Archosaur footprints (cf. *Brachychirotherium*) with unusual morphology from the Upper Triassic Fleming Fjord Formation (Norian–Rhaetian) of East Greenland

HENDRIK KLEIN1*, JESPER MILÄN2,3, LARS B. CLEMMENSEN3, NICOLAJ FROBØSE3, OCTÁVIO MATEUS4,5, NICOLE KLEIN6, JAN S. ADOLFSSEN2, ELIZA J. ESTRUP7 & OLIVER WINGS8

1 Saurierwelt Paläontologisches Museum, Alte Richt 7, D-92318 Neumarkt, Germany
2 Geomuseum Faxe/Østsjællands Museum, Østervej 2, DK-4640 Faxe, Denmark
3 Department for Geosciences and Natural Resource Managements, University of Copenhagen, Øster Voldgade 10, DK-1350 Copenhagen K, Denmark
4 Department of Earth Sciences, GeoBioTec, Faculdade de Ciências e Tecnologia, FCT, Universidade Nova de Lisboa, 2829-516 Caparica, Portugal
5 Museu da Louriña, Rua João Luís de Moura 95, 2530-158 Lourinhã, Portugal
6 Staatliches Museum für Naturkunde Stuttgart, Rosensteinstr. 1, 70191 Stuttgart, Germany
7 Geocenter Møns Klint, Stengård Svej 8, DK-4791 Borre, Denmark
8 Niedersächsisches Landesmuseum Hannover, Willy-Brandt-Allee 5, 30169 Hannover, Germany

*Corresponding author (e-mail: Hendrik.Klein@combyphone.eu)

Abstract: The Ørsted Dal Member of the Upper Triassic Fleming Fjord Formation in East Greenland is well known for its rich vertebrate fauna, represented by numerous specimens of both body and ichnofossils. In particular, the footprints of theropod dinosaurs have been described. Recently, an international expedition discovered several slabs with 100 small chirotheriid pes and manus imprints (pes length 4–4.5 cm) in siliciclastic deposits of this unit. They show strong similarities with *Brachychirotherium*, a characteristic Upper Triassic ichnogenus with a global distribution. A peculiar feature in the Fleming Fjord specimens is the lack of a fifth digit, even in more deeply impressed imprints. Therefore, the specimens are assigned here tentatively to cf. *Brachychirotherium*. Possibly, this characteristic is related to the extremely small size and early ontogenetic stage of the trackmaker. The record from Greenland is the first evidence of this morphotype from the Fleming Fjord Formation. Candidate trackmakers are crocodylian stem group archosaurs; however, a distinct correlation with known osteological taxa from this unit is not currently possible. While the occurrence of sauropodomorph plateosaurs in the bone record links the Greenland assemblage more closely to that from the Germanic Basin of central Europe, here the described footprints suggest a Pangaea-wide exchange.

Supplementary material: Three-dimensional model of cf. *Brachychirotherium* pes–manus set (from MGUH 31233b) from the Upper Triassic Fleming Fjord Formation (Norian–Rhaetian) of East Greenland as pdf, ply and jpg files (3D model created by Oliver Wings; photographs taken by Jesper Milàn) is available at https://doi.org/10.6084/m9.figshare.c.2133546

During the Geocenter Møns Klint Dinosaur Expedition in July 2012, a new site with archosaur footprints was discovered by one of the authors (OW) in the lowermost part of the Ørsted Dal Member of the Upper Triassic Fleming Fjord Formation in mountain slopes facing the Carlsberg Fjord in East Greenland (Fig. 1). These archosaur footprints add to the knowledge of vertebrate life in the ancient lake and steppe system, and supplement previous descriptions of tetrapod footprints and other vertebrate trace fossils from the Upper Triassic deposits (Jenkins et al. 1994; Clemmensen et al. 1998; Milàn et al. 2004, 2006, 2012; Milàn & Bromley 2006; Niedźwiedzki et al. 2014; Sulej et al. 2014). They increase the diversity of tetrapod footprint assemblages from the Fleming Fjord Formation, reflecting a typical Late Triassic community with dinosaur and stem-crocodylian
archosaurs. This is in congruence with the osteological record, which has an equivalent in assemblages of the Germanic Basin of central Europe (Jenkins et al. 1994). The new material, consisting of abundant and well-preserved imprints of identical shape, shows an anatomically based morphological peculiarity, and therefore requires a detailed discussion of the ichnotaxonomy and the possible trackmaker.

Geology and the depositional environment

The Jameson Land Basin lies in central East Greenland at 72° N (Fig. 1). The basin contains a seemingly complete (up to 300 m thick) Upper Triassic succession of lake deposits (Clemmensen et al. 1998). Stratigraphically, these sediments belong to the Fleming Fjord Formation, which is composed of a lower Edderfugledal Member, a middle Malmros Klint Member and an upper Ørsted Dal Member (Fig. 2) (Clemmensen 1980a, b). The latter is divided into a lower unit of red mudstones and sandstones (the Carlsberg Fjord beds), and an upper unit of light greyish dolomitic limestones and variegated mudstones (the Tait Bjerg Beds) (Clemmensen et al. 1998).

When the sediments of the Fleming Fjord Formation were deposited, the Jameson Land Basin lay at about 40° N, at the margin of the dry interior of the supercontinent Pangaea (Kent & Clemmensen 1996; Clemmensen et al. 1998; Kent & Tauxe 2005). Lake deposition was likely to have been influenced by seasonal, as well as orbitally controlled, precipitation changes (Clemmensen et al. 1998). The lake sediments record a gradual, long-term (a few million years) change in climate from semi-arid to humid, probably reflecting a slow northwards drift of the continent in latest Triassic times (Clemmensen et al. 1998; Kent & Tauxe 2005).

Deposits of the Fleming Fjord Formation are particularly well exposed in mountain slopes facing...
the Carlsberg Fjord. The tracks in question were found in the lowermost part of the Carlsberg Fjord beds in the Ørsted Dal Member at the Lepidopteriselv and Tait Bjerg sections (Fig. 2). The Carlsberg Fjord beds are here composed of red mudstones repeatedly interbedded by thin silt- and sandstones, with wave and current ripples. The sedimentary characteristics of these deposits indicate deposition in a relatively shallow lake that frequently dried up (Clemmensen et al. 1998). The base of the unit contains two fluvial sheet sandstones and one lenticular-shaped channel sandstone. The tracks from Lepidopteriselv locality were found in loose blocks at the base of the fluvial channel sandstone, and the sedimentary characteristics of the blocks indicate that they belong to the uppermost part of the composite channel sandstone (Figs 2 & 3).

Magnetostratigraphical investigation of the Fleming Fjord Formation suggests that the tracks from the basal part of the Ørsted Dal Member are about 210 myr old and, hence, of Norian age (Clemmensen et al. 1998). A diverse assemblage of fossil vertebrates also indicates that the main part of the Ørsted Dal Member is of Norian age (Jenkins et al. 1994).

**Material and methods**

Three slabs with footprints have been recovered (Fig. 1). Two of them have been found at a locality (Lepidopteriselv) at the eastern margin of the Jameson Land Basin in East Greenland (Fig. 1). These are loose blocks (consisting of several pieces), but sedimentological data indicate that they originate from the topmost part of a fluvial channel sandstone in the lowermost part of the Ørsted Dal Member of the Fleming Fjord Formation (Fig. 3). The third sample was collected at Tait Bjerg (Fig. 1). These footprints were seen *in situ* at the top of a fluvial sheet sandstone in the lowermost part of the Ørsted Dal Member. The material is stored in the collection of the Natural History Museum of Denmark (Copenhagen) (MGUH) under catalogue numbers MGUH 31233a–c, MGUH 31234 and MGUH 31235. Concave imprints were outlined with chalk, drawn onto transparency film and digitized using Adobe Illustrator CS3 software. Silicone moulds...
and plaster casts were taken from the best-preserved footprints. Photographs were taken under artificial and natural light from both original slabs and casts, and all measurements were made from the original material only (Table 1).

Untextured 3D models from the slabs and high-resolution models from the best-preserved tracks were produced with photogrammetry. Digital photographs were subsequently processed using the methods described by Mallison & Wings (2014) using the photogrammetric software Agisoft PhotoScan Professional 1.0.4 build 1847 (64 bit) with generally high settings.

Tetrapod footprints

Formerly known material

The Ørsted Dal Member of the Fleming Fjord Formation has provided a rich tetrapod ichnofauna, including trackways of theropods with tridactyl footprints up to 28 cm long that can be assigned to the Eubrontes–Anchisauripus–Grallator plexus (Olsen et al. 1998; Gatesy et al. 1999; Milan et al. 2004, 2006; Lucas et al. 2010). Large imprints of a quadruped, up to 53 cm in length, were also described (Jenkins et al. 1994; Clemmensen et al. 1998; Lockley & Bromley 2000).

Recently, Niedźwiedzi et al. (2014) and Sulej et al. (2014) reported new discoveries of trackways of large quadrupeds (up to 50 cm pes length) from the Malmros Klint Member of the Fleming Fjord Formation and attributed them to sauropodomorph dinosaurs. These large tracks from Greenland have been compared with Eosauropus, a purported sauropodomorph ichnotaxon known from Upper Triassic deposits of North America and Europe (Lockley & Meyer 2000; Lockley et al. 2006, 2011; Niedźwiedzi et al. 2014). Until now, they were figured without details or only symbolically on a map of the track surface (Jenkins et al. 1994; Sulej et al. 2014). Details of their morphology and documentation using close-up photographs and sketches were not given by these authors, and their identification as large chirotheres, therefore, cannot presently be excluded.

Jenkins et al. (1994, p. 19) also mentioned trackways of a smaller quadruped from the Ørsted Dal Member and small (2.5–5 cm) tetradactyl imprints showing ‘four relatively short digits with rounded ends’ from the Malmros Klint Member. The latter they consider as probable amphibian tracks. Again, no figures were provided and these authors did not indicate the whereabouts of the material. Probably, these footprints were left in the field.

New discoveries

The material described here consists of about 100 distinct pes and manus imprints made by similarly sized individuals distributed on three main slabs (Figs 4–10). The dimensions of the largest slab (broken into three pieces) are $93 \times 76 \times 4$ cm, displaying about 60 imprints (Figs 4, 5, 7a & 8) with generally high settings.

Table 1. Measurements of cf. Brachychirotherium footprints described in this study compared with selective data from characteristic Upper Triassic Brachychirotherium of different localities

<table>
<thead>
<tr>
<th>Ichnotaxa</th>
<th>cf. Brachychirotherium (this study)</th>
<th>B. hassfurtense*</th>
<th>B. thuringiacum†</th>
<th>B. parvum‡</th>
</tr>
</thead>
<tbody>
<tr>
<td>pl</td>
<td>4.0–4.5</td>
<td>24.1</td>
<td>18.0–19.0</td>
<td>13.5</td>
</tr>
<tr>
<td>pw</td>
<td>4.0–4.5</td>
<td>25.6</td>
<td>18.0–19.0</td>
<td>14.0–17.0</td>
</tr>
<tr>
<td>ml</td>
<td>2.0–2.2</td>
<td>–</td>
<td>5.8–9.7</td>
<td>6.0–8.0</td>
</tr>
<tr>
<td>mw</td>
<td>2.0–2.2</td>
<td>–</td>
<td>7.3–11.0</td>
<td>8.0–9.0</td>
</tr>
<tr>
<td>p I–IV</td>
<td>44</td>
<td>50</td>
<td>40–55</td>
<td>74</td>
</tr>
<tr>
<td>SL</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>– 92</td>
</tr>
<tr>
<td>PA</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>– 153</td>
</tr>
</tbody>
</table>

*†From the Hassberge Formation (Carnian) of northern Bavaria, Germany (Brachychirotherium type locality and horizon) (from Karl & Haubold 1998).
‡From the Redonda Formation (Chinde Group, Norian–Rhaetian) of New Mexico, USA (from Lucas et al. 2010). Notice all imprints being as wide as long, or wider than long.

pl, pes length (in cm); pw, pes width (in cm); ml, manus length (in cm); mw, manus width (in cm); p I–IV, divarication between pedal digits I and IV (in °); SL, stride length (in cm); PA, pace angulation (in °).

§Mean values. Length and width of imprints refers to anterior digit group I–IV only, except values in bold that include digit V.
belongs to the same track level as MGUH 31233a–c but does not match the footprints on the slabs with the concave traces. Pes imprints are tetradactyl with moderate spread digits (the angle between digits I and IV is 44°). The digits are relatively broad and short, and show rounded phalangeal pads and small thin claws (Figs 4, 5, 10). Their lengths and widths are within the same size range (i.e. at 4–4.5 cm). Digit proportions are III > II > IV > I. A distinct metatarsophalangeal area is visible, sometimes showing well-developed rounded pads (Figs 4b, c & 10a). The proximal margin can be variable, appearing as posteriorly concave, convex or straight owing to different preservation and imprint depth. The smaller manus imprints are 2–2.2 cm in length and width, respectively, and mostly show only three (II–IV), but occasionally four (I–IV), digits (Fig. 10a–c, e). Digit proportions of the manus are similar as in the pes, with III > II > IV > I. Even in deeper (5 mm) imprints, all pes and manus tracks lack digit V. The manus imprints are inwardly rotated relative to the pes. The position of the manus relative to the pes is variable, appearing anteromedial, anterior or even antero-lateral (Fig. 10a–c, e). The well-preserved specimen from slab MGUH 31233b in Figures 4b, c and 10a shows a lateral overstep of the manus by the pes. Some imprints display structures related to scale impressions.

**Fig. 4.** Archosaur footprints cf. *Brachychirotherium* from the Ørsted Dal Member of the Fleming Fjord Formation (Upper Triassic, Norian–Rhaetian) of eastern Greenland. (a) Overview of slab MGUH 31233a–c (broken into three pieces: 31233a is the lower, 31233b the middle and 31233c the upper part) with numerous small pes and manus imprints of cf. *Brachychirotherium* preserved as concave epi-reliefs and outlined with chalk. (b) & (c) Detail of a pes–manus set from the middle part of the slab (MGUH 31233b): (b) photograph of natural mould and (c) a 3D model.
and/or scale drag marks (MGUH 31235: Fig. 6b). These are indistinctly preserved polygons and striations on the lateral sides of digits and along the sole surface, reflecting the dynamics of the pes when being pushed into the substrate and slipping backwards. However, no distinct scale pattern can be observed.

**Discussion**

Morphologically, the imprints described here show a strong similarity with chirotheriid footprints, in particular with the ichnogenus *Brachychotherium* Beurlen, 1950 from the Late Triassic of Europe (Beurlen 1950; Karl & Haubold 1998, 2000; Petti et al. 2009), North America (Baird 1957; Silvestri & Szajna 1993; Lockley & Hunt 1995; Szajna & Silvestri 1996; Szajna & Hartline 2003; Klein et al. 2006; Hunt & Lucas 2007a, b; Lucas et al. 2010; Lucas & Heckert 2011), South America (Melchor & De Valais 2006), North Africa (Hminna et al. 2013) and southern Africa (Ellenberger 1972; D’Orazi Porchetti & Nicosia 2007). The short and broad digits with small, thin claws, and the digit proportions with digit IV < digit II in the Greenland tracks are diagnostic of *Brachychotherium* (Karl & Haubold 1998).

Problematic is the lack of a fifth digit in all (about 100) imprints. Even more deeply impressed footprints show no digit V impression.
This is unusual for Brachychirotherium and for all chirotteriids, especially in the pes imprint. Brachychirotherium occasionally lacks a trace of digit V due to substrate conditions and pes posture (Karl & Haubold 1998), but the large number of tracks from Greenland suggests a relationship of this feature to the pes anatomy of the trackmaker, which obviously had a functionally tetradactyl pes. However, the compact digit group I–IV in the Greenland tracks is typically chirotteriid. The only non-chiroteriid tracks of a quadruped that are similar to Brachychirotherium but show a functionally tetradactyl pes imprint is Batrachopus, mainly known from the Lower Jurassic but also from the Late Triassic (Fig. 11g). Batrachopus is considered to be a crocodylomorph track (Olsen & Padian 1986; Klein & Lucas 2010b). However, the manus imprint of Batrachopus is strongly rotated outwards relative to the pes imprint: whereas, in the material from Greenland, the manus imprint is inwards or slightly rotated outwards with respect to the pes imprint (cf. Fig. 11f & g). In typical Brachychirotherium tracks from the Late Triassic, the manus imprint is turned slightly more outwards than the pes (Fig. 11b–d).

Considering the overall congruence of the Greenland tracks with the ichnogenus Brachychirotherium, as well as the morphological peculiarity and lack of a (diagnostic) fifth digit trace, we tentatively refer the material described here to archosaur footprints cf. Brachychirotherium. Possibly, the lack of digit V generally characterizes juvenile Brachychirotherium (see below), requiring an emendation of the diagnosis of the ichnogenus. Late Triassic Brachychirotherium footprints of small size (<5 cm) are hitherto known only from a few imprints with a preserved fifth digit from a single locality in Argentina, South America (Fig. 11e). Small ‘Brachychirotherium’ tracks from the Middle Triassic of Germany (Haubold 1967, 1971a, b) also show a fifth digit, but the correct identification of the ichnogenus in these strata (originally Late Triassic) is doubtful (Karl & Haubold 1998; Klein & Haubold 2004). Alternatively, the tracks from Greenland could represent a new ichnotaxon, but this cannot be demonstrated here with confidence. Establishing a new ichnotaxon would require the presence of a juvenile Brachychirotherium and early ontogenetic stage to be excluded, and, instead, the presence of a small adult form to be proven. However, a new ichnotaxon would be based weakly on the lack of a fifth digit only.

Trackmakers

Brachychirotherium footprints have most probably been left by stem-crocodylian archosaurs (sensu Brusatte et al. 2010; Nesbitt 2011). The pes of the latter is distinguished from that of dinosaur bird-line archosaurs (Avemetatarsalia sensu Benton 1999) by the broader configuration of the metatarsals. This feature is reflected in the typically broad pes imprints of Brachychirotherium. An exception to this rule is the pes of the stem-crocodylian Poposaurus from the Late Triassic of North America, which has a slender shape developed convergent with that of dinosauromorphs (Farlow et al. 2014). Brachychirotherium has been attributed to rauisuchians, crocodylomorphs and aetosaurs by different authors (Klein et al. 2006; Avanzini et al. 2010; Heckert et al. 2010; Lucas & Heckert 2011; Desojo et al. 2013).

Small Brachychirotherium from Upper Triassic deposits are uncommon, and the tracks from Greenland thus far could represent the smallest known. A detailed study of different-sized specimens and
their morphologies has never been undertaken. Chi-
rotheriid footprints reflecting juvenile individuals
have been described from the Middle Triassic (Ani-
sian) of Germany (Haubold 1967, 1971a, b). Avan-
zini & Lockley (2002) gave a detailed analysis
of different Isochirotherium size classes from the
Middle Triassic of the Southern Alps (Italy), with
some specimens having a pes length of 2.7 cm.
However, these very small imprints show all fea-
tures diagnostic of the ichnogenus Isochirotherium
and of chirotheriids in general, including an impres-
sion of the fifth pedal digit. The Greenland tracks
represent the second occurrence of small-sized cf.
Brachychirotherium footprints from Upper Triassic
deposits. They could possibly represent juveniles
and ontogenetic stages with a still less-developed
(not impressed) digit V or functionally tetractyly
pes. The wide variation in the position and orienta-
tion of the manus imprint (Fig. 10) might reflect the
still unconsolidated gait of juvenile individuals of
basal archosaurs. Possibly, juveniles also walked
with a more digitigrade posture of the pes, where
the short fifth digit did not touch the ground and
therefore left no impression. During growth and
increase in body weight, this might have changed
towards a semi-plantigrade or plantigrade posture,
leaving footprints with distinct sole and digit V
impressions. Allen et al. (2009, 2010) demonstrated
the shift of the centre of mass (COM) during ontog-
ey in extant crocodylians and birds. Crocodylians
show a craniodorsal shift of the COM and an
increase in loading on the hip joints that essentially
For extant crocodylians, Farlow & Britton (2000)
showed that during growth the autopodia become
shorter relative to the glenoacetabular length. How-
ever, comparing the lengths of the toe region and
the plantar/palmar portion of the feet relative to
total leg length, the digits show slightly negative
allometric growth, whereas values for the sole indi-
cate isometric growth (Farlow & Britton, 2000).
This would mean that the sole becomes slightly lon-
ger relative to the digit portion. For the theropod
dinosaur Allosaurus, Foster & Chure (2006) showed
negative allometric growth of metatarsal IV relative
to the tibia. In any case, the increase in weight on the

Fig. 7. Pes and manus imprints of cf. Brachychirotherium: (a) plaster replica taken from slab MGUH 31233a with
two overprinted pes impressions; (b) & (c) natural casts from slab MGUH 31235; and (d) natural mould from a
different slab (MGUH 31234).
hind limbs, combined with the relative shortening of metatarsals, as in some digitigrade bipeds, or the elongation of the palmar/plantar portion in more plantigrade quadruped forms, affect the biomechanics and function of the pes. Whether the same patterns observed in extant archosaurs were present

Fig. 8. Overview sketch of slab MGUH 31233a–c with archosaur footprints of cf. Brachychotherium. Letters (a, b, d & f) indicate the position of footprints outlined in Figure 10.

Fig. 9. Overview sketch of slab MGUH 31235 with archosaur footprints of cf. Brachychotherium. Letters (c & e) indicate the position of footprints outlined in Figure 10.
in the Greenland trackmakers cannot be proved with certainty, but the degree of contact of pedal digit V with the substrate, and the preservation of its impression, was probably controlled by differential growth of limb parts.

The osteological tetrapod record of the Fleming Fjord Formation thus far yields temnospondyls, possible sphenodonts and lepidosaurs, turtles, phyto-
saurs, a possible rauisuchian, aeotosaurs, pterosaurs, prosauropods, theropods, and mammals (Jenkins et al. 1994; Mateus et al. 2014; Sulej et al. 2014). From these, only stem-crocodylian archosaurs (rauisuchians, aeotosaurs) might be cautiously considered as potential trackmaker candidates (Karl & Haubold 1998; Heckert et al. 2010; Lucas & Heckert 2011). An affinity of *Brachychirotherium* to aeotosaurs was suggested by Lucas & Heckert (2011), who found several synapomorphies of footprints and pes skeletons. Skeletal remains of aeotosaurs are known from the Fleming Fjord Formation by the taxa *Aetosaurus ferratus* and *Paratypothorax andressi* (Jenkins et al. 1994). The size of the pes of *Aetosaurus* is in the range of the tracks described (Schoch 2007): however, the skeleton shows a relatively long digit IV in the pes and manus, whereas the tracks have a short pedal and manual digit IV (IV < II). Unfortunately, no trackway is preserved in the chirother-iid material from Greenland, and a comparison with the pattern of *Brachychirotherium* trackways from other localities is not possible. It is important to mention that the occurrence of body fossils and footprints in the same stratigraphic unit does not necessarily indicate a relationship. For example, the Olenekei–Anisian (Lower–Middle Triassic) Moenkopi Formation/Group is dominated by am-
phibian body fossils, whereas the tetrapod footprint record shows only a few amphibian, but abun-
dant reptile, tracks (Peabody 1948; Klein & Lucas 2010a). This is probably due to alternating ecologi-
cal conditions with different precipitation and water supply under a semi-arid climate and/or a tapho-
monic effect due to the different conditions under which skeletons or footprints can be preserved.

**Biostratigraphy, palaeobiogeography, palaeoecology**

If the footprints from Greenland represent the ichnogenus *Brachychirotherium*, then they match characteristic tetrapod footprint assemblages from Upper Triassic deposits in the global record (Klein & Haubold 2007; Klein & Lucas 2010c). Klein & Lucas (2010c) proposed a *Brachychirotherium* biochron for Carnian–Rhaetian assemblages that, similar to those from Greenland, contain abundant tridactyl footprints of the *Grallator–Eubrontes* type (see above).

Following Lucas (1998, 2010), the vertebrate body fossil assemblage in Greenland belongs to the Revueltian land-vertebrate faunachron (LVF), which can be cross-correlated with the Carnian–Norian as it is similar to typical associations in Central Europe (e.g. from the German Keuper). In particular, the occurrence of the prosauropod *Plateosaurus* resembles assemblages from the Germanic Basin, whereas the Triassic bone record from North America lacks prosauropods. Against the background of the close proximity of what is today Greenland and North America in Late Triassic times, this can possibly be explained by the existence of larger barriers such as mountain ranges and/or climatic obstacles. In a reconstruction of climate belts on the Late Triassic supercontinent Pangaea, Kent et al. (2014) placed the Germanic Basin and the Jameson Land Basin in a relatively humid climate belt, while the North American
localities are situated in a relatively arid climate belt. This climatic zonation may have controlled faunal variation on the continent. However, tetrapod footprints of the ichnogenus *Eosauropus*, originally described from the Upper Triassic of North America and possibly present in the Flemingfjord assemblage (see above), have been attributed to sauropodomorphs (Lockley *et al.* 2001, 2006, 2011). Interestingly, the Late Triassic record of the Argana Basin of Morocco, which was positioned close to eastern North America prior to the Atlantic rifting, also lacks sauropodomorph body fossils. Purported prosauropods from Morocco (*Azendohsaurus*: Jalil 1999) have more recently been referred to basal archosauromorphs (Flynn *et al.* 2010). In any case, the footprint record, and the presence of the characteristic and widely distributed Upper Triassic morphotype *Brachychirotherium* in the Fleming Fjord Formation of Greenland, indicates a distinct faunal exchange of trackmakers within Triassic Pangaea.

In Late Triassic times, the Jameson Land Basin lay at 40° N in a transition zone between the relatively dry interior of the supercontinent Pangaea and the more humid peripheral part of this continent (Kent & Tauxe 2005). Much of the interior of the continent was influenced by monsoonal rain, and shallow lakes were formed. The archosaur footprints were found in a fluvial channel sandstone in the lowermost part of the Carlsberg Fjord beds during a period when the overall climate of the basin was relatively dry (Kent *et al.* 2014). The

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**Fig. 11.** (a)–(e) Sketches of *Brachychirotherium* pes and manus from different Upper Triassic localities. (a) *Brachychirotherium hassfurtense* (type ichnospecies, lectotype) from the Hassberge Formation (Late Carnian, Upper Triassic) of northern Bavaria, Germany. (b) *B. thuringiacum* from same stratum and locality as (a). (c) *B. parvum* (holotype) from the Passaic Formation (Norian, Upper Triassic) of New Jersey, USA. (d) *B. parvum* from the Redonda Formation (Norian–Rhaetian, Upper Triassic) of New Mexico, USA. (e) *B. isp.* from the Los Colorados Formation of NW Argentina. (f) cf. *Brachychirotherium* from the Ørsted Dal Member of the Fleming Fjord Formation (Upper Triassic, Norian–Rhaetian) of eastern Greenland. (g) *Batrachopus* from the Newark Supergroup (Lower Jurassic) of North America. From Klein & Lucas (2010a, b).
formation of a relatively thick channel sandstone may record intense monsoonal rain transporting sand into the marginal part of the lake system (Clemmensen et al. 2015).

The lake deposits have been shown to contain a high-resolution record of orbitally forced climate change controlling the intensity of monsoonal rain with time (e.g. Kemp & Coe 2007; Volmer et al. 2008). Previous studies of sedimentary cycles in the Late Triassic lake deposits of the Jameson Land Basin suggest that this lake system too was influenced by orbitally forced climate change (Clemmensen et al. 1998). Detailed studies of facies, colour variation, measurements of magnetic susceptibility, gamma-ray intensity and carbonate content were used in a recent study by Frobose & Clemmensen (2014) to detect statistical cyclicity in the uppermost part of the Carlsberg Fjord beds. These new data indicate, in agreement with Clemmensen et al. (1998), the existence of a composite cyclicity in the Carlsberg Fjord beds, with significant cycles. This cyclicity matches well the theoretically expected cyclicity in a lake system that experienced orbitally controlled variation in precipitation and lake environment (Clemmensen et al. 2015).

The track-bearing Carlsberg Fjord beds form part of a lake succession that indicates a long-term shift in climate from semi-arid to humid at the transition from the Norian to the Rhaetian (Clemmensen et al. 1998, 2015). This long-term shift in climate has been explained by the slow northwards drift of the continent in the Late Triassic (Clemmensen et al. 1998; Kent & Tauxe 2005).

Conclusions and perspectives

Clarification of the question of whether the herein-described tracks represent a very unusual morphotype of Brachychirotherium, possibly an early ontogenetic stage or a new ichnotaxon, must await additional data. Studies of ontogenetic growth in the autopodia of basal archosaurs based on footprints and ontogenetic scaling of limb anatomy in Alligator mississippiensis. Journal of Anatomy, 216, 423–445.


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