## The earliest phyllolepid (Placodermi, Arthrodira) from the Late Lochkovian (Early Devonian) of Yunnan (South China)

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**Abstract** – *Gavinaspis convergens*, a new genus and species of the Phyllolepida (Placodermi: Arthrodira), is described on the basis of skull remains from the Late Lochkovian (Xitun Formation, Early Devonian) of Qujing (Yunnan, South China). This new form displays a mosaic of characters of basal actinolepidoid arthrodires and more derived phyllolepids. A new hypothesis is proposed concerning the origin of the unpaired centronuchal plate of the Phyllolepida by a fusion of the paired central plates into one single dermal element and the loss of the nuchal plate. A phylogenetic analysis suggests the position of *Gavinaspis* gen. nov. as the sister group of the Phyllolepidiae, in a distinct new family (Gavinaspidiae fam. nov.). This new form suggests a possible Chinese origin for the Phyllolepida or that the common ancestor to Phyllolepida lived in an area including both South China and Gondwana, and in any case corroborates the palaeogeographic proximity between Australia and South China during the Devonian Period.

Keywords: Devonian, China, Placodermi, phyllolepids, biostratigraphy, palaeobiogeography.

### 1. Introduction

The Phyllolepida are a peculiar group of the Arthrodira (Placodermi), widespread in the Givetian-Famennian of Gondwana (Australia, Antarctica, Turkey, South America: e.g. Janvier, 1983; Long, 1984, 2003; Ritchie, 1984, 2005; Young, 1984, 1988, 1991, 2005a,b,c; Young & Moody, 2002; Young & Long, 2005; Young, Long & Turner, 1993; Young, Moody & Casas, 2000) and in the Famennian of Euramerica (North America, Greenland, West Europe, Russia: e.g. Agassiz, 1844; Denison, 1978; Heintz, 1930; Lane, Cuffey & Daeschler, 2001; Leriche, 1931; Lohest, 1888; Newberry, 1889; Rohon, 1900; Stensiö, 1934, 1969). They are characterized by a dorsoventrally flattened dermal armour, the lack of rostral and pineal plates, and the presence of a large unpaired centronuchal plate, for which no consensus exists as to its origin (either due to the fusion of nuchal and central plates, or to the loss of central plates). The centronuchal plate is surrounded by a 'ring' of perinuchal plates. There is no external foramen for the endolymphatic duct on the paranuchal plates. Post-marginal plates are lacking, and anterior and/or posterior median ventral plates are reduced. The ornamentation mainly consists of ridges lacking semidentine.

First considered as belonging to the 'coelacanths' (Agassiz, 1844), the 'crossopterygians' (Zittel, 1887–90), the Placodermi (Newberry, 1889) and the Heterostraci (Woodward, 1915), the Phyllolepida were finally assigned with certainty to the Placodermi (Stensiö,

1934). Subsequently, they were considered as either sharing an immediate common ancestor with the Arthrodira (Denison, 1978), belonging to the Actinolepidoidei (Long, 1984), or being of indetermined position within the Arthrodira (Goujet & Young, 1995). They were considered as close to *Wuttagoonaspis* (Long, 1984; Young, 1980; Young & Goujet, 2003) and/or to Antarctaspis (Denison, 1978; Long, 1984). More recently, a sister-group relationship between the Phyllolepida (as a crownward member of the top of an 'actinolepidoid' paraphyletic ensemble) and the Phlyctaenioidei ('Phlyctaenii' plus Brachythoraci) was proposed, and Antarctaspis and Wuttagoonaspis are the most inclusive taxa among the Arthrodira (Dupret, 2004). However, the latter hypothesis was not followed by Young (2005a,b,c), who argued that the ridged ornamentation of the Phyllolepida and Wuttagoonaspis is homologous and therefore is an important synapomorphy of this ensemble.

Because the phyllolepid remains were abundantly encountered in the eastern margin of Gondwana before the Famennian, and because the earliest known and most primitive forms were found in Australia, it was commonly accepted that this group originated from Australia. However, the most plesiomorphic phyllolepids so far known (*Cobandrahlepis* Young, 2005*c*, and *Yurammia* Young, 2005*c*, both dated as Givetian) already possess many phyllolepid diagnostic features (e.g. the presence of the centronuchal plate). As a consequence, the phylogenetic relationships between the Phyllolepida and the other groups of Placodermi have long been debated, without leading to a consensus.

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Table 1. Abbreviations used in text and figures

Abbreviation	Definition								
Anatomical structure									
a.po.p	anterior postorbital process								
ald.c	canal for the lateral dorsal aorta								
сс	central sensory line groove								
Ce	central plate								
CeN	centronuchal plate								
d.end	endolymphatic duct								
d.end.e	external foramen for the endolymphatic duct								
d.end.i	internal (proximal) foramen for the endolymphatic duct								
d.mc	cucullaris fossa								
f.s.PaN	sub-paranuchal fossa								
ioc	infraorbital sensory line groove								
lc	main lateral sensory line groove								
М	marginal plate								
mpl	middle pitline								
N	nuchal plate								
oa.ADL	overlap area for the anterior dorsolateral plate (sliding dermal craniothoracic articulation)								
oa.CeN	overlap area of the paranuchal plates for the centronuchal plate								
oa.RC	overlap area for the dermal rostral capsule								
occ	occipital cross commissure								
occ.p	posterior occipital process								
o.n	orbital notch								
p.po.p	posterior postorbital process								
PaN	paranuchal plate								
Pi	pineal plate								
pmc	postmarginal sensory line groove								
ppl	posterior pitline								
PrO	preorbital plate								
PtO	postorbital plate								
R	rostral plate								
r.csa	ridge impression of the anterior semi-circular canal on the dorsal surface of neurocranium								
r.csp	ridge impression of the posterior semi-circular canal on the dorsal surface of neurocranium								
r.nc	impression of the neural canal								
SOC	supraorbital sensory line groove								
spio.a.b.c	foramina for spino-occipital nerves								
sv.p	supravagal process								
Tree statistics									
n	number of equally parsimonious trees								
L	tree length								
CI	consistency index								
RI	retention index								
SC	strict consensus								

It is also noteworthy that despite the close connection between South China and Australia during the Devonian Period (after the Pragian: Zhu, 2000; Zhu & Zhao, 2006; Zhu, Wang & Wang, 2000), no phyllolepid remains (or closely related forms) have been found in China. The new form *Gavinaspis convergens* described here throws some new light onto the systematic and palaeogeographic origin of the Phyllolepida.

### 2. Material and methods

All of the referred specimens are housed in the IVPP (Institute of Vertebrate Palaeontology and Palaeoanthropology, Beijing, People's Republic of China). They were collected by the Early Vertebrate Research Group of IVPP during recent field trips (1998–2006) to Quijng, Yunnan, and have been prepared mechanically.

Abbreviations for placoderm dermal bones and other structures, and phylogenetic abbreviations as used in the text and figures, are listed in Table 1.

### 3. Geological setting

The specimens described herein come from the middle part of the Xitun Formation at a locality close to Xitun village in the suburb of Qujing (Yunnan, South China, Figs 1, 2; Diabolepis-Nanpanaspis macrovertebrate assemblage II of Zhu, Wang & Wang, 2000; Late Lochkovian; only one locality). The matrix of the specimens is a light-grey clayey limestone. The Xitun Formation of Qujing yields remains of the Galeaspida Eugaleaspis changi, Nanpanaspis microculus, Laxaspis qujingensis, Cyclodiscaspis ctenus (Liu, 1965, 1975), Microholonaspis microthyris, Hyperaspis acclivi (Pan, 1992), the Thelodonti Parathelodus scitulus, P. asiatica, P. catalatus, P. trilobatus, P. cornuformis (Wang, 1997), the Chondrichthyes Gualepis elegans, Changolepis tricuspidus, Peilepis solida, Ohiolepis? xitunensis, the Acanthodii Nostolepis sp., Youngacanthus gracili (Wang, 1984), the Sarcopterygii Youngolepis praecursor (Chang & Yu, 1981), Diabolepis speratus (Chang & Yu, 1984), Psarolepis romeri



Figure 1. Position of Qujing, Yunnan, South China.

(Yu, 1998; Zhu, Yu & Janvier, 1999), Achoania jarviki (Zhu, Yu & Ahlberg, 2001), Styloichthys changae (Zhu & Yu, 2002), and Meemannia eos (Zhu et al. 2006), the Antiarcha Yunnanolepis chii (Liu, 1963), Y. parvus, Y. porifera, Phymolepis cuifengshanensis, P. guoruii, Zhanjilepis aspratilis, Chuchinolepis gracilis, C. qujingensis, C. sulcata, C. robusta (Chang, 1978; Zhang, 1978, 1984; Zhu, 1996), the Arthrodira Szelepis yunnanensis (Liu, 1979, 1981) and the Petalichthyida (Zhu, 2000; pers. obs.).

### 4. Systematic palaeontology

PLACODERMI McCoy, 1848 ARTHRODIRA Woodward, 1891 PHYLLOLEPIDA Stensiö, 1934

*Diagnosis.* 'Actinolepidoid' Arthrodira in which an unpaired centronuchal plate is present and no paired central plates are identified. Centronuchal plate is surrounded by four or five paired bones including preorbital, postorbital, marginal and paranuchal plates. Post-marginal plates are absent.

*Remarks.* The diagnosis above and systematic rank are modified after Young (2005*b*, p. 175) to incorporate new data on the Chinese phyllolepid. Despite the presence of an unambiguous centronuchal plate, we consider that the skull roof pattern and the outline of the associated neurocranium of the new form described herein do not fit the recently emended diagnosis of the family Phyllolepididae Woodward, 1891 (non rank-order Phyllolepida Stensiö, 1934) proposed by Young (2005*b*,*c*) or by Ritchie (2005). Moreover, the phylogenetic analysis attempted herein clearly indicates a sister-group relationship between *Gavinaspis* gen. nov. and the Phyllolepididae. Other characters mentioned in



Figure 2. Stratigraphic column of the Lower Devonian deposits in Yunnan, South China. The fish icon represents the stratigraphic position of *Gavinaspis convergens* gen. et sp. nov. Modified after Zhu & Zhao, 2006.

Young's (2005*b*) diagnosis are still applicable for the family Phyllolepididae Woodward, 1891 (all Phyllolepida except Gavinaspididae fam. nov.): centronuchal plate as broad as or broader than long, and surrounded by five smaller paired bones; rostral and pineal plates absent from skull roof; posterolateral plate absent from trunk armour; median dorsal plate without an inner keel, and anterior dorsolateral plate with a narrow elongate area; posterior ventrolateral plate of triangular shape, lacking a lateral lamina; dermal ornament mainly consisting of smooth concentric ridges, with some tubercles and tubercle rows.

#### Family GAVINASPIDIDAE fam. nov.

*Diagnosis.* Same as for the type genus and its type species, by monotypy.

Type genus. Gavinaspis gen. nov.

Genus Gavinaspis gen. nov.

Diagnosis. Same as for the type species, by monotypy.



Figure 3. Schematic reconstruction of the skull roof of *Gavinaspis convergens* gen. et sp. nov. Tectonic dextral shear distortion has been corrected by means of computerized assistance (torsion  $5^{\circ}$ ). (a), dorsal view; (b) ventral view. Scale bar 1 cm. For key to abbreviations, see Table 1.

*Etymology.* To acknowledge the major contribution to the study of phyllolepid placoderms by Dr Gavin C. Young, Australian National University, Canberra.

Type species. Gavinaspis convergens gen. et sp. nov.

### Gavinaspis convergens gen. et sp. nov. Figures 3–7

*Holotype.* Associated postethmoid part of a skull roof in ventral and dorsal views (IVPP V 15085-1, 2), with its neurocranium attached (in dorsal view, IVPP V 15085-2).

*Other material.* Three left paranuchal plates (in internal and external views; IVPP V 15086.1-3); thoracic material referred to this taxon is unknown.

*Etymology.* Suggesting the convergence of middle and posterior pitlines in the midline, as well as the fusion of paired central plates into a single unpaired unit.

*Locality and horizon.* From a locality close to the Xitun village in the suburb of Qujing, Yunnan, southeastern China, Xitun Formation, Late Lochkovian, Early Devonian.

*Diagnosis.* Phyllolepid with the occipital portion of neurocranium very elongated and slender; dermal rostral capsule not fused to the postethmoid part of skull roof; postorbital plate contacting the paranuchal plate mesially to infraorbital and main lateral sensory line grooves (at least on the internal side of skull roof); marginal plate not extending mesially to infraorbital and main sensory line grooves in internal view; ornamentation consisting of elongated tubercles, sometimes merging into ridges.

*Description.* The specimen is tectonically deformed by a dextral shear (the right side is posteriorly displaced relative to the left side).

*Neurocranium.* This is only visible in dorsal view, but is poorly preserved in its pre-vagal portion, consisting of a natural mould of the overlying radiating fibres of the dermal plates and sensory line grooves. No trace of an antorbital or a supraorbital process is visible. The anterior postorbital process (a.po.p, Fig. 4a) is barely visible and situated around a crack in the specimen; nevertheless, it seems to be as large as that in non-phyllolepid 'actinolepidoids'. The posterior postorbital process (p.po.p, Figs 4a, 5a), though its outline is hardly distinguishable, is well developed and bifid, as in all 'Dolichothoraci'. The supravagal process (sv.p, Figs 4a, 5a) is poorly developed, and its tip is not preserved. In the otic region, two shallow ridges corresponding to the underlying anterior and posterior semicircular canals are visible (r.csa, r.csp, Figs 4a, 5a). In the mid-plane of the otic region, a quite large longitudinal ridge may be interpreted as the impression of the underlying neural canal surrounding the medulla oblongata (r.nc, Figs 4a, 5a). The most surprising feature of the neurocranium is the postvagal (occipital) region, which is very long and narrow, and thus resembles more that of some Petalichthyida (e.g. Macropetalichthys in Stensiö, 1969, Fig. 21) than that of the Arthrodira or the Placodermi in general. A vascular plexus is visible at the limit between the neurocranium and the overlying dermal bones. The subparanuchal fossa (f.s.PaN, Figs 4a, 5a) is well marked. In the occipital part, the lateral walls of the perichondral bone of neurocranium are poorly preserved on the ventral side of the skull roof but show at least three foramina for the spino-occipital nerves (spio.a-c, Figs 4c, 5c); since the occipital region is very elongated, it is possible that there were more than three pairs of spino-occipital nerves. The posterior wall of the neurocranium, though poorly preserved, shows the two posterior occipital processes (occ.p, Figs 4c, 5c), separated by a shallow embayment. A poorly preserved oblique tube, along the lateral side of the occipital part of the neurocranium, is interpreted as the canal for the lateral dorsal aorta (ald.c, Figs 4a, 5a).

*Skull roof.* The dermal rostral capsule (usually composed of the pineal, rostral and postnasal plates) is missing, as is the case in most 'actinolepidoids', but its position is indicated by an anterior embayment of the preorbital plates, and a slight overlap area on its anterior margin (oa.RC, Figs 4b, 5b).

The postethmoid region of the skull roof is well preserved and is longer than wide (Figs 3, 4b–c, 5b–c). The sensory line system is more conspicuous on the internal side (broad thickenings) than on the external side (very narrow grooves, when visible). The very tiny and dense tuberculated ornamentation on the external side of the skull roof obscures bone sutures.



The preorbital plates (PrO, Figs 3, 4a–c, 5a–c) are broad and very short paired plates. Their anterior edge shows a shallow embayment for the insertion of the dermal rostral capsule. The supraorbital sensory line groove (soc, Figs 3–5) extends back only to the radiation centre of the preorbital plate. The contact between the preorbital and postorbital plates seems to be a butting contact, with no overlap. A low thickening suggests that the centronuchal plate slightly overlaps the preorbital plates. The orbital notch (o.n, Figs 4c, 5c) is shared by the edges of the preorbital and postorbital plates.

The radiation centre of the postorbital plates (PtO, Figs 3, 4a-c, 5b-c) is situated at the junction of the infraorbital (ioc, Figs 3, 4a-c, 5b-c, 6) and central (cc, Figs 3, 4b-c, 5b-c) sensory line grooves. The central sensory line groove does not reach the radiation centre of the centronuchal plate. Mesially to the infraorbital sensory line groove, the postorbital plate shows a long and very slender posterior process contacting the similarly slender anterior process of the paranuchal plate (PaN, Figs 3, 4a-c, 5b-c). The junction of these two processes (along the infraorbital sensory line groove) prevents the centronuchal (CeN, Figs 3, 4b-c, 5b-c, 6) from contacting the marginal plates (M, Figs 3, 4a-c, 5b-c, 6), at least on the internal side of the skull roof. This character is a derived feature shared by the superfamily Kujdanowiaspidoidea (sensu V. Dupret, unpub. Ph.D. thesis, Muséum National d'Histoire Naturelle, 2003; Dupret, Goujet & Mark-Kurik, 2007) and by the derived Phyllolepida (Austrophyllolepis and Phyllolepis; Cowralepis displays a fenestra rather than a real contact between the postorbital and paranuchal plates; see Ritchie, 2005, Fig. 20). The condition in Placolepis (Ritchie, 1984) and in Cobandrahlepis (Young, 2005c) resembles that in the family Actinolepididae and in the Phlyctaenioidei, where the marginal plate separates the postorbital from the paranuchal plates. It is noteworthy that the postorbital plates contact the paranuchal plate in Wuttagoonaspis, but via neither a blade nor a process (Ritchie, 1973).

The marginal plates (M, Figs 3, 4a-c, 5b-c, 6) are large elements, gently convex laterally, and forming the lateral edge of the skull roof, although their anterior and posterior boundaries are rather unclear. The radiation fibres on the internal side of the skull roof clearly show that their radiation centre is situated at the junction between the infraorbital, postmarginal (pmc, Figs 3, 4b-c, 5b-c, 6) and main (lc, Figs 3, 4a-c, 5b-c, 6) sensory line grooves, and that the plate does not extend mesially to the infraorbital and main sensory line grooves. This is a derived character displayed by the family Kujdanowiaspididae (sensu V. Dupret, unpub. Ph.D. thesis, Muséum National d'Histoire Naturelle, 2003 and Dupret, Goujet & Mark-Kurik, 2007, non Berg, 1955, 1958), but unknown in the Phyllolepida. It is noteworthy that the outline of the marginal plate is poorly known in the early Arthrodira Wuttagoonaspis (Ritchie, 1973) and Antarctaspis (White, 1968, non Denison, 1978). The postmarginal sensory line (pmc, Figs 3, 4b–c, 5b–c, 6) groove is very large on the internal side of the plate. The postmarginal plates are clearly absent, as there is no overlap area (nor bone suture) for them on the marginal or on the paranuchal plates.

Since the median unpaired element is huge and the central plates are not individualized, we propose to term this median dermal component of the skull roof the centronuchal plate

Figure 4. Head of *Gavinaspis convergens* gen. et sp. nov. (a–c, holotype). (a) Dorsal view of the neurocranium and impression of skull roof, no. IVPP V 15085-2; (b) dorsal (external) view of

the skull roof, no. IVPP V 15085-1; (c) ventral (internal) view of the skull roof, no. IVPP V 15085-1. All scale bars 1 cm. For key to abbreviations, see Table 1.



Figure 5. Head of *Gavinaspis convergens* gen. et sp. nov. (a–c, holotype). (a) Dorsal view of the neurocranium and impression of skull roof, holotype no. IVPP V 15085-2; (b) dorsal (external)

(CeN, Figs 3, 4b–c, 5b–c), by homology with the pattern displayed by the Phyllolepididae. This plate is much longer than wide and its radiation centre is anteriorly placed (at the level of the first third of the plate length). It contacts the preorbital plates anteriorly, without separating them. A low thickened area around this contact suggests a possible overlap of the preorbital plates by the centronuchal plate, as is the case in the known phyllolepids.

The external side exposes particularly well the posterior end of the middle pitline and the anterior end of the posterior pitline (mpl, ppl, Figs 3a, 4b, 5b); both are located at the radiation centre of the plate. The anterior and posterior ends of the posterior pitlines are not connected (that is, they do not form a continuous groove on the dermal bones), thereby implying a superficial course. The central sensory line groove does not reach onto the radiation centre. A thin, posteromesially directed groove that runs from almost the posterior end of the supraorbital groove toward the radiation centre of the centronuchal plate is interpreted here as an ornamentation artefact, and not as the continuity of the supraorbital sensory line groove.

The paranuchal plates (PaN, Figs 3, 4b-c, 5b-c, 6) are also large elements of the skull roof. Their radiation centre is anteriorly situated and close to the external foramen for the endolymphatic duct (d.end.e, Figs 3a, 4b, 5b), as is the case in the non-phyllolepid Actinolepidoidei; in the Phyllolepididae, the radiation centre of the paranuchal plate is close to the posterior edge of the plate. Nevertheless, it is noteworthy that, as in the Phyllolepididae, the radiation centre is at the level of the junction between the curved main lateral line groove and the posterior pitline, which roughly corresponds to the concave anteromesial margin of the paranuchal plate and to the level of the proximal end of the median oblique process of Young (2005c, Fig. 4A) and Ritchie (2005, Fig. 7F-G) in the Phyllolepididae. The paranuchal plates are widely overlapped mesially by the centronuchal plate as in the Phyllolepididae, although in the latter the posterior part of the paranuchal plate is much shorter. Posteriorly to the radiation centre on the internal side, there is a thick, long and posterolaterally tapering ridge that slightly bends laterally from the endolymphatic duct (d.end, Figs 3b, 4c, 5c, 6) toward the posterior part of the main sensory line groove. This divides the plate into two almost equal parts. At midlength of the paranuchal plate, the oblique ridge divides in two parts: one straight and tapering posterolaterally bears the main sensory line canal; the other one is more laterally directed and does not seem to have any particular function, unless perhaps an attachment area for the levator muscles of the head, as it is situated along the cucullaris fossa (d.mc, Figs 4c, 5c). The posterolateral edge of the paranuchal plate is smooth, corresponding to a sliding neck-joint type of dermal craniothoracic articulation (oa.ADL, Figs 4c, 5c), as is the case in the 'Actinolepidoidei' (and in the Phyllolepida).

Externally, the occipital cross commissure (occ, Figs 3, 4b, 5b, 6) and the posterior end of the posterior pitline (ppl, Figs 3a, 4b, 5b) are in the normal location, at the level of the plate radiation centre. The occipital cross commissure is far in front of the posterior skull roof margin, so it could not have run posteriorly to the skull roof (in a 'nuchal gap' or on an extrascapular element as in some other placoderms), but rather had a transverse course.

view of the skull roof, no. IVPP V 15085-1; (c) ventral (internal) view of the skull roof, no. IVPP V 15085-1. All scale bars 1 cm. For key to abbreviations, see Table 1.



Figure 6. Three left paranuchal plates of *Gavinaspis convergens* gen. et sp. nov. (a) dorsal (external) view of left paranuchal plate, specimen no. IVPP V 15086.1; (b, c) ventral (internal) view of left paranuchal plate, specimen no. IVPP V 15086.2 (b) and specimen no. IVPP V 15086.3 (c). All scale bars 1 cm. For key to abbreviations, see Table 1.



Figure 7. Tuberculated ornamentation of Gavinaspis convergens gen. et sp. nov. (specimen no. IVPP V 15085-1). Scale bar 1 cm.

*Ornamentation.* On the skull this mainly consists of minute elongated tubercles, some of which merge into short ridges reaching a few millimetres in length. The tubercles do not resemble those of the Phyllolepididae (e.g. *Austrophyllolepis in* Long, 1984, Figs 5, 19), since in the Gondwanan forms the tubercles are clearly rounded. Nevertheless, the ridge formation seems to be same in *Gavinaspis* and the Phyllolepididae, except for the fact that, in the latter, ridges are osseous and do not bear dentine-like tissues.

*Restoration.* Computerized assistance, using a distortion algorithm of Adobe<sup>®</sup> Photoshop<sup>®</sup>, was used for the restoration of *Gavinaspis convergens* gen. et sp. nov. diplayed in Figure 3. The midline axis is defined by the position of the supraorbital sensory line grooves on the preorbital plates, the radiation centre of the centronuchal plate and the

endolymphatic ducts on the paranuchal plates. A transverse axis is indicated by the endolymphatic ducts (ventral view). These axes intersect at an angle of  $85^{\circ}$ , digitally restored to  $90^{\circ}$  to remove the oblique distortion. An inverse  $5^{\circ}$  distortion was made for the ventral view.

### 5. Discussion

### 5.a. The endolymphatic duct in the Phyllolepida

The endolymphatic duct is a tubular structure that connects the labyrinth to the external environment. It opens externally by a foramen, and internally into the saccula of the inner ear by a proximal foramen. The endolymphatic duct is long and oblique in the

Arthrodira (except for the Phyllolepididae), as in Gavinaspis. An endolymphatic duct is unknown in the Phyllolepididae (except possibly for Cowralepis; see below), where an external foramen on the paranuchal plate has never been found. Consequently, some characters referring to these structures in the previous phylogenetic analyses were coded as not applicable (Dupret, 2004; Dupret, Goujet & Mark-Kurik, 2007). Optimization of these 'missing data' by the software supported a close relationship between the Phyllolepida and the Phlyctaenioidei (Dupret, 2004; Dupret, Goujet & Mark-Kurik, 2007), but Young (2005b) considered this evidence of relationship as artificial (along with other character discussions mentioned below). Nevertheless, the discovery in Cobandrahlepis (Young, 2005c, Figs 3A-C, 4A-B, m.pr) and in Cowralepis (Ritchie, 2005, Figs 7F-G, mp.PNu) of an anterior oblique internal mesial process on the paranuchal plate could be interpreted as a vestigial endolymphatic duct in the Phyllolepida. A very short 'craniospinal process ridge' on the internal side of a paranuchal plate of Austrophyllolepis youngi was also described (Long, 1984, Fig. 19A). Ritchie (2005) pointed out the lack of an external foramen for the endolymphatic duct on the paranuchal plate in Cowralepis, and suggested that the opening might have been situated in the gap between the paranuchal and centronuchal plate. However, the actual absence of an endolymphatic external pore in all other phyllolepidids suggests that its function was progressively lost, with a vestigial stage of a blind endolymphatic duct possibly present in Cobandrahlepis and Austrophyllolepis.

### 5.b. Which definition for the central plates?

The central plates are conspicuous in most placoderms except for the Antiarcha. As for the Phyllolepida, there is no consensus about the absence of these plates, whether they fused to the nuchal plate, or are completely lost and replaced by the nuchal plate. A third hypothesis, suggested here by Gavinaspis, has never before been proposed: the central plates are fused in a single element and the nuchal plate is lost. In order to attempt an explanation, it is necessary to consider the definition and homologies of the central plate. The central plates may be identified on the basis of three major criteria: the position of the plate, the presence of a central sensory line groove/canal, and the presence of the middle and posterior pitlines. (1) The topographic *position* of the plate on the skull roof may be used but though convenient, it is not accurate enough. (2) The presence of a central sensory line groove/canal is applicable in most cases but some taxa show a more or less long central sensory line groove that does not reach the central plate (e.g. the 'Phlyctaenii' Dicksonosteus arcticus with a very short groove, see Goujet, 1984; in the Antarctaspidae with a rather elongated groove as in Toombalepis tuberculata Young & Goujet, 2003,

or Yujiangolepis liujingensis, Wang, Pan & Wang, 1998 (see also Young & Goujet, 2003), Antarctaspis mcmurdoensis White, 1968 (non Denison, 1978); this condition has led Denison to believe in the absence of central plates in Antarctaspis mcmurdoensis). In the Macropetalichthyidae Lunaspis broilii Gross, 1937 and the Quasipetalichthyidae Eurycaraspis incilis Liu, 1991, this groove/canal is simply absent. (3) The presence of the middle and posterior pitlines (when visible) is the most accurate character that can be used to identify the central plates, at least in the Arthrodira. Indeed, in closely related groups like the Macropetalichthyidae, the posterior pitline can be located on the anterior paranuchal plate (e.g. Lunaspis broilii in Gross, 1937). It is also noteworthy that some Quasipetalichthyidae display a second posterior pitline on the central plate (e.g. Eurycaraspis incilis Liu, 1991). Nevertheless, in the known Antarctaspididae (basal Arthrodira), the anterior ends of the posterior pitlines are located around the radiation centre of the nuchal plate (see Young & Goujet, 2003). In the latter case, only the topographic position of the central plate is used as a criterion. In other words, we face a circular argument situation, even though we consider that the most accurate definition refers to the position of the middle and posterior pitlines.

The hypothesis that the central plates have fused into a single median element in the new taxon may be compared with the family Actinolepididae (genera Bollandaspis and Actinolepis; see Schmidt, 1976; Mark-Kurik, 1973, 1985), in which the preorbital plates are fused in a single unit. This may be the case in the Phyllolepida, if we refer to the position of the posterior pitlines, but it is also challenged (yet not refuted) by the antarctaspidid skull roof pattern. Nevertheless, Graham-Smith (1978), based on the numerous abnormal specimens of the antiarch Bothriolepis, concluded that the sensory lines could have become anchored to different combinations of the bone rudiments ('primordium' of Stensiö, 1947) at an early stage of the skeletogenesis, and that during later growth they were consequently drawn along different courses. Such a variable feature may then become fixed following a speciation event, and this scheme is proposed for Antarctaspididae herein: the pitline terminations would have been anchored to the nuchal plate primordium, therefore the adult pattern does not show any association between the pitlines and the central plates.

# 5.c. Hypotheses about processes leading to an unpaired centronuchal plate pattern (Fig. 8)

Here we consider the hypothetical transformation of the skull roof pattern from an actinolepid ancestor (*Kujdanowiaspis podolica* Brotzen, 1934) to a phyllolepid pattern (*Placolepis budawangensis* Ritchie, 1984).



Figure 8. Four hypothetical morphological transformations from a classical actinolepid condition (*Kujdanowiaspis podolica*) to a basal phyllolepid condition (*Placolepis budawangensis*), depending on the initial homology assumption about the centronuchal plate (external view). (a) The most likely hypothesis (plain line), where the process involves a fusion of the paired central plates and the loss of the nuchal plate. The other three hypotheses (dashed lines) are: (b) the process involves the loss of the central plates; (c) the process involves the fusion of central and nuchal plates; (d) the so-called postnasal plate of phyllolepids would be homologous with the central plate of other arthrodires (Young, 2005c). Not to scale. For key to abbreviations, see Table 1.

In the hypothesis favoured here (Fig. 8a), the central plates fuse altogether and the nuchal plate is lost. Ideally, the posterior part of the centronuchal plate would become rather narrow or perhaps not form the posterior edge of the skull roof, the paranuchal plates would become the most important components of the posterior part of the skull roof, and the centre of radiation of the unique central plate would be anteriorly placed as are the ends of the middle and posterior pitlines. This almost perfectly fits the pattern displayed by *Gavinaspis*.

If we consider a loss of the central plates (Fig. 8b), all the peri-central plates would become larger (not only the nuchal plate), the centre of radiation of the nuchal plate should be at the geometric centre of the plate, the posterior part of the nuchal plate is not narrowed, and the middle and posterior pitlines may be lost (lacking anchoring points, if not attached to the nuchal primordium). This does not fit the pattern of *Gavinaspis convergens* gen. et sp. nov.

If we consider the fusion of the central and nuchal plates (Fig. 8c), the posterior part of the centronuchal plate is not narrowed, and the radiation centre coincides with the geometric centre of the plate. This again does not fit the pattern displayed by *Gavinaspis convergens* gen. et sp. nov.

Recently, Young proposed a possible fourth homology, according to which the phyllolepid so-called 'postnasal plate' could be homologous to the arthrodiran central plate (Fig. 8d), 'because interpretation as the central plate of other placoderms seems equally likely using the two criteria of relationship to adjacent bones and possession of sensory grooves' (Young, 2005*c*, p. 262). This hypothesis, though alluring, is not confirmed by the pattern in *Gavinaspis convergens* gen. et sp. nov.

#### 5.d. Remarks concerning the ornamentation

There are different hypotheses of homology about this character, all leading to interesting discussions. Before the description of Wuttagoonaspis (Ritchie, 1973), its ridged plates were attributed to phyllolepid remains (e.g. Rade, 1964, on the authority of Prof. Tør Orvig). Despite the presence of ridges in *Wuttagoonaspis*, Ritchie considered this genus only distantly related to the Phyllolepida. Dupret (2004) regarded the ridged ornament as a non-homologous character; the alternative hypothesis (e.g. Long, 1984; Miles, 1971; Young, 1980; Young & Goujet, 2003) is that the ridged ornamentation is one of the synapomorphies shared by *Wuttagoonaspis* and the Phyllolepida. However, the Phyllolepida and Wuttagoonaspis are not the only placoderms that show a ridged pattern. In many groups of Arthrodira or related forms, at least one genus displays this kind of ornamentation (possibly together with tubercles): Actinolepis



Figure 9. Phylogenetic relationships among Arthrodira, and classification of the Phyllolepida.

(Mark-Kurik, 1973, 1985) and Baringaspis (Miles, 1973) for the non-phyllolepid 'Actinolepidoidei', Diadsomaspis (Gross, 1937) for the phlyctaeniids, Holonema (e.g. Newberry, 1889) for the Brachythoraci, and Lunaspis for the Petalichthyida (Gross, 1961). It is also noteworthy that the tissular structures are different in the selected groups; in some Phyllolepida, the ridges are osseous and lack dentinous tissue, whereas, as in the forms cited above, the ridges are composed by bone and semi-dentine like any tubercle. Nevertheless, the histological structure of the tubercles of Austrophyllolepis and Placolepis remains unknown (see Long, 1984, Fig. 19B; Ritchie, 1984, Figs 6B, 7A, C, E-G, K, M). Moreover, the dermal ridges in *Wuttagoonaspis* display a different shape than those of phyllolepids: in the former the ridges are thicker and display small nipple-like structures all along.

### 5.e. Phylogenetic analysis

The ingroup is composed of 38 taxa, among which are the best known actinolepids (nineteen species), some phlyctaeniids (seven species), brachythoracids (three species), basal arthrodires (*Wuttagoonaspis fletcheri* and *Antarctaspis mcmurdoensis*) and phyllolepids (five species, including *Gavinaspis convergens* gen. et sp. nov. and *Cowralepis mclachlani* Ritchie, 2005). The outgroup is composed of the petalichthyids *Lunaspis broilii* Gross, 1937 and *Eurycaraspis incilis* Liu, 1991. The complete list of the 63 characters is given in Appendix 1 (character state codings do not indicate any *a priori* primitive or derived condition); these are scored for the 38 taxa in the data matrix in Appendix 2.

In order better to accommodate the various hypotheses about the centronuchal plate homologies, three different coding strategies have been attempted. (1) The centronuchal plate consists of the fused central plates and the nuchal plate is lost; in this coding, every character referring to the nuchal plate is coded as not applicable for phyllolepids and *Gavinaspis*; there are thus 63 characters in the matrix. (2) The centronuchal plate consists of the nuchal plate and the central plates are lost; in this coding, every character referring to the nuchal plate and the central plates are lost; in this coding, every character referring to the central plates is coded as not applicable for phyllolepids and *Gavinaspis*; there are thus 62 characters in the matrix. (3) The centronuchal plate consists of the fused central and nuchal plates; there are thus 62 characters in the matrix.

Each data matrix was treated with Nexus Data Editor 0.5.0 (Page, 2001), and the analysis performed by P.A.U.P. 4.0.b10 (Swofford, 1989–1997). The heuristic search logarithm was used, because of the large number of taxa. All characters were unordered and unpolarized *a priori*, and the trees were rooted with the two petalichthyid taxa as outgroup. Wagner optimization was used because it accepts both convergences and reversions. The optimization of missing data was carried out using ACCTRAN (favouring reversions).

The resulting trees are different depending on the chosen coding for the centronuchal plate (coding 1: n = 37, L = 152, CI = 0.4211, RI = 0.7381; coding 2: n = 38, L = 149, CI = 0.4228, RI = 0.7346; coding 3: n = 37, L = 152, CI = 0.4145, RI = 0.7351). It is nevertheless noteworthy that the three strict consensus trees obtained have the same topology. In this respect, the results discussed below only refer to the strict consensus tree obtained with the first coding (Fig. 9; L = 154, CI = 0.4156, RI = 0.723).

In the strict consensus tree, the 'Actinolepidoidei' are paraphyletic (nodes 1 to 22) but their internal



Figure 10. (a–i) The nine possible intra-Phyllolepida topologies encountered among the 37 equally parsimonious trees obtained from the entire data matrix used here. (j) Ritchie's (2005) phylogeny.

relationships are different from those published earlier (Dupret, 2004; Dupret, Goujet & Mark-Kurik, 2007). The Arthrodira are monophyletic (node 1). Antarctaspis and Wuttagoonaspis are still the successive most inclusive arthrodire taxa (nodes 1 to 2). The genus Aethaspis appears here as monophyletic (node 4). The family Actinolepididae (node 19) is still phylogenetically independent of other actinolepids (closer to Bryantolepis and the Phlyctaenioidei than to other actinolepids), and Aleosteus is the sister group of Simblaspis in this new scheme (node 8). The Phlyctaenioidei are a monophyletic group (node 23), but the 'Phlyctaenii' appear paraphyletic, because the family Phlyctaeniidae (Phlyctaenius acadicus and Pageauaspis russelli) shares many symplesiomorphies with the more basal actinolepids (nodes 23 to 25).

The Phyllolepida are monophyletic (node 16), and in this new consensus tree they are not closely related to the Phlyctaenioidei. It is noteworthy that the deletion of both *Gavinaspis* and the family Phlyctaeniidae yields a close relationship between

the Phyllolepida and the Phlyctaenioidei, similar to that proposed by Dupret (2004). Except for the sister-group relationship between Gavinaspis and the family Phyllolepididae, the internal relationships of the Phyllolepididae are not resolved (polytomy node 17); this might be due to the large number of characters involved in this analysis. Among the 37 equiparsimonious trees, nine intra-Phyllolepida topologies are encountered (Fig. 10a-i), but none of them corresponds to Ritchie's phylogeny hypothesis for the group (Ritchie, 2005, p. 225, Fig. 20B; Fig. 10j). Ritchie considered that there was a gradual cline between the basal *Placolepis* and the more derived *Phyllolepis*, and that the intermediate forms of this cline (Cowralepis and Austrophyllolepis) illustrate a progressive modification of the paranuchal plate (and subsequently the centronuchal plate) shape and/or of the main lateral sensory line groove, inducing a progressive loss of contact between the centronuchal and marginal plates (primitive feature displayed by *Placolepis*) that is replaced by a contact between the



Figure 11. Summary of known and supposed stratigraphic ranges (Lochkovian–Famennian) for the Wuttagoonaspida and Phyllolepida (Arthrodira) of South China (S.CH), Gondwana and Euramerica (for Gondwana: Amad. – Amadeus Basin; Ant. – Antarctica; Georg. – Georgina Basin; N.S.W. – New South Wales; SE. Aust. – southeastern Australia; Turk. – Turkey; Venez. – Venezuela; Vict. – Victoria. For Euramerica: Balt. – Baltic States; Belg. – Belgium; Green. – Greenland; N. Am. – Northern America (Pennsylvania, USA); Scot. – Scotland (UK). Supposed stratigraphic ranges indicated by dashed lines with a question mark. Australian stratigraphic range after Young, 2005*a*,*b*; Young & Long, 2005) (updated after Young, 1993, 1999). Conodont zonation from Zhu, Wang & Wang, 2000 (Lochkovian) and Talent *et al.* 2000 (Pragian–Famennian). Lochkovian macrovertebrate assemblages (MAV I–III, left sided of the column) after Zhu, Wang & Wang, 2000; Pragian–Famennian macrovertebrate assemblages (MAV2–15, right side of the column), miospore (GH, GF, VCo, LN, LV) and conodont zone is approximate (modified after Young, 1996; Young & Turner, 2000; after Young, 2005*a*,*b*; Young & Long, 2005).

postorbital and paranuchal plates (*Austrophyllolepis*) and *Phyllolepis*). Even though our topologies do not mirror this cline, we agree with Ritchie's hypothesis, and we suggest that a phyllolepid data matrix at the species level should be attempted (thus avoiding homoplasic interferences from other taxa). Nonetheless, the clear sister-group relationship between *Gavinaspis* and the Phyllolepididae, as well as the peculiar skull roof pattern of the former, support the erection of a new distinct, but monogeneric, family for *Gavinaspis*: the Gavinaspididae fam. nov.

## 5.f. Palaeobiogeographic and palaeogeographic implications

It is commonly considered that the Devonian palaeoglobe was divided into two major landmasses by the Rheic and Palaeo-Tethys oceans, with a northern Laurussian and a southern Gondwanan landmass group. One of the exceptions consists of the main body of the China palaeocontinent, belonging to neither Gondwana, nor Laurussia, and hence being termed the Pan-Cathaysian landmass group (Zhu & Zhao, 2006). As a consequence, a high level of vertebrate endemism is observed in South China during Early Devonian times. In the Early Devonian global reconstructions of Scotese (1997), the Gondwana and Pan-Cathaysian landmass groups are placed well apart from each other.

Since the hitherto most ancient known Phyllolepida have been encountered in eastern Gondwana (*Placolepis harajica* Young, 2005*b* from the Early Givetian of Amadeus Basin, central Australia; *Placolepis tingeyi* Young & Long, 2005 and *Austrophyllolepis quiltyi* Young & Long, 2005 from the Early Givetian of the Aztec siltstones, southern Victoria, Antarctica; see Figs 11, 12), a Gondwanan origin for the Phyllolepida was proposed (Young, 2005*a*). Since the new genus *Gavinaspis* is the oldest non-Gondwanan representative of the Phyllolepida (see phylogenetic analysis), and the Pan-Cathaysian landmass is isolated from the



Figure 12. Global distribution pattern for the Phyllolepida and Wuttagoonaspida, plotted on a Devonian palaeogeographic reconstruction (after Zhu & Zhao, 2006). Gavinaspis is indicated by a black star; the Wuttagoonaspis assemblages by white stars; Givetian-Frasnian Phyllolepida by closed circles; Famennian Phyllolepida (genus Phyllolepis) by open circles. Chinese Lochkovian locality: 1 - Qujing (Yunnan, South China); Gondwanan Givetian-Frasnian localities: 2 - Townsville area (Queensland, Australia: Young, 2005b); 3 - Carnarvon Basin (Western Australia: J. A. Long, unpub. data, in Young, 2005b); 4 - Amadeus Basin and 5 - Georgina Basin (central Australia: Young, 2005a,b; Young & Goujet, 2003); 6 – West-central New South Wales (Australia: Hills, 1931, 1936; Ritchie, 1973; Young, 1993, 1999); 7 - Braidwood-Pambula-Mount Howitt (south east Australia: Long, 1984; Ritchie, 1984; Young, 1983); 8 -Transantarctic Mountains, southern Victoria land (Antarctica: Young & Long, 2005); 9 - Upper Antalya Nappe (Western Lycian Taurus, Turkey); 10 - Sierra de Perija (Venezuela: Young & Moody, 2002; Young, Moody & Casas, 2000); Euramerican Famennian localities: 11 - Virginia-Pennsylvania (USA: Daeschler, Frumes & Mullison, 2003; Lane & Cuffey, 2005); 12 - Scotland (UK: Agassiz, 1844; Woodward, 1915); 13 - Wallonia (Belgium: Leriche, 1931; Lohest, 1888); 14 -East Greenland (Heintz, 1930; Stensiö, 1934, 1939); 15 - Baltic States (Vasiliauskas, 1963); 16 - Timan (Russia: Esin et al. 2000). ANT - Antarctica; AR - Arabia; ARM - Armorica; AU - Australia; BAL - Baltica; G - Greenland; IN - India; KAZ -Kazakhstan; N AM – North America; NC – North China Block; SC - South China Block; SOUTH AM - South America; TAR - Tarim.

Gondwanan one (before the Emsian), this hypothesis seems untenable. This also challenges the hypothesis of a more or less large ocean between the South China block and the northern Gondwanan margin proposed in most palaeogeographic reconstructions (e.g. Cocks & Torsvik, 2002; Torsvik & Cocks, 2004), at any rate from the Givetian (Middle Devonian), as far as Phyllolepida are concerned. When studying other early vertebrate groups (e.g. Sarcopterygii), a shallow marine connection between South China and northeastern Gondwana may have occurred as late as the Pragian– Emsian boundary (E'Em bioevent). Indeed, before the E'Em bioevent, the Placodermi and other early vertebrate taxa (e.g. Galeaspida) show a well-marked endemism in South China. Most of them became extinct during and after this episode, most probably because of the arrival of, and the subsequent competition with, eastern Gondwanan forms (Zhu, 2000). This event is proposed for dating the southward migration of Phyllolepida into eastern Gondwanan margins.

Although the Wuttagoonaspida and Phyllolepida are not closely phylogenetically related, it is remarkable that the most ancient '*Wuttagoonaspis* assemblage' occurrence in Australia is dated as end-Pragian or Early Emsian (Fig. 11; Young, 2005b, Fig. 4; Young, 2005a, Fig. 5). Nevertheless, a recent fieldtrip in Zhaotong (Pragian, Yunnan, South China) yielded an incomplete skull roof that could belong to the Wuttagoonaspida; this specimen is presently under study. In other words, both the Phyllolepida and Wuttagoonaspida could have originated from South China, and would have invaded Gondwana during the E'Em bioevent.

If one can consider that the Chinese origin for the Phyllolepida is settled, the dispersal process of this group is more problematic. It can nevertheless be divided into four steps.

- (1) Invasion into Gondwana took place during the E'Em event (without evidence of a later return into China since this group is still unknown in younger Chinese strata; the latter remark can be explained by biological considerations, *Gavinaspis* being a rather bigger and non-flattened organism than Gondwanan phyllolepids and hence being probably more nektonic than sub-benthic, a 'come-back' to China was perhaps impossible for the latter).
- (2) Eastward (to Venezuela, Fig. 12) and westward (to Turkey, Fig. 12) dispersal occurred in Gondwana until the Late Frasnian. It is also possible, though without fossil evidence, that the Turkish forms dispersed more westwardly to South America (Fig. 12).
- (3) An invasion northward into Laurussia/Euramerica occurred during the Frasnian-Famennian boundary. The only hitherto known Euramerican Phyllolepida are dated as Famennian, and all belong to the genus Phyllolepis Agassiz, 1844. It is noteworthy that not only the Phyllolepida invaded Euramerica at the end of the Frasnian or the beginning of the Famennian. A northward invasion by the Groenlandaspididae (Placodermi, 'Phlyctaenii'), the Megalichthyidae and the Rhizodontida (Sarcopterygii) is coeval with a southward dispersal of the genera Asterolepis (Placodermi, Antiarcha) and Holoptychius (Sarcopterygii) (Dupret, Clément & Janvier, 2005). The discovery of similar Frasnian vertebrate faunas in Turkey (Western Lycian Taurus: Janvier, 1983; Janvier, Clément & Cloutier, 2007), in Colombia (Cuche Formation, Department of Boyacá: Janvier & Villarroel, 2000)

and in Venezuela (Sierra de Perijá: Young & Moody, 2002; Young, Moody & Casas, 2000) sheds light on two possible dispersal routes. The main question is then which dispersal route may have been followed for this invasion, that is, whether it is through a Middle Eastern or a South American route. Comparisons between the phylogenies of the different vertebrate groups involved should be considered in order to supply an answer, although it is noteworthy that only the Colombian fauna includes some Euramerican forms (Asterolepis and Holoptychius); moreover, the Turkish groenlandaspidids may be endemics and would not have led to any descent outside this area (Dupret, Clément & Janvier, 2005). Consequently, a South American route is preferred here. A last, but unlikely, possibility would be a circum-Rheic Ocean migration between Turkey and South America slightly before and during the Frasnian-Famennian faunal interchange.

(4) The Famennian phyllolepid invasion into Euramerica is confusing (Figs 11, 12). Indeed, it is noteworthy that the earliest Euramerican phyllolepids are encountered in the Baltic States (close to the palaeoequator), whereas the latest are encountered in Pennsylvania (close to the southern palaeotropic). Paradoxically, the supposedly most primitive forms (morphologically close to *Placolepis* and *Austrophyllolepis*) are encountered in Belgium (Young, 2005*a*, p. 207) and North America (Lane & Cuffey, 2005).

### 6. Conclusions and summary

The new form Gavinaspis convergens gen. et sp. nov. demonstrates the systematic and palaeogeographic origins of the suborder Phyllolepida. It is dated from the Late Lochkovian of Yunnan (South China) and provides some interesting anatomical characters, intermediate between a classical actinolepid arthrodire and a more derived phyllolepid. Its peculiar centronuchal plate permits the suggestion of a new hypothesis concerning the formation of this dermal element, that is, the fusion of the central plates into a single element along with with the loss of the nuchal plate. Its phylogenetic relationship with other phyllolepids leads to the erection of the new family Gavinaspididae, the sister family to the Gondwanan and Euramerican Phyllolepididae. Its early age suggests a possible South Chinese origin for the Phyllolepida, rather than the Gondwanan origin previously proposed. The invasion of Gondwana by the Phyllolepida is likely to have occurred at the end of the Pragian, during the E'Em bioevent, together with other early vertebrate migrations.

Further investigations of the Late Frasnian vertebrate faunas and the Euramerican *Phyllolepis* species records are needed, in order to elaborate more precise dispersal processes. Acknowledgements. We thank Zhao Wenjin and Jia Liantao (IVPP, P.R. China) for the help in the field and with photography, and Xiong Cuihua for the preparation. Edouard Poty (Université de Liège, Belgium) permitted V. D. to access the Famennian phyllolepid collections. Wang Junqing (IVPP) added interesting remarks concerning the Early Devonian Arthrodires of Yunnan. John Long (Museum Victoria, Australia) and Gavin Young (Australian National University, Canberra ACT, Australia) contributed motivating discussions concerning Phyllolepida and basal Arthrodira during the IPC 2006 in Beijing. Two anonymous reviewers greatly enhanced the quality of the manuscript. This work was supported by the Major Basic Research Projects (2006CB806400) of MST of China, the Chinese Foundation of Natural Sciences (40332017 and 40602005), and UNESCO-IGCP 491. The authors thank the two anonymous reviewers for their constructive remarks and corrections.

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### Appendix 1. List of characters

- 1. Link between the two neurocranial components (ethmoid and postethmoid parts):
  - 0. no link ('loose nose' fishes)

1. fusion (by means of either osseous trabecles, or complete fusion)

2. Position of the foramen for the hyomandibular branch of the facial nerve (fVIIHm) in relation to the anterior postorbital process:

0. foramen in the distal part of the anterior postorbital process

1. foramen in a proximal and posterior position to the process

- 3. Neurocranial supraorbital process:
  - 0. absent
  - 1. present
- 4. Neurocranial basal process:
  - 0. absent
  - 1. present

- 5. Rostral, pineal or rostropineal plates:
  - 0. absent 1. present
- 6. Pineal or rostropineal plate separates the preorbital plates:0. no
- yes
   Rostral and pineal plates fused into a single rostropineal component:
  - 0. no
  - 1. yes
- 8. Preorbital plates show an embayment for the insertion of the pineal or the rostropineal plate:
  - 0. no, or very shallow
  - 1. yes, very deep
- 9. Pineal or rostropineal plate fused to the skull roof: 0. no
  - 1. yes
- 10. Postnasal plates fused to the preorbital plates:
  - 0. yes
  - 1. no
- 11. Position of the orbits in the skull roof:
  - 0. dorsal
- lateral
   Preorbital plates:
  - 0. separate
  - 1. fused
- 13. External morphology of the sensory line system:
  - 0. canals with external pores
  - 1. grooves
- 14. Supraorbital sensory lines:
  - 0. separate
  - 1. meet posteriorly
- 15. Infraorbital and main sensory lines grooves run along the mesial margin of the marginal plate:
  - 0. no
  - 1. yes
- 16. Central plates:
  - 0. fused into a centronuchal plate
  - 1. individualized paired elements
- 17. Shape of suture between central plates:
  - 0. straight
  - 1. sinuous
- Pineal (or rostropineal) plate contacts the central plates:
   no
  - 1. yes
- 19. Posterior edge of the preorbital plates indents the anterior edge of the central plates:
  - 0. no
  - 1. yes
- 20. Contact between central and preorbital plates:
  - 0. yes
  - 1. no
- 21. Contact between central and marginal plates:
  - 0. yes
  - 1. no
- 22. Contact between postorbital and paranuchal plates: 0. yes
  - 1. no
- 23. Nuchal plate:
  - 0. absent
  - 1. present
- 24. Nuchal plate separates the central plates:
  - 0. no 1. yes
- 25. Contact between orbits and central plates:
  - 0. no
  - 1. yes

26. Preorbital plates are part of the orbital margin: 0. yes

1. no

- 27. Central sensory line groove extends from the postorbital plate to the radiation centre of the central plates: 0. no
  - 1. yes
- 28 Posterior pitline present on both central and paranuchal plates:
  - 0. anterior and posterior ends clearly connected
  - 1. anterior and posterior ends not connected, implying a superficial course
- 29. Central sensory line groove: 0. absent
  - 1. present
- 30. Postmarginal plate:
  - 0. absent
  - 1. present
- 31. Morphology of the anterior external nuchal central plates contact suture:
  - 0. nuchal plate indents the central plates
  - 1. straight suture
- 32. Contact between the pineal and nuchal plates: 0. no
  - 1. yes
- 33. Number of paranuchal plates pairs:0. one pair
  - 1. two pairs
- 34. Occipital cross commissure:
  - 0. on both nuchal and paranuchal plates 1. only on paranuchal plates
- 35. Posterolateral edge of the paranuchal plates: 0. convex
  - 1. concave
- 36. Posterior process of the paranuchal plate behind the nuchal plate (external side):
  - 0. absent 1. present
- 37. Position of the external foramen for the endolymphatic duct:

0. well anteriorly to the posterior edge of the paranuchal plate, or on the anterior paranuchal plate for the Petalichthyida

- 1. near to the posterior edge of the paranuchal plate
- 38. Type of exoskeletal dermal craniothoracic articulation: 0. actinolepid 'sliding neck joint'
  - 1. ginglymoid phlyctaenioid type
  - 2. 'spoon-like' petalichthyid type
- 39. Dermal articular condyle of anterior dorsolateral plates:0. close together
  - 1. well apart
- 40. Ventral keel on the internal side of the median dorsal plate:
  - 0. absent
  - 1. present
- 41. Unornamented (overlapped) area on the anterior margin of the median dorsal plate:
  - 0. absent
  - 1. present
- 42. The unornamented zone on the anterior margin of the median dorsal plate is:
  - 0. simple
  - 1. double
- 43. Extrascapular plate:
  - 0. absent
  - 1. present

- 44. Dorsolateral groove (for an accessory sensory line) on the anterior dorsolateral plate:
  - 0. absent
  - 1. present
- 45. Posterolateral plate:
  - 0. absent
  - 1. present
- 46. Pectoral notch of the anterior ventrolateral plate: 0. shallow
  - 1. deep
- 47. Prepectoral process of the anterior ventrolateral plate: 0. short
  - 1. long
- 48. Anteroventral plates:
  - 0. absent
  - 1. present
- 49. Anterolateral and anterior ventrolateral plates connected behind the pectoral fenestra:
  - 0. no
- yes
   Anterior median ventral plate:
- 0. absent
  - 1. present
- 51 Posterior median ventral plate:
  - 0. absent
  - 1. present
- 52. Mutual overlap of posterior ventrolateral plates: 0. simple overlapping
  - 1. sinusoidal/double overlapping
- 53. Spinelets on the mesial side of spinal plate: 0. absent
  - 1. present
- 54. Postmedian dorsal plates:
  - 0. absent
  - 1. present
- 55. Width/length ratio of the preorbital plates:
  - 0. (W/L) > 0.51. (W/L) < 0.5
- 56. Central plates length ratio to the skull roof length (from the anterior edge of preorbital plates to posteriormost edge of the skull roof):
  0. (LC/LSR) < 45%</li>
  - 0. (LC/LSR) < 45% $1. (LC/LSR) \ge 45\%$
- 57. Length/width ratio of the nuchal plate:
  - $0. (L/W) \le 1.5$
  - 1.(L/W) > 1.5
- 58. Length/width ratio of the median dorsal plate:
  - 0. (L/W) < 1.5
  - $1.(L/W) \ge 1.5$
- 59. Length/height ratio of the anterior dorsolateral plate:
  - 0. (L/H) < 1
  - 1. (L/H)  $\ge 1$
- 60. Length/height ratio of the posterior dorsolateral plate:
  - 0. (L/H) < 2
  - 1.  $(L/H) \ge 2$
- 61. Angle between interolateral and spinal plates: 0. angle  $< 110^{\circ}$ 
  - 1. angle  $\geq 110^{\circ}$
- 62. (LSp beard by AVL/LSp) ratio = RSp:
  - 0. RSp < 60%
  - 1.  $RSp \ge 60\%$
- 63. Length/width ratio of posterior ventrolateral plates:
  - 0. (L/W) < 1.5
  - 1.  $(L/W) \ge 1.5$

## Appendix 2. Data matrix

Taxa	a 2	3	4	5	6	7	8	0	10	11	12	13	14
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### Appendix 2. Continued.

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## Appendix 2. Continued.

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