Dipnoan from the Upper Triassic of East Greenland and remarks about palaeobiogeography of Ptychoceratodus

WOJCIECH PAWLAK, MATEUSZ TAŁANDA, TOMASZ SULEJ, and GRZEGORZ NIEDŹWIEDZKI


Here we present a description of the dipnoan remains collected from the middle to upper Norian (Upper Triassic) of Jameson Land, East Greenland. The specimens consist of isolated tooth plates and skull bones of Ptychoceratodus, the most complete Late Triassic dipnoan material from Greenland. This genus is reported for the first time from the Upper Triassic of Greenland. The studied material belongs to Ptychoceratodus rectangulus previously known from the middle–upper Norian of Germany. It fills the biogeographical gap between the records of the Germanic and the Jameson Land basins. A reconstruction of the skull roof is provided, based on isolated bones collected from the same bone-bed. Their good preservation enables recognition of the sensory line pits, arranged similarly as in the extant Protopterus, suggesting a comparable mode of life. This finding has implications for our understanding of the disparity in Ptychoceratodus dipnoans, as well as the morphology between closely related dipnoans of the Late Triassic ecosystems.

Key words: Dipnoi, Ptychoceratodus, Triassic, Norian, Greenland, Carlsberg Fjord Beds.

Wojciech Pawlak [wojciech.pawlak@student.uw.edu.pl] and Mateusz Talanda [m.talanda@biol.uw.edu.pl], Department of Palaeobiology and Evolution, Faculty of Biology, Biological and Chemical Research Centre, University of Warsaw, Zwikir i Wigury 101, 02-089 Warsaw, Poland.

Tomasz Sulej [sulej@twarda.pan.pl], Institute of Paleobiology, Polish Academy of Sciences, Twarda 31/55, 00-818 Warsaw, Poland.

Grzegorz Niedźwiedzki [grzegorz.niedzwiedzki@ebc.uu.s], Department of Organismal Biology, Evolutionary Biology Center, Uppsala University, Norbyvägen 18A, 752 36 Uppsala, Sweden.

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Introduction

The study of Late Triassic dipnoans (lungfish) has recently undergone renewed interest, improving our understanding of the taxonomy, diversity and palaeobiogeography of this important component of Triassic freshwater and marginal marine ecosystems (e.g., Krzyzcyki 2015, 2016; Agnolin et al. 2017, 2018; Krzyzcyki et al. 2018; Bhat and Ray 2020). The dipnoans are adapted to breathing with both gills and lungs (Wright 1974; Holmes 2008). They are characterized by crushing dental plates instead of true teeth. Despite this specialization to durophagy, the extant lungfishes are omnivorous (Kemp 1986; Greenwood 1986). Dipnoans are an old sarcopterygian fish lineage known from the Early Devonian to recent. Their massive and resistant to erosion dental plates are common remains in the Triassic fossil record.

The fossil record of the post-Devonian lungfishes is biased in that bone elements from cranial or postcranial skeleton are relatively rare, mainly preserved as isolated or fragmented elements, and the most common remains of these fishes are their dental plates. Only a few post-Palaeozoic species are known from well preserved cranial or postcranial skeletal remains (Kemp 1998). This residual fossil record gives remarkably incomplete knowledge about lungfish palaeoecology and evolution. Consequently, previous phylogenetic analyses (Miles 1977; Marshall 1986b; Cavin et al. 2007; Kemp 2017) failed to offer a robust evolutionary scenario explaining the major changes within the post-Palaeozoic dipnoans. The Triassic was a golden age for dipnoans, at least when it comes to the number of genera and species (Martin 1982; Marshall 1986a; Schultze 2004). Such a diversity is probably related to a wide global distribution of warm and semi-dry continental climate probably preferred by these fishes, similarly to their Recent representatives. Comprehension
of the Triassic evolutionary radiation of dipnoans is crucial for the understanding of their later Mesozoic and younger evolutionary history. The Triassic dipnoans are known from freshwater and marginal marine (brackish) environments. There are two genera, which are particularly common nearly worldwide: *Arganodus* and *Ptychoceratodus* (Skrzycki et al. 2018). Only two species of *Ptychoceratodus* are known from a relatively complete cranial skeleton: *Ptychoceratodus serratus* from the late Ladinian Erfurt Formation in Germany (Schultz 1981) and *Ptychoceratodus philippi* from the Cynognathus Zone of the Bregersdorp Formation in the Karoo Basin, dated as late Early Triassic–early Middle Triassic (Kemp 1996).

In East Greenland, fossiliferous Upper Triassic of Jameson Land Basin, such as the Malmros Klint Member or Ørsted Dal Member, are known for rich faunal assemblages (Jenkins et al. 1994; Clemmensen et al. 1998, 2016; Sulej et al. 2014). However, the remains of dipnoans are still poorly known from the Late Triassic record of Greenland. The oldest record of lungfishes from Greenland comes from the Middle and Late Devonian (Lehman 1959; Bendix-Almgreen 1976; Blom et al. 2007; Clack et al. 2018). In younger Carboniferous or Permian deposits of Greenland, dipnoan remains are unknown, but some remains have been collected from the Upper Triassic Fleming Fjord Formation (Jenkins et al. 1994; Clemmensen et al. 1998, 2016). Recently, dental plates and jaw bones belonging to a new species of *Ceratodus* (*C. tunuensis*) have been described from the upper Carlsberg Fjord Beds (Agnolin et al. 2018).

In this paper we present a detailed description of the newly collected ptychoceratodontid skull material and highlight its palaeoecological and palaeobiogeographic significances. We compare it with *Ceratodus tunuensis* reported from the same lithostratigraphical unit (Agnolin et al. 2018). The new material represents fossils which are much better preserved than the other dipnoan remains collected so far from the Ørsted Dal Member. For this reason, the study has required a brief re-study of the isolated elements described by Agnolin et al. (2018) and a comparison with other previously described ptychoceratodontid fossils. The Fleming Fjord Formation also yielded some aestivation burrows, which carry information about the Late Triassic climate of north Pangea and broadens our knowledge about the distribution of the aestivation behavior among the early Mesozoic dipnoans. These trace fossils will be a subject of an additional study.

**Institutional abbreviations.** — BRSUG, University of Bristol, Department of Geology, UK; KNK, Greenland National Museum & Archives, Nuuk, Greenland; NHMD, Natural History Museum of Denmark, Copenhagen, Denmark; SMNS, Staatliche Museum für Naturkunde, Stuttgart, Germany.

**Other abbreviations.** — BC1, distance between the mediolingual joint and the tip of the first ridge; CIC2, distance between the tips of the first and second ridges; BCp, the distance between mediolingual joint and the tip of the last ridge.

**Geological setting**

The Upper Triassic succession of the Jameson Land is composed of about 200–300 m thick continental sediments of the Fleming Fjord Formation (Clemmensen et al. 1998) represented mainly by red or greenish mudstone, siltstone, and sandstone beds. The formation is divided into three members: the lowermost Edderdugledal Member, the middle Malmros Klint Member and the uppermost Ørsted Dal Member (Fig. 1). Integrated magnetostratigraphical, cyclostratigraphical, and some biostratigraphical studies revealed that the Fleming Fjord Formation age is late Carnian–early Rhaetian (Clemmensen et al. 1998), Norian–Rhaetian (Clemmensen et al. 2016), or Norian–early Rhaetian (Andrews et al. 2014). Very rare invertebrate body fossils (e.g., conchostracans, bivalves), composition of the tetrapod faunas/ichnofaunas, and land-derived palynomorphs suggest that the Malmros Klint Member is of late Carnian–middle Norian age, and that the Ørsted Dal Member is of middle Norian–early Rhaetian age (Clemmensen 1980; Clemmensen et al. 1998, 2016; Sulej et al. 2014). A conspicuous feature of these deposits are sedimentation cycles reflecting climate changes, which are developed slightly differentially in every lithostratigraphical subunit (Clemmensen et al. 1998).

The fossiliferous lower part of the Ørsted Dal Member, named the Carlsberg Fjord Beds, is composed of intraformational conglomerate, massive mudstone, mudstone with wave ripple cross-lamination, and sandstone with cross-bedding. Desiccation cracks are common in this interval within both the mudstone and the sandstone beds but there are no signs of pedogenesis (Clemmensen et al. 1998). The Ørsted Dal Member contains also two other lithostratigraphic subunits. In the western and central part of the Basin, the Carlsberg Fjord Beds are replaced by fluvial sediments of the Bjergronerne Beds, and both units are overlaid by carbonate-rich mudstone, dolostone, and clay-rich mudstone of the Tait Bjerg Beds (Jenkins et al. 1994).

Both the Malmros Klint Member and Ørsted Dal Member are abundant in skeletal and trace fossils of terrestrial and aquatic vertebrates (Jenkins et al. 1994; Clemmensen et al. 1998; Sulej et al. 2014; Clemmensen et al. 2016). The existing outcrops indicate that the Fleming Fjord Lakeland had about 3000 km², therefore it was a large but predominantly shallow reservoir with frequent desiccation episodes (Clemmensen et al. 1998). The periodic shallow lake conditions favored bone accumulation, thus the Fleming Fjord Formation infill of the Jameson Land is an interesting taphonomical window of the Late Triassic fauna. These fossiliferous deposits have been searched for vertebrate remains since the late 1980s (e.g., Jenkins et al. 1994; Clemmensen et al. 1998, 2016; Shapiro and Jenkins 2001; Milan et al. 2004; Sulej et al. 2014). The vertebrate fauna of
the Fleming Fjord Formation is relatively abundant and quite diverse, and includes actinopterygian and dipnoan fishes, temnospondyls, sauropodomorph and theropod dinosaurs, basal turtles, phytosaurs, aetosaurs, pterosaurs, cynodonts, and mammaliaforms (Jenkins et al. 1994, 1997, 2001, 2008; Sulej et al. 2014; Clemmensen et al. 2016; Niedźwiedzki and Sulej 2017; Marzola et al. 2017; Agnolín et al. 2018). Numerous tetrapod tracks have been found in several horizons inside the Malmros Klint and Ørsted Dal members (e.g., Jenkins et al. 1994; Sulej et al. 2014; Klein et al. 2016; Niedźwiedzki and Sulej 2017; Marzola et al. 2017; Agnolín et al. 2018). Material and methods

The examined specimens were collected during the Polish-Danish expedition to Greenland in 2014 (Sulej et al. 2014). Eleven skeletal elements and one imprint were found, which belonged to different individuals of various sizes. All specimens are disarticulated but three-dimensionally preserved without any signs of compaction and additional deformations. Skull roof bones have a characteristic radial pattern of points and lines diverging from the ossification center. Specimens KNK 4154–4161, 4163–4165 were found in the locality 1

Fig. 1. Study area map (A) and stratigraphic column (B) with marked outcrops and age of the stratigraphic position of examined material. Based on Bengaard and Henriksen (1984) and Clemmensen et al. (1998).
(71.38283 N; 22.55716 W), and KNK 4162 in the locality 2 (71.37126 N; 22.55610 W). Both localities have very similar lithostratigraphic position: section exposed at locality 1 represents transitional layers between the Malmros Klint and the Ørsted Dal members, it is a new fossiliferous spot with poorly exposed bone-bed enriched in remains of freshwater fauna; section exposed at locality 2 shows lowermost part of the Carlsberg Fjord Beds and is rich in disarticulated elements of temnospondyls (Jenkins et al. 1994; Marzola et al. 2017) (Fig. 1). In previous studies this site was named Macknight Bjerg Quarry (see Jenkins et al. 1994). There are no signs of postmortal mechanical wearing on the studied elements. Surfaces, cusps, or crests are well preserved. Some observed changes on the bone surfaces are the effects of recent weathering. Almost all of the skeletal elements were enveloped by a slightly heterogeneous mudstone matrix, which was subjected to dissolving and softening in 10% solution of the acetic acid. Some of the specimens were prepared mechanically when the chemical method did not work.

Collected specimens are catalogued in both NHMD and KNK collections. Correspondence between KNK and NHMD numbers is as follows: KNK 4154 = NHMD 658404, KNK 4155 = NHMD 658405, KNK 4156 = NHMD 657408, KNK 4157 = NHMD 657409, KNK 4158 = NHMD 657410, KNK 4159 = NHMD 657411, KNK 4160 = NHMD 657412, KNK 4161 = NHMD 657413, KNK 4162 = NHMD 657414, KNK 4163 = NHMD 657415, KNK 4164 = NHMD 657416, KNK 4165 = NHMD 657417. In this main text of this article we use only the KNK numbers.

Several terminologies of lungfish skull bones exist in the literature. The reason is unclear homology between skull roof bones of lungfish and other vertebrates. In this paper we use nomenclature of bones and sensory lines after Kemp (1998).

Dental plates morphometry was based on the method proposed by Vorobyeva and Minikh (1968). Measured dimensions are explained on the Fig. 2. Biometric method based on the angles between ridges and sections between characteristic points was criticized by Kemp (1997), however, it is used in this paper only as a support (Kemp 1997; Skrzycki 2015). All the measurements of the specimens originally described herein were taken directly from original fossils. Measurements of *Ceratodus tunuensis* are based on the figures presented in Agnolín et al. (2018). All measurements collected in this study are presented in Table 1. This set of biometric measurements of dental plates was confronted with the measurements from Skrzycki (2015).

Principal Component Analysis was performed in PAST 3.23 software. To normalize the data we transformed the measurements by dividing them by standard deviation. We excluded incomplete specimens. The analysis was performed in order to visualize variation occurring in dental plates of different *Ptychoceratodus* species. Lower and upper dental plates were analyzed separately.

Most of the specimens were imaged using Nikon Digital Camera D5300. KNK 4158 was imaged using Keyence Digital Microscope VHX-900F. Surface of the KNK 4165 is figured as untextured high-detailed surface scan. We used Shining 3D EinScan Pro 2X 3D scanner fixed on a tripod with EXScan Pro 3.2.0.2 software.

### Results

Examined dental plates have typical ptychoceratodont morphology. They are high crowned, with less than 7 ridges.

<table>
<thead>
<tr>
<th>Specimen</th>
<th>BCp</th>
<th>α</th>
<th>δ₁</th>
<th>δ₂</th>
<th>δ₃</th>
<th>BC1</th>
<th>BC2</th>
<th>C1C2</th>
<th>C2C3</th>
<th>C1Cp</th>
<th>C2Cp</th>
</tr>
</thead>
<tbody>
<tr>
<td>KNK 4154</td>
<td>31.0</td>
<td>105.5</td>
<td>92.2</td>
<td>51.5</td>
<td>29.6</td>
<td>22.4</td>
<td>24.1</td>
<td>10.8</td>
<td>10.0</td>
<td>34.4</td>
<td>24.8</td>
</tr>
<tr>
<td>KNK 4155</td>
<td>24.9</td>
<td>109.1</td>
<td>79.5</td>
<td>46.4</td>
<td>27.0</td>
<td>14.4</td>
<td>16.1</td>
<td>8.8</td>
<td>7.1</td>
<td>26.4</td>
<td>18.1</td>
</tr>
<tr>
<td>KNK 4156</td>
<td>24.9</td>
<td>103.4</td>
<td>67.5</td>
<td>35.1</td>
<td>14.3</td>
<td>16.1</td>
<td>17.2</td>
<td>9.3</td>
<td>8.00</td>
<td>23.2</td>
<td>14.0</td>
</tr>
<tr>
<td>KNK 4157</td>
<td>15.1</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>NHMD 115910</td>
<td>50.3</td>
<td>110.7</td>
<td>80.8</td>
<td>49.4</td>
<td>27.9</td>
<td>–</td>
<td>36.5</td>
<td>–</td>
<td>14.9</td>
<td>–</td>
<td>38.4</td>
</tr>
</tbody>
</table>
Ridges originate anteriorly, which results in their fan-like arrangement. Crests meet in the medial margin. Number of the ridges in the examined specimens (5 in the upper and 4 in the lower dental plates) is reduced compared to Ptychoceratodus serratus (6 in the upper and 5 in the lower dental plates). Particularly distinctive feature is the flattening of the last ridge, present in Ptychoceratodus concinnus and Ptychoceratodus rectangulus (Linck 1936). However, dental plates of P. concinnus have a significantly different geometry (wider α angle and more robust first ridge expressed by generally higher BC1/BCp ratio in the upper and higher C1C2/BCp ratio in the lower dental plates) (see Skrzycki 2015; Fig. 3). In scatter plots presented on Fig. 3 the upper dental plates described herein group in close neighborhood of P. rectangulus dental plates, whereas the lower dental plates follow conspicuous ontogenetic trend (Fig. 3). However, the sample in both cases is very small.

Systematic palaeontology

Class Osteichthyes Huxley, 1880
Subclass Sarcopterygii Romer, 1955
Order Dipnoi Müller, 1945
Family incertae sedis
Genus Ptychoceratodus Jaekel, 1926
Type species: Ptychoceratodus serratus (Agassiz, 1838); Aargau, Switzerland; Keupersandstein.

Remarks.—Ptychoceratodus is not assigned to any family because of disagreement between researchers about taxonomy of Mesozoic lungfishes. Ptychoceratodontidae diagnosed on dental plates is commonly used by various authors (Kemp 1998; Martin et al. 1999; Richter and Toledo 2008). However, objections to taxonomy based on the dental plates are also present (Schultze 1992). Taxonomic difficulties are well outlined in Skrzycki (2015). We consider cranial features as a better taxonomic tool, however, because most dipnoan findings are isolated dental plates, there is no other and better possibility to distinguish the species or genera.

Ptychoceratodus rectangulus (Linck, 1936)
Figs. 3–5, 7.
1936 Ceratodus rectangulus sp. nov.; Linck 1936: 1–23, pl. 4: 1–5, pl. 5: 1–4.
1938 Ceratodus rectangulus Linck, 1936; Linck 1938: 1–11, text-figs. 1–3, 5.
1963 Ceratodus rectangulus Linck, 1936; Linck 1963: 1–9, pl. 20, text-fig. 1.
1981 Ceratodus rectangulus Linck, 1936; Martin 1981: 11, text-fig. 5a.
1981 Ptychoceratodus rectangulus (Linck, 1936); Schultze 1981: 21, text-fig. 13.
1998 Ptychoceratodus rectangulus (Linck, 1936); Cuny et al. 1998:7, text-fig. 5j.
2018 Ceratodus tumuensis sp. nov.; Agnolín et al. 2018: 1–6, text-figs. 3–5.

Material.—Right upper dental plate with attached almost complete pterygopalatine bone: KNK 4154; right upper dental plate: KNK 4155; right lower dental plate: KNK 4156; right lower dental plate: KNK 4157; right KLMN+3 bone: KNK 4158; right XYZ bone: KNK 4159; right operculum: KNK 4160; incomplete IJ? bone: KNK 4161; E bone: KNK 4162; left XYZ bone: KNK 4163; incomplete dental plate: KNK 4164; KLMN+3 bone imprint: KNK 4165. All from middle–upper Norian, locality 1, despite KNK 4162 from locality 2: Macknight Bjerg, Greenland. Upper dental plates: NHMD 115910–115912; lower dental plates: NHMD 115913–115916; prearticular bone: NHMD 115917. All from upper Norian, Lepidopteriselv, Greenland. Unidentified dental plates: BRSUG 26352, from Norian, Grozon, France. KLMN+3
Fig. 4. Skull roof bones of dipnoan *Ptychoceratodus rectangulus* (Linck, 1936) from the Carlsberg Fjord Beds, Fleming Fjord Formation (Upper Triassic), Macknight Bjerg, East Greenland. A. KNK 4158, KLMN+3 bone in dorsal (A1, A2) and ventral (A3, A4) views. B. KNK 4159, XYZ bone in lateral (B1, B2) and medial (B3, B4) views. Arrows indicate sutures for corresponding bones.
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**Diagnosis.**—Dental plates high-crowned, lingual margin about two times longer than the mesial. Angle between the medial and lingual margin about 90° or more. Upper dental plates with five ridges, the first ridge initially curved anterolaterally, then slightly twisted posteriorly, the last ridge flattened. Lower dental plates with four ridges, the last ridge flattened on the lingual side. Prearticular sulcus is double (like in other ptychoceratodontids except *P. serratus*), but the posterior sulcus is deeper than the anterior. Pterygopalatine ascending process above the second ridge, slightly shifted toward the third ridge, similarly to *P. serratus*. Thick descending process of KLMN+3 bone with wide basement, confluent with relatively high and wide anterior crest.

**Description.**—**Skull roof:** The largest specimen of the Greenland material is KNK 4158, interpreted as an incomplete right KLMN+3 bone because of characteristic lateral line grooves on the dorsal side, and descending process and crests on the ventral side (Fig. 4A). The bone is 36 mm long and 44 mm wide. The posterior part is damaged, whereas other edges are natural. The lateral part of the bone is clearly downturned. The lateral edge is concave. Posteriorly, the bone expands laterally and forms conspicuous lateral process, which gives KNK 4158 trapeze-like outline from dorsal view. The dorsal surface of the bone is relatively smooth, whereas the lateral and anterior margins display cancellous structure. The supraorbital sensory canal groove

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**Fig. 5.** Skull roof bones and opercle of dipnoan *Ptychoceratodus rectangulus* (Linck, 1936) from the Carlsberg Fjord Beds, Fleming Fjord Formation (Upper Triassic), Macknight Bjerg, East Greenland. A. KNK 4165, KLMN+3 bone imprint, in dorsal view. B. KNK 4160, opercle in lateral (B1) and medial (B2) views. C. KNK 4162, E bone in dorsal view. Arrows indicate sutures for corresponding bones.
angle is not preserved. The anterior part of the groove is shallow, whereas the lateral part is deep and widens toward the bone’s margin. The indentation for the medial process of the XYZ bone is visible posterior to the lateral process. The ventral surface is predominantly cancellous, being smooth only posteriorly. Two crests are visible in the central region of the bone: anterior and posterior (Fig. 4A₁, A₃). The posterior crest is directed straight to the lateral margin of the bone, and forms a slight bulge, more conspicuous laterally. The anterior crest is higher and slenderer than the posterior one, and directed anteriorly. The angle between the crests is roughly right. The anterior crest passes smoothly to the descending process of the KLMN+3 bone. The process is partially broken. However, it is clearly leaned towards the medial side. The groove for the ascending process of pterygoid appears on the medial side of the descending process base. The protrusion for the E bone is visible on the medial margin of the KLMN+3 bone. The morphology of the posterior part of the KLMN+3 bone is visible on the imprint of the same individual KNK 4165 (Fig. 5A). The posterior margin has an indentation for the J bone, however, there are no visible differences between the indentation and rest of the posterior margin. It suggests that the I bone and the J bone were merged into a single IJ bone. Three shallow grooves begin from the inclined openings on the dorsal surface of both specimens. They probably represent blood vessel ducts. Shape of the complete KLMN+3 was roughly rectangular, which differs from anteroposteriorly elongated KLMN+3 of Arganodus atlantis (see Kemp 1998). Two complete XYZ bones were found. KNK 4159 is 52 mm long and belonged to a large fish, whereas KNK 4163 is only 29 mm and represents evidently smaller individual. KNK 4159 is a flat bone, geometrically composed of three distinct regions: a flat posterior process, slightly convex but still relatively flat body, and curved articular process (Fig. 3B). The dorsal margin of XYZ has elongated protrusion for the KLMN+3 bone along the edge. The outer surface of the articular process differs from the other bone parts by its cancellous character. Radial marks cover the outer surface of the bone body and the posterior process, converging at the center of ossification. A temporal sensory line groove goes from the posterior to the anterior margin and is shallowing posteriorly. The split of the temporal line is absent. Thus, probably it was located among soft tissues. Five conspicuous elliptical openings appear in the equal distances inside the temporal line groove. The dorsal and anterodorsal margins articulated the XYZ with the KLMN+3 and IJ bones. The anterodorsal suture is relatively short, what is different from YZ bone in the skull of Ceratodus sturii (see Kemp 1998). The posteriormost part of the dorsal margin and the posterior process were joined with the IJ bone. The articular process, which was joined with the lower jaw is clearly thicker than other parts of the XYZ bone. The specimen KNK 4163 is slightly incomplete. Morphology of the bone surface is similar to the larger KNK 4159, but the distinctive feature of this specimen is bone geometry. Whole KNK 4163 is relatively flat, there is no curvature between the bone body and the articular process, unlike in KNK 4159. The delta-shaped bone (KNK 4162) is surrounded by hard sediment impossible to remove without damaging the specimen, thus only one side of the bone is visible (Fig. 5C). The bone is asymmetrical. A conspicuous convex outer side of the bone is probably its dorsal surface. There are visible openings for the blood vessels and a damaged, anteroposteriorly oriented crest along the entire bone. The inner surface is only partially exposed. There is visible an indentation matching the KLMN+3 protrusions. This is why KNK 4162 is interpreted as the E bone.

**Operculum**: The operculum (KNK 4160) is small, flattening posteriorly (Fig. 5B). Only the ventral half of the operculum is preserved. The posterior edge is very thin. Outer surface of KNK 4160 is significantly eroded, but the original surface was probably smooth. In contrast, the inner surface is cancellous. A conspicuous articulation with the XYZ bone is visible on the anterior margin. The cross section revealed a blood vessel canal that goes along the anterior margin.

**Jaw bones and dental plates**: One pterygopalatine bone was found with attached upper dental plate. KNK 4154 is an almost complete pterygopalatine missing only the ascending process (Fig. 6A). Surface of the pterygopalatine is fibrous above the dental plate, and smooth in the posterior part. The basis of the pterygopalatine process is located between the second and third ridge of the upper dental plate. Prearticular bones are represented only by fragments directly attached to the lower dental plates (KNK 4156, 4157). The ventral surface of the KNK 4156 prearticular bone is fibrous and has a distinctive double sulcus (Fig. 6B₃, B₄). The anterior sulcus is isometric and significantly smaller and shallower than the posterior, which is posteriorly elongated and deeper. Five dental plates were found. KNK 4154 and KNK 4155 are palatal plates of moderate size (31.3 and 22.1 mm long, respectively), whereas KNK 4156 and KNK 4157 are smaller prearticular plates (23.9 and 15.1 mm long, respectively). The dental plates are slightly elongated and high crowned with fan-like pattern of ridges (Fig. 6A–C). Crests of the three anterior ridges are meeting at the medio-lingual junction. The specific feature of the dental plates is flattening of the last ridges.

The upper dental plates have five ridges, and slightly convex occlusal surfaces in the posterior part. The last ridges are almost completely flat. KNK 4155 has conspicuous cusps on the three last ridges. The fifth ridge of the smaller specimen (KNK4154) is more confluent with the fourth ridge than in the bigger (KNK4155).

The lower dental plates have four ridges and the occlusal surface is about flat. The last ridges of lower dental plates are composed of medial crest and flattened lingual part. Cusps are visible on all preserved ridges. KNK 4157 has slightly separated fifth ridge with low crest (Fig. 6C).

First crest is differently oriented in the upper and lower plates. In the upper plates it follows mesial margin, but in the lower it is in distance from the margin (Fig. 6B₁, B₂, C).
This pattern enables occlusion. KNK 4164 represents an incomplete dental plate of a large individual (Fig. 6D). Only first two ridges are preserved. The surface is heavily worn and the ratio of thickness to length is relatively small. It is difficult to establish if it is a palatal or prearticular dental plate because no identifiable bone part is attached to the specimen.

Remarks.—The first description of Ptychoceratodus rectangulus in Linck (1936) was based on eight poorly preserved dental plates from the middle Löwenstein Formation of Ochsenbach (middle Norian, Germany). It was initially assigned to genus Ceratodus, until Schultz (1981) transferred it to the genus Ptychoceratodus. Skull bones were also found in the same formation. Linck (1936, 1938, 1963) made bio-
metrical studies, which were repeated in Skrzycki’s (2015) analysis according to current standards. Full morphological description or biometric measurements were not possible for some of the dental plates due to preservation (Skrzycki 2015). A holotype was never designed for *P. rectangulus*. Cuny et al. (1998) assigned to this species one minute dental plate from the Norian Marnes de Châlins Formation in eastern France. The tooth plates described here are not as numerous as the material from Ochsenbach but are better preserved, what makes more detailed morphological or histological studies possible.

The dental plates described by Agnolin et al. (2018) as *Ceratodus tunuensis* came also from the Carlsberg Fjord Beds. They were classified as ceratodontid due to robust and short ridges having broad occlusal surface with rounded outline. However, all described specimens are worn and incomplete (except the complete but worn NHMD 115912). *Ptychoceratodus rectangulus* specimens from Ochsenbach show similar degree of erosion, however, they are mostly complete. High degree of incompleteness of the *C. tunuensis* dental plates may indicate that they were subject of transport and their wear is partially result of post-mortem erosion. This makes the proper evaluation of features like roundness of the occlusal surface, ridges’ and crests’ length, or depth of the inter-ridge furrows, indicated as diagnostic features of *C. tunuensis*, difficult. Other features of *C. tunuensis* (less vulnerable to erosion) are common for *P. rectangulus*. These features are: the $\alpha$ angle dimension, curvature of the first ridge, dental plates elongation, number of ridges and prearticular sulcus shape. Measurements of the *C. tunuensis* holotype (NHMD 115910) show geometrical similarity to ptychoceratodontids (Fig. 3). On these bases we suppose that *C. tunuensis* is a younger synonym of *P. rectangulus*. It is impossible to determine if the differences in dimensions of the upper dental plates from Greenland and Germany (e.g., wider $\alpha$ angle and narrower $\beta_2$ angle in the German specimens) are caused by an intraspecific variability or evolutionary changes related to isolation in two distant basins (Fig. 7).

**Stratigraphic and geographic range.**—Germany, Ochsenbach region; Löwenstein Formation (Stubensandstein), Middle Keuper, middle–upper Norian. Greenland, Macknight Bjerg Quarry; lower part of the Carlsberg Fjord Beds (Ørsted Dal Member), Fleming Fjord Formation, middle–upper Norian. France, Grozon region; Marnes de Châlins Formation, Norian.

**Discussion**

The five skull bones, from the same strata of the German Triassic as the dental plates, were originally interpreted incorrectly by Linck (1963). They were reinterpreted by Schultze (1981) with a reconstruction of the skull roof, and mentioned by Martin (1981). However, preservation of these skull bones is comparably bad as that of the associated dental plates. All German specimens are imbedded in a rock matrix and only external surfaces of the bones are accessible for study. Most of them are incomplete, with broken or invisible edges, and one specimen was destroyed during preparation (Schultze 1981). The set of the Linck’s (1963) skull roof bones includes, according to Schultze (1981): the KLMN+3 bones left (SMNS 19705) and right (SMNS 19706), the incomplete right I bone (SMNS 19707), the E bone (destroyed) and an unidentified poorly preserved element (SMNS 19708).

In the German material the best preserved are the KLMN+3 bones, which resemble the Greenland specimens. Supraorbital line turning at roughly right angle, and deep indentation for the medial process of the XYZ on the lateral margin are particular similarities. Clear difference between the German and Greenland KLMN+3 bones is the lack of the indentation for the J bone on the posterior margin in SMNS 19705 and conspicuous indentation in KNK 4165 (in SMNS 19706 and KNK 4158 the posterior margins are not preserved or not visible). It implies that the J bone could be very small.
(Schultze 1981), or merged variably with neighboring bones (as suggested by Schultze 1981 concerning \textit{P. serratus}). The J bone in \textit{P. serratus} is placed between the KLMN+3, I, and AB bones. Both specimens interpreted as the I bone (SMNS 19707 and KNK 4161) are highly fragmentary, however SMNS 19707 is much more complete than KNK 4161. Schultze 1981:21 comments: “relation of the I bone and KLMN+3 bone is uncertain, if treated like in \textit{P. serratus} or \textit{Arganodus}, there is no space for the J bone, or the J bone is very small”. Indeed, the I bone has a protrusion interpreted in Schultze 1981 as the joint with the AB bone, thereby there is left only a little space for the J bone. Actually, the KLMN+3 imprint (KNK 4165) seems to confirm merging of the I and J bones into the IJ bone (Fig. 8) because of smooth transition between articulations with the I and J bones.

The delta-shaped KNK 4162 is interpreted as the E bone because its posterolateral edges match the KLMN+3 anteromedial margin. Badly preserved SMNS 19708 can represent the same skull element. In Schultze’s (1981) reconstruction, the destroyed bone was interpreted as the E bone (Linck 1963: fig. 3) on the basis of ragged edge in the posterior part, which matches the anteromedial margin the of the KLMN+3 (SMNS 19705, 19706). These interpretations are not contradictory because the destroyed specimen revealed only the ventral surface (Martin 1981), and KNK 4162 only the dorsal. Both specimens had similar size, therefore the shape differences are not an effect of allometric growth, but according to Schultze (1981), the E bone shape can be variable, or specimens are just incomplete. We used the delta-shaped bone as the E bone in the reconstruction, however, it needs confirmation by new material (Fig. 8). Morphology of the minute bone fragment, described in Linck (1936) as the operculum (SMNS 17973), indicates that it is rather the descending process of the XYZ bone.

The skull roof bone pattern of \textit{Ptychoceratodus rectangulus} (Linck, 1936) is generally similar to other ptychoceratodontids with known skull anatomy: \textit{Ptychoceratodus serratus} (see Schultze 1981), \textit{Ptychoceratodus philipsi} (see Kemp 1996), \textit{Ferganoceratodus jurassicus} Nessov and Kaznyshkin, 1985, and \textit{Ferganoceratodus martini} Ferganoceratodus martini Cavin, Suttethorn, Buffetaut, and Tong, 2007. Unique features of the \textit{P. rectangulus} skull among other ptychoceratodontids are robust ventral crests of the KLMN+3 bone, similar rather to \textit{Arganodus atlantis} (see Martin 1981), and possibly the delta-shaped E bone with dorsal ridge. Right angle curvature of the supraorbital sensory line is similar rather to \textit{Ferganoceratodus} spp. than \textit{Ptychoceratodus} spp. \textit{F. martini} and \textit{F. jurassicus} have reduced number of bones in the medial and mediolateral series in comparison with \textit{P. serratus}. However, number of bones in the medial and mediolateral series in \textit{P. rectangulus} is uncertain. Presence of only two bones at least in the mediolateral series is highly probable, as discussed above, and such a state is illustrated in the reconstruction (Fig. 8).
Ptychoceratodus rectangulus has regularly spaced, relatively large openings in the grooves of temporal line. They correspond to localization of sensory canal pits in Recent lungfishes. Extant Propterus has similarly developed pattern of these openings (compare Webb and Northcutt 1997). Importance of mechanoreception or electrorception for Propterus (Bullock et al. 1983; Orlov et al. 2015) suggest similar mode of life for P. rectangulus.

Ptychoceratodus rectangulus coexisted in the Jameson Land Basin with large temnospondyls like the capitosaurid Cyclotosaurus naraserluuki (see Marzola et al. 2017) and especially common plagiosaurid Gerrothorax pulcherinus (see Jenkins et al. 1994, 2008; Schoch and Witzmann 2012; Sulej et al. 2014). Occurrence of the same dipnoan species in the middle–upper Norian sediments of Germanic and Jameson Land basins indicates that at least then the areas were connected. In the Late Triassic both basins had been situated relatively close (Fig. 7), until the opening of the northernmost part of Atlantic Ocean in the Cretaceous, which split Europe and Greenland (Rowley and Lottes 1988). The Mesozoic rifting favored origin of sedimentation basins, which could have been temporally connected with the Germanic Basin and enabled freshwater fauna exchange (McKie and Williams 2009; Ziegler 1998).

Conclusions

The record of Ptychoceratodus from the Fleming Fjord Formation in East Greenland shows a close correlation between the Late Triassic Jameson Land Basin fish fauna and the fauna of the coeval Germanic Basin. Studied Ptychoceratodus rectangulus remains were found in the transitional layers between the Malmros Klint and the Ørsted Dal members and in the lower part of the Carlsberg Fjord Beds (middle–upper Norian). The species was previously known from the Löwenstein Formation (middle–upper Norian) in Germany. Geographical distribution of P. rectangulus shows at least temporary connection between the two basins during middle–late Norian. Recently described Ceratodus tumenensis from the uppermost part of the Carlsberg Fjord Beds is probably conspecific with P. rectangulus. Ptychoceratodus rectangulus had sensory line pit pattern at least similar to that of the extant Propterus, which suggests a similar mode of life. The Late Triassic dipnoan freshwater fish faunas of the northern Pangea are characterized by a frequent occurrence of Ptychoceratodus.

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References

Clemmensen, L.B. 1980. Triassic lithostratigraphy of East Greenland be-
Clemmensen, L.B., Kent, D.V., and Jenkins F.A. 1998. A Late Triassic lake system in East Greenland: facies, depositional cycles and palaeo-


Kemp, A. 2017. Evolutionary history of lungfishes with a new phylo-


tion of East Greenland and a reassessment of head lifting in temnophi-


Niedźwiedzki, G. 2017. Tetrapod track assemblages from the Upper Tri-

assic Fleming Fjord Formation, East Greenland. *Palaeontologia Afri-

cana* 52: 179.


Schoch, R.R. and Witzmann, F. 2012. Cranial morphology of the plagio-

saurid *Gerrothorax pulcherimus* as an extreme example of evolution-


Shapiro, M.D. and Jenkins, F.A. 2001. Cynodont from the Upper Triassic of East Greenland: tooth replacement and double-rootedness. *Bulle-


Martin, M. 1982. Nouvelles données sur la phylogénie et la systématique des Dipnées postpaléozoïques, conséquences stratigraphiques et paléogéogra-


Martin, M., Barbieri, L., and Cuny, G. 1999. The Madagascan Psychocerat-

donid. *Amphibia, Temnospondyls* from the Fleming Fjord Formation of the Jameson Land Basin (East Greenland). *Journal of Vertebrate Paleon-

ology* 37: e1303501.

Mckie, T. and Williams B. 2009. Triassic palaeogeography and fluvial dis-


ology* 37: e1303501.
Skrzycki, P. 2015. New species of lungfish (Sarcopterygii, Dipnoi) from the Late Triassic Krasiejów site in Poland, with remarks on the ontogeny of Triassic dipnoan tooth plates. *Journal of Vertebrate Paleontology* 35: e964357.


