A Silurian ‘marrellomorph’ arthropod

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Xylokorys chledophilia, a new arthropod with three-dimensionally preserved soft tissues, is described from the Herefordshire (Silurian) Lagerstätte of England. The head and trunk are covered by a relatively featureless ovoid carapace, which comprises a domed central part and a flange-like border. The head bears five pairs of appendages. The first is uniramous, with dorsal and ventral projections distally. Appendages two to four are biramous and each endopod terminates in two projections. Appendage five is possibly biramous. The hypostome is very long and rectangular in outline. There are approximately 35 pairs of biramous trunk appendages. Each exopod comprises a long slender shaft bearing numerous fine filaments; each endopod comprises a ribbon-like shaft bearing paddle-like endites. Morphological comparisons and cladistic analyses of X. chledophilia indicate affinity with Vachonisia rogeri from the Lower Devonian Hunsrück Slate, within the marrellomorphs, but assignment to Marrellomorpha is provisional pending revision of other members of this clade. Xylokorys is the first ‘marrellomorph’ to be reported from the Silurian. It is interpreted as a benthic particle filter feeder, which may also have consumed prey items.

Keywords: Arthropoda; exceptional preservation; Herefordshire Lagerstätte; Marrellomorpha; Silurian; Wenlock Series

1. INTRODUCTION

A new arthropod, Xylokorys chledophilia, is described from the Herefordshire Lagerstätte of the Welsh Borderland (Briggs et al. 1996; see Siveter et al. in press and references therein), in which a wide variety of invertebrates of mid-Silurian, Wenlock Series age (approx. 425 Myr ago) are exceptionally preserved in three dimensions. Arthropods form one of the main components of this fauna, including a stem-group chelicerate (Orr et al. 2000b; Sutton et al. 2002), two myocarid ostracodes (Siveter et al. 2003, 2007), a phyllocarid (Briggs et al. 2004), a pycnogonid (Siveter et al. 2004), a barnacle (Briggs et al. 2005) and a stem-group crustacean (Siveter et al. 2007). Xylokorys chledophilia is placed provisionally within Marrellomorpha pending revision of other taxa assigned to this inadequately known arthropod group; it provides the most complete three-dimensional data on any ‘marrellomorph’ species. The four previously recorded marrellomorph genera are known from a few localities of Cambrian, Ordovician and Devonian age in, respectively, Canada and China (Marrella), Bohemia and Morocco (Furca) and Germany (Mimetaster and Vachonisia). Furca is known from approximately 55 specimens (Chlupaˇcˇ 1999; P. Van Roy 2007, personal communication), Mimetaster and Vachonisia from, respectively, approximately 75 and 20 (Stürmer & Bergström 1976; Bartels et al. 1998; G. Oltmann 2007, personal communication), whereas over 24 000 specimens have been recorded for Marrella (Garcı´a-Bellido & Collins 2006). Xylokorys is the first example reported from the Silurian: just two specimens are known, comprising approximately 0.05% of the recorded number of specimens from the Herefordshire Lagerstätte.

Marrellomorphs have been considered by many authors as basal to trilobites and their ‘arachnomorph’ relatives, the latter comprising many fossil taxa as well as living and fossil chelicerates; however, in some investigations, they have resolved basal to all other schizoramian arthropods, including the crustaceans (see §4).

2. MATERIAL AND METHODS

The fossils of the Herefordshire Lagerstätte are preserved in three-dimensional form as calcitic void infills in early diagenetic carbonate concretions within a volcaniastic horizon (Orr et al. 2000a). They retain fine morphological detail of external surfaces, and some also show the remains of the gut. In order to make them available for study, the specimens are serially ground, digitally photographed and then rendered in the round as ‘virtual fossils’ by computer (Sutton et al. 2001a,b, 2002). One of the Xylokorys specimens (OUMNH C.29604) has been reconstructed following serial grinding at 30 μm intervals, and then reconstruction at 60 μm intervals except for detailed reconstruction of the trunk appendages, which used every 30 μm ‘slice’. Prior to grinding, this specimen was traversed by

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Electronic supplementary material is available at http://dx.doi.org/10.1098/rspb.2007.0712 or via http://www.journals.royalsoc.ac.uk.

Received 26 May 2007
Accepted 21 June 2007

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Figure 1. *Xylokorys chledophilia*. (a–l) Almost complete specimen, OUMNH C.29604, ‘virtual’ reconstructions. (a,b,g,k) Dorsal stereo-pair, lateral view, anterior and ventral stereo-pairs, ×1.6. (f) Posteroventral view, ×2.4. (h) Anterior oblique stereo-pair, ×1.7. (c,d,l) Carapace removed, lateral view, dorsal and anterior oblique stereo-pairs, ×1.6. (e) Hypostome with surrounding appendages removed, ventral stereo-pair, ×2.7. (i,j) Gut infill, ventral and lateral stereo-pairs, ×3.4. A1, appendage 1; A2, limb base of appendage 2; A2en, appendage 2 endopod; A2ex, appendage 2 exopod; A3, limb base of appendage 3; A3en, appendage 3 endopod; A3ex, appendage 3 exopod; A4, limb base of appendage 4; A4en, appendage 4 endopod; A4ex, appendage 4 exopod; A5, limb base of appendage 5; A5ex, appendage 5, presumed exopod; AP?, anterior section of pharynx ?; BR, border ridge; CB, carapace border; Co, constriction; Fu, furrow; Hy, hypostome; In, intestine; Mo, mouth; MR, median ridge; Oe, oesophagus; Ph?, pharynx ?; Ten, trunk appendage endopod; Tex, trunk appendage exopod.
three parallel cuts with a 300 μm saw. The first traverses the anterior extremity of the specimen, passing through the carapace on the right-hand side as well as the distal part of the first pair of appendages; the second occurs at the junction of the head and trunk appendages; and the third is near the posterior margin. The virtual fossil was studied on-screen using interactive stereo-capable viewing software, supplemented by hard copy stereo-pair images.

3. SYSTEMATIC PALAEONTOLOGY

Phylum: Arthropoda

? Class: Marrellomorpha Beurlen, 1934

Genus: Xylokorys gen. nov.

Derivation of name. Greek, xylos (pith) + korys (helmet), after the fancied resemblance of the carapace to the favoured tropical headgear of European military personnel during colonial times.
Diagnosis. Carapace covering head and trunk, ovoid in outline, with a domed central part and a flange-like border indented by a narrow anterior notch and a wide posterior embayment. The head bears five pairs of appendages. The first is short, uniramous, with dorsal and ventral projections distally, the dorsal one (at least) ending in an ear-like process. Appendages two to four are large and biramous. Each endopod bears two projections distally and each exopod bears a fringe of setae distally. The exopod of the fourth appendage is very long. Appendage five is possibly biramous. The hypostome is very long and subrectangular in outline. There are approximately 35 pairs of biramous trunk appendages. Each exopod comprises a long slender shaft bearing approximately 25 fine filaments. Each endopod comprises a ribbon-like shaft with one or two endites proximally, four paddle-like endites along its length and a similar terminal process.

Species: Xylokorys chledophilia sp. nov.

Derivation of name. Greek, chledos (mud) + philia (fondness)

Holotype. As for the genus.

Other material. OUMNH C.29604 (figure 2k), a complete specimen; median length approximately 32 mm.

The datasets from serial grinding are housed in the Oxford University Museum of Natural History (OUMNH). The description is based on OUMNH C.29604.

(a) Carapace

The carapace (figures 1a, h–k, f–h, k and 2lm) is ovoid in outline, the length approximately 1.5 times the width. In frontal profile, it is dome-like, the height approximately 0.5 times the width; each side is gently convex dorsally and slopes laterally at approximately 45°. The maximum height is very close to the anterior margin (figure 1b). In lateral view, the median ridge approximates a straight line with a slight concavity posterior of the mid-length. The anterior margin slopes steeply anteroventrally curving abruptly into the convex ventral margin that converges gradually and then more rapidly with the dorsal margin in the posterior third of its length, the two presumably merging in the carapace border. The first appendage emerges through a rounded median notch that is approximately 0.25 times the width of the carapace. Posteriorly, the carapace margin is very broadly extended laterally (preserved on left-hand side only); its shape mediolaterally is unknown. A median ridge runs the length of the carapace; it is very narrow anteriorly, widens slightly almost to the mid-length and narrows slightly to the posterior margin. There is a pronounced carapace border, which is narrowest anteriorly, becoming wider and more distinct posteriorly where the weak border furrow becomes broader. At the mid-length, the border is approximately 0.1 times the width of the carapace. On the posterior part of the border, particularly on the right-hand side, two weak parallel ridges are evident oriented obliquely to the margin. The lateral border is thicker than the main part of the carapace, especially near the two ridges (figure 2m). The border is slightly concave ventrally here and its inner margin is most sharply defined. The carapace inside the border is featureless apart from the median ridge and a slightly flattened ovoid area that flanks the median ridge and at maximum is 25% as wide as the carapace, and is from approximately 10 to 70% of the carapace length from its anterior end. This area appears to reflect the internal attachment area of the body.

(b) Head

Eyes are absent. The first appendage is uniramous; it originates immediately anterior to the lateral shoulder of the hypostome and extends just beyond the carapace margin (figures 1a–d, f–h, k, l and 2a, d, k, l). Five pairs are evident, presumably corresponding to podomeres. The most proximal podomere is stout and directed anteriorly. The appendage flexes anteroventrally at the articulation between podomeres one and two and anteriorly again between three and four. The third podomere bears a long, fine seta ventrolaterally. The appendage terminates in a dorsal and a ventral projection; the former comprises a laterally flat, ear-like process, and the latter expands for a short distance proximally but is unknown distally. One of these is presumably a projection of the penultimate podomere (similar to the structure of the great appendage in, for example, Alalcomenaeus (see Briggs & Collins 1999), or the fixed finger of a chela).

The second appendage is biramous (figures 1a–d, f–h, k, l and 2a, e, f). The elongate proximal part of the limb base bears a spine-like extension posteriorly that is directed strongly posteriorly and slightly inwards near the mouth. The more distal part of the limb base is laterally compressed with subrectangular inner and outer faces, the inner of which is weakly divided by a shallow groove into anterior and posterior portions that continue distally into an exopod and endopod, respectively. The proximal part of the endopod (? a single podomere) is directed subanteriorly and is laterally compressed. The second part (? two subequally short podomeres) flexes subventrally and is shorter, and bears several setae distally on its outer margin. In the third part, also subventrally directed, the endopod divides into a long outer and a slightly shorter inner projection that both bear radiating setae distally, and the inner branch has three discrete setae (corresponding to three podomeres) along its inner margin. One of the projections is presumably that of the penultimate podomere, as in appendage one. The proximal part of the exopod (? a single podomere) is directed anteriorly, the second (? two podomeres) flexes subventrally and the third (? four or five short podomeres) flexes posteriorly to be subparallel to the carapace margin, and it bears two rows of setae that project normal to the axis. All three parts are laterally compressed.

The third appendage (figures 1a–d, f–h, k, l and 2a, f, l) inserts posteroventral to the second and is very similar to it, except that it is slightly larger and the spine-like extension of the limb base trends more inwards.

The fourth appendage is biramous (figures 1a–d, f–h, k, l and 2a, g, d, m). It inserts at the posterior margin of the hypostome, posteroventral to the third appendage. In contrast to the spine-like extension in appendages two and three, a relatively large, transversely elongate gnathobase-like structure is present in the same position, comprising a sharp medial flange and a small posteromedial spine. The endopod is similar in structure to that of the second appendage but is much smaller. The first two parts of the exopod are similar to those of the second
appendage. The third part extends backwards from the
second part with weak geniculation, is twice the length of
its second appendage homologue and bears six setal tufts
on its outer margin and terminates in one or two setae
(probably indicating seven podomeres in total).

The fifth appendage is incompletely preserved and
comprises a limb base that inserts immediately posterior
of the fourth appendage and a single ramus (figures 1c,dklj
and 2a,h). The ramus trends sinuously anterolaterally
(separate podomeres are unresolved), is antenna-like and
thickened slightly proximally. It is laterally compressed in
cross section and shows no evidence of setae. The
articulation to the limb base is unknown, but this ramus
is interpreted as an exopod owing to its general similarity
to that of the fourth appendage. A small endopod may
have originally been present, and subsequently 'lost'
within the preservation gap between head and trunk.

There is a long, narrow, subrectangular hypostome
(length to width ratio 3.3 : 1) subdivided at a pointed
lateral shoulder just posterior to mid-length (figure 1e,k).
The anterior part is aligned subparallel to the dorsal ridge
on the carapace, and in transverse profile has a pitched,
roof-like form with a long, apically rounded median ridge.
The anterior border appears to be subrhomboidal in form
and is attached to the carapace border as part of the
carapace notch, and there is a small subtriangular anterior
wing. The posterior part of the hypostome is flexed
ventrally at approximately 15°; the median ridge on this
part is less pronounced. The posterior margin of the
hypostome is gently convex.

Sections of the alimentary canal are preserved as a
sedimentary infill (figures 1c,dl,ij,l and 2a,l,m). The
presumed mouth lies dorsal to the posterior hypostomal
margin. It opens into an anteriorly expanding tube-like
section (buccal cavity/anterior section of pharynx) above
the posterior section of the hypostome, followed by a very
short constriction that is triangular in cross section (apex
dorsal) above the juncture of the anterior and posterior
parts of the hypostome (interpretation uncertain).
Anteriorly, the canal widens into a short section compris-
ing six longitudinal ridges and furrows, which may
represent (? pharyngeal) muscle; alternate ridges (ventral
and dorsolateral) are broader. Further anteriorly, it
becomes a simple tube again (presumed oesophagus),
subcircular in cross section, and bends dorsally and
sharply backwards at the front of the head. Three traces
of the intestine are present in the trunk region; the most
anterior shows constrictions; the tiny, most posterior trace
constrains the position of the anus to near the end of the
trunk where it was probably terminal.

(c) Trunk
The trunk is narrow, tapers very gradually posteriorly and
bears at least 32 biramous appendages; the most posterior
are not fully resolved and there are probably approxi-
mately 35 in total (figures 1d,lf,jl and 2a–c). The proximal
part of each limb is small, the preservation poor and the
detailed form is obscured. The endopods are imbricated
and collectively form a cone-like basket that in outline
flares slightly outwards anteriorly. The shaft of each
endopod is broad and ribbon-like. One or two smaller
endites occur on the most proximal part of the endopod
but details here are not well resolved. More distally, there
are four large, flat, paddle-like endites and the shaft
terminates in a similar fifth process. The exopods are
confined within the domed part of the carapace. Those of
the more anterior appendages extend outwards and
slightly downwards from their attachment, before curving
posteroventrally; in the posterior part of the trunk, they
become progressively more tightly flexed. The main shaft
of the exopod is narrow and dorsoventrally flattened
(podomeres cannot be discerned). On its posterior
margin, it bears very fine, straight, discrete filaments that
trend posteriory and overlap those of the preceding
exopod. At mid-trunk position, there are approximately
25 such filaments per exopod. They are longest at about
one-third from the base of the main shaft; distally from
here they gradually shorten to nothing and proximally they
shorten to approximately one-quarter their maximum
length. The filaments appear as dots at the limit of
resolution in section images; they are likely to be
subcircular to oval in cross section. A minute section of
the trunk extends posteriorly beyond the last appendage
pair, but it is insufficiently resolved for discrimination of
possible segment boundaries or telson.

4. DISCUSSION
(a) Affinities
A cladistic analysis was performed using the arthropod
morphological character matrix of Wills et al. (1995,
1998), with the addition of Xylokorys and the inclusion of
a reinterpreted appendage morphology of Vachonisia (see
below and electronic supplementary material). Xylokorys
falls in Marrellomorpha, forming a basal sister taxon to a
(Vachonisia (Marrella + Mimetaster)) clade, in 35% of the
most parsimonious trees. The marrellomorphs resolve as
stem-group crustaceans in 87% of the most parsimonious
trees. Although our analysis supports the association of
Xylokorys with Marrellomorpha, we treat the recovered
position within the marrellomorphs and the relationship of
this clade to other arthropod groups as merely indicative:
other studies of the marrellomorphs are pending and a
strictly morphological comparison suggests a close
relationship between Xylokorys and Vachonisia (see
below). An identical stem-group crustacean position for
marrellomorphs was recovered in a small subset of the
trees obtained by Wills et al. (1998), but marrellomorphs
mainly resolved within the schizoramian (archanomorph
and crustacean) stem in their analysis (see also Cotton &
Braddy 2004). Many other positions for Marrellomorpha
have been argued: an isolated position within Arachno-
morpha (Stürmer & Bergström 1976); a basal position
within Lamellipedia (Hou & Bergström 1997) or within a
resurrected Trilobitomorpha (Bergström & Hou 2003);
and as a stem group to other schizoromorphs (archan-
morphs and parvancorinomorphs, the latter possibly
including Vachonisia; Lin et al. 2006).

A rigorous reassessment of the relationships of the
marrellomorphs is beyond the scope of this paper. Even
though the Wills et al. (1998) study was a landmark in
using morphological data from fossils and living taxa, it
has been somewhat superseded by recent molecular and
total evidence approaches (e.g. Giribet et al. 2001;
Mallat & Giribet 2006) and new fossil discoveries. New
data on Marrella from the Burgess Shale (Garcia-Bellido &
Collins 2006), Vachonisia from the Hunsrück Shale
(Oltmann et al. in preparation) and Furca from the

Orдовик Morocco and Bohemia (Van Roy 2006a,b; see also Chlupáč 1999 and Bergström & Hou 2003), will need to be integrated into a new assessment. *Marrella* and *Mimetaster* have always been regarded as closely related (e.g. Stürmer & Bergström 1976; Wills et al. 1995, 1998) and the carapace of *Furca* is very similar to that of *Marrella* (e.g. Chlupáč 1999). The relationship of *Vachonisia* to this clade has proved more problematic, and several authors have regarded its marrellomorph affinities as at best uncertain (e.g. Stürmer & Bergström 1976; Hou & Bergström 1997; Lin et al. 2006; Van Roy 2006a,b). Cladistic analyses that placed *Vachonisia* together with *Marrella* and *Mimetaster* (Wills et al. 1995, 1998) used data matrices of morphological characters largely without ordering or weighting. This may place undue emphasis on the large number of trunk appendages that decrease in size posteriorly, a condition that may be plesiomorphic. The cephalic appendages of *Xylokorys* and *Vachonisia*, on the other hand, are very different from those of *Marrella* and *Mimetaster*. *Xylokorys* and *Vachonisia* are also united by the large carapace that covers body and appendages.

Stürmer & Bergström’s (1976, figure 10) investigation of the head region of *Vachonisia* identified a pair of uniramous antennae followed by three pairs of uniramous appendages that lack exopods, the posterior two bearing strong endites with a spinose tip. The morphology of *Xylokorys* suggests that the three post-antennal uniramous appendages reconstructed by Stürmer & Bergström (1976, figure 10) in *Vachonisia* are a composite of the exopods and endopods of appendages two to four (Oltmann et al. in preparation). The endites reconstructed by Stürmer & Bergström are distal projections of the endopods of appendages two and three (see also Bartels et al. 1998, figure 99).

*Xylokorys* differs from *Vachonisia* in the following ways: in lacking a median rostrum within the anterior carapace notch; in the lack of postolateral projections of the carapace; in the relatively narrower carapace border; in the longer antenna that is divided distally; in the presence of fewer (from approx. 35 as opposed to approx. 80) trunk appendages (though note that the number of appendages in *Vachonisia* was estimated in the holotype (see Lehmann 1955; Stürmer & Bergström 1976), which is approximately 1.7 times as long as OUMNH C.29604); and in its strongly geniculate endopods of appendages two and three may have raked the sediment and brought food into suspension. The trunk appendages may have captured edible particles on the endites from suspension and transported them anteriorly to the mouth. The proximal parts of the head appendages are robust, arranged in overlapping fashion, and project towards the mouth, where they could have been involved in the mechanical breakdown of food. The sharply edged, flange-like extension of the fourth appendage may have served as an incisor. *Xylokorys* does not appear to have ingested large items of food: the lack of space between carapace and oesophagus (figures 1bc and 2) precludes an expanded stomach. Thus, *Xylokorys* is interpreted as essentially a benthic particle filter feeder, which may also have consumed prey items (the feeding habits of the living branchiopod *Triops* may be analogous; see Brusca & Brusca 2003).

The Leverhulme Trust (F/08581/E) and English Nature are thanked for their support; K. Saunders and N. Francis for technical assistance; R. Fenn, T. Hall and J. Sinclair for their general assistance, and G. Oltmann, J. Rust and J. Bergström for discussion of *Vachonisia*.

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