Anhanguera*<sup>27</sup> (Fig. 3), *Hamipterus*<sup>48</sup> (Fig. 4), and *Caupedactylus*<sup>38</sup> (Fig. 3h).

# The palate in non-pterodactyloids

There are not many specimens of non-pterodactyloid pterosaurs that allow the observation of the palate<sup>13,16,24,25,40,44,49,50</sup> (Table 1). Among the best-preserved materials are BML R 2786<sup>18</sup> and CM 11434<sup>33</sup> (Fig. 5a) representing *Rhamphorhynchus*, and *Cacibupteryx* (IGO-V 208<sup>36</sup> Fig. 5b), two Jurassic rhamphorhynchids. Our basic reconstruction follows the one published by Ösi et al.<sup>13</sup>, but with different interpretations of bones and palatal openings (Fig. 6a). According to our interpretation, the palatine has two rami, which form the medial and posterior margin of the maxillopalatine fenestra. Therefore, the opening identified by Ösi et al.<sup>13</sup> as the

Fig. 3 | The similar structure on the palatal region in pterodactyloids. a A piece of skull of Hamipterus tianshanensis (left side) IVPP V 18943.3. b CTrendered bone corresponding to the (a) in dorsal and ventral view. c A close-up shot of the palatine of Ludodactylus SMNK PAL 3828 in dorsal view. d the posterior part of the palate of Hongshanopterus lacustris IVPP V 14852 in ventral view, modified from ref. 35. e CL-scan results corresponding to the circle area in (d). f CL-scan results corresponding to the circle area in (g). g The posterior part of palate of Kunpengopterus sinensis IVPP V 23674 in dorsal view, modified from ref. 41. h The posterior part of the palate of Caupedactylus ybaka MN 4726-V in dorsal view, modified from ref. 38. i The posterior part of the palate of Tupuxuara leonardii IMCF 1052. i The posterior part of the palate of Gnathosaurus subulatus SOS 4580 (cast) in ventral view. k The posterior part of the palate of Pteranodon, modified from ref. 64. l-r Seven skulls of Anhanguera focus on the palate region, and they are RGM 401880 (lateral-dorsal view on the right side, photographed by Dr. T. Rodrigues), AMNH 22555 (lateral view on the right side), SNSB-BSPG 1982 I 89 (lateral view on the left side), AMNH 25555 (dorsal-lateral view on left side, CT-rendered model modified from ref. 33, SAO 16494 (ventral view, photographed by Dr. T. Rodrigues), SNSB-BSPG 1982 I 89 (ventral view, photographed by Dr. T. Rodrigues), MN 4805-V (ventral view), respectively. s The posterior part of palate of Tropeognathus mesembrinus BSP 1987 I 46 (cast) in ventral view. Scale bar:  $\mathbf{a}$ ,  $\mathbf{d}$ ,  $\mathbf{e}$ ,  $\mathbf{g} = 10 \text{ mm}$ ;  $\mathbf{b} = 4.2 \text{ mm}$ ;  $\mathbf{j} = 20 \text{ mm}$ ; f = 1 mm; h, i, k, l, m, p, q, r = 50 mm; n = 100 mm; s = 85 mm.



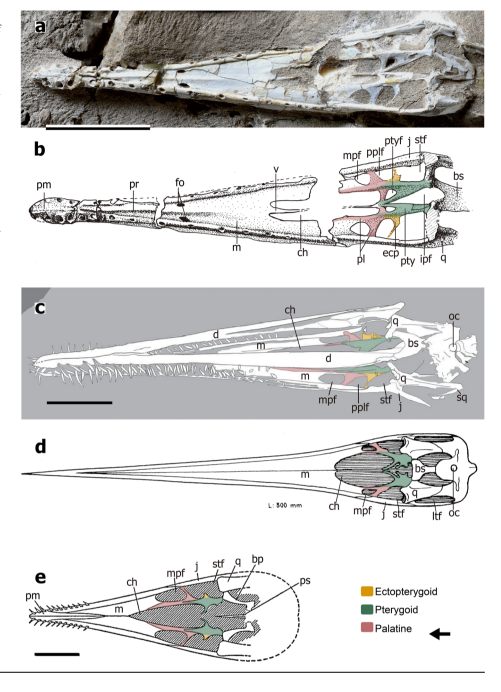
suborbital fenestra and pterygoectopterygoid fenestra should be redesignated as the maxillopalatine and postpalatine fenestrae. Furthermore, Ösi et al. <sup>13</sup> have identified the palatine as being a long, flattened element positioned lateral to the choanae. If, indeed, Ösi et al. <sup>13</sup> are correct, then the palatal configuration of *Dorygnathus*, which also shows an apertura maxillopremaxillaris not recognized in other non-pterodactyloids, might differ from what we present here.

Another member of the Rhamphorhynchidae with a well-preserved palate that, albeit still preserved in the matrix, was observed through CT-scan<sup>51</sup> is *Dearc sgiathanach*. Based on our analysis of the CL-scan of *Kun-pengopterus*, the "y" shaped element observed on the dorsal part that was interpreted as being the ectopterygoid<sup>51</sup> is here regarded to be the palatine. This bone is positioned posterior to the margin of the most anterior opening -the maxillopalatine fenestra-. Both palatines meet at the midline and

contact the vomer. The original openings designated as the suborbital fenestra and pneumatic foramen (perhaps they meant the pterygoid-ectopterygoid fenestra) are, according to our interpretations, the maxillopalatine fenestra and the postpalatine fenestra.

There are at least two other quite distinctive palate configurations present in non-pterodactyloids. The most extreme, as has been already pointed out before, is found in anurognathids where most elements are reduced to rodlike structures<sup>32,50,52-54</sup>. The other distinctive palate has been reported in the wukongopterid *Kunpengopterus* with different relation of the palatal openings, particularly the huge size of the postpalatine fenestra<sup>41</sup>. Based on our study, what originally has been identified as the postpalatine and the secondary subtemporal fenestrae are actually the maxillopalatine and the postpalatine fenestrae, respectively (Fig. 6b). Furthermore, we here reinterpret the ectopterygoid and the lateral process of the

Fig. 4 | The nearly complete palatal region from the other four pterodactyloids. a Photo of a skull of Hamipterus tianshanensis IVPP V 18943.1 seen on the ventral side, and **b** is the sketch drawing of it. c Line drawings of the ventral side of *Liaodactylus* primus PMOL-AP00031, modified from ref. 42. d Reconstruction of the ventral side of Nyctosaurus gracilis FM P 25026, modified from ref. 12. e Reconstruction of the ventral side of Aurorazhdarcho micronyx BSP 1936 I 50, modified from ref. 23. Abbreviations: bp basipterygoid, bs basisphenoid, ch choana, d dentary, ecp ectopterygoid, fo foramen, ipf interpterygoid fenestra, j jugal, ltf lower temporal fenestra, m maxilla, mpf maxillopalatine fenestra, oc occipital condyle, pm premaxilla, pl palatine, pplf postpalatine fenestra, pr palatal ridge, pty pterygoid, ptyf pterygoid fenestra, ps parasphenoid, q quadrate, sq squamosal, stf subtemporal fenestra, v vomer. Scale bars: a, b = 50 mm, c = 20 mm, e = 5 mm.



pterygoid of the original description<sup>41</sup> as the palatine and the ectopterygoid, respectively (Fig. 6b).

Previous reconstructions of the palate of other non-pterodactyloid taxa are also reinterpreted here. The sole known specimen of *Parapsicephalus* (GSM 3166)<sup>16,44</sup> is not very well preserved and shows only the partial right side of the palate that might follow the same pattern (Fig. 5c) as we report for the rhamporhynchids discussed previously. In *Campylognathoides* (CM 11424)<sup>24,49</sup> the bone observable through the orbit and identified as the ectopterygoid<sup>24</sup> is here the same as our interpretation (Fig. 5e). Lastly, in the palate of *Scaphognathus* (GPIB 1304)<sup>25,40</sup> the bones identified as the ectopterygoid might be the palatine (Fig. 5f).

#### The palate in pterodactyloids

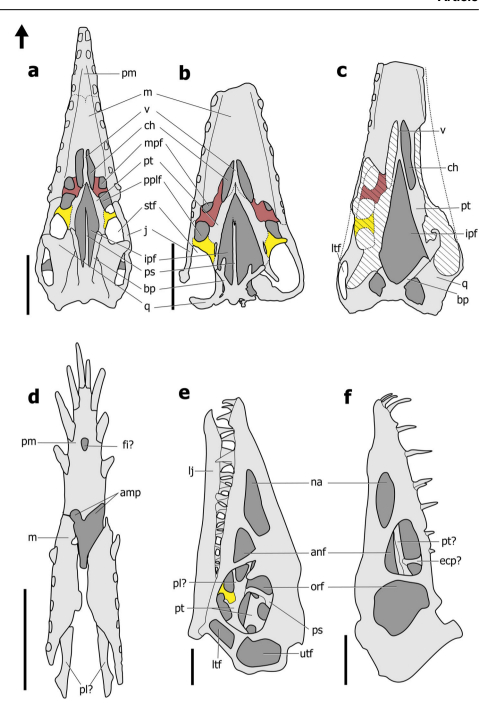
The new interpretation of the palatal structure in *Dsungaripterus* (Dsungaripteridae) can also be applied to other pterodactyloid clades. CT-scans and CL-scans of *Hongshanopterus*<sup>35</sup> (Istiodactylidae) (Fig. 3d, e) and

*Hamipterus*<sup>48</sup> (Hamipteridae) allowed the examination of the dorsal side of the palate, indicating the presence of a "y" shape palatine structure as in *Dsungaripterus*. In several other pterodactyloids such as *Anhanguera*<sup>10,27,33,34</sup> (Anhangueridae, Fig. 3d, j, q), *Ludodactylus*<sup>55</sup> (Ornithocheiridae, Fig. 3c) and probably also in *Caupedactylus*<sup>38</sup> (Tapejaridae, Fig. 3h), also present a similar presence of a "y" shape palatine.

Regarding palatal openings, several pterodactyloid taxa show three main palatal openings positioned lateral to the choanae. Based on our study, the most anterior of these openings should be the maxillopalatine fenestra. This appears to be the case of the tapejarid *Tupuxuara*<sup>14</sup>, and archae-opterodactyloid ctenochasmatids *Gnathosaurus*<sup>12</sup> and *Liaodactylus*<sup>42</sup> (Figs. 3j, i, 4c).

Some other pterodactyloids present only two openings lateral to the choana, such as *Pteranodon*<sup>31</sup> (Pteranodontidae), *Tropeognathus*<sup>26</sup> and *Anhanguera*<sup>27,56</sup> (Anhangueridae), *Nyctosaurus*<sup>19</sup> (Nyctosauridae), and *Aurorazhdarcho*<sup>57</sup> (Ctenochasmatidae or *Pterodactylus*<sup>23</sup>, Pterodactylidae).

Fig. 5 | Line drawings of the palatal region in nonpterodactyloid. a Rhamphorhynchus muensteri CM 11434, modified from ref. 33. b Cacibupteryx caribensis IGO-V 208, modified from ref. 36. c Parapsicephalus purdoni GSM 3166, modified from ref. 16, and the skull length is about 140 mm. d Dorygnathus banthensis SMNS 18969, modified from ref. 13. e Campylognathoides liasicus CM 11424, modified from ref. 49. f Scaphognathus crassirostris GPIB 1304 (Nr. 109), modified from ref. 25. The red represents the palatine and the yellow represents the ectopterygoid. Abbreviations: amp apertura maxillo-premaxillaris, anf antorbital fenestra, bp basipterygoid, ch choana, ecp ectopterygoid, fi foramen incisivum, ipf interpterygoid fenestra, j jugal, lj lower jaw, ltf lower temporal fenestra, m maxilla, mpf maxillopalatine fenestra, na naris, orf orbital fossa, pl palatine, pm premaxilla, pplf postpalatine fenestra, ps parasphenoid, pt pterygoid, q quadrate, stf subtemporal fenestra, utf upper temporal fenestra, v vomer. Scale bar: a, f = 20 mm; b, d = 50 mm; e = 10 mm.



The most anterior one, as in other pterodactyloids, is the maxillopalatine fenestra, followed by the subtemporal fenestra (Figs. 3k–s, 4d, e). This suggests that they lack a postpalatine fenestra. The main reason for this is the possible lack of the development of a lateral process of the ectopterygoid, that in the other studied pterodactyloid separates postpalatine fenestra from the subtemporal fenestra (Fig. 6c). In some specimens of *Anhanguera* (SNSB-BSPG 1982 I 89), *Hongshanopterus*, *Gnathosaurus*, *Aurorazhdarcho*, the ectopterygoid shows a blunt incipient process directed into the subtemporal opening.

As observed in non-pterodactyloids, some palatal elements might also be reinterpreted in pterodactyloids. The palatine as presented in previous studies<sup>12,23,26-28,31,38,43,48</sup> is here reinterpreted as the median maxilla process (Fig. 6c). Also the bone interpreted as the ectopterygoid in some taxa<sup>28,31,35,38,42,43,48,58,59</sup> are here considered as being the palatines, which shows two rami as observed in the CT-scans of *Dsungaripterus*.

The diverse shape of the maxillopalatine fenestra observed in non-pteradactyoids and pterodactyloids (Figs. 2, 3, 4) is influenced by several factors, including the shape and extension of the palatine process of the maxilla, the angle between the two rami of palatine, and the contact of the palatine lateral ramus with bony bar formed by maxilla and jugal. The angle between the two rami of the palatine varies from ~30° in *Dsungaripterus*, *Caupedactylus*, *Tupuxuara*, *Nyctosaurus gracilis*, and *Pteranodon*, ~45° in *Hongshanopterus*, *Aurorazhdarcho*, *Liaodactylus primus*, and *Gnathosaurus*, and 45°–60° in *Hamipterus* and *Anhanguera*.

In the region between the choanae and the interpterygoid fenestra, a pair of small openings are present and vary within pterodactyloids in size. For example, in *Dsungaripterus* these openings are small, forming a pterygoid foramen, while in *Anhanguera*, *Hamipterus*, and *Caupedactylus* they are large, forming a pterygoid fenestra (Figs. 3, 4). There seems to be some variation in the sizes of these openings, that could be observed in some

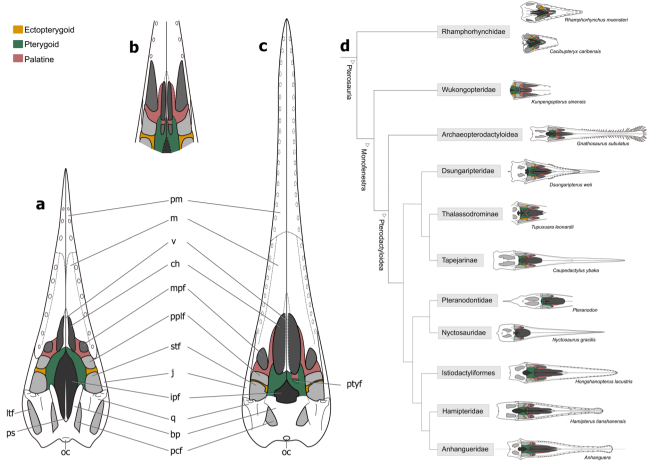


Fig. 6 | The line drawings of three hypothetical models of the palate in ventral view and its evolution in pterosaurs. a palate model based on *Rhamphorhynchus* and *Cacibupteryx*. b Palate model of *Kunpengopterus*. c Palate model of pterodactyloids. d The cladogram was simplified and modified from ref. 47. Line drawings of the palates of *Rhamphorhynchus*, *Cacibupteryx*, *Gnathosaurus*, *Caupedactylus*, *Pteranodon*, *Nyctosaurus*, *Hongshanopterus*, and *Anhanguera* were

reconstructed and modified from previous publications  $^{12,27,33,35,36,38,64}$ . Abbreviations: bp basipterygoid, ch choana, ipf interpterygoid fenestra, j jugal, ltf lower temporal fenestra, m maxilla, mpf maxillopalatine fenestra, oc occipital condyle, pcf posterior cranial fenestra, pm premaxilla, pplf postpalatine fenestra, ps parasphenoid, ptyf pterygoid fenestra, q quadrate, stf subtemporal fenestra, v vomer.

specimens of *Anhanguera*, being more developed in some (SNSB-BSPG 1982 I 89, MN 4805-V, BSP 1987 I 46, and SAO 16494) and smaller in others (Fig. 3, RGM 401 880, AMNH 25555, SNSB-BSPG 1982 I 90, and AMNH 22555).

Although an extensive comparison of the pterosaur palate with other diapsids is beyond the scope of this paper, there are a few comments and differences that can be highlighted. The most significant is the presence of a maxillopalatine fenestra, which is absent in other diapsids 58,60,61 and might turn out to be a synapomorphy of Pterosauria. In crocodyliforms, the suborbital fenestra is anteriorly formed by the maxilla and the palatine, and posteriorly by the pterygoid and ectopterygoid, lacking any maxillopalatine fenestra 1. In prolacertiformes (e.g., ref. 63) and lepidosauromopha 58,60, the internal naris is positioned at the lateral side of the palate, followed by a postpalatine fenestra, and lacks any extra opening between them. But the postpalatine fenestra is similar to the one observed in pterosaurs, since it is formed anteriorly by the palatine and posteriorly by ectopterygoid.

To conclude, the new relation established here between the palatine, ectopterygoid, maxilla, and pterygoid suggest some reinterpretation of the main palatal openings (Fig. 6d). Although it has to be acknowledged that there is still much work on the pterosaur palate necessary to be able to establish stronger evolutionary trends on this portion of the pterosaur skull, six main general remarks can be presented.

1. In some non-pterodactyloids (e.g., *Rhamphorhynchus* and *Cacibupteryx*), the anterior process of the pterygoids is not as developed as it is in pterodactyloids (Fig. 6d).

- 2. The angle between the two rami of the palatine in some non-pterodactyloids are nearly 90°, in *Kunpengopterus* is 45°, and in pterodactyloids are <60° (e.g., ~30° in *Dsungaripterus*, ~60° in *Hamipterus*).
- In pterodactyloids the jugal process of the maxilla extends posteriorly until the ectopterygoid, but not in the non-pterodactyloids (*Rham-phorhynchus* and *Cacibupteryx*).
- 4. The last tooth is posterior to the anterior margin of maxillopalatine fenestrae in non-pterodactyloids (e.g., Dorygnathus, Cacibupteryx, Rhamphorhynchus, and Kunpengopterus), but is anterior to the maxillopalatine fenestrae in toothed pterodactyloids (e.g., Hongshanopterus, Anhanguera, Tropeognathus, Liaodactylus, Gnathosaurus, Aurorazhdarcho, Hamipterus, and Dsungaripterus).
- 5. Pterodactyloids have a pair of pterygoid openings (foramen or fenestra) bordered by the lateral ramus of the palatine and the median process of pterygoid. This opening is present in the nonpterodactyloid *Kunpengopterus* and *Cacibupteryx*, but not in *Rhamphorhynchus*.
- 6. Along with the great variation in the sizes of the palatal openings such as the pterygoid foramen or fenestra, the posterior margin of the choanae moved posteriorly to the maxillopalatine fenestra in *Kunpengopterus* and pterodactyloids (Fig. 6d) and even posterior to the postpalatine fenestra in some pterodactyloids (e.g., *Caupedactylus*, *Tupuxuara*, and *Dsungaripterus*).

# **Data availability**

The authors declare that the main data supporting the findings of this study are available within the article and its Supplementary Information file. Extra data were available from the corresponding author or the first author upon request.

Received: 21 May 2023; Accepted: 2 April 2024; Published online: 12 April 2024

#### References

- Bennett, S. C. The phylogenetic position of the *Pterosauria* within the Archosauromorpha. Zool. J. Linnean Soc. 118, 261–308 (1996).
- Kellner, A. W. A. Pterosaur phylogeny and comments on the evolutionary history of the group *Geol. Soc. Spec. Publ.* 217, 105–137 (2003).
- Unwin, D. M. On the phylogeny and evolutionary history of pterosaurs. Geol. Soc. Spec. Publ. 217, 139–190 (2003).
- 4. Andres, B., Clark, J. M. & Xu, X. The earliest pterodactyloid and the origin of the group. *Curr. Biol.* **24**, 1011–1016 (2014).
- Codorniú, L. S., Paulina-Carabajal, A., Pol, D., Unwin, D. M. & Rauhut, O. W. M. A Jurassic pterosaur from Patagonia and the origin of the pterodactyloid neurocranium. *PeerJ* 4, e2311 (2016).
- Vidovic, S. U. & Martill, D. M. The taxonomy and phylogeny of Diopecephalus kochi (Wagner, 1837) and 'Germanodactylus rhamphastinus' (Wagner, 1851). Geol. Soc. Spec. Publ. 455, 125–147 (2017).
- Ezcurra, M. D. et al. Enigmatic dinosaur precursors bridge the gap to the origin of *Pterosauria*. *Nature* 588, 445–449 (2020).
- Pêgas, R. V., Costa, F. R. & Kellner, A. W. A. Reconstruction of the adductor chamber and predicted bite force in pterodactyloids (*Pterosauria*). Zool. J. Linnean Soc. https://doi.org/10.1093/ zoolinnean/zlaa163 (2021).
- Yu, Y., Zhang, C. & Xu, X. Complex macroevolution of pterosaurs. Curr. Biol. 33, 770–779.e4 (2023).
- Wellnhofer, P. Weitere Pterosaurierfunde aus der Santana-Formation (Apt) der Chapada do Araripe, Brasilien. *Palaeontogr. Abt. A* 215, 43–101 (1991).
- Kellner, A. W. A. et al. Pterodactyloid pterosaur bones from Cretaceous deposits of the Antarctic Peninsula. An. Acad. Bras. Ciênc. 91, e20191300 (2019).
- Wellnhofer, P. Pterosauria. Handbuch der paläoherpetologie Teil 19, x–82 (1978).
- Ősi, A., Prondvai, E., Frey, E. & Pohl, B. New interpretation of the palate of pterosaurs. Anat. Rec. 293, 243–258 (2010).
- Pinheiro, F. L. & Schultz, C. L. An unusual pterosaur specimen (Pterodactyloidea,? Azhdarchoidea) from the Early Cretaceous Romualdo Formation of Brazil, and the evolution of the pterodactyloid palate. PLoS ONE 7, e50088 (2012).
- Marsh, O. C. Principal characters of American Cretaceous Pterodactyls: part 1, the skull of *Pteranodon. Am. J. Sci.* 27, 423–426 (1884).
- Newton, E. T. XIX. On the skull, brain, and auditory organ a new species of pterosaurian (*Scaphognathus purdoni*), from the Upper Lias near Whitby, Yorkshire. *Philos. Trans. R. Soc. Lond. B Biol. Sci.* 179, 503–537 (1888).
- 17. Seeley, H. *Dragons of the Air: An Account of Extinct Flying Reptiles* (Appleton & Co, Methuen & Co, 1901).
- Woodward, A. S. On two skulls of the ornithosaurian Ohampnorhynchus. Ann. Mag. Nat. Hist. ume 9, 1–5 (1902).
- Williston, S. W. On the Skull of *Nyctodactylus*, an Upper Cretaceous Pterodactyl. *J. Geol.* 10, 520–531 (1902).
- von Huene, F. Beitra ge zur Kenntnis des Scha dels einiger Flugsaurier. Geol Pala ontol Abh Neue Folge 13, 57–65 (1914).

- Broili, F. Ctenochasma gracile Oppel. Sond. Geog. Jahresheften 29/ 30, 299–309 (1919).
- Arthaber, G. Studien über Flugsaurier und Bearbeitung des Wiener Exemplares von Dorygnathus banthensis Theod. sp. Akademie der Wissenschaften in Wien. Mathematisch-Naturwissenschaftliche Klasse: Anzeiger 13, 1–2 (1919).
- 23. Wellnhofer, P. Die Pterodactyloidea (Pterosauria) der Oberjura-Plattenkalke süddeutschlands, Bayer. Akademie der Wissenschaften, Mathematisch-Naturwissenschaftliche Klasse, Abhandlungen, Neue folge Heft **141**, 1–133 (1970).
- Wellnhofer, P. Campylognathoides liasicus (Quenstedt), an Upper Liassic pterosaur from Holzmaden: the Pittsburgh specimen. Ann. Camegie Mus. 147, 5–34 (1974).
- Wellnhofer, P. Die Rhamphorhynchoidea (Pterosauria) der Oberjura-Plattenkalke süddeutschlands Teil I. *Palaeontogr. Abt. A* 148, 1–33 (1975).
- Wellnhofer, P. New crested pterosaurs from the Lower Cretaceous of Brazil. Mitteilungen der Bayerischen Staatssammlung für Paläontologie und Historische. Geologie 27, 175–186 (1987).
- Wellnhofer, P. Neue Pterosaurier aus der Santana-Formation (Apt) der Chapada do Araripe, Brasilien. *Palaeontogr. Abt. A* 187, 105–182 (1985).
- Campos, D. D. A. & Kellner, A. W. A. Panorama of the flying reptiles study in Brazil and South America. An. Acad. Bras. Ciênc. 57, 454–466 (1985).
- Kellner, A. W. A. A new edentate pterosaur of the lower Cretaceous from the Araripe Basin, Northeast Brazil. An. Acad. Bras. Ciênc. 61, 439–446 (1989).
- Kellner, A. W. A. & Campos, D. A. A new species of *Tupuxuara* (Pterosauria, Tapejaridae) from the Early Cretaceous of Brazil. *An. Acad. Bras. Ciênc.* 66, 467–474 (1994).
- Bennett, S. C. The osteology and functional morphology of the Late Cretaceous pterosaur *Pteranodon. Palaeontogr. Abt. A* 260, 1–153 (2001).
- 32. Bennett, S. C. A second specimen of the pterosaur *Anurognathus ammoni*. *Paläontol*. *Z*. **81**, 376–398 (2007).
- Witmer, L. M., Chatterjee, S., Franzosa, J. & Rowe, T. Neuroanatomy of flying reptiles and implications for flight, posture and behaviour. *Nature* 425, 950–953 (2003).
- Veldmeijer, A. J. Description of Coloborhynchus spielbergi sp. nov.(Pterodactyloidea) from the Albian (Lower Cretaceous) of Brazil. Scripta. Geologica 125, 35–139 (2003).
- Wang, X., Campos, D. de A., Zhou, Z. & Kellner, A. W. A. A primitive istiodactylid pterosaur (Pterodactyloidea) from the Jiufotang formation (Early Cretaceous), northeast China. *Zootaxa* 1813, 1–18 (2008).
- Gasparini, Z., Fernández, M. & de la Fuente, M. A new pterosaur from the Jurassic of Cuba. *Palaeontology* 47, 919–927 (2004).
- 37. Padian, K. The Early Jurassic pterosaur *Dorygnathus banthensis* (Theodori, 1830). *Spec. Pap. Palaeontol.* **80**, 1–64 (2008).
- Kellner, A. W. A. A new unusual tapejarid (Pterosauria, Pterodactyloidea) from the Early Cretaceous Romualdo Formation, Araripe Basin, Brazil. Earth Environ. Sci. Trans. R. Soc. Edinb. 103, 409–421 (2013).
- Bennett, S. C. The morphology and taxonomy of the pterosaur Cycnorhamphus. Neues Jahrb. Geol. Paläontol. Abh. 267, 23–41 (2013).
- Bennett, S. C. A new specimen of the pterosaur Scaphognathus crassirostris, with comments on constraint of cervical vertebrae number in pterosaurs. Neues Jahrb. Geol. Paläontol. Abh. 271, 327–348 (2014).
- Cheng, X., Jiang, S., Wang, X. & Kellner, A. W. A. New anatomical information of the wukongopterid *Kunpengopterus sinensis* Wang et al., 2010 based on a new specimen. *PeerJ* 5, e4102 (2017).

- Zhou, C. et al. Earliest filter-feeding pterosaur from the Jurassic of China and ecological evolution of Pterodactyloidea. *R. Soc. Open Sci.* 4, 160672 (2017).
- Chen, H. et al. New anatomical information on *Dsungaripterus* weii Young, 1964 with focus on the palatal region. *PeerJ* 8, e8741 (2020).
- O'Sullivan, M. & Martill, D. M. The taxonomy and systematics of Parapsicephalus purdoni (Reptilia: Pterosauria) from the Lower Jurassic Whitby Mudstone Formation, Whitby, U.K. Hist. Biol. 29, 1009–1018 (2017).
- Peters, D. A reexamination of four prolacertiforms with implications for pterosaur phylogenesis. *Riv. Ital. Paleontol. Stratigr.* 106, 293–336 (2000).
- Martin-Silverstone, E. et al. A new pterosaur from the Middle Jurassic of Skye, Scotland and the early diversification of flying reptiles. J. Vertebr. Paleontol. e2298741 https://doi.org/10.1080/02724634. 2023.2298741 (2024).
- Kellner, A. W. A. et al. First complete pterosaur from the Afro-Arabian continent: insight into pterodactyloid diversity. Sci. Rep. 9, 17875 (2019).
- Wang, X. et al. Sexually dimorphic tridimensionally preserved pterosaurs and their eggs from China. *Curr. Biol.* 24, 1323–1330 (2014).
- Padian, K. The Early Jurassic pterosaur Campyloganthoides (Strand, 1928). Spec. Pap. Palaeontol. 80, 69–107 (2008).
- Ji, Q. & Yuan, C. Discovery of two kinds of protofeathered pterosaurs in the Mesozoic Daohugou Biota in the Ningcheng region and its stratigraphic and biologic significances. *Geol. Rev.* 48, 221–224 (2002).
- Jagielska, N. et al. A skeleton from the Middle Jurassic of Scotland illuminates an earlier origin of large pterosaurs. *Curr. Biol.* 32, 1446–1453 (2022).
- Unwin, D. M., Lü, J. & Bakhurina, N. N. On the systematic and stratigraphic significance of pterosaurs from the Lower Cretaceous Yixian Formation (Jehol Group) of Liaoning, China. *Mitt. Mus. Nat.kd. Berl. Geowissenschaftliche Reihe* 3, 181–206 (2000).
- Wang, X., Zhou, Z., Zhang, F. & Xu, X. A nearly completely articulated rhamphorhynchoid pterosaur with exceptionally well-preserved wing membranes and "hairs" from Inner Mongolia, northeast China. *Chin.* Sci. Bull. 47, 226–230 (2002).
- 54. Yang, Z. et al. Pterosaur integumentary structures with complex feather-like branching. *Nat. Ecol. Evol.* **3**, 24–30 (2019).
- Frey, E., Martill, D. M. & Buchy, M.-C. A new crested ornithocheirid from the Lower Cretaceous of northeastern Brazil and the unusual death of an unusual pterosaur. *Geol. Soc. Spec. Publ.* 217, 55–63 (2003).
- Pinheiro, F. L. & Rodrigues, T. Anhanguera taxonomy revisited: is our understanding of Santana Group pterosaur diversity biased by poor biological and stratigraphic control? *PeerJ* 5, e3285 (2017).
- 57. Bennett, S. C. New information on body size and cranial display structures of *Pterodactylus antiquus*, with a revision of the genus. *Paläontol. Z.* **87**, 269–289 (2013).
- Evans, S. E. The skull of lizards and Tuatara. in *Biology of the Reptilia*.
  Volume 20. Morphology H. The Skull of Lepidosauria (eds Gans, C. et al.) Ch. 1, 1–347 (Society for the Study of Amphibians and Reptiles, 2008).
- Wellnhofer, P. The Illustrated Encyclopedia of Pterosaurs (Salamander Books, 1991).
- Benton, M. J. Classification and phylogeny of the diapsid reptiles. *Zool. J. Linnean Soc.* 84, 97–164 (1985).
- Hu, H. et al. Evolution of the vomer and its implications for cranial kinesis in Paraves. figshare https://doi.org/10.6084/M9.FIGSHARE. 7769279.V2 (2019).

- Dollman, K. N., Clark, J. M., Norell, M. A., Xing, X. & Choiniere, J. N. Convergent evolution of a Eusuchian-type secondary palate within Shartegosuchidae. *Am. Mus. Novit.* 3901, 1–23 (2018).
- 63. Sen, K. Pamelaria dolichotrachela, a new prolacertid reptile from the Middle Triassic of India. *J. Asian Earth Sci.* **21**, 663–681 (2003).
- 64. Eaton, G. F. Osteology of Pteranodon. *Mem. Con. Acad. Arts Sci.* **2**, 1–38 (1910).

# **Acknowledgements**

We acknowledge our IVPP research team members Long Xiang, Yan Li, Hongjiao Zhou, Ruijie Wang, Jinkun Mao, Rui Qiu, Yang Li, Xinjun Zhang and Junxia Wang for fieldwork and specimen preparation. The authors would like to thank Dr. Yemao Hou, Pengfei Yin, Xiaoyi Dong, and Dr. Yanxin Gong for their help with the CT and CL scanning and 3D reconstructions in IVPP. We have benefited from discussions with Drs. Xing Xu, Liping Dong, Jun Liu, Rui Qiu, Xin Cheng, Xinjun Zhang, and Yang Li. For granting access and assistance to the study of AMNH 25555, we thank Drs. Mark A Norell (American Museum of Natural History) and Congyu Yu, Qigao Jiangzuo. Also, we would like to thank Prof. Lawrence M. Witmer (Ohio University) for kindly helping with the data file of CM 11434, Prof. Taissa Rodrigues (Universidade Federal do Espírito Santo), Prof. Renan A.M. Bantim (Universidade Regional do Cariri) and Rodrigo V. Pêgas (Universidade Federal do ABC) for providing high-resolution photos Anhangueridae, Maaradactylus kellneri, and Tupuxuara leonardii respectively. We also thank Dr. Thomas Stidham and Dr. Paul Rummy for their constructive comments that highly improve the standards of this manuscript. This work was supported by the following funding: National Natural Science Foundation of China (42288201, 42072028, 41572020, and 42302003), the Strategic Priority Research Program (B) of CAS (XDB26000000), and the Youth Innovation Promotion Association of Chinese Academy of Sciences (2019075). AWAK acknowledges funding from the Conselho Nacional de Desenvolvimento Científico e Tecnológico (CNPg # 308707/2023-0, #406779/2021-0, #406902/2022-4) and the Fundação Carlos Chagas Filho de Amparo à Pesquisa do Estado do Rio de Janeiro (FAPERJ #E-26/ 201.095/2022).

#### **Author contributions**

All authors contributed to the interpretation and discussion of results. H.C. designed the study, collected the data, and wrote the paper. S.J., A.W.A.K., and X.W. performed the research and contributed to the initial discussions.

### **Competing interests**

The authors declare no competing interests.

## **Additional information**

**Supplementary information** The online version contains supplementary material available at https://doi.org/10.1038/s42003-024-06132-6.

**Correspondence** and requests for materials should be addressed to Xiaolin Wang.

**Peer review information** *Communications Biology* thanks David Unwin and the other, anonymous, reviewer(s) for their contribution to the peer review of this work. Primary Handling Editors: Borja Figueirido and Luke R. Grinham.

**Reprints and permissions information** is available at http://www.nature.com/reprints

**Publisher's note** Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons licence, and indicate if changes were made. The images or other third party material in this article are included in the article's Creative Commons licence, unless indicated otherwise in a credit line to the material. If material is not included in the article's Creative Commons licence and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this licence, visit <a href="http://creativecommons.org/licenses/by/4.0/">http://creativecommons.org/licenses/by/4.0/</a>.

© The Author(s) 2024