A new azhdarchoid pterosaur from the Lower Cretaceous of Brazil and the paleobiogeography of the Tapejaridae

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The Tapejaridae were an apparently worldwide distributed clade of edentulous pterosaurs, being a major component of several Lower Cretaceous terrestrial faunas. Despite their distribution across Gondwana and Laurasia, the oldest tapejarid remains were found in Barremian units from Europe, what led to the assumption that the clade originated in Eurasia and later dispersed southwards. Here we present a new tapejarid pterosaur species (Karihidraco dianae gen. et sp. nov.) from the Lower Cretaceous Romualdo Formation of Brazil. The holotype (MPSC R 1056) comprises an incomplete, three dimensionally preserved skull, lower jaw, and cervical vertebrae. It shows a unique combination of features such as unusually tall and comparatively short nasoantorbital fenestrae, as well as a premaxillary crest forming an angle of about 45° with respect to main skull axis. Phylogenetic analyses recover the new taxon as a Tityruxara-related Thalassodrominae, a clade of early-diverging tapejarids that were apparently indigenous to central Gondwana. The inclusion of the new taxon in current phylogenetic frameworks, in addition to similarity cluster analyses of Early Cretaceous tapejarid-bearing pterosaur faunas, indicate Gondwana as the most parsimonious origin center for Tapejaridae, and show that pterosaur communities were affected by large scale tectonic-driven vicariant events.

Key words: Pterosauria, Azhdarchoidea, phylogeny, biogeography, Cretaceous, Romualdo Formation, Araripe Basin, Brazil.

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Introduction

The Romualdo Formation of the Araripe Basin (Lower Cretaceous, Northeastern Brazil) is a well-known fossil Konservat-Lagerstätte, especially notorious for the occurrence of some of the best three-dimensionally preserved pterosaur specimens worldwide, as well as for its diverse pterosaur fauna (e.g., Kellner et al. 2002, 2013; Vila Nova et al. 2011; Bantim et al. 2014; Pinheiro and Rodrigues 2017). A major component of the Santana Group pterosaur fauna, the Tapejaridae, is a peculiar clade of toothless animals that draws attention due the extreme morphology observed in some of its representatives. Tapejarids were so far reported in Lower Cretaceous strata from China, Morocco, Spain, England and Brazil (e.g., Kellner 1989; Wellnhofer and Buffetau 1999; Wang and Zhou 2003; Holgado et al. 2011; Vullo et al. 2012; Kellner et al. 2019; Martill et al. 2020a, b). The tapejarid clade Thalassodromineae is thus far exclusive to the Lower Cretaceous of Northeastern Brazil (Kellner and Campos 2007), whereas the Tapejarinae, the most diverse tapejarid clade, had a widespread distribution in Gondwana and Laurasia (Martill et al. 2020a, b).

The Thalassodromineae were originally defined by Kellner and Campos (2007) as those tapejarids presenting high nasoantorbital fenestra and a premaxillary sagittal crest beginning on the anterior part of the skull and expanding
posteriorly. The Tapejarinae, the other tapejarid subclade, would allocate those pterosaurs with a downturned rostral end of the maxilla, a premaxillary crest constricted above the nasoantorbital fenestra, as well as for having orbits positioned only slightly below the dorsal margin of the nasoantorbital fenestra (Kellner and Campos 2007, Pinheiro et al. 2011). Despite being consistently recovered as a basal tapejarine (e.g., Kellner 2013; Holgado et al. 2019; Kellner et al. 2019), the Brazilian tapejarid *Caupedactylus ybaka* shares nasoantorbital fenestrae, Thalassodrominae a tall, unconstricted premaxillary crest, and orbits positioned very low with respect to the nasoantorbital fenestrae.

Tapejarine pterosaurs have a wider stratigraphic distribution when compared to other tapejarids, and their oldest remains were recovered from Barremian European units (Vullo et al. 2012; Martill et al. 2020a). This led to the conclusion that the Tapejaridae originated in Eurasia, posteriorly spreading southwards to diversify in central Gondwana (Vullo et al. 2012; but see Pêgas et al. 2016).

Here we report a new edentulous pterosaur species from the Romualdo Formation, consisting of an incomplete skull associated to the lower jaws and cervical vertebrae I–IV. Besides adding to the pterosaur diversity of the Araripe Basin, the new taxon contributes with new information on tapejarid biogeography, challenging an Eurasian origin for this clade.

**Institutional abbreviations.**—AMNH, American Museum of Natural History, New York, USA; CPA, Centro de Pesquisas Paleontológicas da Chapada do Araripe, Crato, Brazil; IMCF, Iwaki Coal and Fossil Museum, Japan; MCT, DGM, Museu de Ciências da Terra Rio de Janeiro, Brazil; MN, Museu Nacional, Rio de Janeiro, Brazil; MPSC, Museu de Paleontologia Plácido Cidade Nuvens, Santana do Cariri, Brazil; NMSG, Naturmuseum St. Gallen, Switzerland; SMNK, Staatliches Museum für Naturkunde, Karlsruhe, Germany; UFC, Universidade Federal do Ceará, Fortaleza, Brazil.

**Other abbreviations.**—Char., character; OTU, operational taxonomic unit.

**Nomenclatural acts.**—This published work and the nomenclatural acts it contains have been registered in ZooBank: urn:lsid:zoobank.org:pub:3031C608-123C-422C-820D-249892E8CBA5

## Material and methods

**Phylogenetic analyses.**—Cladistic analyses were performed to test the relationships of *Kariridraco dianae* gen. et sp. nov. within Pterosaurus. The new specimen (MPSC R 1056) was included as an operational taxonomic unit (=OTU) in a modified version of Holgado et al. (2019) dataset, resulting in a data matrix that totalizes 146 characters and 60 OTUs. Updates in Holgado et al. (2019) original dataset are as follows: (i) inclusion of two additional OTUs (*Keresdrakon vilsoni* and *Tupuxuara longicristatus*); (ii) corrections on the original coding of several tapejarids for character 53, following firsthand observation of specimens; and (iii) inclusion of two additional characters (Char. 36: premaxillary sagittal crest, when starting at close or at the anterior portion of the skull, tilt angle [0] under 30°, [1] 30–45°, [2] 45–60°, [3] over 60°, character inapplicable for pterosaurs lacking a rostral premaxillary crest; and Char. 146: palate, surface at mid-length [0] flat or concave, [1] convex). Updated scores for *K. vilsoni* were included following the modified dataset of Holgado and Pêgas (2020). Our preferred analysis excludes a priori this last OTU, as it was detected that its inclusion generates considerable noise during parsing. The data matrix was edited using MESQUITE v.3.51 (Maddison and Maddison 2018), and is included as a SOM, Supplementary Online Material available at http://app.pan.pl/SOM/app66-Cerqueira_etal_SOM.pdf. Heuristic searches were performed using the TNT (Tree analysis using New Technology) version 1.5 (Goloboff and Catalano 2016) software. Searches were conducted using TNT tree bisection reconnection swapping algorithm, with 1000 replications and random seed.

**Faunistic similarity indexes.**—The Jaccard similarity index is a β-diversity method used to estimate similarity between three or more sample sites using presence/absence data and paired groups. In Jaccard index, 0 indicates a total dissimilarity between a pair of sites, and a 1 value means that both sites are equal (Sales et al. 2018).

Here we use the Jaccard similarity index to quantify overall taxonomic similarity between different tapejarid-bearing pterosaur faunas. Only pterosaur data was considered for each unit, being recorded the presence/absence of different groups (using subfamilies as preferred Linnaean taxonomic hierarchy, see SOM). All sampled units contain tapejarid representatives, but our raw sample is not restricted to this clade. Sampled sedimentary units are distributed both in Laurasia and Gondwana, and they are as follows: Crato, Romualdo, and Goio-Erê formations (Brazil); Jiufotang and Yixian formations (China); Kem Kem Group (Morocco); Wessex Formation (England, UK) and La Huérguina Formation (Spain). As the Hungarian Csehbánya Formation, thus far records only one nominal pterosaur taxon, we choose to prune it a priori from our dataset. The list of sampled units and recorded taxa considered in the present study is displayed in the SOM. To calculate Jaccard indexes, we used the clustering analyses included in the software package PAST v. 3.18 (Hammer et al. 2001) and the data was presented as a dendrogram.

## Geological setting

The material described herein comes from a carbonate concretion found in the Romualdo Formation (e.g., Saraiva et al. 2007), one of the units that compose the Araripe Basin’s sedimentary succession. The Romualdo Formation com-
prises a wide range of lithologies and is characterized by conglomeratic sandstone followed by a sequence of green and black shale (Assine 2007), stratified conglomerates, fine-to medium-grained sandstone, laminated limestone, marl, and coquina levels (Assine et al. 2014; Melo et al. 2020). The black shale is occasionally intercalated with several layers where carbonate concretions are abundant (Fara et al. 2005; Saraiva et al. 2007; Vila Nova et al. 2011; Kellner et al. 2013). These layers probably were deposited following pulses of mass mortality, and the lithologies characterize a coastal/marine environment (Melo et al. 2020). The Romualdo Formation was recently considered as Aaptian in age, based on palynomorph, foraminifera and ostracod data (Arari and Assine 2020; Melo et al. 2020). The Romualdo Formation crops out on the slopes of the Araripe Plateau, Northeastern Brazil. The precise locality were the new material was found is, however, unknown.

Systematic palaeontology

Pterosauria Kaup, 1834
Pterodactyloidea Plieninger, 1901
Azhdarchoidea Nesov, 1984 (sensu Unwin 2003)
Tapejaridae Kellner, 1989
Thalassodrominae Kellner and Campos, 2007

Genus Kariridraco gen. et sp. nov.

Zoobank LSID: urn:lsid:zoobank.org:act:77359171-69E4-4C18-A52A-285F1CF4294D

Etymology: After the Kariris, former indigenous people that lived in the area of the Araripe Plateau, combined with the Latin draco, dragon.

Type species: Kariridraco dianae gen. et sp. nov., see below.

Diagnosis.—As for the type species, by monotypy.

Kariridraco dianae gen. et sp. nov.

Zoobank LSID: urn:lsid:zoobank.org:act:D311DC9D-B9ED-4D0A-BAC1-6DC36584B9D5

Etymology: In reference to Diana Prince, alter ego of the DC fictional character Wonder Woman.

Holotype: MPSC R 1056 consists of a fairly complete skull, preserving most of its temporal, orbital, and preorbital portions, part of its premaxillary crest, an almost complete lower jaw and four cervical vertebrae (Fig. 1). Mechanical preparation revealed that the original concretion bearing the fossil was a composite, in which the rostral ends of the premaxillae and dentaries of a second pterosaur specimen was glued to the holotype. In order to avoid misinterpretations, we choose to only illustrate those elements that safely belong to a single individual. A photograph of the composite material is depicted in the SOM: fig. 1.

Type locality: The exact location where MPSC R 1056 was found is, however, unknown, since the specimen was donated to MPSC by local workers. The Romualdo Formation is geographically restricted to the Araripe Plateau, Northeastern Brazil.

Type horizon: Romualdo Formation, Aaptian, Lower Cretaceous.

Diagnosis.—Kariridraco dianae gen. et sp. nov. differs from all other pterosaurs for the following combination of characters (characters marked with * represent autapomorphies): presence of a subtle jugal shelf; comparatively short and tall nasoantorbital fenestra; premaxillary crest forming an angle of about 45° with respect to the maxilla*; lacrimal with an anteriorly-directed deep concavity*, and a single elliptic lateral pneumatic foramen in the left lateral surface of cervical IV centrum.

Description.—The holotype (MPSC R 1056) consists of a partially preserved skull, including the premaxilla, maxilla, premaxillary crest, part of the orbital and temporal regions (jugal, lacrimal, quadratoquadraticus), in addition to an almost complete lower jaw. Associate postcranial elements include the atlas-axis complex and two mid-cervicals. The specimen lacks most of the rostrum anterior to the nasoantorbital fenestra, as well as the premaxillary crest above the temporal region and part of its skull roof.

MPSC R 1056 presents the typical Romualdo Formation preservation, lacking evidence of lateral distortion. The lower jaw has a preserved length of 205 mm and is still in close association to the skull. Both upper and lower jaws are edentulous, a typical feature within Azhdarchoidea (Unwin 2003; Kellner 2004; Witton 2007; Kellner et al. 2019). The most striking features of the new specimen are the exceptionally tall nasoantorbital fenestra and the well-developed, steep premaxillary crest, this latter having its outermost bony layer eroded and exposing trabecular bone. The orbit is piriform, with a subvertical anterior margin limited by the lacrimal ramus of the jugal and the lacrimal itself. The dorsoposterior margin of the orbits, which would be formed by frontoparietales, are absent. In close association to the orbits, K. dianae gen. et sp. nov. preserves a slit-like lower temporal fenestra. Most of the occiput was lost, but a wide occipital plate is visible posterior to the lower temporal fenestra, being detached from its original position. Cervical vertebrae were preserved articulated, still attached to the skull, but were posteriorly isolated through mechanical preparation.

The unusually steep premaxillary crest and the fact that the rostrum anterior to this structure was adulterated by fossil dealers led us to raise concerns about the crest authenticity. Careful examination of the fossil and surrounding matrix revealed that, although there are signs of a breakage close to the base of the crest (where it contacts the main body of the premaxillomaxilla), the two resulting counterparts fit together, with no signs of add-ons or other modifications. Bone surface at the base of the crest was eroded off, so that trabecular bone is exposed at that portion. Additionally, the carbonate concretion remained uniform at the surroundings of the breakage, differing, for instance, from the composite rostrum (Fig. 1, SOM: fig. 1), in which the anterior portion of the upper and lower jaws of a second individual was glued together with MPSC R 1056. This second specimen was preserved in a different, much lighter in color, carbonate matrix. Crest base preparation removed uniform rock throughout the whole procedure, while mechanical removal of matrix in composite specimens fatally menages to reveal add-ons (usually epoxy resin) used to glue pieces together. It
is also noteworthy that, close to the anteroventral border of the nasoantorbital fenestra, the premaxillomaxilla thickens to delineate the rounded outline of the aperture, showing that, MPSC R 1056 indeed had an unusually short nasoantorbital fenestra. This dorsal expansion of the jugal process of the premaxillomaxilla is compatible with the crest's position, steepness and shape.

*Premaxillomaxilla*: The most striking feature of the fused premaxilla/maxilla is the well-developed sagittal crest. As preserved, this structure has a total length of 202 mm. The crest is broken posterodorsally, so that it is not possible to estimate its total original length. Parts of the crest are still embedded in carbonate, as its delicate nature prevented total preparation without further damaging, even using standard techniques for Santana crested pterosaurs (Silva and Kellner 2006). The posteroventral margin of the crest is notably straight and forms the preserved anterior borders of the nasoantorbital fenestrae (Fig. 1). The exact spot where the premaxillary crest would contact the frontoparietals is impossible to estimate. Close to its base, the crest widens slightly, so that the anteroventral end of this structure is deeper than the posterodorsal one.

The anterior-most portion of the premaxillomaxillae is not preserved. The premaxillomaxillae anteriorly border the large and wide nasoantorbital fenestrae, the anterior limit of which is unusually steep (about 45° with respect to the horizontal plane) (Fig. 1A2). Most of the anterodorsal and ventral borders of the nasoantorbital fenestrae are also limited by the premaxillomaxillae. These latter contact the jugals posteroventrally, even though the precise suture between both bones cannot be determined with confidence. Ventrally, the premaxillomaxillae bend medially to form broad palatal plates anterior to the choanae (see Ősi et al. 2010; Pinheiro and Schultz 2012). Incomplete preservation...
prevents the recognition of the entire extent of the premaxillomaxillae, as these bones display an anteroventrally-oriented breakage close to the anterior border of the nasoantorbital opening. Even though the specimen is visible only in the right lateral view, it is clear that the palatal surface of the maxillary palatal plate is strongly convex.

**Jugal:** Despite being difficult to recognize suture lines with the surrounding bones, the right jugal is apparently triradiate, with three well-developed processes (Fig. 1A₂). The jugal is well-preserved (except for a fracture later restored), not showing signs of erosion. Anteriorly, a maxillary process apparently contacts the maxilla close to the ventral midpoint of the nasoantorbital fenestra. The dorsally-oriented lacrimal process detaches itself from the main body of the jugal by the presence of a medial recess, ventrally bordered by a moderately-developed jugal shelf. The lacrimal process very thin and is positioned at an almost right angle with respect to the maxillary process. Posterdorsally, the jugal tapers to form an acute process that makes most of the preserved dorsoventral orbital margin.

**Lacrimal:** The lacrimal is exposed in its right lateral view, being its medial surface still embedded in matrix. The tridimensional conformation of this bone was not altered by preservation. The ventral limit of the lacrimal is positioned at the mid orbital height, and this bone broadens dorssally to make the posterocephalal margin of the nasoantorbital opening. Albeit laterally eroded, the anterior margin of the lacrimal, which forms the ventral margin of the neural canal, is not surrounded by pneumatic foramina anteriorly (Fig. 2A₁, A₂). In posterior view, however, the neural canal is laterally flanked by two small elliptical pneumatic foramina, one in each side (Fig. 2A₂, A₅). In both its left and right lateral surfaces, the atlas-axis complex displays a small rounded pneumatic foramen located in the middle portion of the axial centrum (Fig. 2A₆).

**Quadratojugal:** The quadratojugal is fused to the main corpus of the jugal, forming a thin rod that limits posteroventrally the lower temporal fenestra.

**Quadrate:** The right quadrate is preserved as a thin bone rod that contributes to the posteroventral margin of the lower temporal fenestra, having contact with the squamosal and quadratojugal. Quadrate inclination with respect to the horizontal plane is of about 130°.

**Squamosal:** Only a fragment of the right squamosal is preserved, forming a thin process that reaches the middle posterior margin of the orbit. Together with the quadrate and quadratojugal the squamosal also composes the posterior margin of the lower temporal fenestra.

**Palate:** The palatal surface of *Kariridraco dianae* gen. et sp. nov. is preserved, except for the anterior most portion and the articular portion for the lower jaw. Only the right pterygoid is visible, being still embedded in matrix and not completely preserved. It connects with the maxilla in its anterior portion, where a fibrous joint marks the limits between both bones (Fig. 1). A thin bone rod adjacent to the pterygoid may be a lateral process to the pterygoid main body. The pterygoid has no preserved connection with the ectopterygoid or jugal. A small portion of the ectopterygoid is also preserved, still attached to the jugal. The proximal portion of this bone rod is, however, still covered by matrix, obscuring the connection between the former and the pterygoid.

**Lower jaw:** The lower jaw is almost complete and still occluded to the skull. It only lacks the rostral-most portion (anterior to the nasoantorbital fenestra). Bone fusion, however, precludes the exact identification of the posterior mandibular bones. Posterior to the jaw junction, a ventrally-sloping retroarticular process is visible. Albeit displaying a fair preservation, the lower jaw displays a restored fracture spot in its mid-length. There is no evidence of a dentary crest in its preserved portion, but this structure could have been present in the missing rostral part. The lower jaw appears to be completely edentulous (Fig. 1).

**Atlas-axis:** The atlas-axis complex is completely fused, what is usually displayed by adult individuals (e.g., Bennett 1993; Kellner 2015; Fig. 2). The atlas is fused to the axis by its annular components, located at the base of the vertebrae. Above the annular components is the round-shaped neural canal, which is not surrounded by pneumatic foramina anteriorly (Fig. 2A₁, A₂). In posterior view, however, the neural canal is laterally flanked by two small elliptical pneumatic foramina, one in each side (Fig. 2A₂, A₅). In both its left and right lateral surfaces, the atlas-axis complex displays a small rounded pneumatic foramen located in the middle portion of the axial centrum (Fig. 2A₆). The axis presents a well-developed, almost complete neural spine, displaying some erosion in its dorsal and anterior portions (Fig. 2, 3). It binds the well-developed postzygapophyses, which form an angle of 135° between each other. In lateral view, the neural spine forms an angle of about 10° with the vertical plane close to its base, bending backwards to form an angle of 40° in its superior portion. Posteriorly, the axis has well-developed postzygapophyses. The condyle is located dorsal to the postzygapophyses. It is elliptical in shape, with its transverse length much larger than its height. It is dorsally followed by a concavity that forms the ventral margin of the neural canal. There is a slight slope lateral to this concavity, and the ventral margin of the centrum between the postzygapophyses is convex (Fig. 3A₁, A₅).

**Vertebrae III and IV:** The third cervical vertebra has a total length of 45.79 mm and a maximum width of 44.13 mm. It displays a good preservation, except for its left ventral surface, which shows a circular erosion and slight fragmentation. The vertebra has a subcircular neural canal, flanked by two lateral pneumatic foramina both in its anterior and posterior portions (Fig. 4).

Cervical III centrum apparently lacks lateral pneumatic foramina (Fig. 4A₁, A₃). While its left surface is incomplete, the right one is well-preserved and not pierced by any aperture. The neural spine of the third vertebra is incomplete but was apparently moderately tall. It expands laterally in its base to form the pre- and postzygapophyses (Fig. 5A₁, A₅). These latter are dorsally limited by dorsomedially directed laminae, which connects the tip of the zygapophyses to the
neural spine. Albeit better preserved, the prezygapophyses are less developed than the postzygapophyses.

The hypapophysis is a small bone protuberance positioned ventrally to the cotyle, while the postexapophyses are well developed and almost complete (Fig. 5). Their ventral portions show slight slopes, also observable in lateral view. Such slopes, however, do not interfere with the overall flat ventral surface of the centrum. The condyle is dorsal to the postexapophyses. Its dorsal portion has a concavity that connects to the neural canal, forming its ventral
margin. Its dorsal portion presents slight lateral slopes on both sides. The condyle is directed horizontally with relation to the main axis of the vertebra. The neural canal is flanked by two highly asymmetric pneumatic foramina (Fig. 4A2, A5).

Cervical IV is poorly preserved. It has its dorsal surface almost completely eroded. The neural canal, as well its adjacent structures, is absent. This is also the case for the neural spine, condyle, postzygapophyses and postexapophyses (Fig. 6). There is a small pneumatic foramen on the ventral left surface of the centrum, close to where the left postzygapophysis would originate (Fig. 6A3, A6). The vertebra IV has a slight concavity in its ventral surface (Fig. 6A3, A6), which is more prominent than what is displayed by cervical III. The hypapophysis is fragmented, but reveals to be better developed when compared to what is observed in
cervical III. The right prezygapophysis is better preserved than the left one, which lacks its anterior portion. The right prezygapophysis presents its original conformation and is well pronounced (Fig. 7).

Stratigraphic and geographic range.—Type locality and horizon only.

Phylogenetic results

Our phylogenetic assessment of *Kariridraco dianae* gen. et sp. nov. using the modified version of Holgado et al. (2019) dataset resulted in 6 most parsimonious trees (MPTs) of 345 steps each. The strict consensus tree (Fig. 8) recovers a monophyletic Tapejariidae (sensu Kellner and Campos 2007). *Kariridraco dianae* gen. et sp. nov. is recovered as the sister taxon of *Tupuxuara* (*T. leonardi* + *T. longicristatus*). *Thalassodromus sethi* Kellner and Campos, 2002 stands as the sister taxon to (*Kariridraco + Tupuxuara*) and, as such, Thalassodrominae was recovered as monophyletic. The Tapejarinae was also recovered and, within it, *Caupedactylus ybaka* Kellner, 2013, and *Aymeredactylus cearensis* Pégas, Leal, and Kellner, 2016, are successive sister taxa to a clade formed by *Eopteranodon lii* Lü and Zhang, 2005, “Huaxiapterus” benxiensis Lü, Gao, Xing, Li, and Ji 2007, “Huaxiapterus” corollatus Lü, Jin, Unwin, Zhao, Azuma, and Ji, 2006, *Sinopterus dongi* Wang and Zhou, 2003, *Tapejara wellnhoferi* Kellner, 1989, *Europejara ocladesorum* Vullo, Marugán-Lobón, Kellner, Buscalioni, Gomez, Fuente, and Moratalla, 2012, *Caiuajara dobruskii* Manzig, Kellner, Weinschütz, Fragoso, Vega, Guimarães, Godoy, Liccardo, Ricetti, and Moura 2014, and *Tupandactylus imperator* Campos and Kellner, 1997. Notably, Chinese tapejarines form a monophyletic group, whereas Brazilian representatives of this clade are dispersed in the topology, being either recovered as early-diverging tapejarines (*Caupedactylus ybaka* and *Aymeredactylus cearensis*) or forming a clade together with the Laurasian *Europejara*. The clade formed by *Kariridraco* and *Tupuxuara* species is held by a single unambiguous synapomorphy (Char. 146, palatal surface convex at mid-length). Our second analysis, including *Keresdrakon vilsoni* resulted in 15 equally parsimonious trees of 348 steps. Although the consensus topology dis-
plays Keresdrakon as the sister-taxon to the Tapejaridae (in agreement with Kellner et al. 2019), tapejarid taxa relationships are mainly unresolved. Only the Thalassodrominae (Tupuxuara spp. + Thalassodromeus sethi + Kariridraco dianae gen. et sp. nov.) was recovered as a clade, being placed in a polytomy together with Aymberedactylus cearensis, Caupedactylus ybaka and a clade formed by the remaining tapejarines (Tapejara wellnhoferi, Europejara olcadesorum, Caiuajara dobruskii, Tupandactylus imperator, Sinopterus dongi, “Huaxiapterus” spp., and Eopteranodon lli) (SOM: fig. 2).

Similarity indexes for tapejarid-bearing pterosaur faunas

Our results demonstrate a high similarity index between Wessex and La Huérguina formations (England and Spain, respectively), even though pterosaur specimens are comparatively rare in both units (Fig. 9B). The resulted dendrogram also evidences that the pair formed by La Huérguina and Wessex formations is more akin to the other two Laurasian units sampled here (Yixian and Jiufotang formations) than to the Gondwanan Crato, Romualdo, Goiô-Erê formations (Brazil) and the Kem Kem beds (Morocco). On the other hand, the Kem Kem beds and Crato Formation cluster with the Romualdo Formation, being a higher similarity detected between Crato Formation and the Kem Kem beds. The recovered position of Goiô-Erê Formation may be a sampling bias, as only two pterosaur taxa were so far recovered from this unit.

Discussion

Comparisons.—As preserved, Kariridraco dianae gen. et sp. nov. jaws are entirely edentulous, and empty alveoli are absent. Among Romualdo Formation pterosaurs, the lack of teeth is only displayed by the Tapejaridae (e.g., Kellner 2004). A different condition is seen in the well-represented anhanguerians (Rodrigues and Kellner 2013). Although edentulous chaoyangopterid pterosaurs have been previously reported for the Araripe Basin, they are so far restricted to the lower Crato Formation. The most informative putative chaoyangopterid thus far reported for the Crato beds is Lacusovagus magnificens (Witton 2008), which is represented by a partially complete skull, preserving its anterior rostrum (SMNK PAL 4325). Kariridraco dianae gen. et sp. nov. clearly differs from L. magnificens.
by the presence of a well-developed premaxillary crest. Comparisons between palatal bones are impossible because *L. magnificens* holotype (SMNK PAL 4325) is still partially embedded by matrix. More recently, Leal et al. (2018) described mid-cervical vertebrae (IV–VII) putatively attributable to *L. magnificens* (UFC-721). Direct comparisons between UFC-721 and *K. dianae* gen. et sp. nov. are also not possible because the latter preserves only the first four vertebrae. We should note, however, that *K. dianae* gen. et sp. nov. cervicals display well-developed neural spines, whereas
Leal et al. (2018) describe low neural spines for UFC-721. Also, Leal et al. (2018) report the absence of lateral pneumatic foramina and presence of pronounced postexapophyses. Albeit *K. dianae* gen. et sp. nov. also displays well-developed postexapophyses, lateral pneumatic foramina are present in cervical IV. UFC-721 also differs from *K. dianae* gen. et sp. nov. in having the typical centrum elongation of chaoyangopterids (Zhou 2010; Leal et al. 2018), azhdarchids (Suberbiola et al. 2003; Ösi et al. 2005; Vremir et al. 2013) and ctenochasmatids (Andres and Ji 2008).

*Kariridraco dianae* gen. et sp. nov. displays typical tapejarid features (Fig. 10), such as a nasoantorbital fenestra corresponding to more than 45% of the total skull size (estimated); orbit showing an inverted pear shape; long sagittal crest formed mainly by the premaxilla and extending posteriorly, and a thin subvertical lacrimal process of jugal (Kellner 2004; Kellner and Campos 2007). The monophyly of Tapejaridae was questioned by some authors, such as Unwin and Lü (1997) and Unwin (2003), who agree in that *Tapejaridae sensu* Kellner (2003) would be paraphyletic with respect to *Azhdarchidae*. According to those, *Tupuxuara and Thalassodromeus* would share a series of characters in common with azhdarchids, while absent in *Tapejara*-like taxa. This hypothesis was later supported by, for instance, Martill and Naish (2006) and Lü et al. (2006). Pinheiro et al. (2011), however, recovered Tapejaridae sensu Kellner and Campos (2007) as monophyletic, in agreement with Kellner (2004) and other previous works (Wang et al. 2005, 2008, 2009; Andres and Ji 2008).

Kellner and Campos (2007) divide the Tapejaridae in two independent clades with sister-taxon relationship. The Tapejarinae would include those tapejarids with a downturned rostrum, premaxillary crest constricted above the nasoantorbital fenestra, low posterior expansion of the cranial crest and orbits positioned on a level only slight below the dorsal margin of the nasoantorbital fenestra (Kellner and Campos 2007; Pinheiro et al. 2011; Fig. 10B). Tapejarinae includes *Tapejara wellnhoferi, Tupandactylus imperator, Tupandactylus navigans* Frey, Martill, and Buchy 2003, *Sinopterus dongi*, “Huaxiapterus” *jii* Lü and Yuan, 2005, “Huaxiapterus” *corollatus*, “Huaxiapterus” *benxiensis*, *Eopteranodon lii*, *Europejara olcadesorum*, *Caudipteryx ybaka*, *Caiuajara dobruskii*, *Aymberedactylus cearensis*, *Sinopterus lingyuuanensis* Lü, Teng, Sun, Shen, Li, Gao, and Liu, 2016, *Sinopterus atavismus* (Lü, Teng, Sun, Shen, Li, Gao, and Liu, 2016), and *Wightia declivirostris* Martill, Green, Smith, Jacobs, and Winch, 2020. By their turn, the Thalassodromeinae are characterized by having very high nasoantorbital fenestrae and by a high premaxillary crest which is not constricted above the nasoantorbital openings.
such as to the Early Cretaceous of southern Gondwana (Brazil), such as Thalassodromeus sethi, Thalassodromeus oberlii (Headden and Campos, 2015), Tupuxuara longicristatus Kellner and Campos, 1988, Tupuxuara leonardi Kellner and Campos, 1994, and putatively “Tupuxuara” deliradamus. It is noteworthy that, albeit being consistently recovered as an early-diverging tapejarine, Caupedactylus ybaka shares with thalassodromines a high premaxillary crest that is unconstricted above the nasoantorbital openings.

With respect to the Tapejarinae, Kariridraco dianae gen. et sp. nov. differs from Tapejara wellnhoferi by several features, such as the triangular shape of the nasoantorbital fenestra (sub-oval in Tapejara wellnhoferi, based in the specimen AMNH 24440) and the overall morphology of the sagittal crest (Kellner 1989; Wellnhofer and Kellner 1991). It also differs from Tupandactylus by the triangular shape of its nasoantorbital fenestra, the presence of a jugal shelf and an angle of 150° between the quadrate with respect to the jugal bar. Tupandactylus has an elliptical nasoantorbital fenestra, apparently lacks a jugal shelf and has an anteriorly high sagittal crest that constricts itself above the nasoantorbital openings (based on specimens MCT 1622-R and CPCA 3590) (Campos and Kellner 1997; Pinheiro et al. 2011). Kariridraco dianae gen. et sp. nov. resembles Caupedactylus ybaka in having a steep anterior margin of the nasoantorbital fenestra (steeper in K. dianae gen. et sp. nov.); orbits positioned low in the skull; an unconstricted premaxillary crest (the last two characters being also displayed by thalassodromines); and a jugal shelf (better developed in Caupedactylus ybaka). However, K. dianae gen. et sp. nov. is easily distinguishable from C. ybaka for having a much lower premaxillary crest, a convex palate below the nasoantorbital fenestrae (mainly flat in C. ybaka), a more vertical lacrimal process of the jugal and a much less developed jugal shelf.

As for the Thalassodrominae, Kariridraco dianae gen. et sp. nov. shares with Thalassodromeus sethi (holotype DGM 1476 R), a somewhat triangular nasoantorbital fenestra, a 90° angle between the lacrimal and the maxillary processes of the jugal and a jugal shelf. Despite that, the nasoantorbital fenestra is higher in K. dianae gen. et sp. nov. than what is displayed by T. sethi (Pégas et al. 2018), having also a much steeper anterior margin (about 45° with respect to the horizontal plane in its anterior portion). Additional differences between the new taxon and T. sethi include its palatal morphology, strongly concave posteriorly in T. sethi (Kellner and Campos 2007; Pégas et al. 2018) and distinctly convex in K. dianae gen. et sp. nov. We do not have any elements to directly compare K. dianae gen. et sp. nov. with Thalassodromeus oberlii, as the latter is only known by its holotype (NMSG SAO 251093), a mandibular symphysis. We, however, agree with Pégas et al. (2018) in that NMSG SAO 251093 can be confidently attributed to a species of Thalassodromeus and, thus, probably came from a very distinct animal with respect to K. dianae gen. et sp. nov.

The near triangular-shaped nasoantorbital fenestra, a 90° angle of the lacrimal process with respect to the jugal bar and a convex palate are features shared between Kariridraco dianae gen. et sp. nov. and Tupuxuara leonardi (IMCF 1052). They differ, however, by the lack of the jugal-shelf in this later, as well as much shorter nasoantorbital openings and a steeper sagittal crest in K. dianae. Although known only by a fragmentary skull (holotype MN-6591-V), Tupuxuara longicristatus differs from K. dianae gen. et sp. nov. in having a much less steep premaxillary crest and apparently longer nasoantorbital openings. In addition, the dorsal edge of T. longicristatus premaxillary crest is slightly convex, differing from both K. dianae gen. et sp. nov. and T. leonardi. “Tupuxuara” deliradamus (SMNK PAL 6410) has an angle of 150° between quadrate and the maxillary process of the jugal, which is similar to the condition displayed by Caupedactylus ybaka (MN 4726-V, Kellner, 2013, Fig. 10B), differing from K. dianae gen. et sp. nov. Further comparisons between K. dianae gen. et sp. nov. and “Tupuxuara” deliradamus are hindered by the poor preservation of the holotype of the latter (SMNK PAL 6410), precluding the estimation of nasoantorbital fenestrations and shape and size (Witton 2009). We should note that “Tupuxuara” deliradamus is a highly problematic taxon, the validity of which will be assessed in a future contribution.

The total preserved cervical vertebrae complex (from atlas-axis to cervical IV) of K. dianae gen. et sp. nov. has a set of features only displayed by Thalassodrominae, such as well pronounced postexapophyses, reduced hypapophysis, and pronounced, hatchet-like, neural spines (Aires et al. 2014; Vila Nova et al. 2015). The presence of two foramina adjacent to the neural canal and the flat ventral surface of the vertebrae is also peculiar to thalassodromines (Vila Nova et al. 2015). Regarding the atlas-axis complex, K. dianae gen. et sp. nov. is also similar to taxa of Thalassodrominae in having a flat ventral surface and two foramina adjacent to the neural canal. However, the atlas-axis shows a small single lateral foramen on both sides of the element, a feature previously observed in Tapejara wellnhoferi (SMNK PAL 1135; Eck et al. 2011). The presence of adjacent foramina in the fourth vertebra is impossible to confirm due to its poor preservation. Eck et al. (2011) mentions the presence of a single foramen on either side of the centrum of Tapejara wellnhoferi cervicals. The presence of a single foramen in both sides of K. dianae gen. et sp. nov. cervical IV may result from individual variation.

Biogeography of Tapejaridae.—The paleobiogeography of pterosaurs is an underappreciated topic, and geographic distribution patterns of tapejarids have been accessed solely by seldom and punctual discussions (e.g., Vullo et al. 2012; Pégas et al. 2016; Martill et al. 2020a). An attempt was proposed by Sayão et al. (2012) when presenting a comparison of the distribution patterns of pterosaurs in light of the phylogenies existing at the time. It was concluded that the phylogenies available were based on taxa from a few lo-
cations, specially from Fossillagerstattën, which prevented the application of cladistic and event-based biogeographic methods (Sayão et al. 2012).

Previous works sometimes relied on ambiguous evidence to determine origin centers for the Tapejaridae. As an example, a Eurasian origin was proposed for this clade based on the recovery of slightly older taxa in Europe and China, when compared to South American representatives (Vullo et al. 2012). The sole occurrence of older remains of a specific taxon is, however, well-established as a fragile criterion to determine origin centers (Cain 1944), especially when dealing with taxa that are notorious for their patchy and biased fossil record, as is the case of pterosaurs (Sayão et al. 2012; Butler et al. 2013).

The results of our faunistic similarity assessment of tapejarid-bearing units evidence a clear large-scale provincialism between Gondwanan and Laurasian pterosaur communities (Fig. 9A, B). This was already briefly argued for tapejarines by Martill et al. (2020a) but note that our results are based in all the taxa recovered from each sampled locality. Also, similarity indexes respect geographic distance in Laurasian faunas, with La Huérguina and Wessex formations having a much higher average similarity when compared to the Chinese Yixian and Jiufotang formations. The similarity between pterosaur communities from Northeastern Brazil and Morocco agrees with the results of Sales et al. (2018), who detected a similar pattern for dinosaurian faunas. We should note, however, that the recovered similarity may (or may not) be confirmed by future works, since comparisons between Brazilian and Moroccan pterosaur faunas are hindered by the still incipient pterosaur discoveries in Morocco (Kellner and Mader 1997; Mader and Kellner 1999; Jacobs et al. 2020). In spite of that, our results suggest that, as occurred for strictly terrestrial faunas, pterosaur communities were also affected by large scale tectonic-driven vicariant events.

Although pterosaurs were probably more capable to overcome geographical barriers when compared with terrestrial, non-flying, animals (Sayão et al. 2012), it should also be taken into account that smaller-scale phenomena such as high rates of oceanic spread, volcanism, excessive atmospheric CO₂ heating the surface temperature (Weissert et al. 2003) possibly also contributed to the framework that would explain pterosaur distribution patterns.

When analyzed together, tapejarid phylogeny is in perfect agreement with their geographic distribution. The earliest-diverging clade (Thalassodrominae) is restricted to...
Gondwanan deposits, whereas the specialized tapejarines are widely spread throughout Gondwana and Laurasia, even though they seem to respect smaller scale provinces, as is suggested by the monophyletic Chinese tapejarines (Fig. 9C). The apparent restriction of early-diverging forms to Gondwana, allied to the fact that the sister-taxon to the Tapejaridae (*Keresdrakon vilsoni*) was recovered from Southern Brazil, suggests a Gondwanan origin for the clade, what is in disagreement with the proposition of Vullo et al. (2012). We should note that similar conclusions as we propose here have already been suggested by Pêgas et al. (2016) who, based on a similar topology as we recovered, defied Vullo et al. (2012) Eurasian origin for the Tapejaridae. The presence of slightly older tapejarid remains in Europe and China (Vullo et al. 2012; Martill et al. 2020a) can be readily explained by the strong “Lagerstätten effect” that affects pterosaur fossil record, resulting in severe sampling biases that can obscure large-scale diversity patterns (Butler et al. 2009, 2013; Sayão et al. 2012).

Conclusions

*Kariridraco dianae* gen. et sp. nov. (Fig. 11) has a set of diagnostic features that distinguishes it from other azhdarchoid pterosaurs, such as the presence of a concavity in the upper portion of the lacrimal and an angle of about 45° between the maxilla and the premaxillary crest. *Kariridraco dianae* gen. et sp. nov. was consistently recovered as a thalassodromine tapejarid, closely related to the well-known genus *Tupuxuara*. Our assessment of tapejarid paleobiography recovered Gondwana as the most parsimonious origin center for the Tapejaridae, also suggesting that, similarly to strictly terrestrial animals, pterosaur distribution is marked by tectonic-driven provincialism.

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References


