

# Evolution and identity of synapsid carpal bones

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To date there is little information on carpal bone homology in late Palaeozoic and Mesozoic Synapsida. Crucial to the understanding of homology in synapsid carpal elements is the fact that different nomenclatures are used for the carpals of non-mammalian Synapsida (Gegenbauer's canonical nomenclature) and Mammaliaformes (mammalian nomenclature). The homologies of the carpals of non-mammalian synapsids and mammals were established early last century and have not been reviewed since then. Here we provide a detailed study of the carpal bones of synapsids ranging in age from the early Permian to Late Cretaceous. The mammalian lunate, previously considered the homologue of the intermedium of non-mammalian synapsids, is interpreted here as homologous to their lateral centrale. We interpret the single mammalian centrale as a homologue of the medial centrale of non-mammalian synapsids. In some synapsid specimens, we found that one or two centralia are fused to the radiale (e.g., the gorgonopsian *Arctognathus* and tritylodontid *Bienotheroides*), supporting a digging habit. A third centrale is present in the therapsid *Theriongnathus*, very likely an abnormal duplication. An additional medial bone in a biarmosuchian was interpreted as a prepollex/sesamoid. A cartilaginous prepollex/sesamoid may also have been present in several non-mammalian synapsids, which have an open space proximal to distal carpal I. Distal carpal V is completely lost in dicynodonts and it is mainly fused to distal carpal IV in the adult stage of most other therapsid groups, but showed a delayed development in most non-mammalian cynodonts. In mammalian forms, distal carpal V is not present. Our observations provide an updated revision of synapsid carpal homologies, mainly on the basis of position and anatomical contacts and also taking into account the results of embryological studies.

**Key words:** Synapsida, carpus, intermedium, lunate, manus, homology, Permian, Mesozoic.

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## Introduction

The homologies of carpal bones in late Palaeozoic and Mesozoic Synapsida have not been extensively researched. Homology is a key concept for different disciplines such as taxonomy, systematics, and morphology. In some cases, a deep understanding of homologies for the interpretation of evolutionary changes can only be provided by fossils. An example of this is the proposed homology of the amniote astragalus with four tarsal bones in anamniote tetrapods (O'Keefe et al. 2006), or the established homologies of the

different phalanges in mammals and non-therapsid synapsids (Hopson 1995). Under these circumstances, providing reliable hypotheses of homology is a fundamental palaeontological task. The synapsid carpus is a complex structure consisting of many small bones, and homologizing them in fossils spanning over 170 million years is challenging. Here we investigate the homology and evolutionary change of the synapsid carpals from the early Permian to the end of the Cretaceous on the basis of currently accepted synapsid systematics.

The carpus is the proximal part of the manus (see expanded concept in SOM, Supplementary Online Material

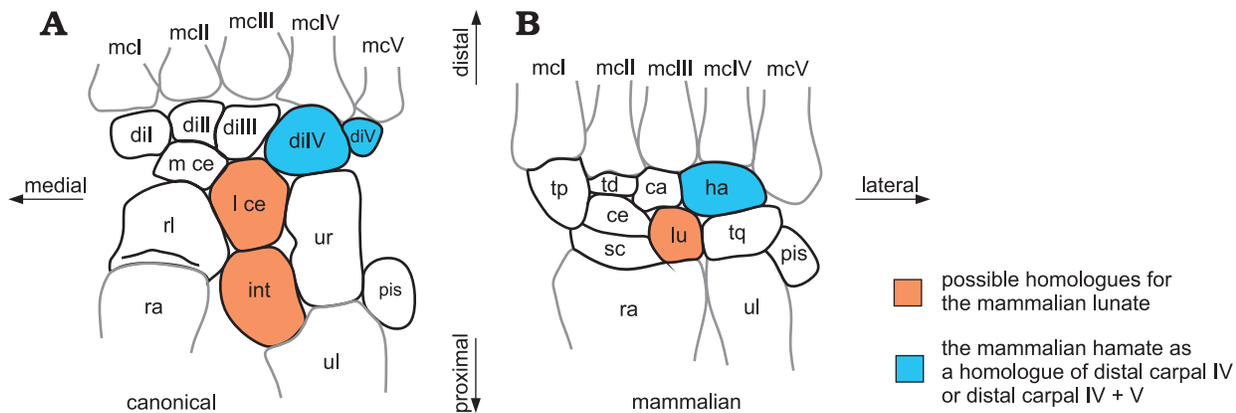


Fig. 1. Schematic diagrams of a non-therapsid synapsid (“pelycosaur”) carpus (A) and a mammaliaform carpus (B), labelled using the canonical and mammalian nomenclatures. Abbreviations: ca, capitate; ce, centrale; di, distal carpal; ha, hamate; int, intermedium; l ce, lateral centrale; lu, lunate; mc, metacarpal; m ce, medial centrale; pis, pisiform; ra, radius; rl, radiale; sc, scaphoid; td, trapezoid; tp, trapezium; tq, triquetrum; ul, ulna; ur, ulnare.

available at [http://app.pan.pl/SOM/app65-Kummell\\_et\\_al\\_SOM.pdf](http://app.pan.pl/SOM/app65-Kummell_et_al_SOM.pdf)). Traditionally, the carpals of mammals were designated using a different nomenclature than that of reptiles and amphibians. These terms were known as the “mammalian nomenclature” (sensu Shubin and Alberch 1986) and the “canonic nomenclature” (sensu Čihák 1972), respectively. The canonic nomenclature (henceforth «canonical nomenclature») is not only used for reptiles and amphibians, but also for the non-mammaliaform members of the clade Synapsida (Fig. 1A; e.g., Broom 1904; Jenkins 1971; Liu et al. 2017), whereas the carpal bones of Mesozoic Mammaliaformes are designated with the mammalian nomenclature (Fig. 1B, Table 1; e.g., Kielan-Jaworowska 1977; Ji et al. 2002; Luo et al. 2003). These different naming conventions can lead to confusion in anatomical descriptions. The use of different nomenclatures for the carpal bones of mammaliaforms and the remaining synapsids is an historical artefact. It probably arose because of the early classification of non-mammaliaform synapsids within the class Reptilia (“mammal-like reptiles”), whereas those specimens now termed Mesozoic Mammaliaformes were, as a whole, considered to be members of Mammalia in former times.

The nomenclature of the mammalian carpus was first established for the human carpals in the 17<sup>th</sup> and 18<sup>th</sup> centuries (Lyser 1653; Monro 1726; Albinus 1726; see McMurrich 1914) and later extended to describe the mammalian carpus in general. A combination of the three slightly differing nomenclature systems of Lyser (1653), Monro (1726), and Albinus (1726), together with the term central bone (centrale) are still in use to designate carpal bones of modern mammals and Mesozoic mammaliaforms (Table 1; e.g. Kielan-Jaworowska 1977; Ji et al. 2002; Luo et al. 2003). In 1864, Gegenbaur introduced the canonical nomenclature for the carpus, which was more generally applicable to all classes of vertebrates. He established homologies of the different carpal bones in all vertebrate clades and homologized the reptilian and mammalian carpals (Table 1).

The first descriptions of the carpus in non-mammaliaform synapsids were provided by Seeley (1888, 1895)

and Bardeleben (1889). These used the mammalian nomenclature more broadly and also applied it to describe the carpals of the Permo-Triassic fossil synapsids. In contrast, Broom (1901) established a homology of the Permo-Triassic synapsids and the mammalian carpals adopting the canonical and the mammalian systems to describe the proximal and central carpals of the dicynodont “*Udenodon gracilis*” (= *Dicynodontoides recuvidens*; Angielczyk et al. 2009). However, in his later publications, he only used the canonical nomenclature to describe the carpus of Permo-Triassic synapsids (Broom 1904, 1907, 1913, 1930). Broom (1901) homologized the canonical and mammalian nomenclature, following Gegenbaur (1864), but he misinterpreted the medial centrale as distal carpal I, thus identifying only one centrale in “*Udenodon*”. However, later on, he described two centralia in non-mammaliaform synapsids (Broom 1904, 1907). The carpus in non-mammaliaform synapsids has five distal carpals and two centralia, two carpals more than extant, basal mammals, which indicates that two carpal bones were lost along the evolutionary transition to mammals (Fig. 1). Broom (1901, 1904, 1907) identified one of the lost carpals as distal carpal V and proposed the second to be a centrale when he homologized the mammalian lunate (= lunar) with the intermedium.

Table 1. Homology of the reptilian and mammalian carpals after Gegenbaur (1864).

Canonical nomenclature	Mammalian nomenclature
radiale	scaphoideum
intermedium	lunatum
ulnare	triquetrum
pisiforme	pisiforme
centrale	centrale
carpale 1	mutungulatum majus/trapezium
carpale 2	mutungulatum minus/trapezoides
carpale 3	capitatum
carpale 4	hamatum
carpale 5	

The homology of the canonical and mammalian nomenclature erected by Gegenbaur (1864; see Table 1) and adopted by Broom (1901, 1904) is followed up to the present day (e.g., Ihle et al. 1927; Romer and Parsons 1977; Salomon et al. 2005; Kivell 2016) and has not been revised, even in the light of new palaeontological and embryological discoveries.

From this historical perspective, the following questions arise: do the homologies proposed by Gegenbaur (1864) between the carpals of reptiles and mammals and by Broom (1901, 1904, 1907) between non-mammalian synapsids and mammals still hold? Are there newer interpretations possible in the light of more recently collected specimens? Here we do not discuss Gegenbaur's (1864) homologies between the carpals of reptiles and mammals because of the limitations of our fossil sample. Instead, we focus only on the question of homology between the carpals of non-mammalian synapsids and basal mammals and the proposal put forward by Broom (1901, 1904). We studied carpal bones from Permian to Cretaceous Synapsida to understand carpal homology from a palaeontological perspective. In particular, we were interested in following the loss of the two carpal bones in the evolution towards mammals (Fig. 1).

The loss of distal carpal V in fossil synapsids was addressed by Hopson (1995). He suggested two variants of bone loss: non-ossification in Dicynodontia and gomphodont Cynodontia, and fusion to distal carpal IV in Biarmosuchia, Gorgonopsia, Therocephalia, and Mammalia. Here we provide new information on this topic based on additional material.

The loss of one centrale during the transition to mammals and the proposed homology between the intermedium and lunate (Broom 1901, 1904) can be questioned for two reasons: firstly, according to its position and bone contacts, the lunate must be interpreted as the lateral centrale of non-mammalian synapsids and the intermedium as the lost carpal element (Fig. 1).

The second reason for the uncertainty of intermedium-lunate homology emerges from studies of mammalian embryology. Carpals appear as chondrogenic foci in early ontogeny and may fuse or disappear later in development, providing clues about the identity of the carpal bones. There is an ongoing debate among embryologists about the identity of the mammalian lunate. Some of them interpret the mammalian lunate as a homologue of the reptilian intermedium (Steiner 1942; Schmidt-Ehrenberg 1942; Shubin and Alberch 1986; Milaire 1978; Heppleston 2010) as did Gegenbaur (1864). Others homologize the lunate with a reptilian centrale and propose that the intermedium was lost or fused to another bone of the forelimb (Holmgren 1933, 1952; Kindahl 1941, 1942a, b, 1944; Slabý 1967, 1968; Čihák 1972). This disagreement between embryologists prompts further scrutiny of each argument.

From studies on prenatal development in modern mammals, embryologists proposed that distal carpal V fused to another manual cartilaginous anlage in early ontogeny. So the loss of this bone in some mammals is indeed related to

ontogenetic fusion (Milaire 1978; Holmgren 1952; Čihák 1972; Slabý 1967). Thus, in addition to paleontological investigation, we refer to recent embryological studies for further data on carpal bone loss and putative homologies.

As well as approaching the issue of the loss of carpals in major synapsid clades, we describe carpal bone additions and losses in single species or individuals. Some synapsid fossils possess an additional central bone. This is the case in the therocephalian *Theriognathus* NHMUK R 5694 (Boonstra 1934: 260, fig. 34) and in a Russian biarmosuchian PIN 1758/320 (Chudinov 1983: 55–56, figs. 3–6). Boonstra (1934) and Chudinov (1983) interpreted the additional bone as a third centrale. In other cases, the most medial or preaxial bone was interpreted as a prepollex, as in the cases of *Theriodesmus phylarchus* NHMUK 49392 (probably a biarmosuchian, FA personal observation; Bardeleben 1889), in “*Opisthoctenodon agilis*” (Broom 1904; which represents most likely the dicynodont *Pristerodon*; Keyser 1993; Angielczyk et al. 2005) and as a probable prepollex in *Zhangheotherium* (Hu et al. 1998) and *Asioryctes* (Kielan-Jaworowska 1977; Kielan-Jaworowska et al. 2004).

Other authors described an open space medial to the central row in non-mammalian synapsids (e.g., Romer and Price 1940; Case 1907), and interpreted it as a lacuna previously occupied by a medially situated, cartilaginous sesamoid. To date, the evolutionary history of the third centralia, prepollices, and sesamoids have not been systematically researched in fossil synapsids. Here, we investigated if there were more cases of synapsid fossils with more than two centralia. We also sought evidence of a prepollex, sesamoid, or an open space on the medial side of the carpus, where these elements could have been present. Because it was not possible to distinguish between prepollices and radial sesamoids in fossils, we used the term “prepollex/sesamoid” bones for such additional preaxial bones (SOM).

In this paper, we use the canonical nomenclature for both non-mammalian synapsids and mammalian forms. This is also in accordance with the fundamental embryological studies of Schmidt-Ehrenberg (1942), Holmgren (1933, 1952), and Shubin and Alberch (1986). The only exception is the lunate of mammalian nomenclature whose identity within synapsids needs clarification. Thus, we continue using the term “lunate” in mammalian forms. The following list clarifies the terminology used here: radiale, scaphoid; ulnare, triquetrum; intermedium, intermedium of non-mammalian Synapsida; lunate, lunate of Mammalian forms; medial and lateral central, centralia in non-mammalian Synapsida; central, single centrale in Mammalian forms; distal carpal I, trapezium; distal carpal II, trapezoid; distal carpal III, capitate; distal carpal IV ( $\pm$  distal carpal V), hamate.

*Institutional abbreviations.*—AM, Albany Museum, Grahamstown, South Africa; AMNH, American Museum of Natural History, New York, USA; BP, Evolutionary Studies Institute, University of the Witwatersrand (formerly Bernard Price Institute for Palaeontological Research), Johannesburg,

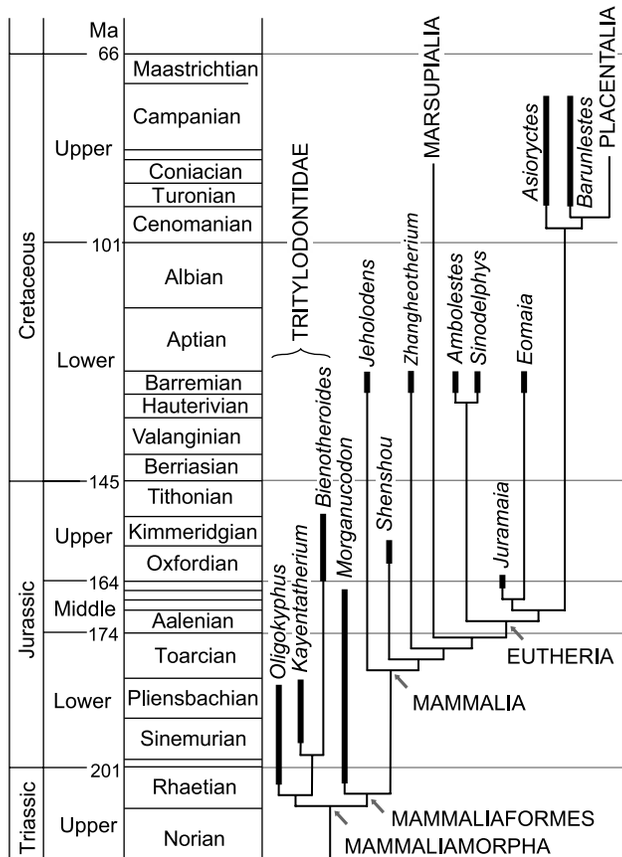
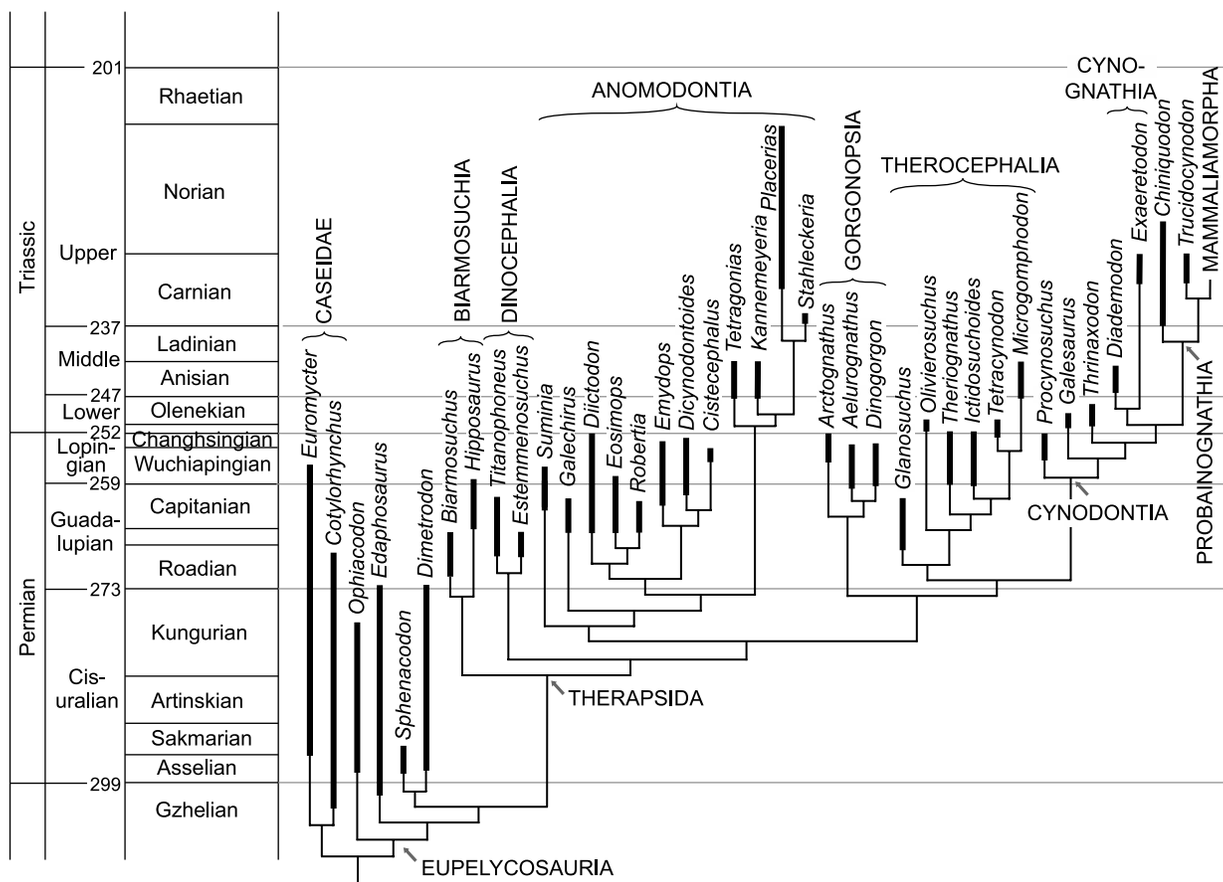


Fig. 2. Cladograms of fossil Synapsida. The cladograms are compromise trees in which different phylogenetic hypotheses were considered. Because of small space, some age names are not included in the cladogramms. These are: Wordian in the Guadalupian; Induan in the Lower Triassic; Hettangian in the Early Jurassic; Bajocian, Bathonian and Callovian in the Middle Jurassic and Santonian in the Late Cretaceous. Geological timescale after Cohen et al. (2013, updated), correlations after Schneider et al. (2020). **A.** Non-therapsid Synapsida (after Reisz 1986; Modesto et al. 2011; Reisz et al. 2011), Dinocephalia (after Kammerer 2011), Anomodontia (after Maisch 2001; Vega-Dias et al. 2004; Fröbisch and Reisz 2011; Kammerer et al. 2011; Angielczyk and Rubidge 2013), Gorgonopsia (after Kammerer 2016; Kammerer and Masyutin 2018; SK personal communication with Christian Kammerer 2019), Terocephalia (after Huttenlocker and Smith 2017), non-mammaliomorph Cynodontia (after Abdala 2007; Ruta et al. 2013). **B.** Mammaliomorpha (after Kielan-Jaworowska et al. 2004; Watabe et al. 2007; Luo et al. 2011; Ruta et al. 2013; Bi et al. 2014, 2018; Velazco et al. 2017).

South Africa; CAGS, Chinese Academy of Geological Sciences, Beijing, China; CGS, Council for Geosciences, Pretoria, South Africa; FMNH, Field Museum of Natural History, Chicago, USA; GMV, National Geological Museum of China, Beijing, China; GPIT, Paleontology Department and Museum, Institute of Geosciences, Eberhard Karls University, Tübingen, Germany; IVPP, Institute of Vertebrate Paleontology and Paleoanthropology, Beijing, China; MCZ, Museum of Comparative Zoology, Harvard University, Cambridge, USA; MNHN.F, Muséum national d'Histoire naturelle, collection de Paléontologie, Paris, France; NMQR, National Museum, Bloemfontein, South Africa; NHMUK, Natural History Museum, London, UK; OMNH, Oklahoma Museum of Natural History, Norman, USA; OUMNH-TSK, Oxford University Museum, T.S. Kemp Collection, Oxford, UK (material now deposited in the NHMUK); PIN, Paleontological Institute, Russian Academy of Sciences, Moscow, Russia; PVL, Colección Palaeontología de Vertebrados Lillo, Universidad Nacional de Tucumán, Argentina; RC, Rubidge Collection, Wellwood, Graaff-Reinet, South Africa; SAM, Iziko South African Museum, Cape Town, South Africa; TM, Northern Flagship Institution, Transvaal Museum, Pretoria, South Africa; TMM, Texas Memorial Museum, Austin, USA; UFRGS, Universidade Federal do Rio Grande do Sul, Porto Alegre, Brazil; UMZC, University Museum of Zoology, Cambridge, UK; USNM, National Museum of Natural History, Washington D.C., USA; WCW, Wucuiwan field collection, housed at the IVPP.

## Material and methods

Sixty-four synapsid specimens from 39 genera were studied. Most of the specimens, including some casts, were examined directly. The rest of the data was obtained from photographs, scans or from publications, shown in the following list. Here, we usually use the generic name for specimens. The specimen number (museum accession number) is provided, when detailed observations were made on just one specimen of a genus or species with several specimens. Accession numbers are also listed, when the identity of the fossil was uncertain. The taxonomy of the Gorgonopsia is based on the research of Christian Kammerer (SK personal communication 2019).

Caseidae: *Euromycter rutenus* MNHN.F.MCL-2 (Sigogneau-Russell and Russell 1974: fig. 18), *Cotylorhynchus romeri* OMNH 00655 (Stovall et al. 1966: fig. 13).

Non-therapsid Eupelycosauria: *Ophiacodon retroversus* FMNH UC 458, MCZ 1203, *Ophiacodon mirus* FMNH UC 671 (cast), *Edaphosaurus boanerges* NHMUK R 9204 (cast), *Sphenacodon ferox* CM 76895 (Henrici et al. 2005: photo, fig. 1), *Dimetrodon milleri* MCZ 1365 (cast).

Biarmosuchia: Biarmosuchidae indet. PIN 1758/320, *Hipposaurus major* SAM-PK-9081.

Dinocephalia: *Titanophoneus potens* PIN 157/1, *Estemmenosuchus uralensis* PIN 1758/23.

Anomodontia: *Suminia getmanovi* PIN 2212/62 (Fröbisch and Reisz 2011: figs. 9, 12), *Galechirus scholtzi* SAM-PK-1068, AMNH 5516, *Eosimops newtoni* BP/1/6674, *Robertia broomiana* SAM-PK-11885a, b, *Diictodon feliceps* CGS FL186, TM 4991, UMZC T 420, GPIT/RE/7193, SAM-PK-K10699, CGS RMS214, CGS T72, SAM-PK-K10636, *Cistecephalus microrhinus* BP/1/2124, BP/1/2915, *Kannemeyeria simocephalus* NHMUK R 3741, *Stahleckeria potens* MCZ 1688.

Gorgonopsia: *Arctognathus curvimola* SAM-PK-3329, *Aelurognathus tigriceps* SAM-PK-2342, cf. *Cynariops robustus* SAM-PK-K10000, *Dinogorgon rubidgei* BP/1/2190, Gorgonopsia indet. BP/1/1210.

Therocephalia: *Glanosuchus macrops* SAM-PK-K7809, SAM-PK-12051, CGS RS424, *Olivierosuchus parringtoni* BP/1/3973, BP/1/3849, *Theriognathus microps* NHMUK R 5694, *Ictidosuchoides longiceps* CGS CM86-655, *Ictidosuchoides longiceps* or *Ictidosuchops intermedium* BP/1/2294, *Tetracynodon darti* AM 3677, BP/1/2710, *Microgomphodon oligocynus* SAM-PK-K10160.

Non-mammalian Cynodontia: *Procynosuchus delaharpeae* BP/1/591, NHMUK PV R 37054 (formerly OUMNH TSK 34), RC92, *Galesaurus planiceps* BP/1/2513, SAM-PK-K10465, SAM-PK-K10468, *Thrinaxodon liorhinus* BP/1/1737, BP/1/7199 (CT-scan), *Diademodon tetragonus* NHMUK R-3581, USNM 23352, Cynognathia indet. BP/1/4534, *Exaeretodon argentinus* PVL 2554, *Trucidocynodon riograndensis* UFRGS PV-1051T.

Basal Mammalianmorphea: Tritylodontidae indet. WCW-06A-34, *Bienotheroides wanhsienensis* IVPP V 7905, *Kayentatherium wellsi* TMM 43690-5.136 (scan, Eva Hoffman, see also Hoffman and Rowe 2018: supplement).

Mammaliaformes: *Jeholodens jenkinsi* GMV 2139a (original and cast), *Zhangheotherium quinquecuspidens* IVPP V7466, *Eomaia scansoria* CAGS01-IG-1a (cast and Luo et al. 2003: fig. 2).

Mammalian embryological studies were considered alongside paleontological observations to provide a thorough basis for testing homologies. Carpal bone homologies were identified in the articulated fossil carpi on the basis of position, relationship of elements to each other and sequential order. Other features (e.g., relative size and shape) played a subordinate role in homology assessment. In disarticulated carpi, relative size, shape and articular facets were used for bone identifications. Because of the high morphological variation of carpal elements, when possible we used articulated, complete to nearly complete carpi with elements in their original position. Using these specimens, we assessed the major evolutionary changes in carpal bones in synapsids. Fused carpals were recognized by fusion lines and/or irregular shapes with set-back angles and indentations. Open spaces in the articulated skeletons may represent unfossilised cartilaginous precursors and were also considered in our interpretations.

To assess the evolution of characters, we mapped them onto phylogenetic trees using Mesquite 3.61 (Maddison and Maddison 2019; see Discussion and SOM: figs. 1–3).

## Description of the synapsid carpus

The descriptions of the bones are mostly presented in dorsal view. However, the dorsal surfaces of some of the fossils studied were not exposed. Such specimens are described in ventral view. The description is made in zero position (sensu Kümmell and Frey 2014b). In zero position the carpus is flat without a transverse arch and the rays are longitudinally aligned and not spread (ray = digit and corresponding metacarpal; Biesecker et al. 2009).

Carpal bones in synapsids sometimes show true joints with a great range of mobility, but usually they are connected by amphiarthroses, implying only a small mobility range. Joints were not exposed in all the partially prepared specimens; therefore we do not distinguish between plane contacts, real articular facets and amphiarthroses.

The carpus of Tritylodontidae resembles that of Mammaliaformes more than that of non-mammalian Synapsida. Thus, we describe the tritylodontid carpus together with species of Mammaliaformes. Tritylodontidae and Mammaliaformes form the clade Mammalianomorpha (Rowe 1988; Luo 2011), and our descriptions discriminate between non-mammalian synapsids and mammalianomorphs.

**Non-mammalian Synapsida.—Radiale:** In dorsal view, the radiale appears either square, irregularly rectangular (transversely orientated) or irregularly pentagonal (Figs. 3A–8A; SOM: table 1: a). In the pentagonal radiale, the lateral and medial margins are parallel to each other, followed distally by two bevelled edges. The radiale shows a wide proximal facet, which occupies the whole distal facet of the radius (SOM: table 1: b). In some Dicynodontia, Gorgonopsia and Therocephalia, the radiale is slightly convex proximally. In species with a quadrangular radiale, the radiale contacts the medial centrale on its distal border. In species with a pentagonal radiale, the bevelled distal edges are the facets for the medial centrale distomedially to distally and the lateral centrale distolaterally (Figs. 4A, B, 6D; SOM: table 1: c–e). The lateral centrale usually lies distolaterally or laterally to the radiale (SOM: table 1: e, f) and distally to the intermedium, which usually protrudes proximally beyond the proximal border of the radiale. In non-mammalian Synapsida, the lateral centrale always contacts the lateral margin of the radiale (Fig. 7; SOM: table 1: f).

**Intermedium:** In non-therapsid Synapsida, the intermedium can be broad and square (e.g., *Euromycter* and *Ophiacodon* MCZ 1203; Fig. 3), rectangular with a proximodistal elongation (e.g., *Ophiacodon* FMNH UC 458, FMNH UC 671) or pentagonal (e.g., Sphenacodontidae; SOM: table 2: a). As reported previously, the “pelycosaurian” intermedium is thin dorsoventrally, convex dorsally and concave ventrally (Romer and Price 1940; Henrici et al. 2005). The “pelycosaurian” intermedium is larger than that of non-mammalian therapsids, i.e., wider in relation to its length (square or pentagonal) and/or longer in relation to the radiale (SOM: table 2: a, e). In “pelycosaurs”, the intermedium is either lon-

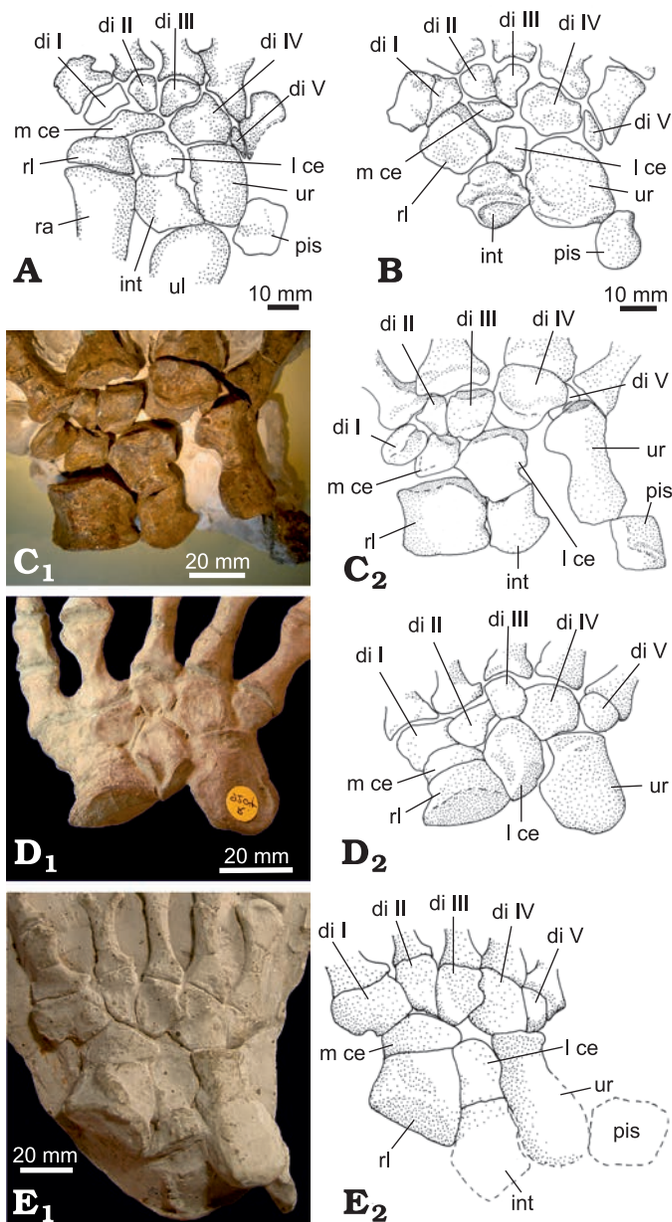


Fig. 3. Carpus of non-therapsid Synapsida. **A.** *Euromycter rutenus* (Sigogneau-Russell and Russell, 1974), MNHN.F.MCL-2, Valady, France, Sakmarian, left carpus (reversed), dorsal view (redrawn from Sigogneau-Russell and Russell 1974: fig. 18). **B.** *Cotylorhynchus romeri* Stovall, 1937, OMNH 00655, Navina, USA, Kungurian, left carpus (reversed), dorsal view (redrawn from Stovall et al. 1966: fig. 13, left). **C.** *Ophiacodon retroversus* Cope, 1878, MCZ 1203, Rattlesnake Canyon, USA, Wichita Group, Cisuralian, right carpus, dorsal view. **D.** *Edaphosaurus boanerges* Romer and Price, 1940, NHMUK R 9204 (cast), Geraldine, Archer County, USA, Wichita Group, Cisuralian, left carpus (reversed), dorsal view. **E.** *Dimetrodon milleri* Romer, 1937, MCZ 1365 (cast), Archer, USA, Putnam Formation, Cisuralian, right carpus, dorsal view. Photographs (C<sub>1</sub>–E<sub>1</sub>) and interpretative drawings (C<sub>2</sub>–E<sub>2</sub>). Abbreviations: di, distal carpal; int, intermedium; l ce, lateral centrale; m ce, medial centrale; pis, pisiform; ra, radius; rl, radiale; ul, ulna; ur, ulnare.

ger or the same length as the radiale, whereas in therapsids, it is the same length or shorter than the radiale. The subadult therapsid *Diictodon* CGS FL186 is the only exception known

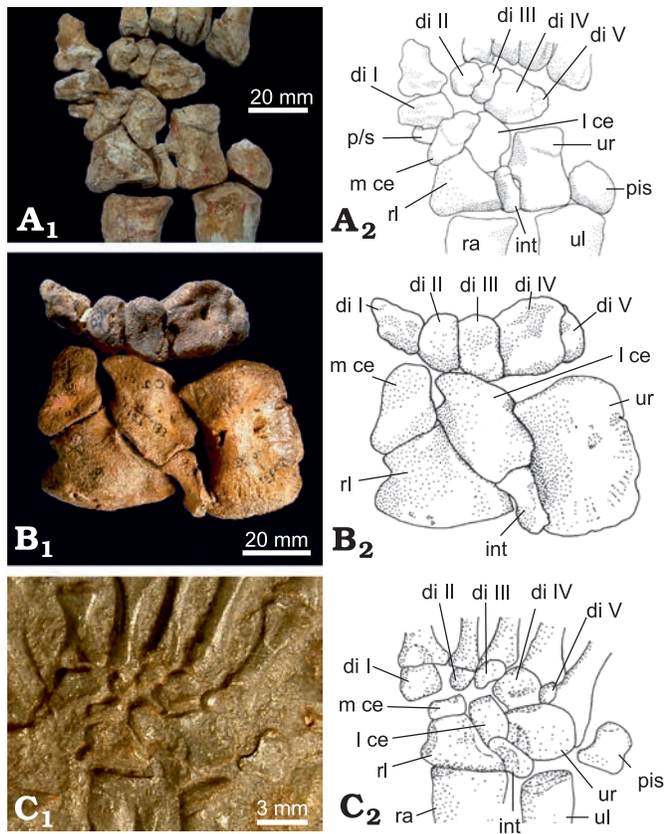


Fig. 4. Carpus of basal Therapsida. **A.** A biarmosuchian PIN 1758/320, Eshovo, Russia, Roadian, Guadalupian, right carpus, slightly disarticulated, dorsal view. **B.** Dinocephalian *Titanophoneus potens* Efremov, 1940, PIN 157/1, Ishevo, Russia, Capitanian, left carpus (reversed), dorsal view. **C.** Basal anomodontian *Galechirus scholtzi* Broom, 1907, SAM-PK-1068, Victoria West, South Africa, Capitanian, right carpus, dorsal view (impression). Photographs (A<sub>1</sub>–C<sub>1</sub>) and interpretative drawings (A<sub>2</sub>–C<sub>2</sub>). Abbreviations: di, distal carpal; int, intermedium; l ce, lateral centrale; m ce, medial centrale; pis, pisiform; p/s, prepollex/sesamoid; ra, radius; rl, radiale; ul, ulna; ur, ulnare.

to us where the intermedium is longer than the radiale (SOM: table 2: e). The non-mammalian therapsid intermedium is lateromedially narrow and dorsoventrally deep. It is a proximodistally oriented, rectangular to hourglass-shaped slender bone (Figs. 4A–8A), or bean-shaped in *Kannemeyeria* and *Galesaurus* BP/1/2513 (Fig. 7C; SOM: table 2: a). Proximally, the intermedium articulates with the ulna (SOM: table 2: b). Distally it contacts the lateral centrale (SOM: table 2: c). Laterally, it is articulated with the ulnare and medially with the radiale and/or the lateral side of the distal end of the radius (SOM: table 2: d). In a few cases, it only contacts the radiale medially (Biarmosuchidea indet. PIN 1758/320, Fig. 4A, *Estemmenosuchus*, *Stahleckeria*, *Glanosuchus* SAM-PK-12051 and CGS RS424, *Theriognathus* and *Ictidosuchoides* CGS CM86-655, Fig. 6D). In Caseidae (Fig. 3A, B) and some single specimens of other groups (*Edaphosaurus*, *Procynosuchus* RC92, Fig. 7A, *Exaeretodon*, and probably *Cistecephalus*), the intermedium is situated even further proximally than in the other synapsids and lies laterally to the radius (SOM: table 2: d).

*Ulnare*: The ulnare is the longest bone of the non-mammalian synapsid carpus, only in some Dicynodontia (*Cistecephalus* and *Stahleckeria*; Fig. 5C), it is approximately the same size as the radiale. In most specimens, it is approximately rectangular and proximodistally elongated (Figs. 4A–8A). Sometimes it is proximally rounded (e.g., in “pelycosaurs”, Fig. 3). In most Theroccephalia it is hourglass shaped (Fig. 6B; SOM: table 3: a). The lateral margin of the ulnare is dorsoventrally thin, while the medial margin is dorsoventrally thicker and usually curved laterally. The ulnare has a complex, mainly convex articular surface on the medial side, which articulates with the intermedium and the lateral centrale. These articulations are often covered by adjacent bones or matrix, and therefore usually not visible. However, in some fossils (the biarmosuchian PIN 1758/320, *Titanophoneus*, *Glanosuchus* CGS RS424, *Procynosuchus* NHMUK PV R 37054, and *Thrinaxodon* BP/1/7199), the facet is exposed, showing a medioventrally pointing triangular process close to the mid-point of the bone’s medial margin. In *Cistecephalus* and one *Diictodon* specimen

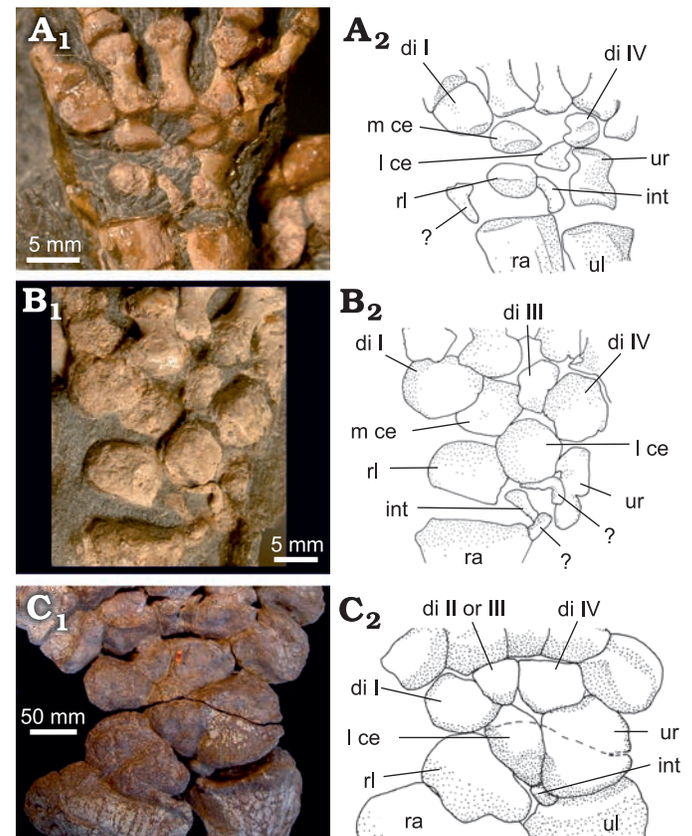


Fig. 5. Carpus of Dicynodontia. **A.** *Diictodon feliceps* (Owen 1876), CGS FL186, Jasfontein, Victoria West, South Africa, *Tropidostoma* Assemblage Zone, Wuchiapingian, right carpus, dorsal view. **B.** *Eosimops newtoni* Broom, 1921, BP/1/6674, Somerfontein, Philipolis District, South Africa, *Pristerognathus* Assemblage Zone, Capitanian–Wuchiapingian, left carpus, ventral view. **C.** *Stahleckeria potens* von Huene, 1935, MCZ 1688, Candelaria, Brazil, Santa Maria Formation, Carnian, right carpus, dorsal view. Photographs (A<sub>1</sub>–C<sub>1</sub>) and interpretative drawings (A<sub>2</sub>–C<sub>2</sub>). Abbreviations: di, distal carpal; int, intermedium; l ce, lateral centrale; m ce, medial centrale; pis, pisiform; ra, radius; rl, radiale; ul, ulna; ur, ulnare. Dotted line, fracture.

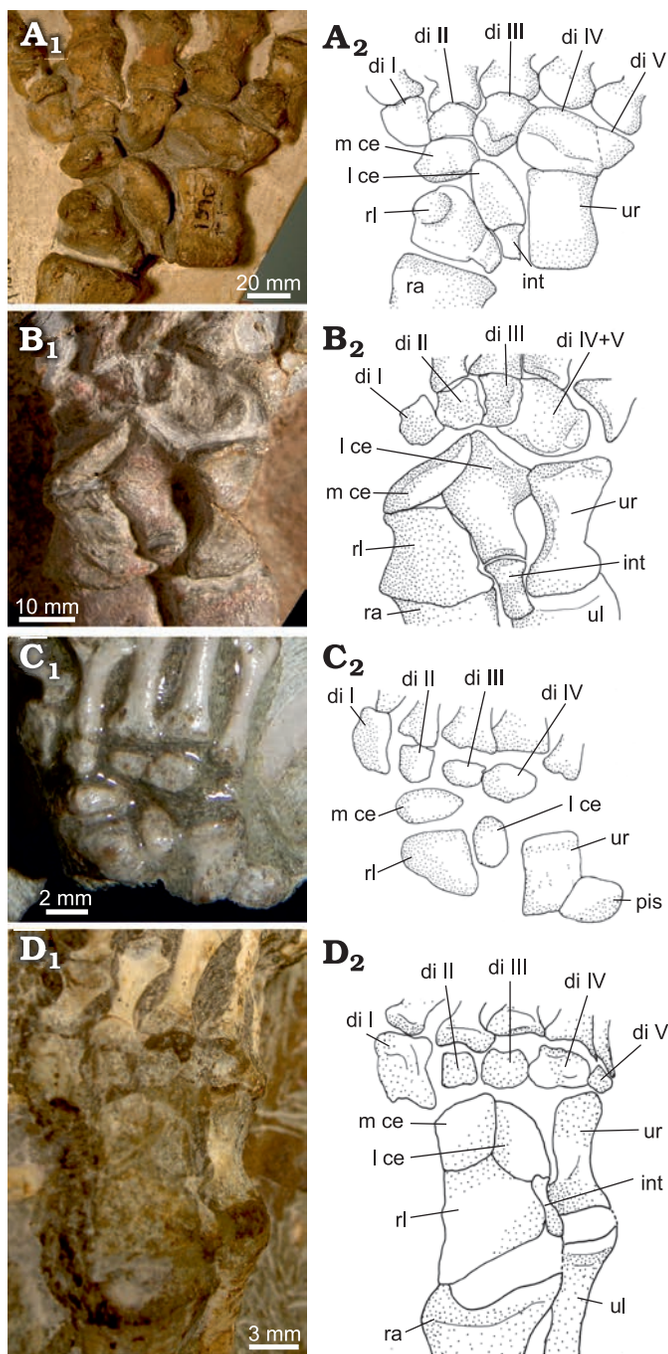


Fig. 6. Carpus of Gorgonopsia and Therocephalia. **A.** Gorgonopsia indet. BP/1/1210, Hoeksplaas, Murraysburg, South Africa, *Daptocephalus* Assemblage Zone, Changhsingian, left carpus (reversed), dorsal view. **B.** Basal therocephalian *Glanosuchus macrops* Broom, 1904, SAM-PK-K7809, La-de-da, Beaufort West, South Africa, *Tapinocephalus* Assemblage Zone, Capitanian, left carpus (reversed), dorsal view. **C.** Therocephalian *Tetracyonodon darti* Sigogneau, 1963, AM 3677, farm Carlton Heights, Pixley ka Seme District, South Africa, *Lystrosaurus* Assemblage Zone, Early Triassic, right carpus, dorsal view (photo courtesy of Gabriela Fontanarrosa). **D.** Therocephalian *Ictidosuchooides longiceps* Broom, 1920, CGS CM86-655, Secretaris Kraal 19, Murraysburg, South Africa, *Daptocephalus* Assemblage Zone, Changhsingian, right carpus, dorsal view. Photographs (A<sub>1</sub>–D<sub>1</sub>) and interpretative drawings (A<sub>2</sub>–D<sub>2</sub>). Abbreviations: di, distal carpal; int, intermedium; l ce, lateral centrale; m ce, medial centrale; pis, pisiform; ra, radiale; rl, radiale; ul, ulna; ur, ulnare.

(CGS T72), the medial border of the ulnare only contacts the intermedium (SOM: table 3: c). Distally and sometimes distomedially, the contacting surface of the ulnare receives the distal carpal IV. Distally, in some cases it also receives distal carpal V (SOM: table 3d, e). Distolaterally, the ulnare contacts distal carpal V (e.g., most “pelycosaur”, and *Galechirus* SAM-PK-1068, *Ictidosuchooides* CGS CM86-655, *Procynosuchus* NHMUK PV R 37054, *Thrinaxodon* BP/1/7199), metacarpal V (usually in Dicynodontia, *Glanosuchus* CGS RS424, *Procynosuchus* RC92), or leaves an open space between its distolateral margin and metacarpal V (many cases of non-mammalian therapsids; SOM: table 3: f). Proximally the ulnare has a broad articulation area contacting the ulna (SOM: table 3: g).

**Pisiform:** The pisiform is often missing in synapsid fossils, especially in most Dicynodontia, Gorgonopsia and Therocephalia. From the 49 non-mammalian synapsid carpi studied here, in which the typical place of the pisiform lateral of the wrist joint is well exposed, 26 lack an associated pisiform (SOM: table 4: a). However, in all synapsid groups, at least some specimens possess a pisiform, suggesting that it is usually present (Figs. 3–5, 7, 8A), but probably easily lost during fossilisation. The pisiform is a subcircular to oval bone, but can be square-shaped in some basal synapsids (*Euromycter*, *Ophiacodon*, and *Dimetrodon*), or sickle-shaped as in the cynodont *Trucidocynodon* (SOM: table 4: a). It is usually positioned close to the proximolateral border of the ulnare and the distolateral margin of the ulna (SOM: table 4: b).

**Lateral centrale:** The outline of the lateral centrale can be square or rectangular in proximodistal orientation, subcircular or oval and rhomboid. Nearly all forms occur in most therapsid groups (Figs. 3A–8A; SOM: table 5: a). In non-mammalian Cynodontia, it is usually longer than the radiale, only in *Procynosuchus*, it has the same length (SOM: table 5: b). Proximally it has a facet for the intermedium, distally it contacts distal carpal III, sometimes also partly distal carpals II or IV. Only in the Caseidae *Euromycter* and *Cotylorhynchus*, it is distally articulated to the medial centrale (Fig. 3A, B; SOM: table 5: c). Distomedially, the lateral centrale articulates with the medial centrale and/or distal carpal II (SOM: table 5: d). Distolaterally it articulates with distal carpal IV (SOM: table 5: e). Proximomedially or medially, the lateral centrale is bordered by the radiale (SOM: table 1: e, f) and laterally by the ulnare (SOM: table 3: c).

**Medial centrale:** The outline of the medial centrale is mostly irregularly oval, rectangular or rhomboid (Figs. 3–7). In *Trucidocynodon*, it is triangular (Fig. 8A; SOM: table 6: a). Proximally, or occasionally proximolaterally, it is articulated with the radiale, proximolaterally or laterally with the lateral centrale. Only caseids show a proximal contact to the lateral centrale (SOM: table 6: b–d). In non-mammalian cynodonts, the radiale is always proximal to the medial centrale as is also the case with the single centrale in mammalian forms (SOM: table 6: b; see below). The medial

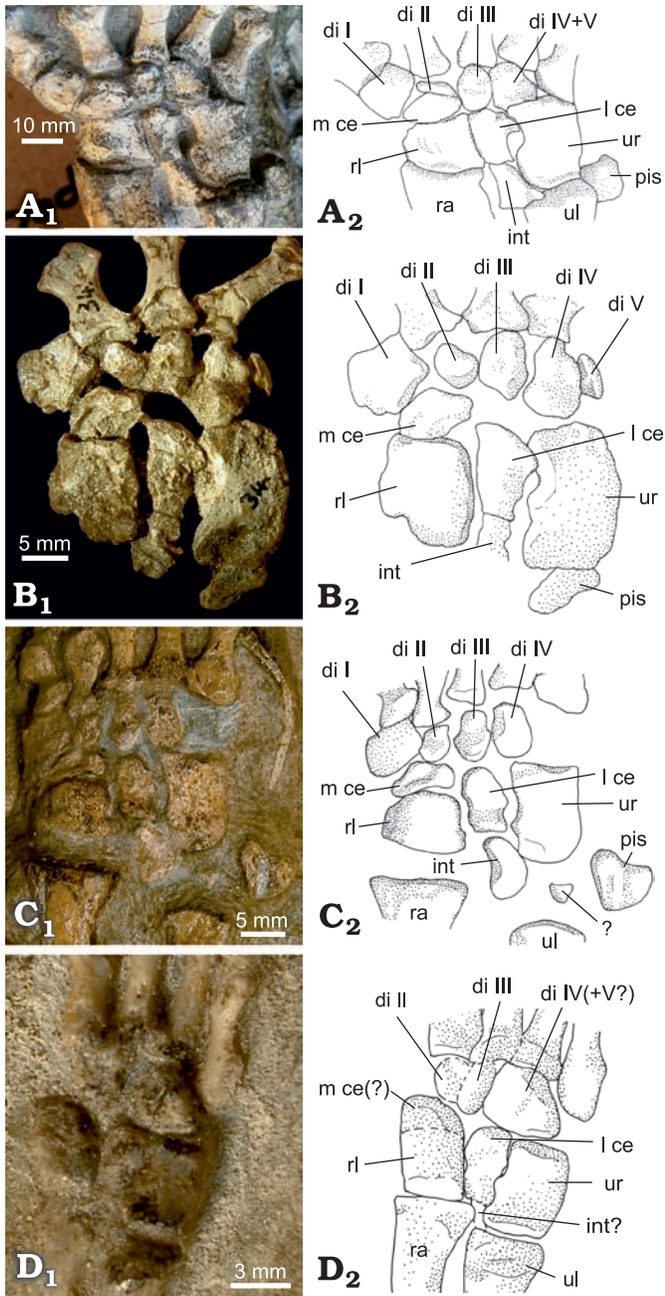


Fig. 7. Carpus of non-mammalian cynodonts. **A.** *Procynosuchus delaharpeae* Broom, 1937, RC92, Doornkloof, Graaff-Reinet, South Africa, *Cistecephalus* Assemblage Zone, Lopingian, right carpus, dorsal view. **B.** *Procynosuchus delaharpeae* Broom 1937, NHMUK PV R 37054, Middle Luangwa Valley, Zambia, *Daptocephalus* Assemblage Zone, Changshingian, right carpus, dorsal view. **C.** *Galesaurus planiceps* Owen 1860, BP/1/2513, Honingkrans, Burgersdorp, South Africa, *Lystrorhynchus* Assemblage Zone, Early Triassic, left carpus (reversed), dorsal view. **D.** *Cynognathia* indet. BP/1/4534, Hugoskop 620, Roxville, South Africa, *Cynognathus* Assemblage Zone, Olenekian–Anisian, left carpus (reversed), dorsal view. Photographs (A<sub>1</sub>–D<sub>1</sub>) and interpretative drawings (A<sub>2</sub>–D<sub>2</sub>). Abbreviations: di, distal carpal; int, intermedium; l ce, lateral centrale; m ce, medial centrale; pis, pisiform; ra, radius; rl, radiale; ul, ulna; ur, ulnare.

centrale is usually adjacent to the distal carpals II and I in non-mammalian synapsids, but often extends to meet distal carpal III at its distolateral edge (SOM: table 6: e).

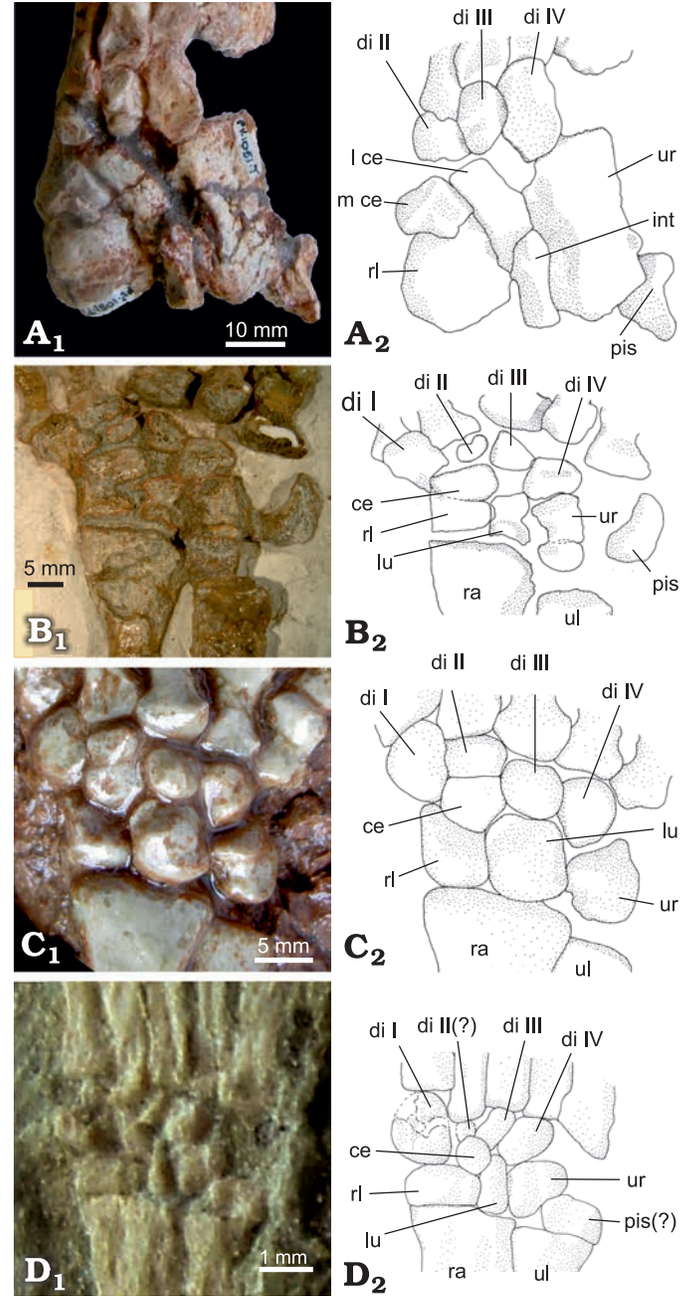


Fig. 8. Carpus of a non-mammalian cynodont and three Mammalia-morpha. **A.** *Trucidocynodon riograndensis* Oliveira, Soares, and Schultz, 2010, UFRGS PV-1051T, Sitio Janner, Rio Grande do Sul, Brazil, Santa Maria Formation, Carnian, right carpus, dorsolateral view. **B.** *Bienotheroides wanhsienensis* Young, 1982, IVPP V 7905, Sichuan, China, Middle to Late Jurassic, right carpus, dorsal view. **C.** Tritylodontidae indet. WCW-06A-34, Wucaiwan, Junggar Basin, northwestern China, Shishugou Formation, probably Oxfordian, right carpus, dorsal view. **D.** *Jeholodens jenkinsi* Ji, Luo, and Ji, 1999, GMV 2139a, Sihetun, Liaoning Province, China, late Barremian, left carpus (reversed), dorsal view. Photographs (A<sub>1</sub>–D<sub>1</sub>) and interpretative drawings (A<sub>2</sub>–D<sub>2</sub>). Abbreviations: di, distal carpal; int, intermedium; l ce, lateral centrale; m ce, medial centrale; pis, pisiform; ra, radius; rl, radiale; ul, ulna; ur, ulnare.

*Three centralia:* Besides the two centralia, a third bone is present in the central row of the theroccephalian *Theriongnathus microps* NHMUK R 5694 (Fig. 9). We do not know

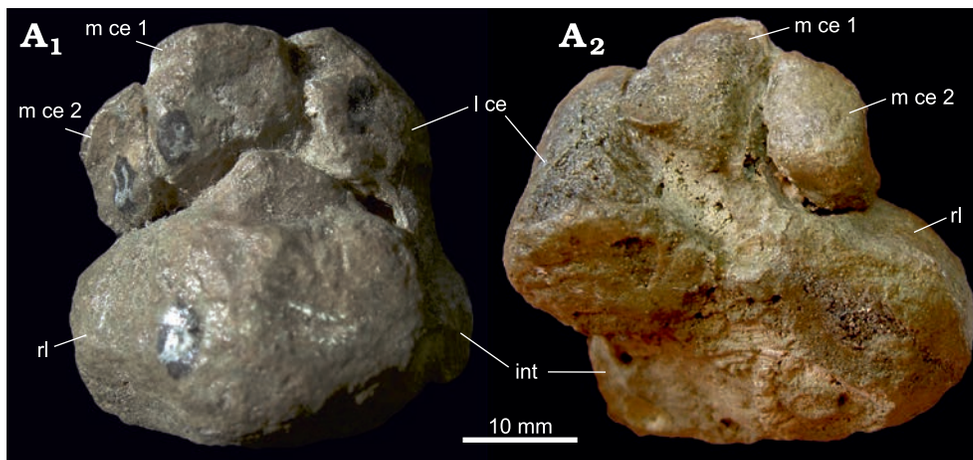


Fig. 9. Medial part of the carpus of *Theriognathus microps* Owen, 1876, NHMUK R 5694, Thaba 'Nchu, South Africa, *Cistecephalus* Assemblage Zone, Lopingian, with three centralia. Dorsal (A<sub>1</sub>) and distoventral (A<sub>2</sub>) views. Abbreviations: int, intermedium; l ce, lateral centrale; m ce, medial centrale; rl, radiale.

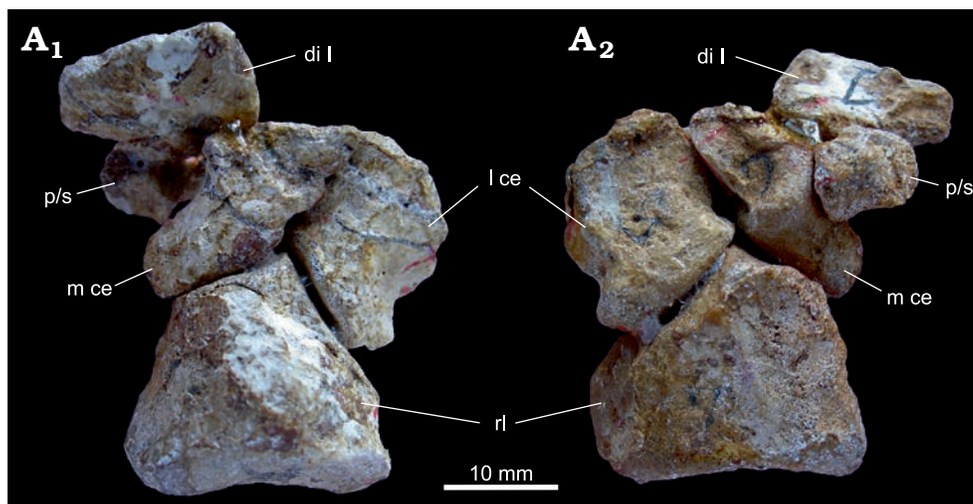


Fig. 10. Prepollex/sesamoid on the carpus of the biarmosuchian PIN 1758/320, Eshovo, Russia, Roadian. Dorsal (A<sub>1</sub>) and ventral (A<sub>2</sub>) views. Abbreviations: di I, distal carpal I; l ce, lateral centrale; m ce, medial centrale; rl, radiale; p/s, prepollex/sesamoid.

if this condition is an individual variant or common also in other specimens of the species *Theriognathus microps*. The lateral centrale is located at its common position, proximal to distal carpal III and distal to the intermedium. Medial to the lateral centrale are two small bones, closely connected to each other with tight fitting articular surfaces (Fig. 9). Together, both medially situated central bones have the general oval outline of the medial centrale and occupy the same position distal/distomedial of the radiale and proximal to distal carpals II and to a small portion of distal carpal I. All three central bones are in articulation with the radiale. There is an empty space medial to the medialmost central bone, between the medial half of the proximal margin of distal carpal I and the distomedial corner of the radiale.

**Prepollex/sesamoid:** There are three central bones present in the biarmosuchian PIN 1758/320. As in *Theriognathus* NHMUK R 5694, the lateral centrale is in the usual position, between distal carpal III distally and the intermedium proximally. The position of the two medially situated central bones is, however, different from that in *Theriognathus* NHMUK R 5694. Adjacent to the lateral centrale is a normal medial centrale occupying the whole distal/distomedial facet of the radiale. It is distally connected to distal carpal

II and the lateral part of the proximal margin of distal carpal I. The third central bone in PIN 1758/320 is oval with an approximate lateromedial orientation of its long axis. It does not articulate with the radiale, but lies on the junction of distal carpal I and the medial centrale, slightly ventral to both, so that its lateral and distolateral border underlies these carpal bones (Figs. 4A, 10). The medial section shows a free ending. In our view, this represents a prepollex/sesamoid, interpreted as additional preaxial bone (see SOM).

In many synapsid fossils, there is an open space proximal/proximomedial to distal carpal I. Usually only the proximolateral side of distal carpal I is articulated with the medial centrale (see section "Distal carpal I" below). The open space is at the position of the prepollex/sesamoid in the biarmosuchian PIN 1758/320, leaving open the possibility that this space was occupied by a cartilaginous prepollex/sesamoid in these species.

**Loss or fusion of centralia:** In the dicynodont *Stahleckeria* MCZ 1688, only one centrale is present (Fig. 5C). This centrale contacts the intermedium proximally and is placed distolateral to the radiale and medial to the ulnare. Distally, distolaterally and distomedially it is connected to distal carpal I, the central distal carpal (II or III) and distal carpal IV

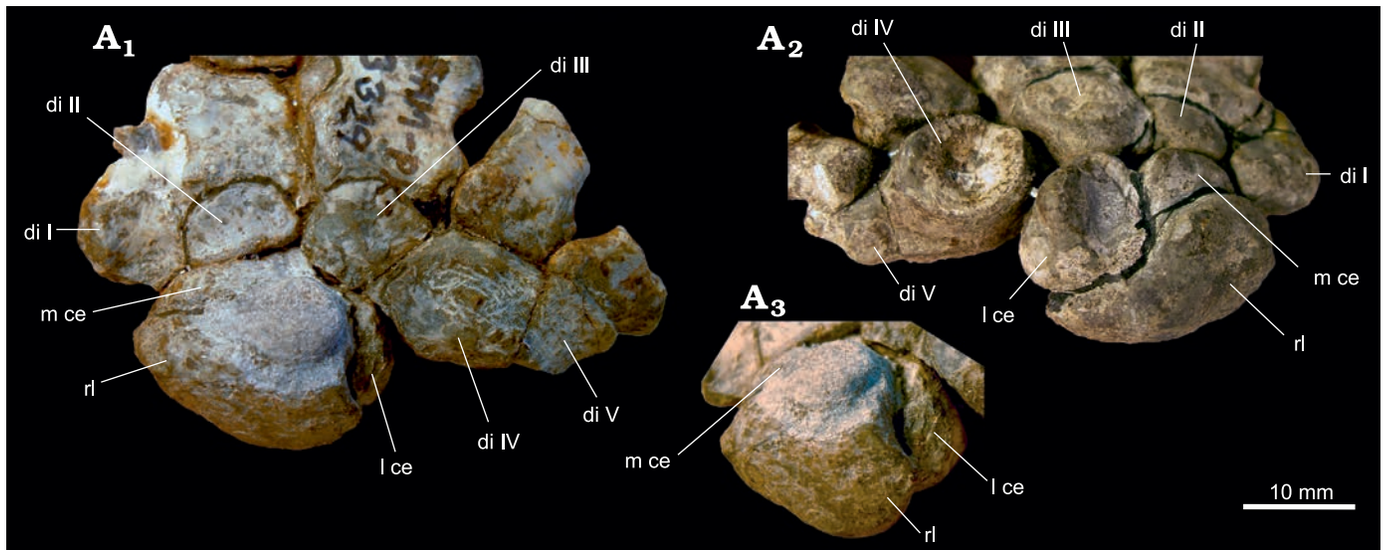


Fig. 11. Part of the carpus of *Arctognathus curvimola* (Owen, 1876), SAM-PK-3329, Oudeberg, Graaff-Reinet, South Africa, *Cistecephalus* Assemblage Zone, Lopingian. Dorsal (A<sub>1</sub>), ventral (A<sub>2</sub>), and dorsolateral (A<sub>3</sub>) views, showing the fusion lines between the radiale and the centralia. Abbreviations: di, distal carpal; l ce, lateral centrale; m ce, medial centrale; rl, radiale.

and can be unequivocally identified as the lateral centrale. The distal border of the radiale is connected to distal carpal I, leaving no space for the medial centrale.

In the gorgonopsian *Arctognathus curvimola* SAM-PK-3329, the centralia are fused to the radiale (Fig. 11). The fusion lines between the centralia and the radiale are clearly present and there are small set-back angles in the junctions between the different fused bones. In dorsal view, the centralia are proximodistally short, but widen ventrally (compare Fig. 11A<sub>1</sub> and A<sub>2</sub>).

In an unidentified cynognathian cynodont BP/1/4534, only the lateral centrale is present as a separate bone (Fig. 7D). The medial centrale is probably fused to the radiale. This fusion is visible as a faint lateromedially orientated line of coalescence. The radiale (in probable fusion with the medial centrale), is approximately the same length as the ulnare. This is uncommon among non-mammalian synapsids, where the ulnare is usually longer than the radiale (Fig. 7). This is an additional suggestion that the medial centrale is fused to the radiale in this specimen. In *Thrinaxodon* BP/1/1737, the medial centrale was also probably fused to the radiale. However, in *Thrinaxodon* BP/1/7199, the bones were separate in both manus. So, the fusion in BP/1/1737 could represent an individual variant.

**Distal carpal I:** The first distal carpal is usually square, oval or rectangular, with a mostly mediolaterally orientated long axis (Figs. 3–7; SOM: table 7: a). In non-therapsid Synapsida and in the anomodont *Galechirus*, distal carpal I is aligned with the row of distal carpals. In some therapsids, however, it lies more distally, medial to the proximal portion of the row of metacarpals as in the dicynodonts *Robertia* and *Diictodon* (Fig. 5A; SOM: table 7: b). In most other non-mammalian synapsids, distal carpal I has an intermediate position. Its proximal portion is aligned with the row of distal carpals and the bone protrudes distally on the medial side of

metacarpal II (SOM: table 7: b; Kümmell and Frey 2014b; Fontanarrosa et al. 2019). The distal carpal I connects the first ray to the carpus. Proximolaterally, or proximally, distal carpal I articulates with the distomedial border of the medial centrale (SOM: table 7: c, d). In *Stahleckeria*, where the medial centrale is absent, distal carpal I articulates with the radiale and lateral centrale instead. Proximomedially and sometimes also proximally, the bone is usually free of connections. There is an empty space between the junction of the distal carpal I and the medial centrale, often extending to the radiale in the carpi with articulated carpals of most non-mammalian synapsids (SOM: table 7: e). In a few fossils, the contact area of distal carpal I to the medial centrale is very short and sometimes nearly absent as in some Gorgonopsia (*Arctognathus*, *Aelurognathus*, and gorgonopsian BP/1/1210; Figs. 6A, 11) and Therocephalia (*Theriognathus*, *Ictidosuchoides* CGS CM86-655, and *Tetracynodon*; Fig. 6C, D). In these cases, there is a large empty space proximal/proximomedial to the distal carpal I. In a few specimens, distal carpal I shows an extensive proximal contact to one or two carpal bones (e.g., *Edaphosaurus*, where the whole proximal side of the bone is articulated with the medial centrale or in the biarmosuchian PIN 1758/320 where it contacts the medial centrale and a prepollex/sesamoid).

**Distal carpals II and III:** Distal carpal II is usually the shortest of the first four distal carpals. However, in many non-therapsid Synapsida, distal carpal I is shorter than distal carpal II (Fig. 3A, C, E) and in a few Therapsida, distal carpal III is the shortest (e.g., *Olivierosuchus* BP/1/3973 and *Tetracynodon* AM 3677; SOM: table 8: a). Distal carpal II is usually oval to quadrangular or triangular (Figs. 3A–8A; SOM: table 8: b). It forms the base for the articulation of ray II with the carpus (SOM: table 8: c). Distal carpal III is mostly oval, sometimes quadrangular or triangular, or

Table 2. State and shape of distal carpals II and III in different *Diictodon* specimens. The specimens are ordered (top to bottom) according to the increasing length of their skulls and/or their long bones.

<i>Diictodon feliceps</i>	Distal carpal II	Distal carpal III
CGS FL186	absent	absent
TM4991	absent	absent
UMZC T 420	absent	absent
GPIT/RE/7193	very small nodule	oval
SAM-PK-K10699	absent	oval
CGS RMS214	very small nodule?	oval
CGS T72	absent	oval
SAM-PK-K10636	small oval nodule	

wedge shaped (SOM: table 9: a). It is associated with the third ray (SOM: table 9: b).

Generally, the carpometacarpal joints II–V form the carpometacarpal line, which is straight or slightly convex distally. In some specimens, however, the carpometacarpal line corresponding to the carpometacarpal joint II is slightly more proximal than the line through the lateral carpometacarpal joints, e.g., in *Edaphosaurus*, *Galesaurus* BP/1/2513 and most Gorgonopsia (Figs. 6A, 7C; SOM: table 8: d).

In *Diictodon* specimens, distal carpal II is usually absent and occasionally distal carpal III is also missing (Fig. 5A, Table 2), having an open space in the articulated carpi at the typical position of these bones, which suggests that these elements were present in cartilage. Distal carpal III is absent in the three smallest *Diictodon* specimens. Distal carpal II is either completely absent or only present as a very small bony nodule in two or three *Diictodon* specimens (Table 2). In *Stahleckeria*, only three distal carpals are visible in dorsal view, but in this case, there is no open space in any position (Fig. 5C). The second distal carpal of *Stahleckeria* is connected to the second ray and the medial side of ray III and the largest, third distal bone connects to rays III, IV and the medial side of V. We are unsure of the identity of the second element, which could be either distal carpal II or III. The position of this bone at the base of ray II and its partial contact with ray III suggests it is more likely distal carpal II. The size and position of the third distal bone suggest it is distal carpal IV. A possible fusion of distal carpal III to distal carpal IV cannot be excluded.

**Distal carpals IV and V:** Distal carpal IV is usually the longest distal carpal in non-therapsid synapsids and dinocephalians, whereas in dicynodonts, therocephalians and cynodonts, it is usually distal carpal I that is the longest (SOM: table 10: a). Distal carpal IV is pentagonal, especially in many non-therapsid synapsids and gorgonopsians. It is usually quadrangular or trapezoidal in biarmosuchians and anomodonts and mostly ovoid in therocephalians and non-mammalian cynodonts (Figs. 3A–8A; SOM: table 10: b). Distal carpal V is small in relation to other carpals and is ovoid, triangular, quadrangular or trapezoidal in outline (SOM: table 10: c).

Five distal carpals is the plesiomorphic condition in synapsids, but in many non-mammalian therapsids, distal

carpal V is lost or fused to distal carpal IV, where a fusion line may be visible (SOM: table 10: d, e).

In “pelycosaur”, distal carpals IV and V articulate with the fourth and fifth rays, respectively (SOM: table 10: f, g). In non-mammalian therapsids, distal carpal IV sometimes extends laterally to articulate with ray V (SOM: table 10: f), especially when distal carpal V is fused to IV. In some specimens with distal carpal V, e.g., *Thrinaxodon*, a lateral extension of distal carpal IV also contacts metacarpal V.

The following states for distal carpal V can be distinguished: (i) Distal carpal V is a separate bone adjacent to distal carpal IV, e.g., in non-therapsid synapsids (Fig. 3), dinocephalians (Fig. 4B), anomodont *Galechirus* (Fig. 4C), and therocephalian *Ictidosuchoides* CGS CM86-655 (Fig. 6D). In a few cases, distal carpal V appears as a small nodule located within a space between metacarpal V and ulnare. This is the case in the gorgonopsian cf. *Cynariops* SAM-PK-K10000, the cynodonts *Procynosuchus* BP/1/591, NHMUK PV R 37054 (Fig. 7B; but not in RC92, Fig. 7A), *Thrinaxodon* and *Diademodon* NHMUK R-3581. (ii) Distal carpal V is not present in the fossil like in the smallest *Diictodon* CGS FL186 (Fig. 5A), therocephalians *Tetracynodon* AM 3677 (Fig. 6C) and *Olivierosuchus* BP/1/3973, cynodonts *Galesaurus* (Fig. 7C), cynognathian BP/1/4534 (Fig. 7D), *Exaeretodon* and *Trucidocynodon* (Fig. 8A, SOM: table 10: h), and there is an open space between metacarpal V and ulnare. (iii) Distal carpals IV and V are fused with a visible fusion line: gorgonopsians *Arctognathus*, *Dinogorgon* and the gorgonopsid BP/1/1210 (Fig. 6A and 11), *Hipposaurus* (see also Boonstra 1965), the biarmosuchian PIN 1758/320 (very faint line dorsally and an indentation denoting the fusion ventrally; see also Chudinov 1983) and therocephalians *Theriognathus* and *Microgomphodon* (SOM: table 10: e). (iv) Distal carpal V is absent or fused with no apparent fusion line and no space between metacarpal V and ulnare: in dicynodonts such as adult specimens of *Diictodon*, in *Stahleckeria* (Fig. 5C) and probably *Cistecephalus*, therocephalians *Glanosuchus* SAM-PK-K7809 and *?Ictidosuchoides* BP/1/2294 and cynodont *Procynosuchus* RC92 (Fig. 7A). However, in *Ictidosuchoides* CGS CM86-655 and in *Procynosuchus* BP/1/591 and NHMUK PV R 37054, the distal carpal V is separated (SOM: table 10: d, e).

**Mammalianomorpha.**—In contrast to the non-mammalian Synapsida, the carpus of Mammalianomorpha is short and more compact in relation to the whole manus. In non-mammalian cynodonts, the ratio of the carpus to the whole manus (measured as carpus + ray III) is approximately 1:3. In tritylodontids it is about 1:4 and in Mesozoic mammaliaforms it is about 1:6–8. This reduction in relative size of the carpus is mainly due to a shortening of the ulnare and the loss of one bone of the proximal or central row and a concomitant elongation of metacarpals and phalanges.

**Radiale:** The radiale is triangular to rectangular in outline in the tritylodontid *Bienotheroides* and the tritylodon-

tid WCW-06A-34, with its apex pointing distomedially and similar, but more rectangular in *Jeholodens* (Fig. 8B–D). It is trapezoid in *Kayentatherium* and short and rectangular in *Zhangheotherium* and *Eomaia* (SOM: table 1: a). In contrast to non-mammalian synapsids, where the proximal side of the radiale contacts the entire distal end of the radius, the mammalian radiale contacts the medial half of the distal facet of the radius (Fig. 8B–D; SOM: table 1: b). The distal or distolateral border of the radiale articulates with the single centrale (SOM: table 1: d, e). The radiale is fused to the centrale in *Bienotheroides*, with a fusion line present between the two bones and a wedge-shaped angle (set-back angle) between the outline of both bones (Fig. 8B). Laterally, the radiale contacts the lunate (SOM: table 1: f). In contrast to most non-mammalian synapsids, there is no empty space between the radiale and distal carpal I, except in *Kayentatherium* (SOM: table 7: e). In the tritylodontid WCW-06A-34 and *Bienotheroides*, the distomedial top of the triangular radiale forms a short process, which articulates slightly with the proximomedial end of distal carpal I (Fig. 8B, C). In *Zhangheotherium*, however, a small nodular bone interpreted as a probable fragment of a prepollex/sesamoid (Hu et al. 1998), intercalates between the distomedial border of the radiale and distal carpal I (Fig. 12; SOM: table 1: c).

**Ulnare:** The mammalian ulnare is pentagonal (tritylodontid WCW-06A-34), rectangular (*Bienotheroides*) or irregularly triangular (*Zhangheotherium*; Figs. 8B, C, 12; SOM: table 3: a). In relation to metacarpal III, it is short compared with the ulnare of non-mammalian cynodonts (ulnare length as a percentage of the length of metacarpal III: non-mammalian cynodonts 70–111% (except cynognathian BP/1/4534 with 57%), tritylodontids 60–62% and Mesozoic mammals 18–29%; SOM: table 3: b). In *Zhangheotherium*, the ulnare is wider than long (Fig. 12). Proximally, the ulnare is articulated with the ulna (SOM: table 3: g). Medially, it contacts the lunate and in the tritylodontid *Bienotheroides*, the unidentified tritylodontid WCW-06A-34 and *Zhangheotherium*, it also contacts the uppermost end of the lateral side of the radius (SOM: table 3: c). It receives distal carpal IV distomedially or distally (SOM: table 3: d, e). Distolaterally there is an open space between metacarpal V, distal carpal IV and the ulnare, and only in *Zhangheotherium*, the ulnare probably articulates with metacarpal V (SOM: table 3: f).

**Pisiform:** The pisiform is sickle-shaped in *Bienotheroides*, subcircular to oval in *Jeholodens* and proximodistally rectangular in *Zhangheotherium* (Figs. 8B, D, 12; SOM: table 4: a). It is relatively long in mammalian forms. The pisiform is usually fossilised on the lateral/proximolateral border of the ulnare, but in *Zhangheotherium*, it lies mainly lateral to the ulna and articulates with the proximal to proximolateral side of the ulnare (Fig. 12; SOM: table 4: b).

**Lunate:** The lunate is square to sub-oval, subcircular or triangular (Figs. 8B–D, 12; SOM: table 5: a) and is longer than the radiale (SOM: table 5: b). The central constriction

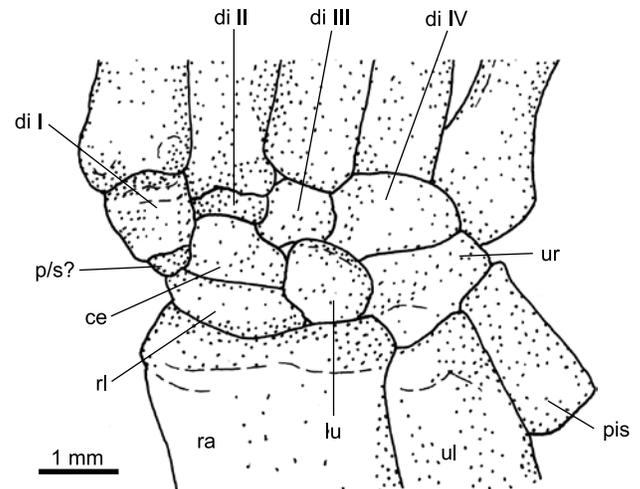


Fig. 12. Carpus of *Zhangheotherium quinquecuspidens* Hu, Wang, Luo, and Li, 1997, IVPP V7466, Jianshangou Valley, Liaoning Province, China, Barremian, right carpus, dorsal view. Abbreviations: di, distal carpal; ce, centrale; lu, lunate; p/s, prepollex/sesamoid, probably a fragment; pis, pisiform; ra, radius; rl, radiale; ul, ulna; ur, ulnare.

of the lunate in *Bienotheroides* is unique in this species. It is prominent and compact in the tritylodontid WCW-06A-34. The lunate articulates distally with distal carpal III, distomedially with the centrale and distolaterally with distal carpal IV (SOM: table 5: c–e). These contacts are identical to those of the lateral centrale in non-mammalian cynodonts, assuming that the single centrale of mammalian forms is homologous with the medial centrale of non-mammalian cynodonts. Between the lunate and distal carpal III, a (small) open space is found in the tritylodontids *Bienotheroides* and *Kayentatherium*, not present in the tritylodontid WCW-06A-34 (Fig. 8B, C). A similar space is also visible distal to the lateral centrale in the cynodonts *Diademodon* NHMUK R-3581 and *Galesaurus* SAM-PK-K10465. In *Diademodon* USNM 23352 it is very small and is absent in *Galesaurus* BP/1/2513 (Fig. 7C), probably becoming obscured by the further growth of the bones. Proximally, the lunate articulates with the lateral portion of the distal facet of the radius (SOM: table 1: b).

**Centrale:** The centrale is irregularly oval or pentagonal (Figs. 8B–D, 12; SOM: table 6: a). Distally, distomedially and -laterally, it contacts distal carpals II, I, and III (SOM: table 6: e). It contacts the radiale proximally and the lunate proximolaterally or laterally (SOM: table 6: b–d). In *Kayentatherium* TMM 43690-5.136, distal carpal II is lost or fused to distal carpal III and the centrale is articulated with distal carpal I and the medial part of the centrally located distal carpal. In *Jeholodens*, the interpretation is difficult, because the presumed distal carpal II, a small bone proximal to metacarpal II, has indistinct edges. A bone approximately the size of distal carpal III lies proximal to this inconspicuous bone, which we interpret here as the centrale (Fig. 8D).

**Prepollex/sesamoid:** A small nodular bone lies medial to the centrale in *Zhangheotherium* and intercalates between the distomedialmost border of the radiale and the proximal part

of distal carpal I (Fig. 12). This is likely to be a fragment of a prepollex/sesamoid, as proposed by Hu et al. (1998).

*Distal carpal I:* Distal carpal I is the longest of the distal carpals in the tritylodontid WCW-06A-34 and in *Bienotheroides* (Fig. 8B, C), whereas in *Kayentatherium* TMM 43690-5.136 and *Zhangheotherium* it is about the same length to distal carpal IV (Fig. 12; SOM: table 10: a). It is square to subcircular in most specimens and rectangular with a proximodistal orientation in *Eomaia* (SOM: table 7: a). In *Jeholodens*, the bone proximal to metacarpal I is damaged and was interpreted as two bones, a distal carpal I and a centrale in Kümmell and Frey (2014b). Here we interpret this bone as a broken distal carpal I in accordance with Ji et al. (2002). A centrale at this position is unlikely because the position of the (medial) centrale in fossil synapsids is very conserved (Fig. 8D). In the tritylodontids WCW-06A-34, *Bienotheroides* and *Kayentatherium* TMM 43690-5.136, distal carpal I extends from the row of distal carpals into the row of metacarpals. In the other mammalian morphs, distal carpal I is situated at the medial end of the row of distal carpals (Fig. 8B, C; SOM: table 7: b). Note that in Mesozoic mammals, the carpometacarpal joint II is shifted slightly proximally (see below), and the distal carpal I is in line with distal carpals III–V, even when distal carpal I is slightly displaced distally compared to ray II (Figs. 8D, 12).

Proximally, distal carpal I contacts the radiale in most mammalian morph species, but connects to a small questionable prepollex/sesamoid in *Zhangheotherium*. In *Kayentatherium*, distal carpal I shows an open space proximally as most non-mammalian morph synapsids (Fig. 12; SOM: table 7: c, e).

*Distal carpals II and III:* Distal carpal II is the shortest of the four distal carpals. It is an oval bone articulating with metacarpal II (Figs. 8B–D, 12; SOM: table 8: b, c). Distal carpal III is also mostly oval, but triangular in *Bienotheroides* (SOM: table 9: a). It articulates with metacarpal III (SOM: table 9: b). The carpometacarpal joint II is shifted proximally in Mesozoic mammals (compared to tritylodontids and most non-mammalian morph cynodonts), so that the carpometacarpal line curves slightly proximally at the position of ray II (Figs. 8D, 12; SOM: table 8: d). In *Kayentatherium* TMM 43690-5.136, only three distal carpals are present. We identified the medial-most distal carpal as distal carpal I and the lateral-most as distal carpal IV. The middle distal carpal is probably slightly turned and is interpreted as distal carpal III because of its size. However, it could also be a fusion of distal carpals II and III. The very small bony structure proximal to metacarpal II of *Jeholodens* is interpreted here as distal carpal II (Fig. 8D).

*Distal carpal IV:* Distal carpal IV is variable in shape: pentagonal, oval or irregularly triangular (Figs. 8B–D, 12; SOM: table 10: b). It articulates with metacarpal IV and usually with the medial part of metacarpal V. In *Zhangheotherium*, it also articulates with the lateral part of metacarpal III (SOM: table 10: f). Between metacarpal V and the ulnare is an open

space in the tritylodontids WCW-06A-34 and *Bienotheroides* and also in the eutriconodont *Jeholodens*. There is a small space between distal carpal IV and ulnare in *Eomaia* (SOM: tables 3: f, 10: h).

## Discussion

This large-scale review of synapsid carpals shows that the position and contacts of single carpal bones are relatively conserved from the Permian to Late Cretaceous, with few exceptions. The different carpal elements usually maintain their positions, with only slight changes in the length or width of individual bones. In the evolution of carpals, not only position and contacts are conserved, but also the relative sizes and width to length proportions of bones. However, the general outlines of the carpal elements are rather variable, with the exception of the mainly rectangular ulnare. Here we assessed homologies according to bone element position, contacts, sequential order and relative size.

Besides the morphological variations of carpal bones related to different locomotory modes, other reasons for shape variation are changes during ontogenetic development (Luo et al. 2003; Stafford and Thorington 1998). Changes in the outline of the carpals do not appear only in early ontogeny, but can develop later, even between subadult and early adult stages, because carpal ossification can occur right up to the latest part of the growing phase (Nesslinger 1956; Oliveira et al. 1998; Stafford and Thorington 1998; Prochel and Sánchez-Villagra 2003; Gilsanz and Ratib 2005; Fröbisch 2008; Wilson et al. 2010). As a general rule, suboval to round shapes occur in the early ontogeny of carpals, whereas complex shapes are found in the mature stages. When other elements of the carpus have a complex outline, round to oval shapes of individual carpal bones suggest a delayed ossification of these elements.

In fossil specimens, fusion lines are important indicators of ontogenetic carpal bone coalescence, whereas open spaces between elements can suggest the persistence of cartilaginous precursors of the bones, which never ossified. However, spaces can also arise from taphonomic distortion of the original bone contacts, e.g., by flattening of the transverse arch during fossilisation.

There are also instances of sudden changes in carpal position and relative size during synapsid evolution, identified using the traditional homology of carpal bones. An example of this is the dimensional change of the intermedium in the transition from “pelycosaurs” to therapsids and from non-mammalian morph cynodonts to mammalian morphs. These sudden changes can arise for several reasons: e.g., a gap in the fossil record or in the sampling used in this study or an inaccurate homologization of the specific carpal bone in the groups before and after the change.

There are significant gaps in the fossil record of synapsids preserving a complete carpus. Besides the poorly sampled “pelycosaurs”, an extensive information gap is found

between basal Sphenacodontidae and the first appearance of therapsids, which cover a time span of around 30 Ma, during the Cisuralian and early Roadian (Sidor and Hopson 1998; Abdala et al. 2008). There is another temporal information gap of about 10 Ma between the first record of Therocephalia and Cynodontia from Wordian to Wuchiapingian (Fig. 2A; Sidor and Hopson 1998). A notable information gap in our sample is present before the emergence of mammalian forms (about 30 Ma, during the Late Triassic; Fig. 2A, B). These information gaps have to be considered when discussing the sudden changes in bone position, contacts, shape and relative size during carpal evolution.

**Position and evolution of radiale and ulnare and the stiffness of the digger's carpus.**—The radiale and ulnare are easily recognized in non-mammalian synapsids, because the radiale occupies the entire distal facet of the radius, and the ulnare articulates with the ulna. The connections of the radiale to the two centralia and the connection of the ulnare to distal carpal IV are both highly conserved. There is a trend towards a shortening of the ulnare in relation to the more elongated metapodium, which emerged in basal mammalian forms (tritylodontids) and became even more pronounced in Mesozoic mammals. The mammalian radiale is narrower in relation to the width of the distal radial facet than in non-mammalian synapsids. It occupies only the medial half of the distal facet of the radius, whereas the lateral half of the distal facet of the radius contacts the lunate. Distally, besides the connection of the radiale to the single centrale of mammalian forms, a new connection of the mammalian radiale to distal carpal I appeared in several specimens of our sample (tritylodontid WCW-06A-34, *Bienotheroides*, probably *Jeholodens*) and some other Mesozoic mammalian forms such as the docodont *Agilodocodon* and the haramiyid *Shenshou lui* (Bi et al. 2014; Meng et al. 2015).

In some specimens, the radiale is fused to the centralia. In the gorgonopsian *Arctognathus curvimola*, both centralia are fused to the radiale. *Arctognathus curvimola* SAM-PK-3329 is interpreted as a highly fossorial animal (Kümmell 2009). This can be deduced from its putative long ungual phalanges in relation to the whole digital length and the compact, stiff carpus (Fig. 11; the tips of the ungual phalanges are broken and their lengths were estimated). Further hints for a digging lifestyle in this species are the short and stout metacarpals and basal and middle phalanges. In addition, the strong basal and middle joints of the digits show near quadrangular facets with low mobility ranges. According to these features, the manus of *Arctognathus curvimola* is midway between that of the scratch diggers *Vombatus* and *Lasiorhinus* and those of the shovel diggers *Talpa* and *Tachyglossus* (Kümmell 2009). Hildebrand and Goslow (2004) suggest that in diggers the carpus is protected against dislocations of the single carpals either by a structural unity of the bones or by the presence of very strong ligaments. In *Talpa*, for example, radiale, lunate and ulnare, though unfused, are tightly bound, so that there

is no mobility between them (Yalden 1966). In *Arctognathus curvimola* the structural unity of the carpus is evident, not only from the fusion of the centralia to the radiale, but also from the compact arrangement of the carpal bones and the very close contact of the distal carpals with the corresponding metacarpals (Fig. 11).

In the undescribed cynognathian cynodont BP/1/4534, the medial centrale probably fused to the radiale as well (Fig. 7D). In contrast to *Arctognathus*, where the structure of the manus suggests that it was an equipped digger, the manual structure of BP/1/4534 suggests it was mainly terrestrial, but could dig sporadically. The length to width-index of the basal phalanx IV was similar (after size corrections) to that of the scratch digger cynodonts *Procynosuchus* and *Chiniquodon* and lower (that means more robust) than that of the extant scratch digger gerbil rodent *Meriones* (Kümmell 2009). Scratch digging therefore, appears to have been possible for the cynognathian BP/1/4534. The fusion of the medial centrale and the radiale would have stabilized the carpus during digging, as in *Arctognathus*. Other Permo-Triassic therapsids may have strengthened their carpus by ligaments rather than by bone fusion, because they show unfused carpals (e.g., *Procynosuchus* and *Diictodon*; Kümmell 2009; Kümmell and Frey 2012).

Fusion of the radiale to other carpal bones also occurred in mammalian forms: a fusion to the centrale in the tritylodontid *Bienotheroides* and to the lunate in the zalambdalestid *Barunlestes* (Kielan-Jaworowska 1978). It is likely that *Bienotheroides* was also capable of scratch digging. The digits are not well preserved; however, the ossified olecranon process of the ulna is very long, 40% of the distal segment of the ulna and the deltopectoral crest of the humerus is prominent and long. These features are thought to be associated with digging abilities in the tritylodontid *Kayentatherium* (Sues and Jenkins 2006).

**Position of the pisiform and its probable sesamoid identity.**—The location of the pisiform in a position approximately distolateral of the ulna and proximolateral of the ulnare is stable throughout fossil synapsids. Slight variations or dislocations are present, so the pisiform is occasionally fossilised just laterally to the ulna or ulnare or (partly) ventrally. The pisiform is not strongly interconnected with other carpal bones, and articulates with the ulna and/or ulnare with short, simple articular surfaces. It is free of contacts laterally, distally and proximally. The contacts to the carpus and its positioning close to the joint between ulna and ulnare, makes the pisiform easy to identify.

The pisiform is often absent in fossil synapsids. However, it is known from specimens of every major lineage of synapsids. Because of this, we consider its frequent absence as taphonomic, arising from the minimal intercalation of the pisiform in the structure of the carpus.

In most placentals for which data are available as well as in the marsupials *Didelphis* and *Monodelphis* (Prochel and Sánchez-Villagra 2003), the pisiform together with the

prepollex/sesamoid are the last carpals to ossify during ontogeny. The late onset of ossification and the minimal contact to the rest of the carpals support the hypothesis that the pisiform is a sesamoid, embedded in the tendon of *m. flexor carpi ulnaris* (Haines 1969; Fabrezi et al. 2007; Fontanarrosa and Abdala 2014, 2016; Amador et al. 2018). Other authors argue for its nature as a true carpal bone (Gillies 1929; Kivell 2016; Diaz and Trainor 2015; Kjosness et al. 2014; Reno et al. 2016; see SOM for further information).

Because of its minimal contacts to the other carpal bones and its free endings distally, laterally and proximally, the synapsid fossil record suggests a sesamoid identity for the pisiform.

**The homology of the mammalian centrale.**—In nearly all non-mammalian synapsids, the medial centrale is articulated distally, distomedially, and distolaterally with distal carpals II and I, often also with distal carpal III. It contacts the radiale proximally or proximolaterally. In some cases it is fused to the radiale (see above). The same connections, with distal carpals I, II, and III and proximally with the radiale, are observed in the single centrale of mammalian synapsids. Because of the relative position towards the medial side of the manus and its anatomical contacts, we interpret the single centrale of mammalian synapsids as homologous to the medial centrale of non-mammalian synapsids.

The mammalian centrale is absent in Monotremata and Marsupialia (Flower 1885; Holmgren 1952; Grassé 1955; Szalay 1994; Flores and Diaz 2009). It is also reported to be lost in some basal eutherians such as *Ambolestes* and *Sinodelphys* (*Sinodelphys* was originally interpreted as metatherian by Luo et al. 2003, but most recently as eutherian by Bi et al. 2018). It is likely that the centrale persisted in the stem lineage of Mesozoic Eutheria (as in *Barunlestes* and probably *Asioryctes*; Kielan-Jaworowska et al. 2004: figs. 13.15 and 13.12), because it appears in many extant placentals, e.g., in *Tupaia* (e.g., Schmidt-Ehrenberg 1942; Stafford and Thorington 1998).

**The question of the lunate homology.**—*Palaeontological and morphological evidence:* In previous anatomical work and textbooks, the intermedium of reptiles and non-mammalian synapsids is homologized with the lunate of mammals (Fig. 13I, A<sub>2</sub>–D<sub>2</sub>; e.g., Gegenbaur 1864; Broom 1901; Ihle et al. 1927; Romer and Parsons 1977; Starck 1979; Salomon et al. 2005; Kivell 2016). Also, Sun and Li (1985) in their description of the basal mammalian tritylodontid *Bienotheroides* designated the bone in the position of the lunate as an intermedium. We argue for the homology of the mammalian lunate with the lateral centrale of non-mammalian synapsids (Fig. 13II, A<sub>2</sub>–D<sub>3</sub>) for the following two reasons. (i) Position: The mammalian lunate articulates distally with the distal carpal III, distolaterally with the distal carpal IV and distomedially with the centrale. It contacts the radiale medially and the ulnare laterally. These contacts are identical to those of the lateral cen-

trale in non-mammalian cynodonts (SOM: tables 1: f, 3: c, 5: c–e). Proximally, the lunate contacts the lateral half of the distal articular surface of the radius. This contact to the radius resembles neither the proximal contact of the lateral centrale, which articulates with the intermedium, nor that of the intermedium, which articulates proximally with the medial section of the distal ulnar facet. So, the articulation of the lunate with the radius is an apomorphy of mammalian synapsids. Thus, in position and contacts, the lunate of mammalian synapsids resembles the lateral centrale of non-mammalian cynodonts more than their intermedium (Fig. 13II, A<sub>2</sub>–D<sub>3</sub>). (ii) In terms of relative size, the mammalian lunate resembles the lateral centrale of non-mammalian cynodonts, as well. The lunate is longer than the corresponding radiale and relatively wide. That is also the case for the lateral centrale and radiale of non-mammalian cynodonts (except the most basal form *Procynosuchus*), whereas the intermedium of non-mammalian cynodonts is very slender, and the same length or shorter than the corresponding radiale (Fig. 13, SOM: tables 2: e, 5: b).

There are two arguments that can be brought against our proposal, which should be discussed here. First, one can argue that during the transition from “pelycosaur” to therapsids, the intermedium altered significantly in form and size (Fig. 13). This would suggest considerable evolutionary plasticity, which could also account for the proposed changes in form at the transition from non-mammalian cynodonts to mammalian synapsids. In our data set, there are temporal information gaps in both transitions: from non-therapsid synapsids to therapsids and from non-mammalian cynodonts to mammalian synapsids (Fig. 2A, B; see above). This leaves open the possibility that large evolutionary changes could have taken place during that time. However, in the transition from non-therapsid synapsids to therapsids, the position of the intermedium and its contacts did not change, but the relative width and dorsoventral depth did change. In non-therapsid synapsids, it is mostly broad, square or pentagonal (Fig. 3A–C) except in two *Ophiacodon* specimens (FMNH UC 671 and FMNH UC 458), where it is longer than in other non-therapsid synapsids (Figs. 3A–C, 13). In therapsids, the intermedium is considerably narrower in dorsal view. The intermedium is dorsoventrally shallow in non-therapsids and deep in therapsids. Because the anatomical position and the relevant contacts remain the same, we propose that the intermedia in both groups are homologous. The change in proportions of the intermedium may be related to the slight rotation of the elbow posteriorly, producing a semi-sprawled posture on the transition to therapsids, which altered the geometry of the wrist (Colbert 1948; Jenkins 1971). However, in the transition from non-mammalian cynodonts to mammalian synapsids, the situation would be different, if the lunate derived from the intermedium, as suggested previously. In this case, the intermedium of non-mammalian cynodonts would not only have changed in relative size and form, but also in position. Such a transformation is unlikely, especially given the fact that this shift must have occurred in the midst

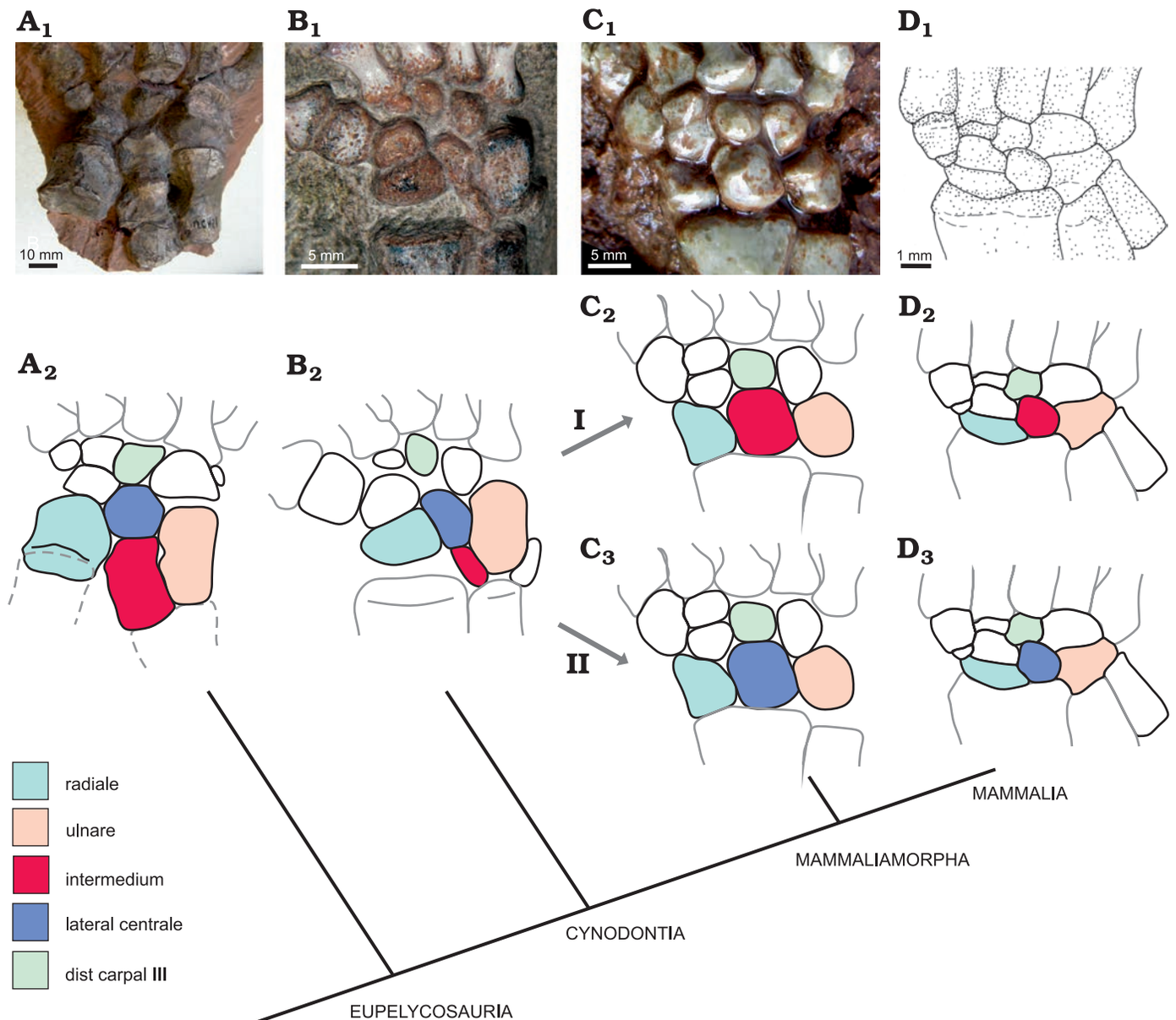


Fig. 13. Homologization of the mammalian morpholunate with the intermedium or lateral centrale of non-mammalian synsids. I. The homologue of the lunate is the intermedium (C<sub>2</sub>, D<sub>2</sub>); this is the traditional view (Gegenbaur 1864; Romer and Parsons 1977; Kivell 2016) and was put forward by Broom (1901) for non-mammalian morph therapsids and extant mammals. II. The homologue of the lunate is the lateral centrale (C<sub>3</sub>, D<sub>3</sub>) as suggested here. A. *Ophiacodon* (FMNH UC 458). B. *Galesaurus* (SAM-PK-K10465). C. Tritylodontid (WCW-06A-34). D. *Zhangheotherium* (IVPP V7466). Photographs (A<sub>1</sub>-C<sub>1</sub>), drawing (D<sub>1</sub>), and explanatory drawings (A<sub>2</sub>-D<sub>2</sub>, C<sub>3</sub>, D<sub>3</sub>). The figures show the right manus in dorsal view, except for A, where the left manus has been lateromedially reversed.

of tight anatomical contacts of the carpal bones. Thus, we argue that the intermedium was lost during the transition from non-mammalian morph synsids to mammalian morphs or otherwise fused to the lateral centrale or another carpal or zeugopodial bone (see below) and that the lunate is the homologue of the lateral centrale.

Secondly, *Kayentatherium* MCZ 8812, which belongs to Tritylodontidae, one of the most basal mammalian morph clades, appears to contradict our proposal at first glance. Sues and Jenkins (2006) described two centralia and one intermedium for each of the partly articulated right and left manus of the specimen. With its square outline, the bone

designated as intermedium resembles the mammalian morph lunate rather than the non-mammalian morph intermedium. Because the tritylodontid *Kayentatherium* is one of the earliest mammalian morphs, the situation in this fossil could indicate that the intermedium changed its form and relative width prior to the evolutionary loss of one carpal bone. In this case, the lunate of mammalian morphs would be homologous to the non-mammalian morph synsids intermedium. But the presence of two centralia and one intermedium in each manus of *Kayentatherium* MCZ 8812 is questionable, because both intermedia are out of place. Another *Kayentatherium* TMM 43690-5.136 was recently described, which show a

carpus fossilised in articulation (Hoffman and Rowe 2018). In this specimen, two centralia or one centrale and one lunate, respectively, are present with no intermedium (according to supplement video 1 in Hoffman and Rowe 2018, and images of scans sent by Eva Hoffman to SK). This articulated carpus of *Kayentatherium* resembles the carpi of the tritylodontid WCW-06A-34 and *Bienotheroides* (Figs. 8B, C, 13). In *Kayentatherium* TMM 43690-5.136, the distal carpal I is comparatively large and resembles the previously designated intermedium in *Kayentatherium* MCZ 8812 (Sues and Jenkins 2006). If the latter were to be distal carpal I, the situation in *Kayentatherium* would not be different from that in other mammalian morphs. A thorough comparison of the two specimens of *Kayentatherium* is necessary to solve the identity of the bones completely.

**Embryological evidence:** An investigation of the early ontogeny of extant mammals may shed light on the question of lunate identity and help to identify the bones that were lost in the transition to mammalian morphs. Despite differing views on lunate identity (intermedium versus centrale), there is general agreement that a mesenchyme string (intermedial string; sensu Schmidt-Ehrenberg 1942) forms in early mammalian ontogeny, which detaches from the ulna, and gives rise to the lunate (Fig. 14A, B; Steiner 1935; Schmidt-Ehrenberg 1942; Holmgren 1933, 1952; Čihák 1972; Shubin and Alberch 1986). While Steiner (1942), Schmidt-Ehrenberg (1942), and Shubin and Alberch (1986) interpret the lunate as homologous to the intermedium, Holmgren (1933, 1952), Kindahl (1941, 1942a, b, 1944), Čihák (1972) and Slabý (1967, 1968) interpret parts of the intermedial mesenchyme string as the homologue of the intermedium and the lunate as homologous to a centrale.

In some mammals (*Tupaia*, *Elephantulus*, *Tarsius*, among many others), Holmgren (1933, 1952) and Kindahl (1942b) observed the chondrification of the mesenchymal remnant (intermedium anlage) of the connection of the lunate anlage to the ulna. This chondrified remnant, interpreted as the intermedium, can persist in these species even when the precursor of the lunate chondrifies (Fig. 14B, C). Holmgren (1952) described that in later ontogenetic stages of mammalian embryos, the intermedium fused to other chondrogenic anlagen: usually to the ulnare (= centrale 4 of Holmgren), sometimes to the lunate (= centrale 1 of Holmgren), and very occasionally to the ulna or the radiale.

The mesenchymal connection between the distomedial corner of the ulna and the lunate, distal to the radius, is the secondary digital arch (sensu Shubin and Alberch 1986), along which the lunate and centrale develop. The proximal part of this axis resembles the intermedium of non-mammalian morph synapsids in shape and position (Fig. 14) and may thus represent the homologue of this bone. In mammalian morphs, it is plausible that the anlage of the intermedium persists through early ontogeny, even though the intermedium in the adult stage is lost. This is because the anlage constitutes the developmental axis of the lunate and centrale in early development.

The embryological studies of Holmgren (1952), Kindahl (1942b), Čihák (1972), and Slabý (1976) present strong evidence that the lunate of mammalian morphs is best homologized with a centrale of non-mammalian morph synapsids. Mammalian morphs thus resemble modern tetrapod taxa such as lizards, crocodiles, and anurans, in which the intermedium is frequently absent in the adult stage (Fabrezi and Alberch 1996; Fabrezi and Barg 2001; Fabrezi et al. 2007). The intermedium may appear as an isolated element in early ontogeny but can also become fused to the radiale in later stages, as in Crocodylia (Müller and Alberch 1990; Müller 1991).

Using the evidence derived from our paleontological investigations and the embryological studies of those mammals, in which both the lunate and the intermedium anlage chondrified, we homologized the mammalian morph lunate with the lateral centrale of non-mammalian morph synapsids. Thus, the mammalian morph lunate should be called “lateral centrale” and the single centrale of mammalian morphs would be the “medial centrale”. For traditional reasons and considering the uncertainty surrounding a probable early ontogenetic fusion of the intermedium to the lateral centrale of mammalian morphs, we argue for the continued use of the term “lunate” and “centrale” in mammalian morphs.

**Third centrale—an abnormal duplication of one centrale?**—Three centralia have been described in the theroccephalian *Theriognathus* NHMUK R 5694 (Boonstra 1934: 260, fig. 34). The two medial bones are in the same position as the medial centrale of many Therapsida. The bones are tightly connected and were previously interpreted as fused (Boonstra 1934). Our study shows they are separate bones, with tight fitting articular surfaces (Fig. 9). We also interpret the three elements as centralia in accordance with Boonstra (1934).

Angielczyk and Rubidge (2013) tentatively identified three centralia in the right carpus of the dicynodont *Eosimops* BP/1/6674, lying lateral and distolateral of the radiale. We interpret the proximalmost of these three bones as the lateral centrale. The medial centrale is, in our opinion, in its normal location distal to the radiale (an element interpreted as the distal carpal II by Angielczyk and Rubidge 2013). The bones of the distal row of carpals are displaced from their natural position and the two distally located bones, interpreted by Angielczyk and Rubidge (2013) as centralia, could also be interpreted as distal elements out of place. The left carpus of the same specimen, however, is fossilised in articulation with only some disturbance at the ulnare. Here, the lateral and medial centralia are present in their usual position. The form, size and proximal articulation of the medial centrale in *Eosimops* is typical for Dicynodontia (Fig. 5B, SOM: table 6: a, b).

A third centrale could arise from “de novo” duplication of an early ontogenetic cartilaginous condensation. Such a case is known from an abnormal duplication of centrale 3 in the turtle *Phrynops hilarii* (Fabrezi et al. 2009). “De novo”

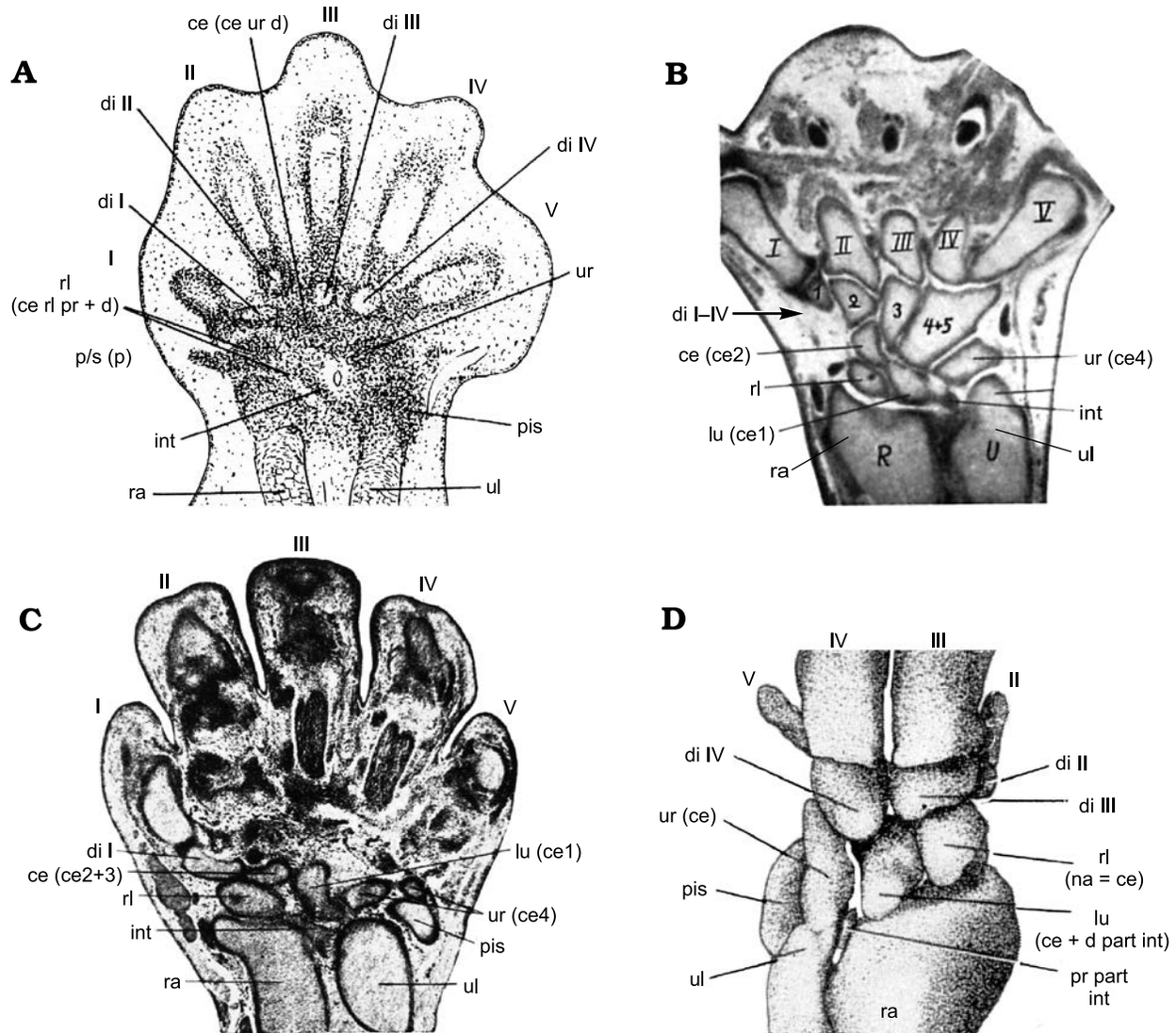


Fig. 14. Interpretation of the cartilaginous anlagen of the embryological carpus of different mammals by different authors. Whenever the radiale, ulnare or lunare have been interpreted previously as homologues of a centrale, the details are given in brackets following the abbreviations in the figures. **A.** *Microcebus myoxinus* Peters, 1852, 10.2 mm neck-rump length (from Schmidt-Ehrenberg 1942: 61, fig. 10, courtesy of Revue Suisse de Zoologie). **B.** *Tarsius spectrumgurskyae* Shekelle, Groves, Maryanto, and Mittermeier, 2017, 20.5 mm body length, (from Holmgren 1952: 80, fig. 93, reproduced with permission of Acta Zoologica, © The Royal Swedish Academy of Sciences, all rights reserved). **C.** *Talpa europaea* Linnaeus, 1758, 15 mm body length (from Kindahl 1942b: 270, fig. 4, courtesy of Sächsisches Staatsarchiv, Staatsarchiv Leipzig). **D.** *Ovis aries* Linnaeus, 1758, 38 mm body length, three-dimensional reconstruction of the left carpus in dorsal aspect (from Slabý 1967: pl. 6: 30). Abbreviations: ce, centrale; d, distal; di, distal carpal; int, intermedium; lu, lunare; na, naviculare; p, prepollex; p/s, prepollex/sesamoid; pis, pisiform; pr, proximal; ra, radius; rl, radiale; ul, ulna; ur, ulnare.

duplication probably occurred in the condensation of the medial centrale in *Theriognathus* NHMUK R 5694, because the two medial centralia occupy the same position as the medial centrale of the other non-mammalian synspsids.

#### Has there been a prepollex/sesamoid in synspsids?—

Prepollices occur in a wide range of tetrapods. They were shown to be present in anurans as a complex of skeletal elements (Fabrezi 2001). They occasionally appear in reptiles (Wagner et al. 2000), birds (Starck 1979) and frequently in mammals (Salesa et al. 2006; Echeverria et al. 2019). However, in many cases the so-called prepollex of mammals was identified as a radial sesamoid (see SOM), so here we designate the prepollex of mammals as “prepollex/ses-

amoid”. A prepollex/sesamoid in extant mammals usually differs from most carpals in being incompletely integrated in the carpus. This bone is often situated at the medial border of the carpus, between radiale and distal carpal I (scaphoid and trapezium) oriented towards the palmar surface, with a free ending pointing medially or distally (Salton and Sargis 2008).

Previously, the three central bones of the Russian biarmosuchian PIN 1758/320 were interpreted as centralia (Chudinov 1983: 55–56). Two of these bones occupy the position of the medial and lateral centrale, whereas the medialmost third central bone lies in the junction of distal carpal I and the medial centrale, in a slightly ventral position. The medial half of this bone shows a free medial margin. This

position and orientation do not support the interpretation as a third centrale and is more consistent with the interpretation of a prepollex/sesamoid instead.

Other non-mammalian synapsids show small bones of uncertain identity close to distal carpal I. These include *Chiniquodon* PVL 3820, *Galesaurus* SAM-PK-K10468 (Kümmell and Frey 2014b), *Suminia* 2212/62 (Fröbisch and Reisz 2011: fig. 9), “*Opisthoctenodon agilis*” (Broom 1904; most likely *Pristerodon*; Keyser 1993, Angielczyk et al. 2005) and *Theriodesmus* NHMUK 49392 (probably a biarmosuchian, FA personal observation; Bardeleben 1889). These bones may also be prepollices/sesamoids.

In most non-mammalian synapsids there is an open space proximal to distal carpal I at the junction of this bone with the medial centrale, sometimes extending to the distomedial rim of the radiale (SOM: table 7: e). This space is at the same position in relation to distal carpal I as the prepollex/sesamoid of the biarmosuchian PIN 1758/320. The proximal contact area of distal carpal I to the medial centrale is usually very short or nearly absent as in some Theriocephalia and Gorgonopsia. In the latter, the first digit is opposable and very manoeuvrable (Kümmell and Frey 2014b), so a strong articular connection to the carpus is likely. Because the free space proximal to distal carpal I appears in both juveniles and adults, its interpretation as representing an unossified, cartilaginous medial extension of the medial centrale is not guaranteed. It is more likely that distal carpal I was supported by a cartilaginous unfossilised prepollex/sesamoid.

Earlier authors proposed the presence of a sesamoid, medial to the medial centrale. Thus, a large sesamoid was proposed to be present proximal to distal carpal I at the junction of distal carpal I and the medial centrale in *Dimetrodon incisivus* (Case 1907) and *Dimetrodon grandis* (Gilmore 1919), and a small sesamoid between the medial centrale and the radiale in non-therapsid Eupelycosauria (Romer and Price 1940: 160). However, recent criticism of these interpretations arose from Henrici et al. (2005), stating that there is no indication of articular surfaces for sesamoids on the medial carpals in the eupelycosaurian *Sphenacodon ferox*. Like Henrici et al. (2005), we did not find any articular structures for a prepollex/sesamoid, similar to those between the other carpals on distal carpal I, medial centrale and radiale in synapsids where the medial portion of the carpus is exposed. However, because of the slightly ventral position of the prepollex/sesamoid in the biarmosuchian PIN 1758/320, its articular structures do not resemble the articular surfaces in other carpal bones. In the biarmosuchian PIN 1758/320, only slight indentations with no special articular surfaces are visible on the first distal carpal and medial centrale (Fig. 10A<sub>2</sub>). Flat, medioventrally facing indentations at the proximal margin of distal carpal I and the distomedial margin of the medial centrale are also present in *Titanophoneus* and *Galesaurus* SAM-PK-K10468, similar to that of the biarmosuchian PIN 1758/320. These indentations do not have a different surface texture that would allow their iden-

tification as true facets. Therefore, a prepollex/sesamoid may have been present in *Titanophoneus* and *Galesaurus* SAM-PK-K10468, but cannot be completely verified. An extensive analysis of these structures in synapsid skeletons was not possible in this study, because the first distal carpal and the medial centrale were not exposed from the ventral side in most fossil Synapsida. Further preparation or CT-scanning would clarify this in the future.

In early non-synapsid amniotes and amphibians, a prepollex/sesamoid often appears in the tracks of these early tetrapods. It was found in the ichnofossil *Amphisauropus*, which presumably represents a track of a seymouriamorph trackmaker (Fig. 15) and also in the earliest diadectomorph tracks *Ichniotherium praesidentis* and the putative tracks of the temnospondyl *Eryops* (Voigt and Ganzelewski 2010; SK personal communication 2016 with Sebastian Voigt; Voigt and Lucas 2017). Despite not being preserved in the fossilised skeleton of *Eryops*, a prepollex was reconstructed by some authors on the basis of the facets of the medialmost centrale and distal carpal I, combined with data from extant amphibians (Steiner 1921; Gregory et al. 1923). In contrast to Gregory et al. (1923), Dilkes (2015) did not reconstruct a prepollex in *Eryops*. To date, a prepollex/sesamoid-like structure was not found in the tracks of synapsids (SK personal communication 2016 with Sebastian Voigt). However, because of the small size of the prepollex/sesamoid found in the biarmosuchian PIN 1758/320, a visible separate prepollex/sesamoid structure within the imprint of a manus would be improbable.

The few instances described here suggest the possibility that in non-mammalian synapsids a cartilaginous prepollex/sesamoid-like structure was present. It probably supported distal carpal I by filling the space on its proximal and proximomedial side. In Mesozoic mammalian morphs, two

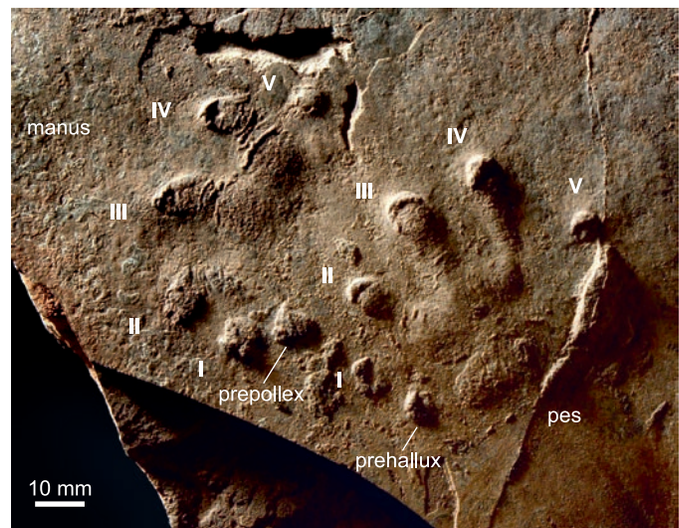


Fig. 15. The track *Amphisauropus* Haubold, 1970, NMMNH P-37922, central New Mexico, USA, Abo Formation, Cisuralian, left manus and pes imprint, convex hyporelief, presumably from a seymouriamorph trackmaker. Photo made by Sebastian Voigt, published with the permission of Sebastian Voigt, all rights reserved (see also Voigt and Lucas 2017: fig. 3F).

putative prepollices have been reported: one in *Asioryctes* (Kielan-Jaworowska 1977; Kielan-Jaworowska et al. 2004) and a fragmentary nodular bone proximal to distal carpal I in *Zhangheotherium* (Hu et al. 1998). Salton and Sargis (2008: fig. 8) reported on a very small rudiment of a prepollex in a specimen of the otter shrew *Potamogale*, intercalated between distal carpal I and the radiale. This bone resembles the nodule in *Zhangheotherium* in both shape and position.

In Mesozoic mammaliaforms, with the exception of *Kayentatherium* TMM 43690-5.136, no free space is present between the junctions of centrale and distal carpal I, centrale and radiale or in the junction of all three bones. Instead, the proximal rim of distal carpal I articulates about its whole width with fossilised bones. Either it articulates with the centrale and the probable prepollex/sesamoid as in *Zhangheotherium* and *Asioryctes* or with the centrale and the radiale. The radiale developed a distomedial process, to connect to distal carpal I in some Mesozoic mammaliaforms without a prepollex/sesamoid. This process is present and short in the tritylodontid WCW-06A-34 and in *Bienotheroides* and is very long in the haramiyidid *Shenshou lui* (Bi et al. 2014). This additional contact between distal carpal I and the radiale of many mammaliaforms integrates distal carpal I further into the carpus adding greater stability and reducing mobility (Kümmell and Frey 2014b).

The prepollex/sesamoid is present in many extant mammals and in some synapsid fossils. The configuration of the fossil carpals suggests it might also have been present in other synapsid taxa. Prepollex/sesamoid impressions in some early tetrapod tracks further suggest that it was fairly widespread among early tetrapods. The systematic framework of the occurrence of prepollex/sesamoid-like structures in most extant mammals and in early tetrapods makes it very likely that such structures were present in synapsids of the Permian and Mesozoic. Indeed, a prepollex/sesamoid was also proposed for stem eutherians by Szalay (1994).

**Position of the distal carpals and their evolutionary loss or fusion.**—In the early ontogenetic development of mammals, the distal carpals and the metacarpals I–IV separate from each other by one of two possible processes. Either the metacarpals segment from their corresponding distal carpal (Shubin and Alberch 1986) or the distal carpal detach from the proximal border of the metacarpal blastemata (Holmgren 1952; Milaire 1978; Johanson et al. 2007). In both cases, the distal carpal and metacarpal are developmentally associated with each other. This is also the case for ray V, where distal carpal V detaches separately from the corresponding metacarpal blastemata in early ontogeny. In later stages, the cartilaginous foci of distal carpals IV and V fuse either to each other (*Mus*, Milaire 1978; *Perameles*, *Dasyopus*, Holmgren 1952) or distal carpal V fuses to metacarpal V (*Homo*, Čihák 1972). Only Slabý (1976, investigating *Procvavia capensis*, *Sus scrofa*, *Ovis aries*, and *Bos taurus*) interpreted distal carpal V as ontogenetically derived from the ulnare. The different manual rays are connected to the carpus by a joint

between the single ray and the corresponding distal carpal, only rays IV and V are both connected to the distal carpal IV of modern mammals, which probably represents a fusion of distal carpals IV and V. The connected developmental derivation and the functional relation through ontogeny makes the association of the distal carpals and the corresponding manual rays very stable in ontogeny. This is also probably the case for phylogeny. Thus, the identity of a single distal carpal is relatively easy to determine by its position at the base of a manual ray.

*Distal carpal I:* Distal carpal I is always the base for ray I, but shows some proximodistal positional changes within fossil synapsids. In “pelycosaurs”, the anomodont *Galechirus* and in Mesozoic mammaliaforms, it lies in the row of distal carpals, whereas in some non-mammalian therapsids (e.g., *Robertia*, *Diictodon*, *Microgomphodon*, *Procynosuchus* RC92, *Exaeretodon*), it is aligned with the row of metacarpals. In all other analyzed non-mammaliaform therapsids, it shows an intermediate position (SOM: table 7: b). In non-mammaliaform synapsids, the distal shift of distal carpal I is connected to the change of the autopodial rolling mode with increasing parasagittal posture, e.g., in Permian dicynodonts (Kümmell and Frey 2014a, b). The autopodia rolled medially to the medial digits in early forms and to the central digits in later forms, facilitated by an increasing length of digit I. During the transition to Mesozoic Mammaliaformes and the tritylodontid *Oligokyphus*, metacarpal I (metacarpal or metatarsal I in *Oligokyphus*) changed its function. In non-mammaliaform synapsids (with the exception of *Galechirus*), metacarpal I functioned as a basal phalanx, whereas in Mesozoic mammaliaforms and *Oligokyphus* it functioned as a true metacarpal. Accordingly, distal carpal I of non-mammaliaform synapsids was more mobile than that of mammaliaformes and *Oligokyphus*, where it is completely integrated in the carpus (Kümmell and Frey 2014b). The situation of the other tritylodontids needs further investigation.

*Distal carpal II and III:* In Dicynodontia, distal carpals II and III are irregularly absent (Fig. 5A) or present. Both bones are absent in the articulated fossil carpus of many specimens of *Diictodon feliceps*, where an empty space occupies their usual position, but they are sometimes present in other specimens of the species. Considering this situation, we interpret the absence of distal carpal II and/or III as due to a heterochronically delayed ossification (paedomorphosis; sensu McNamara 2002) of cartilaginous precursors. Distal carpal III is apparently absent in the smaller, very likely younger forms, but present in the larger, presumably older forms of *Diictodon feliceps*. The ossification of distal carpal II is even more delayed than that of distal carpal III. It is either not present even in the larger and very likely older forms or it is present as a very small bony nodule, as in *Diictodon* GPIT/RE/7193 and in the largest and probably most mature specimen SAM-PK-K10636 (Table 2).

The delayed ossification of distal carpals II and III in *Diictodon* is remarkable, since the animal is known to have

been a burrow-dweller that dug its own burrow (Smith 1987; Ray and Chinsamy 2003). Diggers usually have a strong and stiff, well ossified carpus (see above). As judged from the digits of the manus, *Diictodon* was able to scratch dig (Kümmell 2009). In the manus the second digit is longer than the fourth, a condition typical in scratch diggers for example the extant marsupial *Lasiorhinus latifrons*. The ungual phalanges are relatively short in relation to those of other dicynodonts (the length of the ungual phalanges III in relation to the corresponding digital length is 31%, 34%, 41%, and 49% in four *Diictodon* specimens). These values are typical in extant scratch diggers, as *Meriones shawi* (38%), *Oryctolagus cuniculus* (35%), and *Meles meles* (46%). The robustness of basal phalanx III (median value of the ratio of length to width is 1.65) is similar in other dicynodonts. It is slightly lower (i.e., the phalanges are more robust) than in terrestrial mammals and in the placental scratch digger *Meles meles* (2.48) and resembles that of the marsupial scratch digger *Bettonia pinicillata* (1.78). *Diictodon* shows well ossified joints. It must have been a scratch digger that dug in relatively soft soil (Ray and Chinsamy 2003; Kümmell 2009). The absence of wear facets on the tusks or keratinized frontal part of the snout suggest that these did not play a major role in the digging process of *Diictodon* (Ray and Chinsamy 2003). The digging method of *Diictodon* needs further investigation to explain the paedomorphic evolution of distal carpals II and III in this species.

*Stahleckeria* MCZ 1688 has only three distal carpals. In contrast to the situation in *Diictodon*, there is no empty space in the articulated carpus of *Stahleckeria*, so the distal carpals were lost in evolution or fused to an adjacent bone. The distal carpals lost were distal carpal V (see below) and one of the central distal carpals (II or III), probably III judging by the position of the central distal carpal mainly at the base of ray II. The absence of one central distal carpal or its fusion to another distal carpal was probably associated with the evolution of large sole cushions in that animal, which correlates with reduced carpus mobility (Kümmell 2009). It has also been suggested that distal carpal II was cartilaginous and did not ossify in *Stahleckeria* MCZ 1688 (Romer and Price 1944).

The tritylodontid *Kayentatherium* TMM 43690-5.136 possess only three distal carpals instead of the usual four of mammalianomorphs. Distal carpal II is probably absent or fused to distal carpal III. We analyzed this fossil from images of scans, so a more detailed study of the specimen comparing it with *Kayentatherium* MCZ 8812 (Sues and Jenkins 2006) is needed to investigate the question of the identity of the distal carpals.

*Distal carpal V*: The presence of distal carpal V is pleiomorphic in therapsids and is found in the “pelycosaur” taxa analysed here. During therapsid evolution, it is lost in nearly every group. Hopson (1995) described two processes of losses in therapsids. The first was non-ossification of distal carpal V in Dicynodontia and gomphodont Cynodontia

(with the absence of distal carpal V as an individual variant in some early cynodonts where it probably failed to ossify). The second was fusion to distal carpal IV in Biarmosuchia, Gorgonopsia, Therocephalia, and Mammalia.

Here we propose that the processes leading to the loss of distal carpal V were even more subtle. In the most basal therapsids, the biarmosuchians, distal carpals IV and V are fused. The fusion line of the biarmosuchian PIN 1758/320 is very faint, but clear in *Hipposaurus*. In dinocephalians, distal carpal V is still present. In dicynodonts, distal carpal V is completely lost, as can be inferred from the compact form of distal carpal IV without indentations or set-back angles in its outline and the contact of the proximal rim of metacarpal V with the ulnare in many *Diictodon* specimens and *Stahleckeria* (Fig. 5B, C; SOM: table 3: f). In gorgonopsians, therocephalians and cynodonts, the following states of distal carpal V are present. State 1: An open space is present instead of distal carpal V. Here, distal carpal V was not ossified, but remained in a cartilaginous form (Hopson 1995; Fontanarrosa et al. 2019). This state appears in the therocephalians *Olivierosuchus* BP/1/3973 (Fig. 16A), *Tetracyonodon* AM 3677 (Figs. 6C, 16B) and the cynodonts *Galesaurus* (Fig. 7C), *Exaeretodon* and *Trucidocynodon* (Figs. 8A, 16C). State 2: Ossification of distal carpal V had begun, appearing as a small nodule within an open space between metacarpal V and ulnare, present in the gorgonopsian cf. *Cynariops* SAM-PK-K10000 and the cynodonts *Procynosuchus* BP/1/591 and NHMUK PV R 37054 (Figs. 7B, 16D, E), *Thrinaxodon* and *Diademodon* NHMUK R-3581. State 3: The space is nearly completely filled by the distal carpal V. This state is found in the therocephalian *Ictidosuchoides* CGS CM86-655 (Figs. 6D and 16F). State 4: Distal carpals IV and V are fused with a visible fusion line, which is the case in the gorgonopsians *Arctognathus*, *Dinogorgon* and the gorgonopsid BP/1/1210 (Figs. 6A, 11, 16G, I) and the therocephalians *Theriognathus* (Fig. 16H) and *Microgomphodon*. State 5: Distal carpal IV and V are fused, but no fusion line is present. This state is present in the therocephalians *Glanosuchus* SAM-PK-K7809 (Figs. 6B, 16J; Fontanarrosa et al. 2019), *?Ictidosuchoides* BP/1/2294 (Fig. 16K) and in the cynodont *Procynosuchus* RC92 (Figs. 7A, 16L). The fusion of distal carpal V is deduced from the shape of distal carpal IV with a distal indentation showing the point of fusion of the two bones. A further suggestion advocating a fused state rather than the loss of distal carpal V, is that distal carpal V is also found in other specimens of *Procynosuchus* (state 2).

Thus, in gorgonopsians, therocephalians, and cynodonts, distal carpal V is always present, but appears in different forms: as a cartilaginous precursor, a separate bone or in a fused state. A fused state, in which a fusion line is present or in which there is other evidence suggestive of the fusion of two different bones, provides evidence of two ossification centres, from which two bones grow and fuse during ontogeny. Because carpals usually ossify relatively late compared to the long bones and other bones of the manus (Nesslinger

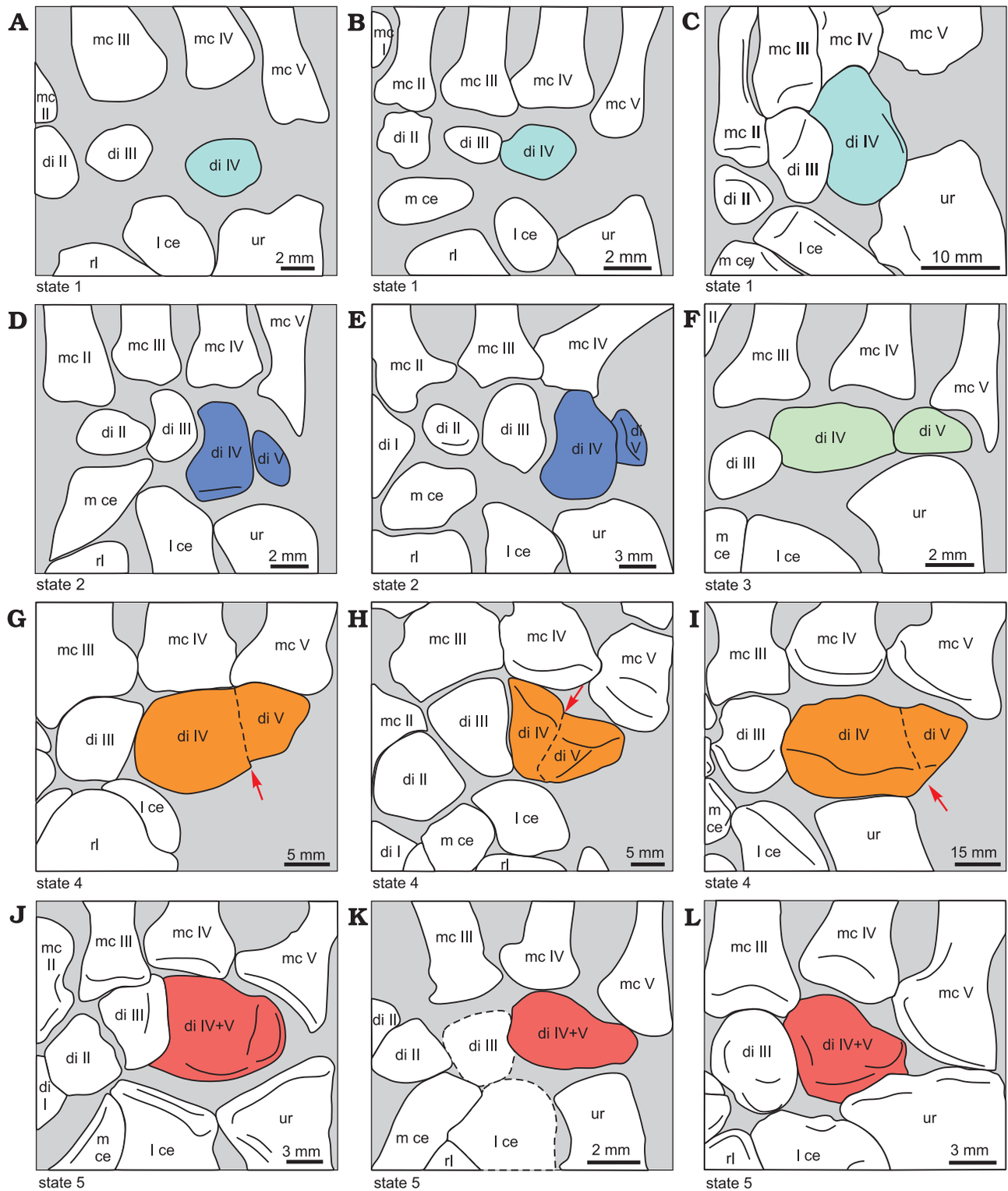


Fig. 16. Selected specimens showing the different states of distal carpal V in gorgonopsians, therocephalians, and cynodonts, all in dorsal view (except A, ventral view). **A.** *Olivierosuchus parringtoni* (Brink, 1965), BP/1/3973, left carpus. **B.** *Tetracynodon darti* Sigogneau, 1963, AM 3677, right carpus. **C.** *Trucidocynodon riograndensis* Oliveira, Soares, and Schultz, 2010, UFRGS PV-1051T, right carpus. **D.** *Procynosuchus delaharpeae* Broom, 1937, BP/1/591, right carpus. **E.** *Procynosuchus delaharpeae* Broom, 1937, NHMUK PV R 37054, right carpus. **F.** *Ictidosuchoides longiceps* Broom, 1920, CGS CM86-655, right carpus. **G.** *Arctognathus curvimola* (Owen, 1876), SAM-PK-3329, right carpus. **H.** *Theriognathus microps* Owen, 1876, NHMUK R 5694, right carpus. **I.** Gorgonopsian BP/1/1210, left carpus (reversed). **J.** *Glanosuchus macrops* Broom, 1904, SAM-PK-K7809, left carpus (reversed). **K.** *Ictidosuchoides longiceps* Broom, 1920, or *Ictidosuchops intermedium* (Broom, 1938), BP/1/2294, right carpus. **L.** *Procynosuchus delaharpeae* Broom, 1937, RC92, right carpus. Colours show the different states of distal carpal IV and V. Arrows show the stippled fusion lines between distal carpals IV and V. Abbreviations: mc, metacarpals; di, distal carpals; l ce, lateral centrale; m ce, medial centrale; rl, radiale; ur, ulnare.

1956; Oliveira et al. 1998; Stafford and Thorington 1998; Prochel and Sánchez-Villagra 2003; Gilsanz and Ratib 2005; Fröbisch 2008; Wilson et al. 2010), the fusion of two ossified carpals may appear in a juvenile or subadult stage. An example of this in extant mammals is found in bats, where the radiale, lunate and centrale fuse in a postnatal stage (Stafford and Thorington 1998). As in any endochondral bone, the ossification of carpals is preceded by a cartilaginous precursor (Holmgren 1952; Stafford and Thorington 1998). So, in early ontogeny (prenatally or postnatally or before or after hatching), there is only a cartilaginous precursor present instead of an ossified distal carpal V. So, it is very likely that individuals with fusion of distal carpals IV and V showed two prior states of distal carpal V earlier in ontogeny: a separate ossified distal carpal V in an older ontogenetic stage and an open space representing the cartilaginous precursor of distal carpal V in a younger ontogenetic stage.

Within our sample, specimens of *Procynosuchus delaharpeae* show such an ontogenetic change in distal carpal V. Two specimens show state 2 and one state 5 (Fig. 16D, E, L). The cynodont *Procynosuchus* BP/1/591 is considered a subadult (see also Brink and Kitching 1951) after the immature growth of carpals, which are more rounded and partly surrounded by open spaces, whereas carpals of adult specimens are not surrounded by open spaces and are more edged and angular. The skull and the long bones of BP/1/591 show smaller sizes than the other fossils of the same species (Table 3). In *Procynosuchus* NHMUK PV R 37054, the carpals are not fully grown. It can be verified as being slightly smaller than *Procynosuchus* RC92 (Table 3). NHMUK PV R 37054 might have been close to being mature, but were probably still developing at the time of death. In contrast, RC92 is considered to be an adult (in fact it is one of the largest specimens of *Procynosuchus*).

Within the *Procynosuchus* specimens, the subadult and the nearly mature form (BP/1/591 and NHMUK PV R 37054) show an immature ossification state in distal carpal V, whereas the mature *Procynosuchus* RC92 shows a fused state without fusion line. That means that fusion of distal carpal IV and V would have occurred close to the adult stage.

An alternative explanation is that the observed differences in distal carpal V in these species are individual variations. Individual variants are known from the fusion of the middle and terminal phalanges in the human fifth toe, which do not appear in all individuals, but in 73% and 77% of the Japanese and Chinese (Hong Kong) populations and in 38–46% of Europeans (Nakashima et al. 1995; Chan et al. 2019). Here fused and unfused phalanges are present in

adults. However, in *Procynosuchus*, the different ontogenetic stages are clearly apparent (Table 3), so the variation is likely to be ontogenetic.

*Ictidosuchooides* CGS CM86-655 and ?*Ictidosuchooides* BP/1/2294 also show different states of distal carpal V (states 3 and 5). The identity of BP/1/2294 is not fully established, and that specimen could also belong to *Ictidosuchops* (the taxonomic identity of *Ictidosuchooides* and *Ictidosuchops* needs reevaluation; FA personal observation). BP/1/2294 is slightly bigger than CGS CM86-655 (humerus length difference 8%), but in both specimens the carpals are not completely developed. These specimens show distal carpal V in states 3 and 5 and demonstrate the high variability of that character in immature therocephalians.

In gorgonopsians and therocephalians, the fused state of the distal carpal V, either with or without fusion line, is the most common state. The fossils with fused distal carpal IV and V (states 4 and 5) in both lineages show fully mature carpals with the exception of ?*Ictidosuchooides* BP/1/2294. Also, the basal cynodont *Procynosuchus* with a fused distal carpal IV and V (RC 92) is interpreted as fully adult. Considering the widespread occurrence of the fused state in adults of gorgonopsians and therocephalians and in the basal adult cynodont *Procynosuchus*, the fusion of distal carpal IV and V is very likely a synapomorphy of Theriodontia. This is even more likely because the basal members of these lineages show fusion (therocephalian *Glanosuchus*, gorgonopsians *Arctognathus* and *Viatkogorgon* (SK personal observation on PIN 2212/61) and adult cynodont *Procynosuchus*). However, some later theriodont adult or nearly mature specimens do not exhibit fusion, but rather an immature state of distal carpal V. This is the case for instance in the gorgonopsian cf. *Cynariops* SAM-PK-K10000 and most non-mammalian cynodonts. Cf. *Cynariops* SAM-PK-K10000 and the cynodont *Thrinaxodon* possess a distal carpal V as a small bone within an open space (state 2) and in the cynodonts *Galesaurus* (Fig. 7C), *Exaeretodon* and *Trucidocynodon* (Fig. 16C) distal carpal V was cartilaginous (state 1). Cf. *Cynariops* SAM-PK-K10000 is an adult judging by the form of its carpals (see also Ray et al. 2004). Both *Thrinaxodon* specimens (BP/1/1737 and BP/1/7199) and *Galesaurus* BP/1/2513 are thought to be adults by Jasinowski et al. (2015), Jasinowski and Abdala (2017), and Brink (1965). The carpals of *Thrinaxodon* BP/1/7199 and *Galesaurus* BP/1/2513 are nearly mature, but probably not completely ossified. The carpal condition of *Exaeretodon* and *Trucidocynodon* indicates that they are fully mature. Variations of the states of distal carpal V in the adult stage

Table 3. Length (in mm) of skull and long bones of *Procynosuchus* specimens and the percentual size differences between the specimens.

<i>Procynosuchus delaharpeae</i>	BP/1/591	NHMUK PV R37054	RC92	Length difference (%)	
	subadult	slightly immature	adult	BP/1/591 and RC92	NHMUK PV R37054 and RC92
Basal skull	82	111	144	43	23
Humerus	48	67	80	40	16
Radius	41		61	33	
Ulna	46		72	36	

of gorgonopsians, therocephalians and cynodonts may be due to changes in the timing of the distal carpal V ontogenetic stages. A delay in the development of distal carpal V could result in an unfused variant of distal carpal V or the presence of a cartilaginous precursor in an adult stage, which shows up as an open space in the fossil. This delayed development of distal carpal V is likely to have occurred in the evolution of the gorgonopsian cf. *Cynariops* SAM-PK-K10000 and most cynodonts. Thus, the development of distal carpal V is heterochronically delayed in many non-mammalian Cynodontia (i.e., the distal carpal V evolved paedomorphically, sensu McNamara 2002).

Only the state of cynognathian BP/1/4534 (Fig. 7D) is ambiguous and similar to many mammalian morphs (see below). It has an open space distolateral to the ulnare, but also a relatively wide distal carpal IV. This bone shows no fusion line, no set-back angle or indentation in the outline and resembles the unfused wide distal carpal IV of *Thrinaxodon*, although it is wider than in *Galesaurus* in which distal carpal V was cartilaginous. So, distal carpal IV could be unfused and distal carpal V could have been cartilaginous. However, it is also possible that distal carpals IV and V were fused in the species represented by this specimen.

In mammalian morphs, distal carpal V is not present. Distal carpal IV often extends towards the medial part of metacarpal V, but leaves a space between the lateral part of metacarpal V and the ulnare. Distal carpal V is either not fossilised and stays cartilaginous or is fused to another forelimb bone, most probably to distal carpal IV. In the case of fusion, it is possible that the cartilaginous precursor became fused to another anlage of forelimb bones in early ontogeny, a situation known to occur in some modern mammals (Miláire 1978; Holmgren 1952; Čihák 1972; Slabý 1976).

**Synopsis on character evolution.**—The key transformations in carpal anatomy of synapsids were a morphological change and reduction in size of the intermedium, the loss of intermedium and distal carpal V, the evolution of the medial and lateral centralia, a positional change of distal carpal I and the overall shortening of the carpal elements, especially of the proximal carpals.

In therapsids, the morphology of the intermedium differs markedly from that of “pelycosaurs”, and it is much reduced in size (in length and/or width) in relation to the radiale (Fig. 17: character 1, SOM: fig. 1, table 2: a, e).

Another putative therapsid synapomorphy is the distal shift of distal carpal I, which became aligned partly with the row of remaining distal carpals and partly with the proximal end of the row of metacarpals (Fig. 17: character 2; SOM: fig. 2, table 7: b). In the anomodont *Galechirus* and the mammals analysed here, the distal carpal I was again aligned in the row of distal carpals (SOM: fig. 2). A further distal shift of distal carpal I producing a complete alignment to the row of metacarpals occurs homoplastically in some dicynodonts (*Diictodon* and *Robertia*), one therocephalian (*Microgomphodon*) and some non-mammalian cyno-

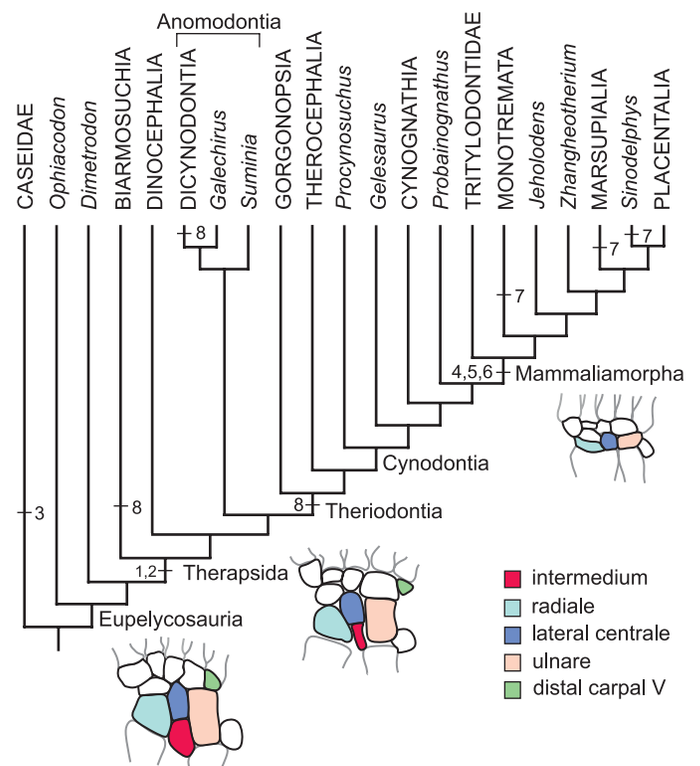


Fig. 17. Phylogenetic tree showing the major evolutionary changes in the carpus during synapsid evolution. For explanation, see text. The phylogeny is based on the same references as Fig. 2A and B.

donts (*Exaeretodon* and *Procynosuchus* RC92; SOM: fig. 2, table 7: b). In caseids, one synapomorphy seems to be a distal contact of the lateral centrale to the medial centrale (Fig. 17: character 3; SOM: table 5: c), whereas the placement of the lateral centrale lateral to the radiale is a synapomorphy of cynodonts (SOM: table 1: f). In mammalian morphs, the intermedium is completely lost or fused to an adjacent carpal (Fig. 17: character 4; SOM: fig. 1). This loss was linked to some important changes in the proximal row of the carpus: the lateral centrale (lunate) became proximally connected to the lateral part of the distal facet of the radius and the radiale became narrower in relation to the radius width, contacting only to the medial side of the radial facet (Fig. 17: character 5; SOM: table 1: b). Simultaneously with these changes, there was a reduction of the relative length of the ulnare (Fig. 17: character 6; SOM: table 3: b). The structure of the proximal row of carpals as a whole, therefore, was reorganized on the transition to mammalian morphs. The size of proximal carpals was further reduced on the line to mammals (SOM: table 3: b), which resulted in a compact and short carpus in Mesozoic mammals.

There is a high degree of plasticity in the condition of the medial centrale, resulting in the loss of the bone by fusion to the radiale, observed in specimens of unrelated lineages: the gorgonopsian *Arctognathus*, the tritylodont *Bienotheroides*, probably in the unnamed cynognathian BP/1/4534 and in one *Thrinaxodon* specimen (BP/1/1737), in which this condition would be an intraspecific variation (SOM: fig. 3).

Loss of the medial centrale with no evidence of fusion to the radiale is also demonstrating a plastic condition observed in the dicynodont *Stahleckeria*, in the evolutionary lines of both monotremes and marsupials, and also in the eutherian *Sinodelphys* (Fig. 17: character 7; SOM: table 3).

The loss of the distal carpal V is an important step towards the reduction of the number of carpal bones, but the record of this loss is widely scattered in punctual records of different therapsids (SOM: table 10: d, e). Absence of distal carpal V or its fusion to distal carpal IV is represented independently in biarmosuchians, dicynodonts and in most adults of theriodonts (as in the basal forms *Viatkogorgon*, *Arctognathus*, *Glanosuchus*, and *Procynosuchus*; Fig. 17: character 8). The condition in theriodonts is, however, complicated by the presence of a separate distal carpal V in the adult gorgonopsian cf. *Cynariops* SAM-PK-K10000, and in the adult non-mammaliaform cynodont *Thrinaxodon*.

With the reduction of two to three carpal bones (intermedium, distal carpal V and in some lineages the medial centrale) and the shortening of the carpus, there is an overall trend of skeletal simplification in the synapsid carpus on the line to mammals, a trend also reported for the cranium of synapsids (Sidor 2001). While the absence or fusion of the medial centrale and the distal carpal V present a highly homoplastic distribution in synapsid phylogeny, the loss of the intermedium occurred only once, as far as can be judged from our sample.

## Conclusions

We analyzed the homology of the carpal bones in fossil representatives of the clade Synapsida from the early Permian to the Late Cretaceous. Our interpretation is in accordance with the traditional view (e.g., Broom 1901, 1904) for the following carpal bones of non-mammaliaform synapsids and mammaliaforms: radiale/scaphoid, ulnare/triquetrum (= cuneiform), pisiform, distal carpal I/trapezium, distal carpal II/trapezoid, distal carpal III/capitate. We homologize the medial centrale of non-mammaliaform synapsids with the single centrale of mammaliaforms. Traditionally, the mammaliaform lunate is considered as a homologue of the intermedium of non-mammaliaform synapsids. We interpret the lunate as the homologue of the lateral centrale of non-mammaliaform synapsids, because of its position and relative proportions, and because of patterns in the embryonal cartilaginous foci of some extant mammals. The proximal contact of the lunate with the radius is an apomorphy of mammaliaforms and the intermedium is lost in this clade. Despite the homologies with the lateral and medial centrale of non-mammaliaform synapsids, we argue for the continued use of the terms “lunate” and “centrale” for the relevant bones in mammaliaform anatomy.

Different carpal elements can be fused to the radiale: in *Arctognathus* both centralia, in *Bienotheroides* the single mammaliaform centrale and in the unidentified

cynognathian BP/1/4534 probably the medial centrale. *Arctognathus* was a fossorial gorgonopsid and the other two taxa also show digging abilities. A digging habit in these taxa would have been facilitated by the stiffening of the carpus through carpal fusion.

In regard to three central bones in *Theriognathus* NHMUK R 5694, we follow Boonstra (1934) and designate them as three centralia on the basis of their positions. In the biarmosuchian PIN 1758/320, we interpret the medial-most of the three central bones as a prepollex/sesamoid, because of its unusual position for a centrale. The open space proximal to distal carpal I in articulated carpi of most non-mammaliaform synapsids, suggests the presence of cartilaginous prepollices/sesamoids at this position during life. The presence of prepollex/sesamoid-like structures in modern mammals and in the tracks of early tetrapods, is a further suggestion that a prepollex/sesamoid could have been present in synapsids during the Permian and Mesozoic.

Distal carpals and metacarpals developmentally derive from each other in ontogeny and remain connected by distinct joints during life, making this connection very stable in ontogeny and phylogeny. Some exceptional situations are: distal carpals II and III highly delayed in development observed in *Diictodon feliceps*, where they evolved paedomorphically. Distal carpal V is lost in Dicynodontia and fused in Biarmosuchia and also in most adults of Gorgonopsia, Therocephalia (including the basal forms of the latter two) and in the adult basal cynodont *Procynosuchus*. In most non-mammaliaform Cynodontia, distal carpal V show a delayed development compared to *Procynosuchus* and basal therocephalians, suggesting distal carpal V evolved paedomorphically. In Mesozoic mammaliaforms, distal carpal V is either cartilaginous or fused, very likely to distal carpal IV.

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## References

- Abdala, F. 2007. Redescription of *Platycraniellus elegans* (Therapsida, Cynodontia) from the Lower Triassic of South Africa, and the cladistic relationships of eutheriodonts. *Palaeontology* 50: 591–618.
- Abdala, F., Rubidge, B.S., and Van den Heever, J.A. 2008. The oldest therocephalians (Therapsida, Eutheriodontia) and the early diversification of Therapsida. *Palaeontology* 51: 1011–1024.
- Albinus, B.S. 1726. *De ossibus corporis humani*. 309 pp. Leidae Batavorum: Apud Henricum Mulhovium, Leiden.
- Amador, L.I., Giannini, N.P., Simmons, N.B., and Abdala, V. 2018. Morphology and evolution of sesamoid elements in bats (Mammalia: Chiroptera). *American Museum Novitates* 3905: 1–38.
- Angielczyk, K.D. and Rubidge, B.S. 2013. Skeletal morphology, phylogenetic relationships and stratigraphic range of *Eosimops newtoni* Broom, a palaecephalid dicynodont (Therapsida, Anomodontia) from the Middle Permian of South Africa. *Journal of Systematic Palaeontology* 11: 191–231.
- Angielczyk, K.D., Fröbisch, J., and Smith, R.M.H. 2005. On the stratigraphic range of the dicynodont taxon *Emydops* (Therapsida: Anomodontia) in the Karoo Basin, South Africa. *Palaeontologia Africana* 41: 23–33.
- Angielczyk, K.D., Sidor, C.A., Nesbitt, S.J., Smith, R.M.H., and Tsuji, L.A. 2009. Taxonomic revision and new observations on the postcranial skeleton, biogeography, and biostratigraphy of the dicynodont genus *Dicynodontoides*, the senior subjective synonym of *Kingoria* (Therapsida, Anomodontia). *Journal of Vertebrate Paleontology* 29: 1174–1187.
- Bardeleben, K. 1889. On the prepollex and prehallux, with observations on the carpus of *Theriodesmus phylarchus*. *Proceedings of the Zoological Society of London* 18: 259–262.
- Bi, S., Wang, Y., Guan, J., Sheng, X., and Meng, J. 2014. Three new Jurassic euharamiyidan species reinforce early divergence of mammals. *Nature* 514: 579–584.
- Bi, S., Zheng, X., Wang, X., Cignetti, N.E., Yang, S., and Wible, J.R. 2018. An early Cretaceous eutherian and the placental–marsupial dichotomy. *Nature* 558: 390–395.
- Biesecker, L.G., Aase, J.M., Clericuzio, C., Gurrieri, F., Temple, I.K., and Toriello, H. 2009. Defining morphology: Hands and feet. *American Journal of Medical Genetics Part A* 149A: 93–127.
- Boonstra, L.D. 1934. A contribution to the morphology of the mammal-like reptiles of the suborder Therocephalia. *Annals of the South African Museum* 31: 215–267.
- Boonstra, L.D. 1965. The girdles and limbs of the Gorgonopsia of the *Tapirocephalus* Zone. *Annals of the South African Museum* 48: 237–249.
- Brink, A.S. 1965. On two new specimens of *Lystrosaurus*-Zone cynodonts. *Palaeontologia Africana* 9: 107–122.
- Brink, A.S. and Kitching, J.W. 1951. On *Leavachia*, a procynosuchid cynodont from the Middle *Cistecephalus* Zone. *South African Journal of Science* 1951: 342–347.
- Broom, R. 1901. On the structure and affinities of *Udenodon*. *Proceedings of the Zoological Society of London* 901: 162–190.
- Broom, R. 1904. The origin of the mammalian carpus and tarsus. *Transactions of the South African Philosophical Society* 15: 89–94.
- Broom, R. 1907. On the origin of the mammal-like reptiles. *Proceedings of the Zoological Society of London* 1907: 1047–1061.
- Broom, R. 1913. On the origin of the mammalian digital formula. *Anatomischer Anzeiger* 43: 230–232.
- Broom, R. 1930. On the structure of the mammal-like reptiles of the suborder Gorgonopsia. *Philosophical Transactions of the Royal Society of London B* 218: 345–371.
- Case, E.C. 1907. Revision of the Pelycosauria of North America. *Carnegie Institution of Washington Publication* 55: 1–176.
- Chan, Y.L.C., Dao, W.H., Yeung, T., and Chow, E.C.S. 2019. Prevalence of toe symphalangism in Hong Kong Chinese population. *Journal of Orthopaedics, Trauma and Rehabilitation* 26 (1): 56–60.
- Chudinov, P.K. [Čudinov, P.K.] 1983. Early therapsids [in Russian]. *Trudy Palaeontologičeskij Institut, Akademiâ Nauk SSSR* 202: 1–229.
- Čihák, R. 1972. Ontogenesis of the skeleton and intrinsic muscles of the human hand and foot. *Advances in Anatomy, Embryology and Cell Biology* 46: 1–194.
- Cohen, K.M., Finney, S.C., Gibbard, P.L., and Fan, J.-X. 2013 (updated). The ICS International Chronostratigraphic Chart. *Episodes* 36: 199–204. [published online, <http://www.stratigraphy.org/ICSChart/ChronostratChart2020-01.pdf>]
- Colbert, E.H. 1948. The mammal-like reptile *Lycaenops*. *Bulletin American Museum of Natural History* 89: 357–404.
- Diaz, R.E. and Trainor, P.A. 2015. Hand/foot splitting and the “re-evolution” of mesopodial skeletal elements during the evolution and radiation of chameleons. *BMC Evolutionary Biology* 15: 184.
- Dilkes, D. 2015. Carpus and tarsus of Temnospondyli. *Vertebrate Anatomy Morphology Palaeontology* 1: 51–87.
- Echeverria, A.I., Abdala, V., Longo, M.V., and Vassallo, A. 2019. Functional morphology and identity of the thenar pad in the subterranean genus *Ctenomys* (Rodentia, Caviomorpha). *Journal of Anatomy* 235: 940–952. [published online, <https://doi.org/10.1111/joa.13049>]
- Fabrezi, M. 2001. A survey of prepollex and prehallux variation in anuran limbs. *Zoological Journal of the Linnean Society* 131: 227–248.
- Fabrezi, M. and Alberch, P. 1996. The carpal elements of anurans. *Herpetologica* 52: 188–204.
- Fabrezi, M. and Barg, M. 2001. Patterns of carpal development among anuran amphibians. *Journal of Morphology* 249: 210–220.
- Fabrezi, M., Abdala, V., and Martínez Oliver, M.I. 2007. Developmental basis of limb homology in lizards. *Anatomical Record* 290: 900–912.
- Fabrezi, M., Manzano, A., Abdala, V., and Zaher, H. 2009. Developmental basis of limb homology in pleurodiran turtles, and the identity of the hooked element in the chelonian tarsus. *Zoological Journal of the Linnean Society* 155: 845–866.
- Flores, D.A. and Diaz, M.M. 2009. Postcranial skeleton of *Glironia venusta* (Didelphimorphia, Didelphidae, Caluromyinae): Description and functional morphology. *Zoosystematics and Evolution* 85: 311–339.
- Flower, W.H. 1885. *An Introduction to the Osteology of the Mammalia*. 375 pp. Macmillan and Co., London.
- Fontanarrosa, G. and Abdala, V. 2014. Anatomical analysis of the lizard carpal bones in the terms of skilled manual abilities. *Acta Zoologica* 95: 249–263.
- Fontanarrosa, G. and Abdala, V. 2016. Bone indicators of grasping hands in lizards. *PeerJ* 4:e1978.
- Fontanarrosa, G., Abdala, F., Kümmell, S.B., and Gess, R. 2019. The manus of *Tetracyonodon* (Therapsida: Therocephalia) provides evidence

- for survival strategies following the Permian extinction. *Journal of Vertebrate Paleontology* 38: 4.
- Fröbisch, J. and Reisz, R.R. 2011. The postcranial anatomy of *Suminia geymanovi* (Synapsida: Anomodontia), the earliest known arboreal tetrapod. *Zoological Journal of the Linnean Society* 162: 661–698.
- Fröbisch, N.B. 2008. Ossification patterns in the tetrapod limb—conservation and divergence from morphogenetic events. *Biological Reviews* 83: 571–600.
- Gegenbaur, C. 1864. *Untersuchungen zur vergleichenden Anatomie der Wirbeltiere. Erstes Heft: Carpus und Tarsus*. 127 pp. Verlag von Wilhelm Engelmann, Leipzig.
- Gillies, C.D. 1929. The origin of the os pisiform. *Journal of Anatomy* 63: 380–383.
- Gilmore, C.W. 1919. A mounted skeleton of *Dimetrodon gigas* in the United States National Museum with notes on the skeletal anatomy. *Proceedings of the United States National Museum* 56: 525–539.
- Gilsanz, V. and Ratib, O. 2005. *Hand bone age. A digital Atlas of Skeletal Maturity*. 106 pp. Springer, Berlin.
- Grassé, P.P. 1955. *Traité de Zoologie. Anatomie. Systématique. Biologie. Mammifère. Les ordres: anatomie, éthologie, systématique*. Tome XVII, 1147–1447. Masson and Co., Paris.
- Gregory, W.K., Miner, R.W., and Noble G.K. 1923. The carpus of *Eryops* and the structure of the primitive chiropterygium. *Bulletin of the American Museum of Natural History* 48: 279–288.
- Haines, R.W. 1969. Epiphyses and sesamoids. In: C. Gans, A.d'A. Bellairs, and T.S. Parsons, (eds.), *Biology of the Reptilia. Volume 1. Morphology A*, 81–115. Academic Press, London.
- Henrici, A.C., Berman, D.S., Lucas, S.G., Heckert, A.B., Rinehart, L.F., and Zeigler, K.E. 2005. The carpus and tarsus of the early Permian synapsid *Sphenacodon ferox* (Eupelycosauria: Sphenacodontidae). The nonmarine Permian. *New Mexico Museum of Natural History and Science Bulletin* 30: 106–110.
- Heppleston, A.C. 2010. *Patterns and Processes of Digit Number Reduction*. Ph.D. Thesis, McGill University, Montreal. Available at [http://digitool.library.mcgill.ca/R/?func=dbin-jump-full&object\\_id=92163&local\\_base=GEN01-MCG02](http://digitool.library.mcgill.ca/R/?func=dbin-jump-full&object_id=92163&local_base=GEN01-MCG02)
- Hildebrand, M. and Goslow, G.E. 2004. *Vergleichende und funktionelle Anatomie der Wirbeltiere*. 709 pp. Springer, Berlin.
- Hoffman, E.A. and Rowe, T.B. 2018. Jurassic stem-mammal perinates and the origin of mammalian reproduction and growth. *Nature* 561: 104–108.
- Holmgren, N. 1933. On the origin of the tetrapod limb. *Acta Zoologica* 14: 185–295.
- Holmgren, N. 1952. An embryological analysis of the mammalian carpus and its bearing upon the question of the origin of tetrapod limb. *Acta Zoologica* 33: 1–115.
- Hopson, J.A. 1995. Patterns of evolution in the manus and pes of non-mammalian therapsids. *Journal of Vertebrate Paleontology* 15: 615–639.
- Hopson, J.A. and Barghusen, H. 1986. An analysis of therapsid relationships. In: N. Hotton, P.D. MacLean, J.J. Roth, and E.C. Roth (eds.), *The Ecology and Biology of the Mammal-like Reptiles*, 83–106. Smithsonian Institution Press, Washington.
- Hu, Y.M., Wang, Y.Q., Li, C.K., and Luo, Z.X. 1998. Morphology of dentition and forelimb of *Zhangheotherium*. *Vertebrata Palasiatica* 36: 102–125.
- Huttenlocker, A.K. and Smith, R.M.H. 2017. New whaitsioids (Therapsida: Therocephalia) from the Teekloof Formation of South Africa and therocephalian diversity during the end-Guadalupian extinction. *PeerJ* 5: e3868.
- Ihle, J.E.W., Kampen, P.N. van, Nierstrasz, H.F., and Versluys, J. 1927. *Vergleichende Anatomie der Wirbeltiere*. 906 pp. Verlag von Julius Springer, Berlin.
- Jasinoski, S.C., Abdala, F., and Fernandez, V. 2015. Ontogeny of the Early Triassic cynodont *Thrinaxodon liorhinus* (Therapsida): cranial morphology. *The Anatomical Record* 298: 1440–1464.
- Jasinoski, S.C. and Abdala, F. 2017. Cranial ontogeny of the Early Triassic basal cynodont *Galesaurus planiceps*. *The Anatomical Records* 300 (2): 353–381.
- Jenkins, F.A., 1971. The postcranial skeleton of African cynodonts. *Peabody Museum of Natural History, Yale University, Bulletin* 36:1–216.
- Ji, Q., Luo, Z.X., Wible, J.R., Zhang, J.P., and Georgi, J.A. 2002. The earliest known eutherian mammal. *Nature* 416: 816–822.
- Johanson, Z., Josse, J., Boisvert, C.A., Ericsson, R., Sutija, M., and Ahlberg, P.E. 2007. Fish fingers: Digit homologues in sarcopterygian fish fins. *Journal of Experimental Zoology Part B* 308: 757–768.
- Kammerer, C.F. 2011. Systematics of the Anteosauria (Therapsida: Dinocephalia). *Journal of Systematic Palaeontology* 9: 261–304.
- Kammerer, C.F. 2016. Systematics of the Rubidgeinae (Therapsida: Gorgonopsia). *PeerJ* 4: e1608.
- Kammerer, C.F. and Masyutin, V. 2018. Gorgonopsian therapsids (*Nochnitsa* gen. nov. and *Viatkogorgon*) from the Permian Kotelnich locality of Russia. *PeerJ* 6: e4954.
- Kammerer, C.F., Angielczyk, K.D., and Fröbisch, J. 2011. A comprehensive taxonomic revision of *Dicynodon* (Therapsida, Anomodontia) and its implications for dicynodont phylogeny, biogeography, and biostratigraphy. *Journal of Vertebrate Paleontology* 31 (Supplement to 6): 1–158.
- Keyser, A.W. 1993. A re-evaluation of the smaller Endothiodontidae. *Memoirs of the Geological Survey of South Africa* 82: 1–53.
- Kielan-Jaworowska, Z. 1977. Evolution of the therian mammals in the Late Cretaceous of Asia. Part II. Postcranial skeleton in *Kennalestes* and *Asioryctes*. In: Z. Kielan-Jaworowska (ed.), Results Polish Mongolian Palaeontological Expeditions. *Palaeontologica Polonica* 37: 65–84.
- Kielan-Jaworowska, Z. 1978. Evolution of therian mammals in the Late Cretaceous of Asia. Part III. Postcranial skeleton in Zalambdalestidae. In: Z. Kielan-Jaworowska (ed.), Results Polish Mongolian Palaeontological Expeditions. *Palaeontologica Polonica* 38: 5–41.
- Kielan-Jaworowska, Z., Cifelli, R.L., and Luo, Z.X. 2004. *Mammals from the Age of Dinosaurs. Origins, Evolution, and Structure*. 630 pp. Columbia University Press, New York.
- Kindahl, M. 1941. Untersuchungen einiger Entwicklungsstadien der Hand und des Fußes von *Erinaceus europaeus*. *Zeitschrift für mikroskopisch-anatomische Forschung* 50: 458–464.
- Kindahl, M. 1942a. Beitrag zur Kenntnis der Entwicklung des Extremitätenskelets bei *Centetes ecaudatus* und *Ericulus setosus*. *Zeitschrift für mikroskopisch-anatomische Forschung* 51: 322–333.
- Kindahl, M. 1942b. Einige Mitteilungen über die Entwicklung der Hand und des Fußes bei *Talpa europaea* L. *Zeitschrift für mikroskopisch-anatomische Forschung* 52: 267–273.
- Kindahl, M. 1944. On the development of the hand and the foot of *Tarsius tarsius* and *Microcebus myoxinus*. *Acta Zoologica* 25: 49–58.
- Kivell, T.L. 2016. The primate wrist. In: T.L. Kivell, P. Lemelin, B.G. Richmond, and D. Schmitt (eds.), *The Evolution of the Primate Hand. Anatomical, Developmental, Functional, and Paleontological Evidence*, 17–54. Springer, New York.
- Kjosness, K.M., Hines, J.E., Lovejoy, C.O., and Reno, P.L. 2014. The pisiform growth plate is lost in humans and supports a role for Hox in growth plate formation. *Journal of Anatomy* 225: 527–538.
- Kümmell, S.B. 2009. *Die Digi der Synapsida: Anatomie, Evolution und Konstruktionsmorphologie*. 424 pp. Ph.D. Thesis, Fakultät für Biowissenschaften der Universität Witten/Herdecke. Shaker Verlag, Aachen.
- Kümmell, S.B. and Frey, E. 2012. What digits tell us about digging, running and climbing in recent and fossil Synapsida. *Fundamental* 20: 117–119.
- Kümmell, S.B. and Frey, E. 2014a. Autopodial rotation as a measure for stance and gait in Synapsida from early Permian to later Cretaceous. In: M. Delfino, G. Carnevale, and M. Pavia (eds.), *EAVP XII Annual Meeting, Abstract Book and Field Trip Guide*, 91. Museo Regionale di Scienze Naturali, Torino.
- Kümmell, S.B. and Frey, E. 2014b. Range of movement in ray I of manus and pes and the prehensibility of the autopodia in the early Permian to Late Cretaceous non-anomodont Synapsida. *PLoS One* 9: e113911.
- Liu, J., Schneider, V.P., and Olsen, P.E. 2017. The postcranial skeleton of

- Boreogomphodon* (Cynodontia: Traversodontidae) from the Upper Triassic of North Carolina, USA and the comparison with other traversodontids. *PeerJ* 5: e3521.
- Luo, Z.X. 2011. Developmental patterns in Mesozoic evolution of mammal ears. *Annual Review of Ecology, Evolution, and Systematics* 42: 355–80.
- Luo, Z.X., Ji, Q., Wible, J.R., and Yuan, C.X. 2003. An early Cretaceous tribosphenic mammal and metatherian evolution. *Science* 302: 1934–1940.
- Luo, Z.X., Yuan, C.X., Meng, Q.J., and Ji, Q. 2011. A Jurassic eutherian mammal and divergence of marsupials and placentals. *Nature* 476: 442–445.
- Lyser, M. 1653. *Culter anatomicus*. 217 pp. Lamprecht, Copenhagen.
- Maddison, W.P. and Maddison, D.R. 2019. *Mesquite: a Modular System for Evolutionary Analysis*. Version 3.61 <http://www.mesquiteproject.org>
- Maisch, M.W. 2001. Observations on Karoo and Gondwana vertebrates. Part 2: A new skull-reconstruction of *Stahleckeria potens* von Huene, 1935 (Dicynodontia, Middle Triassic) and a reconsideration of kannemeyeriiform phylogeny. *Neues Jahrbuch für Geologie und Paläontologie, Abhandlungen* 220: 127–152.
- McNamara, K.J. 2002. What is heterochrony. In: N. Minugh-Purvis, K.J. McNamara (eds.), *Human Evolution through Developmental Change*, 1–4. The Johns Hopkins University Press, Baltimore.
- McMurrich, J.P. 1914. The nomenclature of the carpal bones. *Anatomical Record* 8: 173–182.
- Meng, Q.J., Ji, Q., Zhang, Y.G., Liu, D., Grossnickle, D.M., and Luo, Z.X. 2015. An arboreal docodont from the Jurassic and mammaliaform ecological diversification. *Science* 347: 764–768.
- Milaire, J. 1978. Étude morphologique, histochemique et autoradiographique de développement du squelette des membres chez l'embryon de souris. I. Membres antérieurs. *Archives Biologiques (Brussels)* 89: 169–216.
- Modesto, S.P., Smith, R.M.H., Campione, N.E., and Reisz, R.R. 2011. The last “pelycosaur”: a varanopid synapsid from the Pristerognathus Assemblage Zone, Middle Permian of South Africa. *Naturwissenschaften* 98: 1027–1034.
- Monro, A. 1726. *The Anatomy of the Humane Bones*. 352 pp. Ruddiman, Edinburgh.
- Müller, G.B. 1991. Evolutionary transformation of limb pattern: heterochrony and secondary fusion. In: J.R. Hinchliffe, J.M. Hurlé, and D. Summerbell (eds.), *Developmental Patterning of the Vertebrate Limb*, 395–405. Plenum Press, New York.
- Müller, G.B. and Alberch, P. 1990. Ontogeny of the limb skeleton in *Alligator mississippiensis*: developmental invariance and change in the evolution of the archosaur limbs. *Journal of Morphology* 203: 151–164.
- Nakashima, T., Hojo, T., Suzuki, K., and Ijichi, M. 1995. Symphalangism (two phalanges) in the digits of the Japanese foot. *Annals of Anatomy* 177: 275–278.
- Nesslinger, C.L. 1956. Ossification centers and skeletal development in the postnatal virginia opossum. *Journal of Mammalogy* 37: 382–394.
- O’Keefe, F.R., Sidor, C.A., Larsson, H.C.E., Maga, A., and Ide, O. 2006. Evolution and homology of the astragalus in early amniotes: New fossils, new perspectives. *Journal of Morphology* 267: 415–425.
- Oliveira, C.A. de, Nogueira, J.C., and Bohórquez Mahecha, G.A. 1998. Sequential order of appearance of ossification centers in the opossum *Didelphis albiventris* (Didelphidae) skeleton during development in the Marsupium. *Annals of Anatomy* 180: 113–121.
- Prochel, J., and Sánchez-Villagra, M.R. 2003. Carpal ontogeny in *Monodelphis domestica* and *Caluromys philander* (Marsupialia). *Zoology* 106: 73–84.
- Ray, S. and Chinsamy, A. 2003. Functional aspects of the postcranial anatomy of the Permian dicynodont *Diictodon* and their ecological implications. *Palaeontology* 46: 151–183.
- Ray, S., Botha, J., and Chinsamy, A. 2004. Bone histology and growth patterns of some nonmammalian therapsids. *Journal of Vertebrate Paleontology* 24: 634–648.
- Reisz, R.R. 1986. Pelycosauria. *Encyclopedia of Paleoherpétology*. Part 17A: 1–102. Gustav Fischer Verlag, Stuttgart.
- Reisz, R.R., Maddin, H.C., Fröbisch, J., and Falconnet, J. 2011. A new large caseid (Synapsida, Caseasauria) from the Permian of Rodez (France), including a reappraisal of “*Casea*” *rutena* Sigogneau-Russell & Russell, 1974. *Geodiversitas* 33: 227–246.
- Reno, P.L., Kjosness, K.M., and Hines, J.E. 2016. The role of Hox in pisiform and calcaneus growth plate formation and the nature of the zeugopod/autopod boundary. *Journal of Experimental Zoology Part B* 326: 303–332.
- Romer, A.S. and Parsons, T.S. 1977. *The Vertebrate Body*. 624 pp. Saunders College Publishing, Philadelphia.
- Romer, A.S. and Price, L.I. 1940. Review of the Pelycosauria. *Special Paper Geological Society of America* 28: 1–538.
- Romer, A.S. and Price, L.I. 1944. *Stahleckeria lenzii*, a giant Triassic Brazilian dicynodont. *Bulletin Museum of Comparative Zoology* 93: 465–491.
- Rowe, T. 1988. Definition, diagnosis and origin of mammalia. *Journal of Vertebrate Paleontology* 8: 241–264.
- Ruta, M., Botha-Brink, J., Mitchell, S.A., and Benton, M.J. 2013. The radiation of cynodonts and the ground plan of mammalian morphological diversity. *Proceedings of the Royal Society B: Biological Sciences* 280: 20131865.
- Salesa, M.J., Antón, M., Peigné, S., and Morales, J. 2006. Evidence of a false thumb in a fossil carnivore clarifies the evolution of pandas. *Proceedings of the National Academy of Sciences* 103: 379–382.
- Salomon, F.V., Geyer, H., and Gille, U. 2005. *Anatomie für die Tiermedizin*. 790 pp. Enke Verlag, Stuttgart.
- Salton, J.A., and Sargis, E.J. 2008. Evolutionary morphology of the Tenrecoidea (Mammalia) carpal complex. *Biological Journal of the Linnean Society* 93: 267–288.
- Schmidt-Ehrenberg, E.C. 1942. Die Embryogenese des Extremitätenskelettes der Säugetiere. *Revue Suisse de Zoologie* 49: 33–131.
- Schneider, J.W., Lucas, S.G., Scholze, F., Voigt, S., Marchetti, L., Klein, H., Opluštil, S., Werneburg, R., Golubev, V.K., Barrick, J.E., Nemyrovskaya, T., Ronchi, A., Day, M.O., Silantiev, V.V., Rößler, R., Sabero, H., Linnemann, U., Zharinova, V., and Shen, S.Z. 2020. Late Paleozoic–early Mesozoic continental biostratigraphy. Links to the Standard Global Chronostratigraphic Scale. *Palaeoworld* 29 (2): 186–238.
- Seeley, H.G. 1888. Researches on the structure, organization and classification of the fossil Reptilia. III. On parts of the skeleton of a mammal from Triassic rocks of Klipfontein, Fraserberg, South Africa (*Theriodesmus phylarchus*, Seeley), illustrating the reptilian inheritance in the mammalian hand. *Philosophical Transactions of the Royal Society, Series B* 179: 141–155.
- Seeley, H.G. 1895. Researches on the structure, organization and classification of the fossil Reptilia. Part IX, Section 4. On the Gomphodontia. *Philosophical Transactions of the Royal Society, Series B* 186: 1–57.
- Shubin, N.H. and Alberch, P. 1986. A morphogenetic approach to the origin and basic organization of the tetrapod limb. *Evolutionary Biology* 20: 319–387.
- Sidor, C.A. 2001. Simplification as a trend in synapsid cranial evolution. *Evolution* 55: 1419–1442.
- Sidor, C.A. and Hopson, J.A. 1998. Ghost lineages and “mammalness”: assessing the temporal pattern of character acquisition in the Synapsida. *Paleobiology* 24: 254–273.
- Sigogneau-Russell, D. and Russell, D.E. 1974. Étude du premier caséidé (Reptilia, Pelycosauria) d’Europe occidentale. *Bulletin du Muséum national d’Histoire naturelle* 230: 145–215.
- Slabý, O. 1967. *Die Morphogenese und phylogenetische Morphologie des Carpus der Paarhufer*. 55 pp. Academia Nakladatelství Československé Akademie Věd, Praha.
- Slabý, O. 1968. Wege und Gesetzmäßigkeiten der Evolution in Bezug auf die phylogenetische Entwicklung der Extremitäten. *Acta Universitatis Carolinae Medica. Monographia* 35: 1–137.
- Smith, R.M. 1987. Helical burrow casts of therapsid origin from the Beaufort Group (Permian) of South Africa. *Palaeogeography, Palaeoclimatology, Palaeoecology* 60: 155–170.
- Stafford, B.J. and Thorington, R.W. 1998. Carpal development and morphology in archontan mammals. *Journal of Morphology* 235: 135–155.

- Starck, D. 1979. *Vergleichende Anatomie der Wirbeltiere, auf evolutionsbiologischer Grundlage. Vol. II: Das Skelettsystem. Allgemeines, Skeletsubstanzen, Skelet der Wirbeltiere einschließlich Lokomotionstypen.* 776 pp. Springer Verlag, Berlin.
- Steiner, H. 1921. Hand und Fuss der Amphibien, ein Beitrag zur Extremitätenfrage. *Anatomischer Anzeiger* 53: 513–542.
- Steiner, H. 1935. Beiträge zur Gliedmaßentheorie. Die Entwicklung des Chiropterygium aus dem Ichthyopterygium. *Revue Suisse de Zoologie* 42: 715–729.
- Steiner, H. 1942. Der Aufbau des Säugetier-Carpus und -Tarsus nach neuen embryologischen Untersuchungen. *Revue Suisse de Zoologie* 49: 217–223.
- Stovall, J.W., Price, L.I., and Romer, A.S. 1966. The postcranial skeleton of the giant Permian pelycosaur *Cotylorhynchus romeri*. *Bulletin of the Museum of Comparative Zoology* 135: 1–30.
- Sues, H.D. and Jenkins, F.A. 2006. The postcranial skeleton of *Kayentatherium wellsi* from the Lower Jurassic Kayenta Formation of Arizona and the phylogenetic significance of postcranial features in tritylodontid cynodonts. In: M.T. Carrano, T.J. Gaudin, R.W. Blob, and J.R. Wible (eds.), *Amniote Paleobiology. Perspectives on the Evolution of Mammals, Birds, and Reptiles*, 114–152. The University of Chicago Press, Chicago.
- Sun, A. and Li, Y. 1985. The postcranial skeleton of the late tritylodont *Bienotheroides* [in Chinese with English summary]. *Vertebrata Palasiatica* 23: 133–151.
- Szalay, F.S. 1994. *Evolutionary History of the Marsupials and an Analysis of Osteological Characters.* 481 pp. Cambridge University Press, Cambridge.
- Vega-Dias, C., Maisch, M., and Schultz, C.L. 2004. A new phylogenetic analyses of Triassic dicynodonts (Therapsida) and the systematic position of *Jachaleria candelariensis* from the Upper Triassic of Brazil. *Neues Jahrbuch für Geologie und Paläontologie, Abhandlungen* 231: 145–166.
- Velazco, P.M., Buczek, A.J., and Novacek, M.J. 2017. Two new tritylodontids (Synapsida, Cynodontia, Mammalia) from the Upper Jurassic, Southwestern Mongolia. *American Museum Novitates* 3874: 1–35.
- Voigt, S. and Ganzelewski, M. 2010. Toward the origin of amniotes: Diadectomorph and synapsid footprints from the early late Carboniferous of Germany. *Acta Palaeontologica Polonica* 55: 57–72.
- Voigt, S. and Lucas, S.G. 2017. Early Permian tetrapod footprints from central New Mexico. New Mexico. *New Mexico Museum of Natural History and Science Bulletin* 77: 333–352.
- Wagner, G.P., Chiu, C.H., and Laubichler, M. 2000. Developmental evolution as a mechanistic science: The inference from developmental mechanisms to evolutionary processes. *American Zoologist* 40: 819–831.
- Watabe, M., Tsubamoto, T., and Tsogtbaatar, K. 2007. A new tritylodontid synapsid from Mongolia. *Acta Palaeontologica Polonica* 52: 263–274.
- Wilson, L.A.B., Schradin, C., Mitgutsch, C., Galliari, F.C., Mess, A., and Sánchez-Villagra, M.R. 2010. Skeletogenesis and sequence heterochrony in rodent evolution, with particular emphasis on the African striped mouse, *Rhodomys pumilio* (Mammalia). *Organisms Diversity & Evolution* 10: 243–258.
- Yalden, D.W. 1966. The anatomy of mole locomotion. *Journal of Zoology* 149: 55–64.