Appendages of an early Cambrian metadoxidid trilobite from Yunnan, SW China support mandibulate affinities of trilobites and artiopods

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Abstract – Appendage anatomy contributes crucial data for understanding the evolution and ecology of Euarthropoda. The Palaeozoic trilobites show a great diversity of exoskeletons in the fossil record. However, soft parts, especially appendages, have only been discovered from a few trilobite species. Here we report extraordinarily preserved appendages in the trilobite species Hongshiyuanaspis yiliangensis Zhang & Lin in Zhang et al. 1980 (Redlichiida, Metadoxididae) from a single mudstone layer of the Xiazhuan fossil assemblage within the Hongjingshao Formation (Cambrian Series 2, Stage 3) near Kunming, Yunnan, SW China. The appendages exhibit the common architecture revealed by other trilobites and artiopods by consisting of a pair of uniramous antennae followed by a series of paired homonomous biramous limbs. The antennae in holaspid individuals comprise up to 27 spinous podomeres and their ontogeny occurs by lengthening of the podomeres. The post-antennal biramous limbs are similar to those in other polymerid trilobites and artiopods by having a single-segmented protopodite and an endopodite comprising seven segments, but possess a unique wide tripartite exopodite with long setae. Sophisticated appendage anatomy, including the body–limb junction, fine setae, putative muscle bundles and duct-type tissues, are also revealed. Appendages of trilobites, artiopods and other upper stem-group euarthropods are compared and summarized. The H. yiliangensis appendages highlight the high morphological disparity of exopodites and the conservativeness of endopodites in trilobites and artiopods. This morphological pattern, together with similar body patterning seen in crustaceans but not in chelicerates, supports the mandibulate affinities of trilobites and at least some artiopods.

Keywords: arthropods, trilobites, appendages, Mandibulata, Cambrian, soft-part preservation

1. Introduction

With biominerlized exoskeletons of more than 20 000 species discovered (only inferior to ostracods; Zhang, 2013), the trilobites (=Trilobita) are one of the most diverse extinct groups of Euarthropoda (e.g. Budd & Telford, 2009) that inhabited Palaeozoic seas from the Cambrian explosion (Hollingworth, 2008) to the end-Permian mass extinction (Owens, 2003). However, in contrast to the megadiversity of exoskeletons, the soft-bodied anatomy of trilobites is poorly known, with only ~30 species primarily from Konservat-Lagerstätten showing soft-part preservation, especially appendages (Table 1; also see Hughes, 2003).

To date, all the reported appendages of polymerid trilobites consist of a pair of uniramous deutocerebral antennae and a series of homonomous biramous post-antennal limbs corresponding to each body segment (see Hughes, 2003; Scholtz & Edgecombe, 2005, 2006). It should be emphasized that the morphology of the post-antennal limbs of Agnostus pisiformis is distinct from that of polymerid trilobites (Müller & Walossek, 1987), which makes the supposed trilobite affinity of Agnostida problematic (Walossek & Müller, 1990; Fortey, 2001; Hughes, 2003). Although antennae of exactly similar uniramous multi-segmented architecture are preserved in ~20 of these polymerid trilobite species, complete post-antennal limbs have only been reconstructed in six species from five of the nine polymerid orders of Trilobita, including Eoredlanchia intermedia, Olenoides serratus, Triarthrus eatoni, Cryptolithus bellulus, Cerarurus pleurexanthemus and Chotecops ferdinandi (see Table 1 for details; Fortey, 2001; Hughes, 2003, 2007). The known post-antennal limbs of these different polymerid trilobite species share a biramous architecture, with two rami (endopodite and exopodite) connected to a protopodite (e.g. Hughes, 2003). The protopodite consists of a single segment (e.g. Ramsköld & Edgecombe, 1996) and the endopodite is made up of seven segments (e.g. Hughes,
Table 1. Updated summary of trilobites reported with preserved appendages, supplemented and modified from Hughes (2003)

<table>
<thead>
<tr>
<th>Species</th>
<th>Order</th>
<th>Family</th>
<th>an</th>
<th>en</th>
<th>ex</th>
<th>ds</th>
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<th>Deposits</th>
<th>Preservation</th>
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<td>Chengjiang Burgess</td>
<td>Shu et al. 1995; Ramsköld &amp; Edgecombe, 1996; Chen &amp; Zhou, 1997; Hou &amp; Bergström, 1997; Hou et al. 2008</td>
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<td>Chengjiang Burgess</td>
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<td>++</td>
<td>+</td>
<td>+</td>
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<td>Cam. S3</td>
<td>Chengjiang Burgess</td>
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<td><strong>Hongshiyananus yiliangensis</strong></td>
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<td>++</td>
<td>++</td>
<td>++</td>
<td>++</td>
<td>Cam. S3</td>
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<td>+</td>
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<td>Guanshan Burgess</td>
<td>Hu et al. 2013</td>
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<td>Guanshan Burgess</td>
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<td>o</td>
<td>Cam. S4</td>
<td>Eum Bay Shale Burgess</td>
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<td>++</td>
<td>++</td>
<td>++</td>
<td>Cam. S5</td>
<td>Burgess Shale Burgess</td>
<td>Walcott, 1918, 1921; Raymond, 1920; Stormer, 1939; Whittington, 1975, 1980</td>
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<td>+</td>
<td>o</td>
<td>o</td>
<td>Cam. S5</td>
<td>Burgess Shale Burgess</td>
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<td>Ptychoparidae</td>
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<td>o</td>
<td>o</td>
<td>Cam. S5</td>
<td>Burgess Shale Burgess</td>
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<tr>
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<td>Ptychoparidae</td>
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<td>+</td>
<td>+</td>
<td>+</td>
<td>Cam. S5</td>
<td>Burgess Shale Burgess</td>
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<td>Ptychoparidae</td>
<td>++</td>
<td>o</td>
<td>o</td>
<td>o</td>
<td>Cam. S4</td>
<td>Drumian Wheeler Fm. Burgess</td>
<td>Briggs et al. 2008</td>
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<td>Agnostidae</td>
<td>++</td>
<td>++</td>
<td>++</td>
<td>++</td>
<td>Cam. Guzhangia</td>
<td>Orsten Phosphatic Müller &amp; Walsheke, 1987</td>
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<td>Nileidae</td>
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<td>+</td>
<td>o</td>
<td></td>
<td>Ord. Tremadocian</td>
<td>Feozuata Burgess</td>
<td>Van Roy et al. 2010</td>
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<td><strong>Placoparia cambriensis</strong></td>
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<td>Plomerididae</td>
<td>++</td>
<td>o</td>
<td>o</td>
<td>o</td>
<td>Ord. Darrwilian</td>
<td>Lower Llanviri Mudstone moul</td>
<td>Whittington, 1993</td>
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<td><strong>Cryptolithus bellulus</strong></td>
<td>Asaphida</td>
<td>Trinucleidae</td>
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<td>o</td>
<td></td>
<td>Mid.-Upp. Ord.</td>
<td>Beecher-type's beds, Pyritic, calcitic</td>
<td>Walcott, 1881, 1912; Beecher, 1895; Raymond, 1920; Stormer, 1939; Bergström, 1972; Farrell et al. 2009</td>
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<td>o</td>
<td>+</td>
<td>+</td>
<td>o</td>
<td>Cam. S5</td>
<td>Trenton Gr. Calcitic</td>
<td>Walcott, 1918, 1921; Raymond, 1920; Stormer, 1939, 1951</td>
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<td>Cheiruridae</td>
<td>o</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>Upp. Ord.</td>
<td>Trenton Gr. Mudstone imprint</td>
<td>Billings, 1870; Raymond, 1920</td>
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<td>Calymenididae</td>
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<td>+</td>
<td>o</td>
<td></td>
<td>Upp. Ord.</td>
<td>Trenton Gr. Mudstone imprint</td>
<td>Walcott, 1884, 1918; Raymond, 1920</td>
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<tr>
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<td>o</td>
<td>+</td>
<td>o</td>
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<td>Upp. Ord.</td>
<td>Richmond Gr. Mudstone imprint</td>
<td>Walcott, 1884, 1918; Raymond, 1920</td>
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<td>Odontopleuridae</td>
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<td>o</td>
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<td>Upp. Ord.</td>
<td>Clays Ferry Fm. Silicified Ross, 1979</td>
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<td><strong>Rheopis et. anserinus</strong></td>
<td>Phacopida</td>
<td>Acastidae</td>
<td>++</td>
<td>o</td>
<td>o</td>
<td>o</td>
<td>Dev. Emsian</td>
<td>Hunsrück Slate Pyritic</td>
<td>Stormer, 1939; Stürmer &amp; Bergström, 1984; Bergström &amp; Brassel, 1884; Bartels, Briggs &amp; Brassel, 1998; Bruton &amp; Haas, 1999</td>
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</table>

Orders and families are mainly based on Harrington et al. (1959) and Whittington et al. (1997). Note that the former Elrathina cordillerae from the Burgess Shale is revised as an unnamed new species of Elrathina (Geyer & Peel, 2017). Preservation of complete, incomplete and absent anatomical structures are indicated by '+++', '+', and 'o', respectively. Abbreviations: an – antenna; ds – digestive system; en – endopodite; ex – exopodite.
Nevertheless, the exopodites show considerable morphological variations in different trilobite species (Müller & Walossek, 1987, fig. 27; Shu et al. 1995, fig. 21; also see Bruton & Haas, 1999, fig. 22 and Hou et al. 2008, fig. 14 for revised reconstructions for C. ferdinandi and E. intermedia, respectively). Other exceptional appendages of polymerid trilobites include a pair of antennae-like cerci that has only been found in the pygidium of O. serratus (e.g. Whittington, 1975, 1980).

The limited knowledge of the soft anatomy of trilobites and other stem-group euarthropods, especially appendages, has long constrained our understanding of the internal and external phylogenetic relationships of trilobites (see Scholtz & Edgecombe, 2005, 2006 and references therein). It was not until recently that phylogenetic analyses resolved the trilobites within the Artiopoda Hou & Bergström, 1997, as close relatives to several groups of soft-bodied ‘trilobitomorph’ euarthropods from Cambrian Lagerstätten (e.g. Edgecombe & Ramköld, 1999; Ortega-Hernández, Legg & Braddy, 2013; Stein et al. 2013; Legg, Sutton & Edgecombe, 2013). These ‘trilobitomorphs’, including concilitergans (e.g. Kuamaia and Saperion), nектaspids (e.g. Naraotaia, Livia and Emucaris), xandarelids (e.g. Xandarella and Cindarella) and other problematic taxa, share the common appendage architecture composed of a pair of uniramous antennae and homonomous post-antennal biramous limbs, as well as other synapomorphies, with polymerid trilobites (e.g. Hou & Bergström, 1997; Edgecombe & Ramköld, 1999; Zhang, Shu & Erwin, 2007; Ortega-Hernández, Legg & Braddy, 2013). Nevertheless, phylogenetic analyses have not reached a consensus on the sister group of trilobites and cannot determine whether the entire Artiopoda is closer to the Mandibulata or Chelicera at present (e.g. Budd & Telford, 2009; Ortega-Hernández, Legg & Braddy, 2013; Stein et al. 2013; Legg, 2014). Therefore, further studies on trilobite soft anatomy are essential to deliver arguments for answers to these questions.

Here we describe exceptionally preserved appendages of the polymerid trilobite *Hongshiyanaaspis yiliangensis* Zhang & Lin in Zhang et al. 1980 (Redlichiiida, Metadoxidae) from the Xizhaxian fossil assemblage in the Hongjingshao Formation (Cambrian Series 2, Stage 3) near Kunming, Yunnan, SW China. The new material confirms the basic architecture of trilobite/artiopodan appendages, but also exhibits a morphological disparity in trilobite/artiopodan exopodites, providing new information for comparative anatomy and elucidating the affinities of trilobites and other related arthropods.

2. Geological setting

All specimens of *Hongshiyanaaspis yiliangensis* were recovered from a single yellowish structureless claystone layer intercalated within sandstone layers from the lower part of the Hongjingshao Formation, the lower fossil horizon yielding the Xiazhuang fossil assemblage at Xiazhuang, Chenggong, Kunming, eastern Yunnan, SW China (see Zeng et al. 2014 for detailed information on geography and stratigraphy). The soft-part preservation and various angles of burial of the fossils indicate that the fossiliferous layer was deposited rapidly, probably by a storm event (e.g. Zhu et al. 2001; Hu et al. 2010, 2013). Other euarthropods recovered from the same layer only include a large bivalve euarthropod *Jagatacaris*, whose biramous limbs comprise more than 20 endopodite podomes and are readily distinguishable from the trilobite limbs (Zeng et al. 2014). Although other trilobite species including *Yunnanocephalus yunnanensis*, *Malongocephalus yunnanensis* and *Kuanyangia* (*Sapushania* *granulosa* were also found in the upper fossil horizon (Zeng et al. 2014), none of these species were discovered from the *Hongshiyanaaspis* bed. The age of the Xiazhuang fossil assemblage is approximately identical to that of the lower part of the Xiaoshiba Lagerstätte (e.g. Hou et al. 2017) because both fossil assemblages are from the same stratigraphic interval in the lower part of the Hongjingshao Formation and from the same *Eoredlichia*–*Wutingaspis* Assemblage Zone of the regional Qiongzhushian Stage (Cambrian Stage 3). This fossil zone also yields the renowned Chengjiang biota from the underlying Yu’anshan Formation (Zhu et al. 2001). The similar age and faunal compositions suggest that these two fossil assemblages from the Hongjingshao Formation can be regarded as continuing the Chengjiang biota (Zeng et al. 2014). However, the upper part of the Xiaoshiba Lagerstätte extends into the regional Canglangpuan (Cambrian Stage 3) *Yiliangella* Assemblage Zone represented by *Zhangshania typica* (Hou et al. 2017), an interval which is absent in the section that contains the Xiazhuang fossil assemblage (Zeng et al. 2014).

3. Materials and methods

A total of 106 early to fully grown holaspid specimens of the trilobite *Hongshiyanaaspis yiliangensis* were studied (Field IDs prefixed by HBHY; see online Supplementary Material available at http://journals.cambridge.org/geo). The majority of these specimens are dorsoventrally embedded, with only 11.3 % laterally compressed. Nearly half of them (45.3 %) exhibit preserved soft parts, including antennae, biramous limbs and parts of the digestive system. Owing to the different angles of burial, the shapes of original three-dimensional structures can vary, especially for the biramous limbs, but structures in various positions or on different levels can also reveal additional details of the morphology. Similar to soft parts of the Chengjiang fossils, the appendages of *H. yiliangensis* are preserved mainly as Fe-rich aluminosilicate films with limited organic ingredients (Zhu, Babcock & Steiner, 2005).

All figured specimens are housed at the Nanjing Institute of Geology and Palaeontology, Chinese
Appendages of an early Cambrian trilobite

Figure 1. (Colour online) Complete specimens of *Hongshiyanaspis yiliangensis* Zhang & Lin *in* Zhang *et al.* 1980 preserved with appendages, from the Xiazhuang fossil assemblage, Kunming, Yunnan, SW China. (a) Dorsoventrally compressed normal-sized holaspid with complete antennae, part, NIGPAS 164504A. (b) Dorsoventrally compressed fully grown holaspid with complete antennae and a series of post-antennal biramous limbs, part only, NIGPAS 164503. (c, d) Laterally compressed normal-sized holaspid with a series of post-antennal biramous limbs, NIGPAS 164505. (c) Part, NIGPAS 164505A. (d) Counterpart, NIGPAS 164505B. Abbreviations: an – antenna; en – endopodite; ex – exopodite; gut – gut of digestive tract.

Academy of Sciences (prefixed by NIGPAS). Appendages were prepared manually using blades. Photographs showing overall morphology were taken using a Nikon D300s digital camera with a Nikon AF-S VR105mm f/2.8G macro lens. Detailed anatomy was captured using a Carl Zeiss SteREO Discovery V12 microscope linked to an AxioCam HR3 digital microscope CCD camera. Illumination from various directions and angles was employed in order to show the three-dimensional structures. Line drawings were prepared on the basis of high-resolution pictures. Measurements were conducted on photographs within Adobe Photoshop™ CS6 and statistically analysed in Microsoft Office Excel™ 2013.

We follow most of the standard terminology for trilobites in Whittington *et al.* (1997), including the terms ‘antenna(e)’, ‘endopodite(s)’ and ‘exopodite(s)’, which are also the most commonly used terms in recent literature on fossil and extant euarthropods. However, the neutral term ‘protopodite’ is used rather than the term ‘basis’ or ‘basipodite’ (e.g. Boxshall, 2004), which is equal to the term ‘coxa’ or ‘coxopodite’ in earlier studies (e.g. Whittington *et al.* 1997). For the protopodite in post-antennal biramous limbs of trilobites and other arthropods, the term ‘basis’ was first introduced by Ramsköld & Edgecombe (1996). However, this term implies the evolutionary hypothesis that an undivided protopodite is homologous to the basis/basipodite in a multi-segmented protopodite with other more proximal podomeres such as the coxa or precoxa (see Boxshall, 2004 for discussion). The corresponding evolutionary scenario would be that the origin of other non-basal podomeres occurred by addition of proximal podomeres (e.g. Walossek & Müller, 1998; Haug *et al.* 2013), which rejects an alternative by the subdivision of an originally undivided protopodite podomere (see Boxshall, 2004 for discussion).

4. Description of appendages

4.a. General arrangement of appendages

The appendages of a *Hongshiyanaspis yiliangensis* holaspid consist of a single pair of uniramous antennae (an) followed by a series of homonomous post-antennal biramous limbs (Figs 1–3). The preservation of incomplete cephalic biramous limbs (Figs 4c–e, 5c, d, 6a) and three paired digestive glands (gd) on the second and third glabellar lobes and the occipital lobe (Figs 1a, 2a, 4a, b, 5a, b) suggest the presence of three corresponding pairs of post-antennal limbs underneath the cephalon (ce). Each of the 14 thoracic segments (th1–th14) bears a single pair of biramous limbs (Figs 1b–d, 2b, 3, 4e, f, 6–8), as supported by limb fragments connected to the 14th thoracic segment (unfigured fragmentary specimen HBHY008). Fragments of limbs are connected to the first and only axial ring of the pygidium (pg) (unfigured fragmentary specimen HBHY008). It is unknown whether there are limbs corresponding to the terminal axial piece of the pygidium, including the cerci. Variations in shape, if there are any, are insignificant between the post-antennal biramous limbs.
4.b. Antennae

The paired uniramous antennae are slender and flexible (Figs 1a, b, 2a, b). Each is attached to the corresponding side of the hypostome (hyp) (Figs 1a, b, 2, 4c, e, 5d, 6a) and emerge at the anterior rim of the cephalon ventrally as in the possible life position (Figs 1a, b, 2, 4a–c, e, 5a, b, d, 6a). The lengths of complete antennae exceed ~ 50% of the cephalon’s length (Figs 1a, b, 2), and the relative proportion between the antennae and the complete body length decreases from 36% (Figs 1a, 2a) to 29% (Figs 1b, 2b) from the normal-sized to fully grown holaspids periods. Individual antennae may be curved by up to 90° at their middle (Figs 4a, 5a) or terminal sections (Figs 1a, 2a). Both antennae may be stretched apart laterally with an intersection angle of up to 105° (Figs 4a, 5a).

The antennae are composed of up to 27 rectangular podomeres in fully grown holaspids (Figs 1b, 2b), while an approximately similar number of podomeres is also found in normal-sized holaspids (Figs 1a, 2a). Although antennae are also preserved in early holaspids (Figs 4a, 5a), their maximum numbers of podomeres cannot be determined owing to the difficulties in preparation. The most proximal podomeres are evidently stouter than the distal ones (Figs 1a, b, 2,
4a–c, e, 5a, b, d, 6a, 9a). Each podomere bears at least one sharp spine (sp) close to its distal arthrodial membrane (Figs 1b, 2b, 4e, 6b, 9b). The length of the spine may reach one-third of the podomere length (Fig. 9b).

4.c. Post-antennal limbs

Each post-antennal limb is biramous and consists of a protopodite (pt) comprising a single segment, an endopodite (en) consisting of seven segments and a tripartite exopodite (ex) (Figs 1b–d, 2b, 3, 7a, b, 8a, b, 10, 11a, g, h, 12a–d, 13). The shapes of these limbs are homonomous, but their sizes decrease correspondingly to the sizes of the thoracic segments (Figs 1b–d, 2b, 3, 4e, f, 6–8, 10). The thoracic limbs are longer and wider than the thoracic exoskeleton so that they stretch out from below the exoskeleton and are arranged in an imbricate series (Figs 1b–d, 2b, 3, 4e, f, 6–8, 10).

4.c.1. Protopodite and body–limb junction

The protopodite is robust and has a subrectangular outline, carrying the endopodite and the exopodite, respectively, at its dorsal and ventral distal margins (Figs 1b–d, 2b, 3, 10, 11a, g, h, 12a–d, 13). It is connected to the body by an arthrodial membrane (am) as the body–limb junction (Figs 7c, 8c, 11a, b, i, 12a–g, 13). The body–limb arthrodial membrane can be preserved as a section about half the width of the protopodite (Figs 11i, 12a–g, 13), or indicated by subparallel annulations (Figs 10, 11a, b). The attachment site of the protopodite to the body is the lateral side of the...
subrectangular to hourglass-shaped sternite (sn) (Figs 7c, 8c, 11i). Putative muscle bundles (ms) are preserved as reddish Fe-rich films along the boundary between the protopodite and the arthrodial membrane (Figs 12e, g, 13b). The protopodite extends significantly in a ventral direction, forming a stout endite (Fig. 12a–g). At least three clusters of robust spines (Figs 12a, b, e, f, 13), together with rows of non-clustered fine spines (Figs 12c, d, g, 13), develop along the ventral margin of the protopodite’s endite and form a gnathobase (gs). Numerous fine setae (st) of a few hundred micrometres in length develop on the surface of the protopodite (Figs 12e, f, 13). A shallow transverse furrow is also present on the protopodite (Figs 10, 11a, b). It is most likely to be a real anatomical structure rather than a taphonomic imprint because no corresponding structures are seen on the exoskeleton.

4.c.2. Endopodite

The endopodite composed of seven podomeres is attached to the distal margin of the protopodite by an arthrodial membrane (Figs 1b–d, 2b, 3, 4e, f, 5, 6, 7a, b,...
Figure 5. Line drawings of *Hongshiyanspis yiliangensis* Zhang & Lin in Zhang *et al.* 1980 preserved with appendages, from the Xiazhuang fossil assemblage, Kunming, Yunnan, SW China. (a) Counterpart only, NIGPAS 164514, as in Figure 4a. (b) Counterpart only, NIGPAS 164512, as in Figure 4b. (c) Part, NIGPAS 164513, as in Figure 4d. (d) Part, NIGPAS 164510, as in Figure 4c. Abbreviations as in Figures 1–3.
Figure 6. Line drawings of *Hongshiyanaspis yiliangensis* Zhang & Lin in Zhang et al. 1980 preserved with appendages, from the Xiazhuang fossil assemblage, Kunming, Yunnan, SW China. (a) Counterpart, NIGPAS 164506b, as in Figure 4e. (b) Part, NIGPAS 164506a, as in Figure 4f. Grey areas indicate digestive system. Abbreviations as in Figures 1–3.

8a, b, 10, 11a, b, g, 12a–g, 13), which is also indicated by putative oblique muscle bundles (Figs 12c, g, 13b). The total length of the endopodite accounts for ∼ 80\% of that of the post-antennal limb (Figs 1b–d, 2b, 3, 10, 11a, g, h, 12d, 13a). The first six podomeres are subrectangular in outline, and their sizes decrease from proximal to distal (Figs 1b–d, 2b, 3, 4e–f, 5c, d, 6, 7a, b, 8a, b, 10, 11a, g, h). Each of these six podomeres bears an endite (ed) with thin spines (Figs 1c, 3a, 12a–d, g, h, 13). Tiny reddish Fe-rich dots interpreted as the bases of setae are present on these podomeres (Figs 12b–d, g, 13). The terminal, seventh podomere is extremely short but connected to the sixth podomere by an arthrodial membrane (Figs 1c, d, 3, 4e, f, 6a, b, 7a, 8a, 11e–h, 12d, h, 13a). Three highly sclerotized sharp claws (cw), one prominent in the middle and two subordinate lateral ones, are attached to the seventh podomere (Figs 11e, f, 12d, h, 13a).

### 4.c.3. Exopodite

The exopodite is oblong to subrectangular in outline and composed of a tripartite flattened flap (ex1–ex3) with setae (xs) and lamellae (lm) (Figs 1b–d, 2b, 3, 7, 8, 10, 11a–d, g, h, j, 12a–d, 13). The total length of the exopodite, flap and setae included, is subequal to that of the endopodite (Figs 1b–d, 2b, 3, 10, 11a, g, h, 12a, 13b). The flap is in addition ∼ 50\% wider than the sagittal length of the thoracic segment and almost double the maximum width of the endopodite. The exopodite flap is attached to the dorsal margin of the protopodite by an arthrodial membrane (Figs 7c, 8c, 10, 11a, b). Two joints, one transverse and the other oblique, divide the flattened flap into three parts (Figs 1b–d, 2b, 3, 4e, 6a, 7a, b, d, 8a, b, d, 10, 11a, g, h, j, 12a, c, d, 13). The transverse joint runs through about the distal third of the flap, separating the flap into a bell-shaped distal part (the third lobe, ex3) and a trapezoidal proximal part (Figs 1b–d, 2b, 3, 4e, 6a, 7a, b, d, 8a, b, d, 10, 11a, c, d, g, h, j, 12a, c, d, 13, 14). The oblique joint, which starts at the posterior end of the transverse joint and terminates at the distal end of the protopodite–exopodite junction, separates the proximal part of the flap into two subtriangular lobes (the first and second lobes, ex1 and ex2) (Figs 1b–d, 2b, 3, 4e, 6a, 7a, b, d, 8a, b, d, 10, 11a, c, d, g, h, j, 12a, c, d, 13). Up to 40 long, non-overlapping setae (xs) develop along the distal and posterior margins of the third lobe (Figs 1b–d, 2b, 3, 7a, b, 8a, b, 10, 11a, c, d, g, h, 12a, 13b, 14e–g). The maximum lengths
of the setae are approximately equal to the width of the flap. Flattened imbricate lamellae (lm) develop along the posterior margin of the second lobe (Figs 1b, 2b, 10, 11c, d). They are preserved fragmentarily (Figs 1b, 2b, 10, 11c) or as imprints on the exoskeleton’s surface (Figs 1b, 2b, 11d). A marginal rim runs along the margin of the flap (Figs 10, 12a–d, 13, 14). The anterior section of the marginal rim (ar) (Figs 10, 12a–d, 13, 14a–d) is generally wider than its posterior and distal sections (pr and dr) (Figs 10, 12a–d, 13, 14e–g). The anterior sections of the marginal rims of different limbs are in addition inserted by duct-type soft tissues (dt) preserved as reddish mineral films that merge into a main stem connected to the body (Figs 7d, 8d, 11j).

### 5. Ontogeny of antennae in Hongshiyanaspis yiliangensis

The variations in the trilobite exoskeletons during ontogeny have been studied for a long time and in many taxa. However, little is known about the ontogenetic pattern of trilobite appendages (see Hughes, 2003, 2007 and references therein). The growth of antennae during the ontogeny of trilobites can be performed via two theoretical models: (1) by addition of podomeres; or (2) by stretching of individual podomeres. In order to test these growth models, the number of podomeres, total lengths of the antennae and the average lengths of the podomeres were measured and analysed statistically from our new *Hongshiyanaspis yiliangensis* material, as well as the lengths of the cephalon as quantification of ontogenetic stage (Table 2). A significantly positive correlation ($R^2 = 0.9369$) is found between the average length of the podomeres and the length of the cephalon (Fig. 15). Combined with the similar number of podomeres in nearly complete antennae in normal-sized and fully grown holaspids (Table 2; Figs 1a, b, 2a, b), the growth of antennae is interpreted to occur predominantly by the lengthening of individual podomeres. This may suggest that the number of podomeres increases during the meraspid stages but remains constant during the holaspid.
period, which is similar to the growth of thoracic segments.

6. Comparative anatomy of trilobites and artiopods

Before discussing the evolution of euarthropod limbs and trilobite affinities, it is necessary to make comprehensive anatomical comparisons between *Hongshiyanaspis yiliangensis* and other members of Redlichida, Trilobita and Artiopoda. Detailed discussions organized by structure are given in the subsections below. The results are summarized in Table 3.

6.a. General arrangement of appendages

The appendages of *Hongshiyanaspis yiliangensis* conform to the basic architecture of polymerid trilobite appendages, which are developed as a pair of uniramous antennae followed by a series of homonomous biramous limbs, one pair at each segment (e.g. Hughes, 2003; Scholtz & Edgecombe, 2005, 2006). The attachment sites of the antennae are close to the lateral margins of the hypostome in *H. yiliangensis*, as shown in other trilobite species (Stürmer & Bergström, 1973; Whittington, 1975, 1993; Whittington & Almond, 1987; Hou et al., 2008). The number of pairs of cephalic biramous limbs posterior to the antennae in *H. yiliangensis* is interpreted to be three, which is consistent with the situations in other well-documented trilobite species (e.g. Hughes, 2003), including *Eoredlichia intermedia* (Hou et al., 2008), *Olenoides serratus* (Whittington, 1975), *Triarthrus eatoni* (Cisne, 1975, 1981; Whittington & Almond, 1987), *Rhenops cf. anserinus* (Bartels, Briggs & Brassel, 1998) and *Chotecops ferdinandi* (Bruton & Haas, 1999). The claim of four pairs in *Ceraurus pleurexanthemus* (Størmer, 1939) or a fourth pair overlapping the cephalic/thoracic boundary in *Placoparia cambriensis* (Edgecombe & Ramsköld, 1999) requires further research (Hughes, 2003; Scholtz & Edgecombe, 2005).

The thoracic limbs of *H. yiliangensis* show no significant variation in shape, similar to other trilobites (see...
Appendages of an early Cambrian trilobite

6.b. Antennae

Like the antennae known from other trilobites, the antennae of *Hongshiyanaspis yiliangensis* comprise numerous podomeres with spines with putative sensory function (Whittington, 1975, 1993; Bruton & Haas, 1999; Hou et al. 2008). The proximal podomeres in *H. yiliangensis* are stouter than the distal podomeres, as seen in other trilobites (Raymond, 1920; Stürmer & Bergström, 1973; Whittington, 1975; Bergström & Brassel, 1984; Whittington & Almond, 1987; Shu et al. 1995), probably providing a stronger mechanical force in the proximal section of the antenna to create effective swinging for the distal section. The maximum number of podomeres in *H. yiliangensis* antennae is 27, whereas more than 40 are found in *Chotocops ferdinandi* (Bruton & Haas, 1999), 45 in *Eoredlichia intermedia* (Hou et al. 2008), ~ 50 in *Olenoides serratus* (Whittington, 1975, fig. 3), > 30 in *Triarthrus eatoni* (Whittington & Almond, 1987) and ~ 30 in *Palaeolenus lantenoisi* (Hu et al. 2013). Thus, the numbers of podomeres in antennae are likely to vary among different trilobite species.

6.c. Post-antennal biramous limbs

The post-antennal biramous limbs of *Hongshiyanaspis yiliangensis* are consistent with the basic architecture shown in other polymerid trilobites and artiopods, in having a protopodite composed of a single segment and an endopodite with seven segments composed of six endite-bearing podomeres and a claw-bearing terminal podomere (Fig. 16; Bergström, 1972; Whittington, 1975; Bergström & Brassel, 1984; Whittington & Almond, 1987; Bruton & Haas, 1999; Hou et al. 2008), but the exopodite is unique with its tripartite flap composition.

6.c.1. Protopodite and body–limb junction

The coexistence of both clustered and non-clustered spines on the gnathobase of the protopodite of *Hongshiyanaspis yiliangensis* (Fig. 16) suggests that reconstructions of gnathobases in various polymerid trilobites are possibly incomplete when they only show either clustered or non-clustered spines (e.g. Müller & Walossek, 1987, fig. 27). Putative muscle bundles around the arthrodial membranes of the protopodite in trilobites have not been reported in former studies and provide new information on the musculature of trilobite limbs. Fine setae that are nicely preserved on the protopodite and probably also on the endopodite are most likely to have a sensory function, as suggested in other euarthropods (e.g. Strausfeld, 2016).

The body–limb junction formed by an arthrodial membrane between the protopodite and each thoracic sternite shown in the *Hongshiyanaspis yiliangensis* material supplements the poor record of the body–limb junction in polymerid trilobites. Although the body–limb arthrodial membrane has been shown in several Orsten stem–euarthropods (see Haug et al. 2013 and references therein), the most similar junction to the arthrodial membrane and sternite as in *H. yiliangensis* best documented in the nektaspid *Misszhouia longicaudata* (Ramsköld et al. 1996, fig. 2; Chen, Edgecombe & Ramsköld, 1997, figs 8a, 9a), where curved annihilations of arthrodial membrane and hourglass-shaped sternites are also visible. No indication for either a second proximal podomere or a proximal endite is present on the protopodite of *H. yiliangensis*.
6.c.2. Endopodite

As in other biramous trilobite limbs, the six proximal podomeres of the endopodite have differentiated shapes, especially in their spinous endites, and show a tendency to taper from the proximal towards the distal end (Fig. 16; Bergström, 1972; Whittington, 1975; Bergström & Brassel, 1984; Whittington & Almond, 1987; Bruton & Haas, 1999; Hou et al. 2008). Rather than simply consisting of claws as indicated in other trilobite species, the seventh and terminal podomere has a short rigid base and is connected to the sixth podomere by an arthrodial membrane. Three distal claws are present, and their morphology and arrangements vary owing to the different angles of burial similar to the Hunsrück Shale species.
Figure 11. (Colour online) Details of post-antennal biramous limbs of *Hongshiyanaspis yiliangensis* Zhang & Lin in Zhang et al. 1980, from the Xiazhuang fossil assemblage, Kunming, Yunnan, SW China. (a–d) Partial view, NIGPAS 164503. (a, b) Nearly complete fourth dextral thoracic limb, as in Figure 10a. (a) Image showing the protopodite, endopodite and exopodite. (b) Close-up of the protopodite in (a) showing an arthrodial membrane as the body–limb junction and a transverse furrow (red arrows). (c, d) Close-ups of exopodites. (c) Distal parts of exopodites of the fifth, sixth and eighth dextral thoracic limbs showing the setae along margins and the transverse joints. (d) Exopodite of the second sinistral thoracic limb, showing reddish imprints of lamellae on the exoskeleton. (e, f) Counterpart, NIGPAS 164506, close-ups of the seventh claw-bearing podomere of the endopodite, showing the arthrodial membrane between the sixth and seventh podomereres of the endopodites (yellow arrows) and the three distal claws, as in the first and second thoracic limbs of Figures 4c, 6a. (g, h) NIGPAS 164505, nearly complete thoracic limbs. (g) Part, NIGPAS 164505a, the third sinistral thoracic limb, as in Figures 1a, 3a. (h) Counterpart, NIGPAS 164505b, the sixth sinistral thoracic limb, as in Figures 1b, 3b. (i) Part only, NIGPAS 164509, close-up of the body–limb junctions, as in Figures 7c, 8c. (j) Counterpart, NIGPAS 164511b, close-up of the exopodites and duct-type soft tissues, as in Figures 7d, 8d. Black and white arrows indicate the transverse and oblique joints separating the exopodite lobes. Red arrows indicate the transverse furrow on the protopodite. Yellow arrows indicate the arthrodial membrane between the sixth and seventh podomereres of the endopodites. Abbreviations as in Figures 1–3, 8.
Table 2. Measurements of antennae and cranidia of holaspid specimens *Hongshiyanaspis yiliangensis* Zhang & Lin in Zhang et al. 1980

<table>
<thead>
<tr>
<th>Specimen</th>
<th>Ontogenetic stage</th>
<th>Left/right antenna</th>
<th>Length of cephalon (mm)</th>
<th>Total length (mm)</th>
<th>Number of podomeres</th>
<th>Average length of podomere (mm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>NIGPAS 164514</td>
<td>Early holaspid</td>
<td>Left</td>
<td>1.85</td>
<td>0.65</td>
<td>6</td>
<td>0.11</td>
</tr>
<tr>
<td>NIGPAS 164514</td>
<td>Early holaspid</td>
<td>Right</td>
<td>1.85</td>
<td>2.03</td>
<td>14</td>
<td>0.15</td>
</tr>
<tr>
<td>NIGPAS 164504</td>
<td>Normal-sized holaspid</td>
<td>Left</td>
<td>4.85</td>
<td>8.25</td>
<td>25</td>
<td>0.33</td>
</tr>
<tr>
<td>NIGPAS 164504</td>
<td>Normal-sized holaspid</td>
<td>Right</td>
<td>4.85</td>
<td>6.92</td>
<td>23</td>
<td>0.30</td>
</tr>
<tr>
<td>NIGPAS 164512</td>
<td>Normal-sized holaspid</td>
<td>Left</td>
<td>5.54</td>
<td>2.38</td>
<td>8</td>
<td>0.30</td>
</tr>
<tr>
<td>NIGPAS 164512</td>
<td>Normal-sized holaspid</td>
<td>Right</td>
<td>5.54</td>
<td>2.57</td>
<td>9</td>
<td>0.29</td>
</tr>
<tr>
<td>NIGPAS 164510</td>
<td>Normal-sized holaspid</td>
<td>Left</td>
<td>5.78</td>
<td>7.33</td>
<td>18</td>
<td>0.41</td>
</tr>
<tr>
<td>NIGPAS 164506</td>
<td>Normal-sized holaspid</td>
<td>Left</td>
<td>6.61</td>
<td>4.69</td>
<td>13</td>
<td>0.36</td>
</tr>
<tr>
<td>NIGPAS 164503</td>
<td>Fully grown holaspid</td>
<td>Left</td>
<td>9.40</td>
<td>13.39</td>
<td>23</td>
<td>0.58</td>
</tr>
<tr>
<td>NIGPAS 164503</td>
<td>Fully grown holaspid</td>
<td>Right</td>
<td>9.40</td>
<td>14.01</td>
<td>27</td>
<td>0.52</td>
</tr>
</tbody>
</table>

Figure 12. (Colour online) Two disarticulated post-antennal biramous limbs of *Hongshiyanaspis yiliangensis* Zhang & Lin in Zhang et al. 1980, from the Xiazhuang fossil assemblage, Kunming, Yunnan, SW China, NIGPAS 164508. (a, b) Limb A as part and counterpart, respectively. (c, d) Limb B as part and counterpart, respectively. (e, f) Close-ups of the protopodite of Limb A, as in (a) and (b), respectively. (g) Close-up of the protopodite of Limb B, as in (c). (h) Close-up of the distal podomeres of Limb B, as in (d). Blue and yellow arrows indicate the proximal and distal boundaries of the arthrodial membrane. Black and white arrows indicate the transverse and oblique joints separating the exopodite lobes. Abbreviations as in Figures 1–3, 8.
Table 3. Comparative anatomy between *Hongshiyanaspis yiliangensis* and other redlichiids, trilobites and artiopods

<table>
<thead>
<tr>
<th>Body parts</th>
<th>Characters</th>
<th><em>Hongshiyanaspis yiliangensis</em></th>
<th>Redlichiida</th>
<th>Trilobita</th>
<th>Artiopoda</th>
</tr>
</thead>
<tbody>
<tr>
<td>Arrangement of appendages</td>
<td>Overall morphology</td>
<td>Deutocerebral/hypostomal antennae and homonomous post-antennal limbs</td>
<td>Same</td>
<td>Mostly three pairs; the claims of four pairs in <em>Ceratops</em> and <em>Placoparia</em> are questionable</td>
<td>Mostly three pairs; those interpreted with more are open to question</td>
</tr>
<tr>
<td>Cephalon</td>
<td>Three pairs of biramous limbs</td>
<td>Same</td>
<td>Mostly three pairs; the claims of four pairs in <em>Ceratops</em> and <em>Placoparia</em> are questionable</td>
<td>Mostly three pairs; those interpreted with more are open to question</td>
<td></td>
</tr>
<tr>
<td>Thorax (trilobites) or trunk (artiopods)</td>
<td>One pair of biramous limbs per thoracic tergites</td>
<td>Same</td>
<td>Likely same; unknown for those with differentiated thorax</td>
<td>Various correspondences between numbers of limbs and tergites, extreme cases in those with fusion of trunk tergites, e.g. naraoiids and tegopeltids</td>
<td></td>
</tr>
<tr>
<td>Pygidium (trilobites only)</td>
<td>Micropygous, one pair of biramous limbs in axial ring, multiple pairs in terminal axial piece</td>
<td>Same</td>
<td>Number of limbs almost doubles the number of pygidial tergites in <em>Triarthrus</em>; unknown in others</td>
<td>Various</td>
<td></td>
</tr>
<tr>
<td>Antennae</td>
<td>Rami</td>
<td>Uniramous</td>
<td>Same</td>
<td>Same</td>
<td>Same</td>
</tr>
<tr>
<td>Numbers of podomeres</td>
<td>Maximum ≥ 27</td>
<td>45 in <em>Eoredlichia</em></td>
<td>Various</td>
<td>Same</td>
<td>Various</td>
</tr>
<tr>
<td>Shapes of podomeres</td>
<td>Thick in proximal, slender in distal</td>
<td>Unknown</td>
<td>Likely same; unknown for those with differentiated thorax</td>
<td>Various</td>
<td></td>
</tr>
<tr>
<td>Spines</td>
<td>No less than one per podomere</td>
<td>Same</td>
<td>Multiple in <em>Olenoides</em></td>
<td>Various</td>
<td></td>
</tr>
<tr>
<td>Post-antennal limbs</td>
<td>Biramous</td>
<td>Same</td>
<td>Same</td>
<td>Same</td>
<td>Mostly same; not obvious in some species</td>
</tr>
<tr>
<td>Body–limb junction</td>
<td>Arthrodial membrane attached to sternite</td>
<td>Unknown</td>
<td>Same</td>
<td>Unknown</td>
<td></td>
</tr>
<tr>
<td>Protopodite</td>
<td>Podomere(s)</td>
<td>Single, no differentiation</td>
<td>Same</td>
<td>Same</td>
<td>Same; except for the prebasal endite in <em>Sidneyia</em></td>
</tr>
<tr>
<td>Gnatobase</td>
<td>Clustered and non-clustered spines</td>
<td>Non-clustered only</td>
<td>Likely both types</td>
<td>Various</td>
<td></td>
</tr>
<tr>
<td>Transverse furrow(s)</td>
<td>One</td>
<td>Unknown</td>
<td>Unknown; rejected in <em>Triarthrus</em></td>
<td>Unknown</td>
<td></td>
</tr>
<tr>
<td>Musculature</td>
<td>Present across arthrodial membrane</td>
<td>Unknown</td>
<td>Unknown</td>
<td>Unknown</td>
<td></td>
</tr>
<tr>
<td>Setae</td>
<td>Sensory setae</td>
<td>Unknown</td>
<td>Putative sensory setae in <em>Olenoides</em></td>
<td>Unknown in most species</td>
<td></td>
</tr>
<tr>
<td>Numbers of podomeres</td>
<td>Seven</td>
<td>Same</td>
<td>Questionable for those are not seven</td>
<td>Mostly unknown, likely various</td>
<td></td>
</tr>
<tr>
<td>Shapes of podomeres</td>
<td>Differentiated</td>
<td>Same</td>
<td>Same</td>
<td>Likely same if present</td>
<td></td>
</tr>
<tr>
<td>Endites</td>
<td>Spinous</td>
<td>Same</td>
<td>Likely same if present</td>
<td>Likely same if present</td>
<td></td>
</tr>
<tr>
<td>Claws</td>
<td>Three rooted on a very short podomere</td>
<td>Three</td>
<td>Three</td>
<td>Likely various</td>
<td></td>
</tr>
<tr>
<td>Musculature</td>
<td>Oblique muscle bundles</td>
<td>Unknown</td>
<td>Unknown</td>
<td>Unknown</td>
<td></td>
</tr>
<tr>
<td>Setae/spines</td>
<td>Sensory setae</td>
<td>Unknown</td>
<td>Lateral spines in <em>Olenoides</em></td>
<td>Mostly unknown, likely various</td>
<td></td>
</tr>
<tr>
<td>Exopodite</td>
<td>Composition</td>
<td>A flap of three lobes</td>
<td>Lobs or shafts with various numbers</td>
<td>Lobs or shafts with various numbers</td>
<td></td>
</tr>
<tr>
<td>Numbers of joints</td>
<td>One transverse and one oblique</td>
<td>One transverse</td>
<td>Various</td>
<td>Various</td>
<td></td>
</tr>
<tr>
<td>Lamellae</td>
<td>Imbricate</td>
<td>Same</td>
<td>Likely same if present</td>
<td>Likely same if present</td>
<td></td>
</tr>
<tr>
<td>Distal setae</td>
<td>Non-overlapping</td>
<td>Same</td>
<td>Likely same if present</td>
<td>Likely same if present</td>
<td></td>
</tr>
<tr>
<td>Marginal rims</td>
<td>Present on lobes</td>
<td>Unknown</td>
<td>Unknown</td>
<td>Unknown</td>
<td></td>
</tr>
<tr>
<td>Duct-type structures</td>
<td>Present, merge into a main stem</td>
<td>Unknown</td>
<td>Unknown</td>
<td>Unknown</td>
<td></td>
</tr>
</tbody>
</table>

Note: the specific names of species are omitted for concision. Situations labelled as ‘same’ are compared with the adjacent column on the left side. The Redlichiida is represented by *Eoredlichia intermedia* (Hou et al. 2008; Shu et al. 1995). The Trilobita represent polymerid trilobites, excludingagnostids. See text for detailed discussions and references.
**Figure 13.** Interpretative line drawings of two disarticulated post-antennal biramous limbs of *Hongshiyanaspis yiliangensis* Zhang & Lin in Zhang et al. 1980, from the Xiazhuang fossil assemblage, Kunming, Yunnan, SW China, NIGPAS 164508. (a) Counterpart, NIGPAS 164508a, as in Figures 12b, d. (b) Part, NIGPAS 164508b, as in Figures 12a, c, but horizontally flipped. Abbreviations as in Figures 1–3, 8.

*Chotecops ferdinandi* (Bruton & Haas, 1999, text-fig. 19), whereas the three claws clearly do not merge together into a common base in *H. yiliangensis*. Taking into account the pyritized preservation, the common base of the claws shown in *C. ferdinandi* (Bruton & Haas, 1999, text-fig. 19) may be a taphonomic artefact.

6.c.3. Exopodite

Exopodites have been previously reported from several polymerid trilobites (Størmer, 1939, 1951; Bergström, 1972; Whittington, 1975; Whittington & Almond, 1987; Bruton & Haas, 1999; Hou et al. 2008). The morphology of the exopodite in *Hongshiyanaspis yiliangensis* is unique in possessing a tripartite flattened flap (Fig. 16). Although tripartite exopodite flaps are also found in several other arthropods, such as *Kwa-maia lata* (Hou & Bergström, 1997), *Sidneyia inexpectans* (Stein, 2013), *Emeraldella brocki* (Stein & Selden, 2012) and *Arthroaspis bergstroemi* (Stein et al. 2013), the flap in *H. yiliangensis* differs in having a transverse and an oblique joint, no sharp discontinuity of the margin at the endpoints of the joints, and lobes with subequal widths. The exopodite flap in *H. yiliangensis* is almost double the maximum width of the endopodite, which is distinct from those described from almost all other lobate exopodites of trilobites (Whittington, 1975; Bruton & Haas, 1999; Hou et al. 2008) but comparable to those in *K. lata* (Hou & Bergström, 1997), *Squamacula clypeata* (Zhang et al. 2004), *Kwanyinaspis maotianshanensis* (Zhang & Shu, 2005) and *Naraoia spinosa* (Zhang, Shu & Erwin, 2007) (also see Ortega-Hernández, Legg & Braddy, 2013, fig. 4 for reconstructions). These wide exopodite flaps in *H. yiliangensis* may be an adaptation to powerful swimming and/or a more effective respiration, with the joints possibly directing water currents. Non-overlapping setae and imbricate lamellae develop along the margins of
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Figure 14. (Colour online) Marginal areas in the exopodites of post-antennal biramous limbs of *Hongshiyanaspis yiliangensis* Zhang & Lin in Zhang et al. 1980, from the Xiazhuang fossil assemblage, Kunming, Yunnan, SW China. (a, b, c–g) NIGPAS 164503. (a, b) Anterior margins of the exopodites in the third and fourth thoracic limbs, respectively. (e–g) Distal and posterior margins of the exopodites in the eighth, sixth and fifth thoracic limbs, respectively. (c, d) NIGAPS 164511b, anterior margins of the exopodites in the fifth and sixth thoracic limbs. Black arrows indicate the transverse joints separating the exopodite lobes. Abbreviations as in Figures 1–3, 8, 10.

the distal (= third) and proximal (= first) lobes of the exopodite flap in *H. yiliangensis*, respectively. This situation is present not only in other polymerid trilobites such as the Cambrian Stage 3 *Eoredlichia intermedia* (Hou et al. 2008) and the Cambrian Stage 5 *Olenoides serratus* (Whittington, 1975), but also in a number of non-trilobite artiopods (see Zhang, Shu & Erwin, 2007; Ortega-Hernández, Legg & Braddy, 2013; Stein et al. 2013 and references therein). Despite the limited knowledge, the overall similarity between the exopodites of *O. serratus* (Order Corynexochida; Whittington, 1975) and *E. intermedia* (Order Redlichiida; Hou et al. 2008) as well as the differences between the almost contemporaneous *E. intermedia* and *H. yiliangensis* (Order Redlichiida) suggests that these anatomical variations are more likely the results of divergence owing to different ecological adaptations of the species rather than the different evolutionary tendencies of trilobite orders.

The exopodite flaps in *Hongshiyanaspis yiliangen sis* exhibit duct-type soft tissues (Fig. 16). These tissues of the different limbs further merge into a main stem in the ventral soft parts of the body. The morphology and positions of these duct-type structures are comparable to those ‘invasive caeca’ or ‘triangular strips’ in a range of stem-group euarthropods from the Burgess Shale (see Aria & Caron, 2015, figs 2, 3, 5, 9). A digestive nature was suggested for those from the Burgess Shale because they are connected to the main alimentary canal (Aria & Caron, 2015). However, the duct-type structures here may be different tissues, since no connection to the gut is observable. Because exopodites have long been interpreted as respiratory organs like gills (e.g. Hou & Bergström, 1997), a putative circulatory nature for these duct-type structures is proposed here. Still, further investigations are required to clarify these alternative possibilities.

6.d. Basic appendage morphology of trilobites and artiopods

By discussing comparative anatomy under the light of the new *Hongshiyanaspis yiliangensis* appendages above, we can conclude in general that all trilobites and other artiopods share the same basic architecture of appendages comprising a pair of deutocerebral/
hystostomal uniramous antennae and a series of paired homonomous biramous limbs (e.g. Hughes, 2003; Scholtz & Edgecombe, 2005, 2006). However, within ascending systematic hierarchies from Redlichiida, Trilobita to Artiopoda, their biramous limbs show a conserved morphology by consisting of seven podomeres in the endopodites and one single segment in the protopodite, but also considerable morphological disparity in the composition of the exopodites.

7. Affinities of trilobites: mandibulate or chelicerate?

Three alternative affinities for trilobites and other artiopods have been hypothesized: as stem-chelicerates, stem-mandibulates or a stem lineage of both Chelicerata and Mandibulata (Budd & Telford, 2009). Comparisons between major upper stem-group euarthropods and the stem and crown groups of Mandibulata and Chelicerata can reveal general evolutionary trends of euarthropod appendages in two aspects, i.e. the arrangement of appendages along the anterior–posterior main body axis, and the composition of limb rami. These trends can be essential for interpreting the affinities of trilobites and artiopods, which are discussed below and summarized in Table 4.

7.a. Arrangement of appendages

Considering the anteriormost (deutocerebral and tritocerebral) appendages, within upper stem-group euarthropods, trilobites and other artiopods are unique in lacking any specialized cephalic feeding appendages (Table 4), such as the tritocerebral specialized post-antennal appendages (SPAs) in large ‘bivalved’ stem-euarthropods (e.g. Legg et al. 2012) and fuxianhuiids (e.g. Yang et al. 2013) and mandibulates (Scholtz & Edgecombe, 2005, 2006). In megacheirans and chelicerates, however, neither the SGAs nor the chelicerae are antenniform but raptorial, and consist of a limited number of podomeres (Chen, Waloszek & Maas, 2004). Therefore, the ‘secondary antennae’ have been regarded as strong evidence supporting the mandibulate affinities of trilobites and other artiopods (Scholtz & Edgecombe, 2005, 2006).

On other hand, all post-tritocerebral limbs are homonomous on both the cephalon and trunk of all the upper stem-group euarthropods mentioned above (e.g. Legg et al. 2012; Ortega-Hernández, Legg & Braddy, 2013; Yang et al. 2013; Aria, Caron & Gaines, 2015). In Mandibulata, although several anterior post-tritocerebral limbs are specialized for feeding and the rest can show various degrees of differentiation or tagmatization, continuous series of undifferentiated homonomous limbs are typically present on the trunk of myriapods and some crustaceans such as Anostraca and Remipedia. By contrast, in Chelicerata (except for pycnogonids), post-tritocerebral limbs on the anterior and posterior body tagma (prosoma and opisthosoma) are always differentiated. In Xiphosura, the post-tritocerebral limbs of the anterior and posterior tagma are differentiated into uniramous legs (except for the last leg with flabellum) and gills, respectively, whereas the posterior limbs have been reduced in other chelicerate crown groups. Therefore, the homonomous pattern of limbs shown in trilobites, artiopods and other upper stem-group euarthropods has been retained in Mandibulata, but was lost by the basic anterior and posterior tagmatization in Chelicerata, providing further evidence supporting the mandibulate concept.

Figure 16. Reconstruction of paired post-antennal biramous limbs connected to sternites. Note that the structures are not in actual life positions, including the orientations of the protopodite–exopodite and body–limb junctions, but are flattened to exhibit the most detailed limb anatomy and their connections, as in the mainstream reconstructions of artiopodan limbs. The proximal lamellae are hypothetical and drawn as dashed lines, and the duct-type soft tissues are in grey. Abbreviations as in Figures 1–3, 8, 10.
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7.b. Biramous limbs

Trilobites and other artiopods are also distinguished by the composition of their post-antennal biramous limbs. Despite that all of these limbs in the upper stem groups of Euarthropoda mentioned above are homonomous, the basic compositions of their endopodites and exopodites are different (Table 4). In fuxianhuiids (e.g. Yang et al. 2013) and large ‘bivalved’ stem-euarthropods (e.g. Legg et al. 2012), the podomeres of the endopodites show no differentiation in shape and their number exceeds ten, and their exopodites are undivided flaps. In megacheirans (e.g. Davidson & Erwin, 2006), endopodite podomeres are also undifferentiated and are at least eight in number (including the distal claw), whereas their exopodites are flaps comprising two lobes. It is worth noting that the morphology of the exopodites is consistent in each of these groups, whereas the number of their endopodite podomeres varies. This indicates that the exopodites are more morphologically conserved than the endopodites in these upper stem-euarthropod groups.

In trilobites and other artiopods, however, endopodites and exopodites exhibit different patterns of morphological disparity. For the endopodites, the number of podomeres is consistently seven (including distal claws) and the morphological differences of the podomeres are geometric rather than qualitative in trilobites and other artiopods with a few questionable exceptions. By contrast, the exopodites vary in composition between any two trilobite and artiopod species. Their exopodites can be a flap comprising one to three lobes, or a shaft, showing a high morphological disparity. Within the two modern euarthropod lineages Mandibulata and Chelicera, this similar disparity of the exopodites is only exhibited in stemgroup and crown-group crustaceans, whereas the exopodites have been reduced or lost in Chelicera.

Developmental biology has shown that endopodites and exopodites are rami originating from the same main proximal–distal axis of the biramous limb bud (Wolff & Scholtz, 2008). However, the striking distinction in patterns of disparity between the endopodites and exopodites seen in stem-euarthropods suggests that their developmental regulatory machineries converged early in the evolution of euarthropod biramous limbs (Davidson & Erwin, 2006). In trilobites, other artiopods and crustaceans, the developmental genetic regulatory programmes were relatively conserved in the endopodites, whereas they were much more flexible in the exopodites. Nevertheless, the situations are reversed in large ‘bivalved’ stem-euarthropods, fuxianhuiids and megacheirans, with higher conservativeness in the exopodites than in the endopodites. The cheliceral concept would thus require abandoning the highly evolved developmental machinery of the exopodites, which is an unlikely case.

Therefore, from a developmental perspective, the pattern of morphological disparity shown by the two rami also supports the mandibulate affinities of trilobites and other artiopods.

7.c. Evolutionary trends of euarthropod appendages

Most recent morphological and molecular phylogenetic frameworks of Euarthropoda have, respectively, revealed a successive appearance of euarthropod...
characters in upper stem groups from ‘bivalved’ stem-arthropods, fuxianhuilids, megacheirans to artiopods, and the relationships of crown groups such as Chelicerata + Mandibulata (Myriapoda + Tetracotyla/Paracrustacea) (e.g. Edgecombe & Legg, 2014). Within the upper stem groups, the early evolution of euarthropod appendages has undergone the origination of deutocerebral (antennae, SGAs) and tritocerebral (SPAs) specialized appendages (Edgecombe & Legg, 2014), reduction in the number of endopodite podomeres from more than ten to the ground plan of seven, and differentiation of endopodite podomeres (Boxshall, 2004, 2013). However, the subdivision of the protopodite, and the differentiation or specialization of post-tritocerebral or trunk appendages should occur during the early evolution of Mandibulata and Chelicerata (Boxshall, 2004, 2013).

7.d. Affinities of trilobites and artiopods

The chelicerate affinity hypothesis for trilobites and artiopods had been the mainstream for a long time until new evidence for mandibulate affinities and conflicts with the chelicerate concept were put forward by Scholtz & Edgecombe (2005, 2006). However, controversies have still been going on since then, as both the mandibulate and chelicerate concepts gained support from different recent phylogenetic analyses (e.g. Ortega-Hernández, Legg & Braddy, 2013; Legg, Sutton & Edgecombe, 2013). The evidence given by Scholtz & Edgecombe (2005, 2006), including the ‘second antennae’ and head segmentation, together with our new arguments based on the homonomous pattern and composition of the biramous limbs, supports the mandibulate concept of trilobites and at least some artiopods. Meanwhile, this evidence proposes the critical character transformations that are required to fit the chelicerate concept, including the loss of antennae, disappearance of delimited cephalon–trunk tagmatization, change of limb patterning along the main body axis and reorganization of developmental machineries in the limb rami. However, we cannot negate the possibility that with new data, some members of the current Artiopoda definition may be closer to the chelicerate lineage. If this is true, subdivision of artiopods would be essential.

8. Conclusion

The appendages of Hongshiyanaspis yiliangensis reinforce the idea that polymerid trilobites share a homonomous arrangement of biramous limbs as well as conserved anatomy in the protopodites and endopodites, but have significant inter-taxa differences in the exopodites. This appendage architecture of trilobites is highly comparable to that of other artiopods. Ontogeny of trilobite antennae is studied for the first time and shows a growth model of lengthening each podomere. By reinvestigating and comparing appendages in upper stem groups and crown groups of Eu-arthropoda, we show similarities in the arrangement of homonomous limbs and patterns of morphological disparity in the endopodites and exopodites between artiopods (including trilobites) and mandibulates. Together with the shared ‘secondary antennae’ and head tagmosis, these new lines of evidence further support the mandibulate affinities of trilobites and at least some artiopods. However, more data on the appendages of trilobites and other stem-group euarthropods are still essential to resolve controversies surrounding the problem of trilobite affinities.

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Supplementary material

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