

## Article

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
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# Ultrastructural evidence shows adaptation to a pelagic lifestyle in Ordovician caryocaridids (Crustacea: phyllocarida)

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**Non-technical Summary**

Caryocaridids are a unique representative of the pelagic arthropod group during the Ordovician and stand out from other arthropods (ostracods, trilobites, etc.) for their remarkable pelagic abilities. Herein, we report on a species of caryocaridids, *Somicaris cedarbergensis*, from the Lower Ordovician in northwestern Xinjiang, NW China, which shows the rare enrolled carapaces with the evidence of cuticular ultrastructure preserved. These caryocaridid specimens from Xinjiang provides the substantial evidence for the presence of caryocaridids in the Central Asian Orogenic Belt. This discovery suggests that *S. cedarbergensis* appeared as early as the Early Ordovician (late Tremadocian) and persisted until the end-Ordovician (Hirnantian) and is the longest-ranging species of known caryocaridids. The cuticle of the carapace in *S. cedarbergensis* is preserved in carbonate-fluorapatite, which can be divided into three mineralized lamellae (outer, middle, and inner). The outer and inner lamellae both consist of three layers, which seem to correspond to the epicuticle, exocuticle, and endocuticle of extant crustacean carapaces, respectively. The particular ultrastructure of the carapace of Ordovician caryocaridids (thin cuticle; thickened inner lamella cuticle; and large, complex oxygen supply system) probably represents an adaptation to the pelagic lifestyle during the Ordovician plankton revolution.

**Abstract**

Caryocaridids are a unique representative of pelagic arthropods from the Ordovician period. They are typically found as flattened carapaces in mudstones and shales. This study reports on a species of caryocaridids, *Somicaris cedarbergensis*, discovered in the Lower Ordovician of northwestern Xinjiang, NW China. The species shows the rare enrolled carapaces with a preserved cuticular ultrastructure. These specimens of caryocaridids from Xinjiang are the first reported in the Yili Block, and provide the substantial evidence that the paleogeographic distribution of caryocaridid phyllocarids could extend to the Central Asian Orogenic Belt. This species existed from the late Tremadocian until the end of the Ordovician (Hirnantian), making it the longest-ranging known species of caryocaridids. The carapace cuticle of *S. cedarbergensis* is composed of carbonate-fluorapatite and can be divided into three mineralized lamellae: outer, middle, and inner. The outer and inner lamellae each consist of three layers that correspond to the epicuticle, exocuticle, and endocuticle of extant crustacean carapaces. Moreover, the polygonal reticulation structure of the carapace in archaeostracans appears to be similar in shape and size to the hemolymph sinuses of leptostracans. This unique ultrastructure of the carapace cuticle in caryocaridids is believed to be better suited for a pelagic lifestyle.

**Introduction**

Phyllocarids are an important group of Malacostraca known for their morphological diversity, prolonged fossil records, and abundant extant relatives (Rolfe 1969, 1981; Schram and Hof 1998; Spears and Abele 1999). However, phyllocarids have received less attention compared with other contemporaneous taxa of the phylum Arthropoda due to their relative rarity and generally poor preservation, which make it difficult to infer their habitat, size, and body structure (Sepkoski 2000). Crustacean cuticles exhibit varying levels of calcification that can differ even among different parts of the carapace of the same organism (Vega et al. 2005). Additionally, diagenetic processes and taphonomy have the potential to alter or even destroy evidence of the original cuticle of crustacean carapaces. Therefore, there have been few studies focused on the ultrastructure and composition of the cuticle of phyllocarids in comparison to ostracods. The known cases are mainly centered on caryocaridids and ceratiocaridids (Rolfe

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1962; Churkin 1966; Miranda 2002; Vannier *et al.* 2003; Whittle *et al.* 2007; Liu *et al.* 2023; Šilinger 2023; Kovář *et al.* 2024).

Most authors have acknowledged the pelagic lifestyle of caryocaridids (e.g., Whittle *et al.* 2007; Collette and Hagadorn 2010; Liu *et al.* 2022, 2023). Ruedemann (1934) suggested that the caryocaridids have a pelagic lifestyle based on their wide paleogeographic distribution and the relative longevity of the genus. Since then, additional scholars have attempted to demonstrate the pelagic habitat of caryocaridids based on accumulated evidence. This includes their cosmopolitan distribution, carapace morphology, abdominal morphology, and association with the shale facies (Størmer 1937; Chlupáč 2003; Vannier *et al.* 2003). However, according to Bulman (1964), the large and heavy carapace of caryocaridids prevents them from living in a pelagic environment. Additionally, caryocaridids often coexist with benthos in certain assemblages. Chlupáč (1970) also suggested that caryocaridids are not strictly part of the plankton and likely attach to floating algal thali near the surface of water. However, Churkin (1966) discovered that the carapace valves of the caryocaridids were relatively large but extremely thin. And no evidence of algal mats has been found in conjunction with caryocaridid remains (Chlupáč 1970, 2003). According to Vannier *et al.* (1997a), the streamlined carapace and flattened furcal rami of caryocaridids are assumed to be adaptation to their active, free-swimming lifestyle.

In extant crustaceans, the cuticular structure of the carapace appears to be closely related to lifestyle (Sohn and Kornicker 1988; Pütz and Buchholz 1991; Yamada 2019). Unfortunately, since Churkin (1966) first studied the cuticular ultrastructure and mineralogical composition of the caryocaridid carapace, no further research has not been conducted, except for Vannier *et al.* (2003), who related the cuticular ultrastructure to the caryocaridid lifestyle. Other authors have provided brief descriptions of the cuticular ultrastructure and mineralogical composition of the carapace in caryocaridids (Miranda 2002; Vannier *et al.* 2003; Whittle *et al.* 2007; Liu *et al.* 2023; Šilinger 2023; Kovář *et al.* 2024).

In recent years, we have investigated the Cambrian–Ordovician transitional strata in the Guozigou area of Xinjiang, NW China. We collected a large number of the Lower Ordovician caryocaridid specimens from the original or nearby sections reported in previous studies (Xiang and Zhang 1984; Zhang 1987). These materials retain the rare enrolled carapace and cuticular ultrastructure of the carapace due to unique taphonomic conditions. We discuss here the carapace characteristics of caryocaridids in three preservation states. Additionally, we studied and evaluated the cuticular ultrastructure of the carapace and provide further evidence that caryocaridids are pelagic arthropods.

## Materials and Methods

The western part of the North Tianshan Mountains in Xinjiang contains relatively continuous Ordovician strata. The most-studied area is in Guozigou (Fig. 1A,B). The Ordovician strata in the Guozigou area are divided into the Guozigou Formation ( $O_1g$ ), the Xinertai Formation ( $O_{1-2x}$ ), the Kekesaleixi Formation ( $O_3kk$ ), the Aketashi Formation ( $O_3a$ ), and the Hudukedaban Formation ( $O_3h$ ) from the bottom to the top (BGXUAR 1987c; XBGMR 1993, 1999; Li 1995; Chen *et al.* 1998). The Xinertai Formation is a collection of darkish-gray and gray siliceous, argillaceous, and carbonaceous siltstones, siliceous rocks, and sandy mudstones, with a few black shales and thin-bedded limestones (XBGMR 1999). It was most likely

deposited in a shelf to slope paleoenvironment (Li 1995; XBGMR 1999). This formation yields a variety of invertebrates, including phyllocarids, graptolites, trilobites, and brachiopods (e.g., Xu and Huang 1979; BGXUAR 1987c; Zhang 1987; Qiao 1989; XBGMR 1993, 1999).

Xiang and Zhang (1984) were the first to report crustacean fossils, probably caryocaridid carapaces, in the Ordovician strata from the Guozigou area. Since then, roughly determined *Caryocaris* sp. has frequently been encountered in the Jiangkunusi, Guozigou, and Kegouqinshan areas in later geological surveys or studies of other fossils (Zhang 1987; Qiao 1989; XBGMR 1993; Zhang 2010; Fig. 1B). However, no systematic description or detailed analysis of caryocaridids has been provided. The phyllocarid materials described in this paper were collected from two stratigraphic sections in the Guozigou area (Fig. 1B,C). Section 1 ( $44^{\circ}28'47.5''N$ ,  $81^{\circ}08'35.8''E$ ) is located on the ridge between the Linkuanggou and Fenggou, where a large number of well-preserved caryocaridid fossils were found in the siliceous siltstones about 1.5 m above the bottom of Bed 6. Section 2 ( $44^{\circ}28'11.4''N$ ,  $81^{\circ}08'4.9''E$ ) is located west of a new highway, approximately 500 m north of the Linkuanggou entrance. Three beds of caryocaridid fossils were collected from the lower part of Bed 10, and the fossils are poorly preserved and scarce (Fig. 1D). The caryocaridid-bearing levels from the Guozigou area cross three graptolite biozones, including the *Adelograptus-Clonograptus* (late Tremadocian, *Adelograptus tenellus* and *Aorograptus victoriae*), *Tetragraptus approximatus* (early Floian), and the *Tshallograptus fruticosus* (early Floian) Biozones, as suggested by the graptolites found in the same bed (Xu and Huang 1979; Qiao 1989; Chen *et al.* 1998; XBGMR 1999; Zhang *et al.* 2004, 2005; Li *et al.* 2018). Our caryocaridid specimens were mainly found in the *Adelograptus-Clonograptus* and *Tetragraptus approximatus* graptolite biozones within the lower Xinertai Formation within the two sections of the Guozigou area. A total of 104 caryocaridid specimens were collected and are currently housed in the State Key Laboratory of Biogeology and Environmental Geology (BGEG) at China University of Geosciences in Wuhan.

The caryocaridid specimens were photographed using an APS-C DSLR camera and illuminated with a microscope cold-light halogen lamp. Images of enrolled specimens were taken using field emission scanning electron microscopes (Hitachi SU 8010) at the State Key Laboratory of Biogeology and Environmental Geology, China University of Geosciences, and Zeiss Axio Zoom V16 at the Nanjing Institute of Geology and Paleontology, Chinese Academy of Sciences. Microzone chemical composition analysis and wavelength dispersive element scanning of fossils were conducted using energy dispersive spectroscopy (EDS; Hitachi SU 8010). Additionally, light photomicrographs of the thin sections were taken using a Zeiss Axioscope A1 polarizing microscope at the China University of Geosciences.

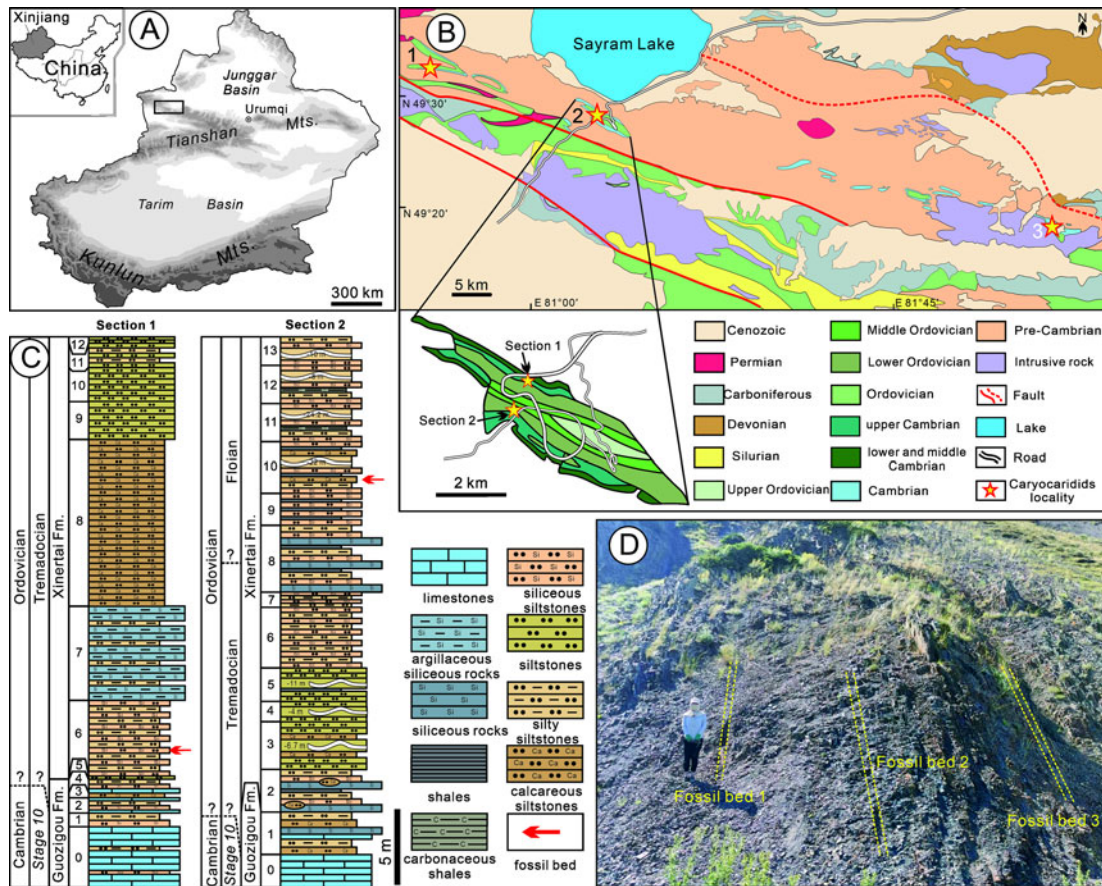
## Morphological Terminology Used

**Lamella/lamellae**, a general expression for a layered appearance of anything; **lamina/laminae**, the fine striations constituting a special structure in the cuticle composed of chitin–protein fibers.

## Results

### Description of Caryocaridids

Out of the caryocaridid specimens collected from the two sections, only 56 could be identified, including 52 incomplete



**Figure 1.** A, Location of North Yili in North Tianshan, Xinjiang, NW China. B, Geologic map of North Yili shown in black box in A (modified from BGXUAR 1987a,b,c): 1, Jiangkunusi; 2, Guozigou; 3, Keguqinshan. C, Stratigraphic column of the Xinertai Formation in the Guozigou area (modified after Zhang 1987; Qiao 1989). D, The outcrop of three fossil beds in Bed 10 of Section 2 (geologist is 165 cm tall).

carapaces, 2 telsons, and 2 furcal rami. The carapace valves of caryocaridids exhibit irregular longitudinal and oblique folds and lines due to the compression deformation. Nearly 40% of them are enrolled. The remaining specimens are highly fragmented and could not be identified.

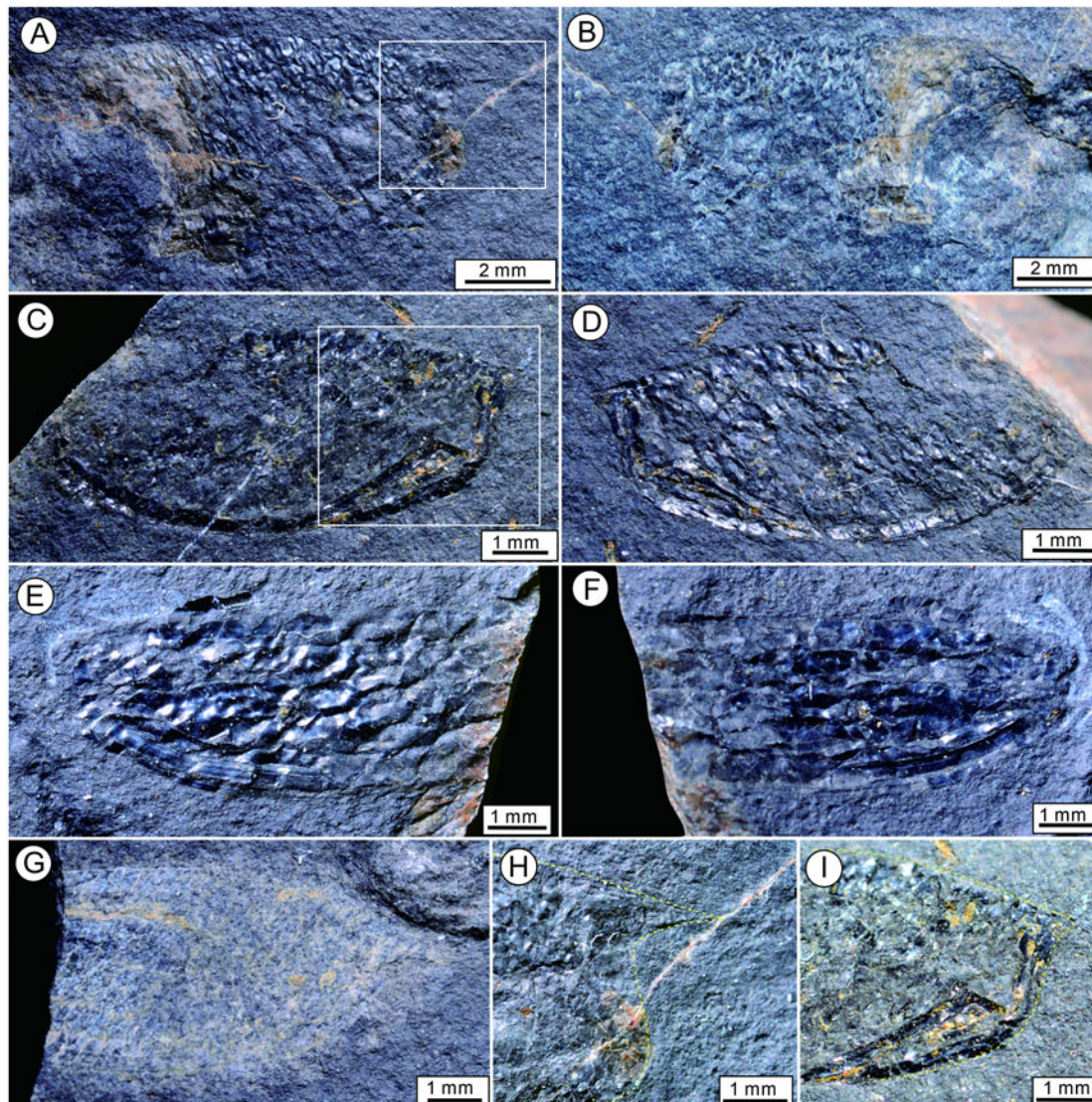
**Detailed Description.** The pod-shaped, bivalved carapace has a length to height ratio of 2.5. The highest point of the carapace is located in the center of the carapace. Dorsal and ventral margins are moderately convex (Figs. 2A–G, 3A,B). The anterior horn of the carapace is sharply pointed and measures 1.5–1.8 mm in length, pointing forward (Fig. 2A,B,G,H). A clear and pronounced ventral doublure with a moderate convexity and constant width of 0.4 mm runs along the ventral border (Figs. 2C–F, 3A,B). The posterior margin of carapace is curved and lacks spinules and a posteroventral spine. The posterodorsal spine is short and pointed (Figs. 2C,D,I, and 3A). The subtriangular telson is dorsoventrally flattened and has a pointed extremity (Fig. 3C,D). The maximum length of the telson is 5.0 mm, with a corresponding width of 1.2 mm near its proximal end. The furcal rami are leaf-shaped, pointed, and longer than the telson. Their outer margins lack spinelike triangular expansion and articulated spines. The maximum width of the furcal rami is located just anterior to its midlength (Fig. 3E,F). A furrow, which runs from almost the base of the spinelike expansion to the axis of the terminal posterior spine, is clearly visible (Fig. 3F). These specimens from the Guozigou area are identified as *Soomicaris cedarbergensis*

(Whittle et al. 2007) based on the carapace outline, which features a long pointed anterior horn and a short pointed posterodorsal spine (Whittle et al. 2007; Racheboeuf and Crasquin 2010).

**Geographic Distribution of Soomicaris**

Caryocaridids are a well-known representative of the pelagic arthropod group during the Ordovician period. Their paleogeographic distribution became global from the Dapingian to Sandbian (Liu et al. 2022). Before the recent description of caryocaridids from the North China Craton (Liu et al. 2023), these arthropods were mainly found in Gondwana, Laurentia, Baltica, Avalonia, Bohemia, and South China (e.g., Chapman 1902, 1934; Woodward 1912; Chlupáč 1970, 2003; Jell 1980; Shen 1986; Racheboeuf et al. 2000, 2009; Whittle et al. 2007; Racheboeuf and Crasquin 2010). *Soomicaris* comprises three known species: *S. cedarbergensis*, *S. ordosensis*, and *S. scanicus*, which are respectively distributed across Clanwilliam in South Africa (late Hirnantian), Inner Mongolia in China (early Sandbian), and Scania in Sweden (late Tremadocian) (Möberg and Segerberg 1906; Whittle et al. 2007; Liu et al. 2023). Freiberger (1947) documented the carapaces of *Caryocaris curvilata* in the Karatau, Kazakhstan. And then, Obut and Zuvtsov (1965) and Abdullaev and Khaletskaia (1970) reported the *Caryocaris baidamtalensis* and *Caryocaris* sp. from the Ordovician strata of Kyrgyzstan and Uzbekistan respectively.





**Figure 2.** The carapaces of *Soomicariscedarbergensis* from the Lower Ordovician Xinertai Formation, Xinjiang, NW China. **A, B**, Incomplete right carapace, BGEG-SJ-01a,b. **C, D**, Incomplete left carapace, BGEG-SJ-03a,b. **E, F**, Incomplete right carapace, BGEG-SJ-06a,b. **G**, Incomplete right carapace, BGEG-SJ-02a. **H**, Enlargement of the white box in **A**, showing the anterior horn; **I**, Enlargement of the white box in **C**, showing the posterodorsal spine.

Tolmacheva *et al.* (2004, 2008) and Nikitina *et al.* (2006) also mentioned the occurrence of some carapace and telson fragments of caryocaridids or phyllocarids in the western Balkhash region and the Kol'denen River area of Kazakhstan. However, these specimens are relatively fragmented and the illustrations are poor or even missing, to the extent that further species identification is not feasible. Undoubtedly, these evidence seems to support the notion that the distribution of caryocaridids could potentially spread to the Central Asian Orogenic Belt (Tolmacheva *et al.* 2021). Therefore, the discovery of *S. cedarbergensis* in the Lower Ordovician of northwestern Xinjiang not only provides more substantial evidence for the presence of caryocaridids in the Central Asian Orogenic Belt but also further extends their distribution range within this geological area.

*Soomicariscedarbergensis*, found in the Xinertai Formation in North Tianshan, is similar to the other two species of this genus co-occurring with graptolites and thin-shelled brachiopods at the same horizon (Möberg and Segerberg 1906; Liu *et al.* 2023). The

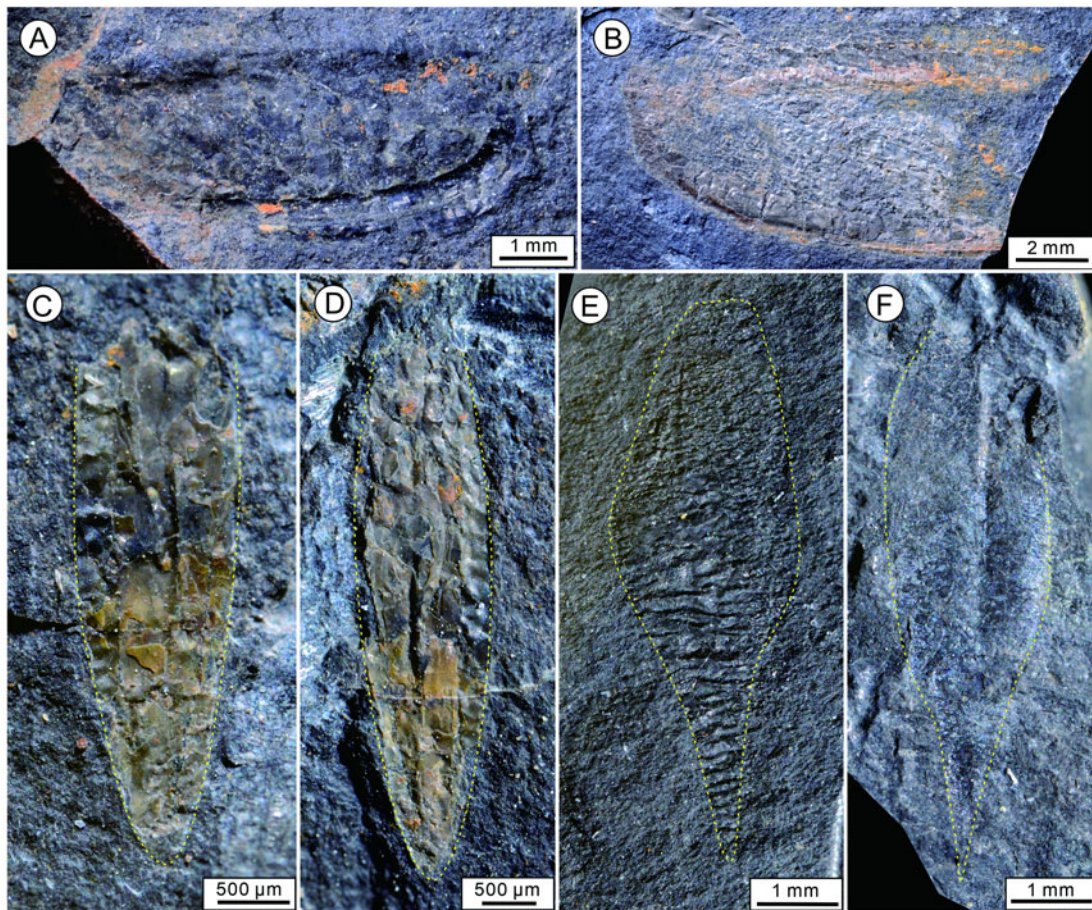
graptolites in the same bed suggest that *S. cedarbergensis* from the Guozigou area of North Tianshan, Xinjiang, is from the late Tremadocian to early Floian, occupying three graptolite biozones. Thus, *S. cedarbergensis* appeared as early as the late Tremadocian and persisted until the end of the Ordovician (Hirnantian), making it the longest-ranging species of known caryocaridids.

## Discussion

### *Taphonomic Types of Carapaces*

The carapaces of the reported specimens of caryocaridids exhibit varying degrees of compression deformation, which can be classified into three types: enrollment, wrinkles, and flattening (e.g., Churkin 1966; Chlupáč 1970; Rushton and Williams 1996; Vannier *et al.* 2003; Fig. 4). Generally, most caryocaridid carapaces are flat in shales, siltstones, and/or siliceous rocks, but complete flattening of carapaces is not common due to their rigid





**Figure 3.** The carapaces, telsons and furcal rami of *Soomicaris cedarbergensis* from the Lower Ordovician Xinertai Formation, Xinjiang, NW China. **A**, Incomplete left carapaces, BGEG-SJ-04. **B**, Incomplete right carapace, BGEG-SJ-10. **C**, **D**, The well-preserved telsons, BGEG-SJ-09a,b. **E**, **F**, Incomplete furcal rami; **E**, BGEG-X-C6, **F**, BGEG-S-C9.

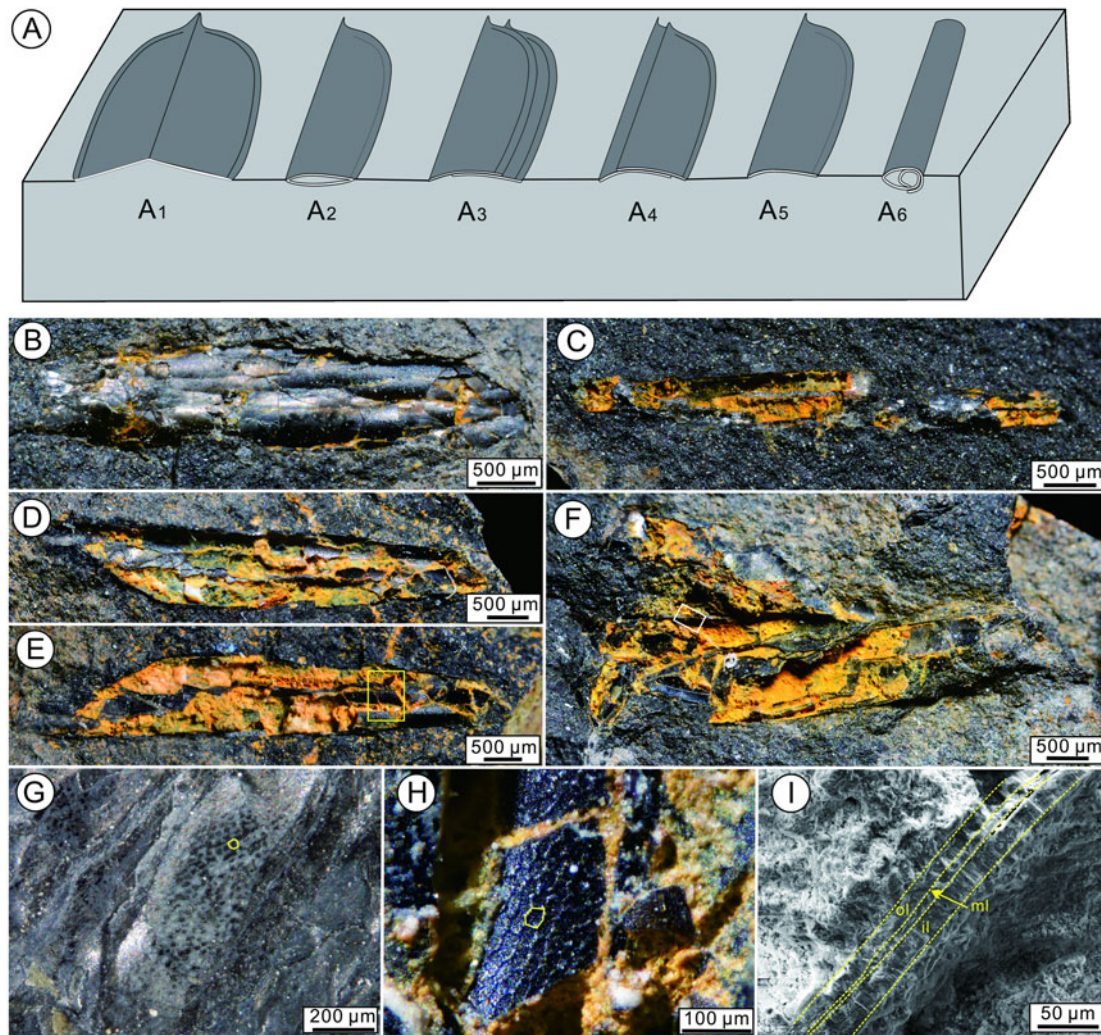
strength (Chlupáč 1970). If the two carapace valves remain jointed after burial, they are eventually preserved in a butterfly shape under vertical burial (e.g., Shen 1986: figs. 3, 4; Wang et al. 2019: fig. 3g; Liu et al. 2023: fig. 2j; Fig. 4A<sub>1</sub>) and in the overlapping form under lateral burial (e.g., Vannier et al. 2003: fig. 5E; Fig. 4A<sub>2</sub>). The two jointed carapace valves may be slightly separated due to bottom currents or bioturbation (e.g., Copeland 1967: plate 162, figs. 1, 2; Jell 1980: fig. 5A; Liu et al. 2023: fig. 2G,M), then compression may result in the impression of the harder border of one carapace into the valve of the opposite carapace, so that four or two parallel grooves following the ventral or dorsal margin of the carapace may be observed (e.g., Chlupáč 1970: plate I, figs. 1, 3, 5; Racheboeuf et al. 2000: fig. 7C; Fig. 4A<sub>3</sub>, A<sub>4</sub>). If the two carapace valves are far apart, they are preserved in a flat, single valve form (e.g., Whittle et al. 2007: fig. 3A, B; Fig. 4A<sub>5</sub>). When the compression deformation is minimal, the carapace valves of caryocaridids show irregular folding and warping, resulting in irregular longitudinal and oblique folds and lines (e.g., Chapman 1934: plates 9, 10; Pillola et al. 2008, plate 1, fig. 4). These folds and lines often cause the carapace valves to break (Chlupáč 1970).

Enrollment is the rarest type, which fully demonstrates the 3D morphology of the caryocaridid carapaces. Before this paper, the only known 3D preserved specimens were *Caryocaris curvilata* from Alaska, North America (Churkin 1966), *Caryocaris wrightii*

from Bohemia (Chlupáč 1970) and the British Isles (Rushton and Williams 1996; Rushton and Ingham 2000; Vannier et al. 2003), and some unknown caryocaridid carapaces from Texas and Nevada, North America (T. A. Hegna personal communication 2024). Approximately 40% of the total number of caryocaridid specimens found in the Guozigou area of Xinjiang are enrolled. These carapace valves are enrolled transversely and are cylindrical, tapering slightly from the middle to the ends, resembling an elongated cigar (Fig. 4A<sub>6</sub>, B–F). All enrolled specimens were partially broken. The largest caryocaridid specimen measures about 5.8 mm in length, with a height of about 0.9 mm and a length-to-height ratio of about 6.5, compared with about 2.5 in associated flattened specimens. The thin black carapace valves, with a thickness of about 40–45 µm (Figs. 5J, 6A), are supported by white calcite or loose limonite deposited between each valve (Fig. 4B–F), or they may adhere to each other to form “thick” carapace valves of two to three layers. This type of carapace is identical to the carapaces described by Churkin (1966) and Chlupáč (1970) for caryocaridids.

The carapace valves are typically flattened and jointed. Before sediment compaction, the carapace valves may undergo partial enrolling and folding, such as folding at the posterior ventral margin of the carapace (Fig. 2I) or complete spiral enrolling (Fig. 4B–F). Chlupáč (1970) discovered that the carapace valves of caryocaridids can be enrolled with either the dorsal or ventral





**Figure 4.** Taphonomic types of caryocaridid carapace and enrolled carapace from the Xinertai Formation, Xinjiang, NW China. **A**, Preservational states of caryocaridid carapace on the surface of bedding plane. **A<sub>1</sub>**, Carapaces preserved in the “butterfly position”; **A<sub>2</sub>**, jointed carapaces buried on one side; **A<sub>3</sub>**, **A<sub>4</sub>**, slightly dislocated carapaces buried on one side; **A<sub>5</sub>**, single carapace buried on one side; **A<sub>6</sub>**, enrolled carapaces. **B–F**, Curled incomplete carapaces, BGEG–SJ–33, 34, 35a, b, 36. **G**, The polygonal reticulation of a carapace, BGEG–SJ–11b; **H**, Enlargement of the yellow box in **E**, showing the polygonal reticulation. **I**, Natural cross-section of the carapace of the white box in **F** showing the three mineralized lamellae. Abbreviations: il, inner lamella; ml, middle lamella; ol, outer lamella.

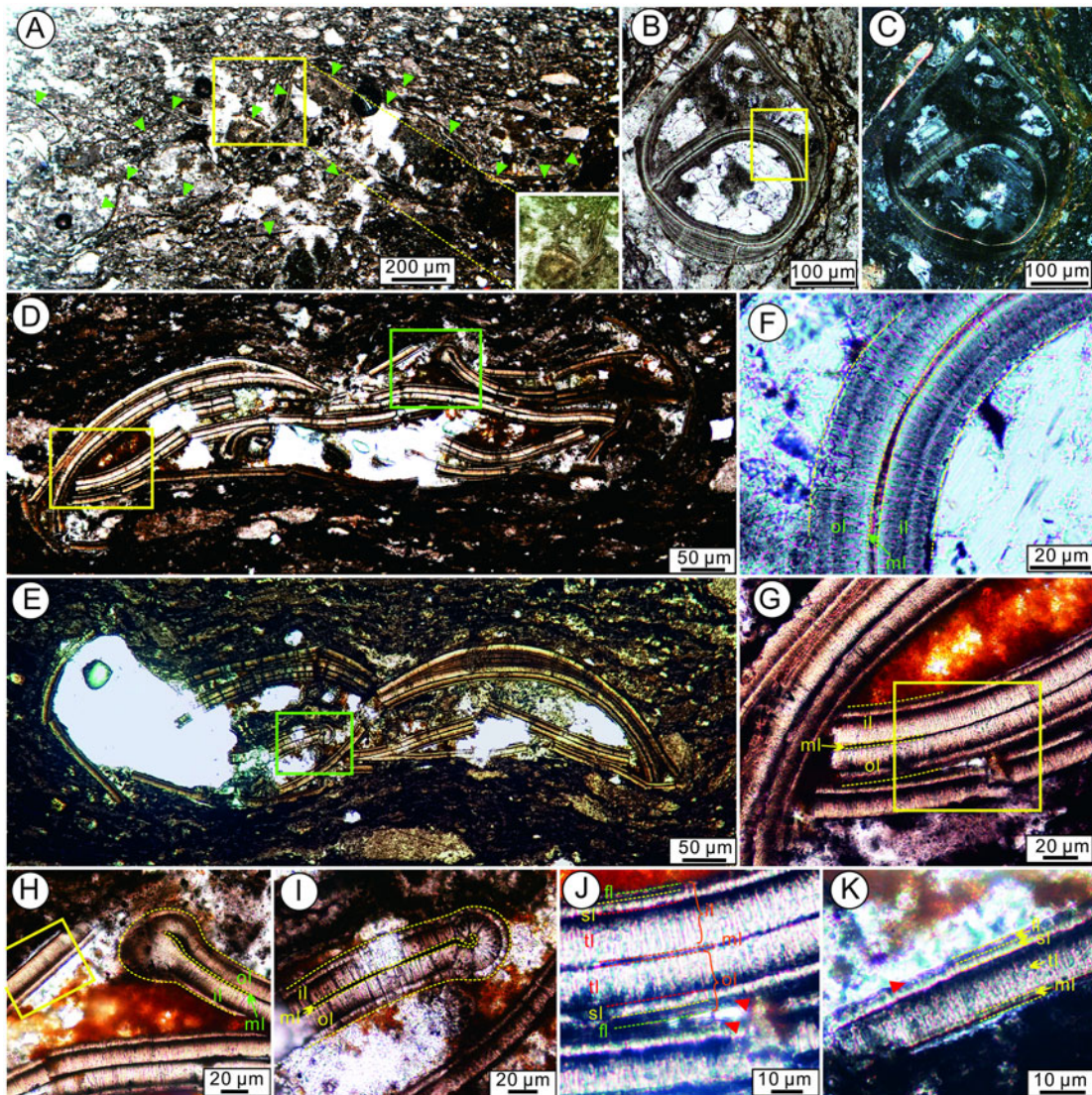
line as the axis, resulting in two types of enrollment: the ventral border is either inside the spiral with the dorsal line remaining outside, or the ventral border remains outside while the dorsal line is enrolled inside the spiral. However, these two types are challenging to observe and distinguish in actual specimens. In our study’s thin-sectioned specimens, the longitudinal section of the carapace valves provides a trace of evidence that the carapace begins to roll along the dorsal hinge line from the ventral margin (Fig. 5B,C), which belongs to the first enrolled type. Racheboeuf *et al.* (2000) proposed that the secondary enrollment of the carapace valves in caryocaridids may be linked to the mechanical properties and architecture of the exoskeleton. Specifically, the thin and flimsy carapace with a high L:H ratio is more susceptible to enrolling. Secondary enrolling of carapace valves typically occurs after the organism’s death or molting, but before compaction and fossilization (Chlupáč 1970). According to Racheboeuf *et al.* (2000), the exuviae or free carapaces are more susceptible to enrolling than the carapaces after natural death. This is because the soft parts of the carapaces in the latter are still attached to the

exoskeleton, which act as a mechanical hindrance to the process of enrolling.

#### *Cuticular Ultrastructure of the Carapace*

The carapace valves of caryocaridids typically exhibit distinct fine lines in thin sections (Fig. 5A). Some of these lines may be bent or flexed, and in some cases, the valves may even fracture along these lines (Fig. 5D,E). Churkin (1966) was the first to report on the ultrastructure of the carapace valves in caryocaridids, and he observed that the carapace was primarily composed of three distinct mineralized lamellae (Fig. 5), which resemble sandwiches. The outer (ol) and inner (il) lamellae are of similar or varying thicknesses and have a yellow-brown to orange-brown color under white polarized light (Fig. 5F–I). They also have weak fibrous structures running perpendicular to the carapace valves in some of the well-preserved specimens (Figs. 5F–I, 6A) and are separated by a black or dark-brown middle lamella (ml). The dense middle lamella can vary in thickness and lacks a





**Figure 5.** Light photomicrographs of the carapace of *Soomnicaris cedarbergensis* from the Xinertai Formation, Xinjiang, NW China. **A**, Light photomicrograph of siliceous siltstones from Bed 6 in Section 1, BGEG-TS-01-1, green triangles indicate the carapace fragments of *Soomnicaris*. **B**, **C**, Photomicrographs of a thin-sectioned specimen through an enrolled carapace, BGEG-TS-01-2. **D**, **E**, Random cross-section across a partially crushed specimen showing a complicated pattern of carapace valves folding, BGEG-TS-03, 04. **F**, **G**, Detailed view showing the characteristic three-layered ultrastructure of the carapace valves from the yellow boxes in **B** and **D**. **H**, **I**, Detailed view showing the closed ventral margin of carapace from the green boxes in **D** and **E**. **J**, **K**, Detailed view showing the seven-layered ultrastructure of carapace of the yellow box in **G** and **H**; red triangles indicate the laminae of the epicuticle of the carapace. Abbreviations: fl, first layer; il, inner lamella; ml, middle lamella; ol, outer lamella; sl, second layer; tl, third layer.

fibrous structure. It has a distinct boundary with the inner and outer lamellae (Figs. 4K, 5F–K, 6A). This is why some carapaces may have a slight lamination or parting. The ventral margin of the carapace features connected inner and outer lamellae that expand into a rounded cross-section, while the middle lamella does not penetrate them (Figs. 5H,I, and 7I–L). This corresponds to the ventral doublure at the ventral margin of the carapace. In the complete enrolled carapace, the sharp end of the carapace may represent the dorsal hinged line, and two connected left and right carapace valves are enrolled along their ventral margins (Figs. 5B,C, and 7A–D). The thickness of the enrolled carapace may vary significantly in different thin sections, and the carapace valves on either side of the supposed dorsal hinge line have variable and unequal thicknesses (Churkin 1966: plate 65, figs. 4–6; Miranda 2002: fig. 6.7,

6.10; Figs. 4B,C, and 6A–D). According to Churkin (1966), this phenomenon is due to the compositional division of the layered carapace valves, rather than an original difference between the two carapace valves.

In addition to the three mineralized lamellae observed in the carapace cuticle of the Xinjiang specimens, there is a microstratification structure consisting of three layers in the inner and outer lamellae of the carapace. The outermost first layer (fl) is yellow-brown and 2–3  $\mu\text{m}$  thick, with some weak horizontal laminae (Fig. 5J,K). The second layer (sl) is yellowish and slightly thicker than the outermost layer, measuring about 4  $\mu\text{m}$  (Fig. 5J,K). The thickest layer is the third layer (tl), which measures 11 to 13  $\mu\text{m}$ . It has a clear boundary between the second layer and middle lamella. The third layer is dark brown to black near the second layer and pale yellow near the middle lamella (Fig. 5J,K). Both

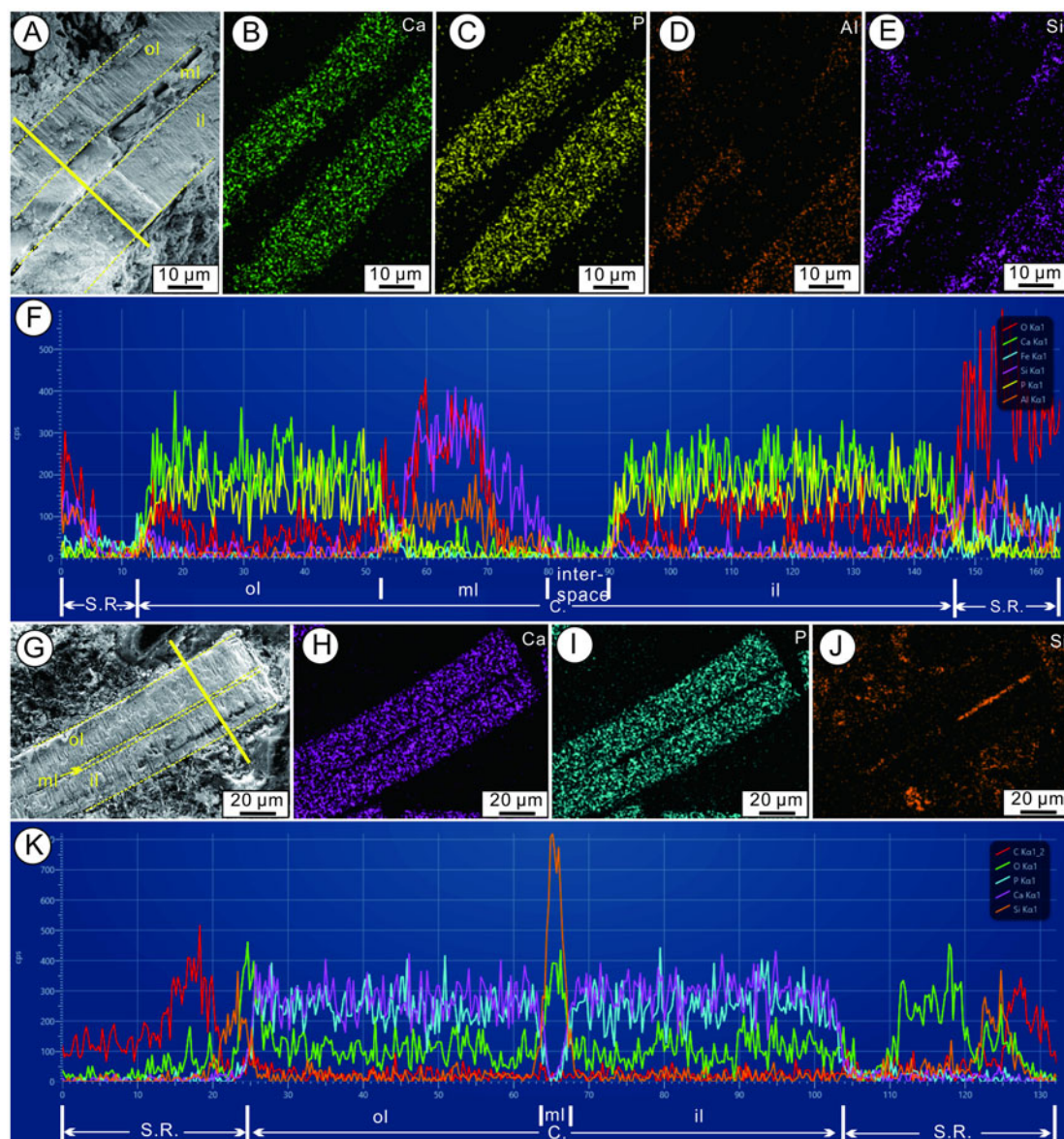


the second and third layers have weak fibrous structures that run perpendicular to the surface of the carapace valves.

The energy spectrum analysis for the natural cross-section and thin section of the carapaces in *S. cedarbergensis* reveals that the main elemental composition of the inner and outer lamellae is similar, as both are rich in calcium, phosphorus, and oxygen. However, the composition of the middle lamella is notably different from that of the inner and outer lamellae, with the main elements being silicon, aluminum, and oxygen (Figs. 6, 7). Therefore, the inner and outer lamellae may consist mainly of  $\text{CaO}$  and  $\text{P}_2\text{O}_5$ , while the middle lamella may consist mainly of  $\text{SiO}_2$  and  $\text{Al}_2\text{O}_3$ . This is consistent with the elemental composition of caryocaridids analyzed in previous studies (Churkin 1966; Miranda 2002; Vannier *et al.* 2003), suggesting that the

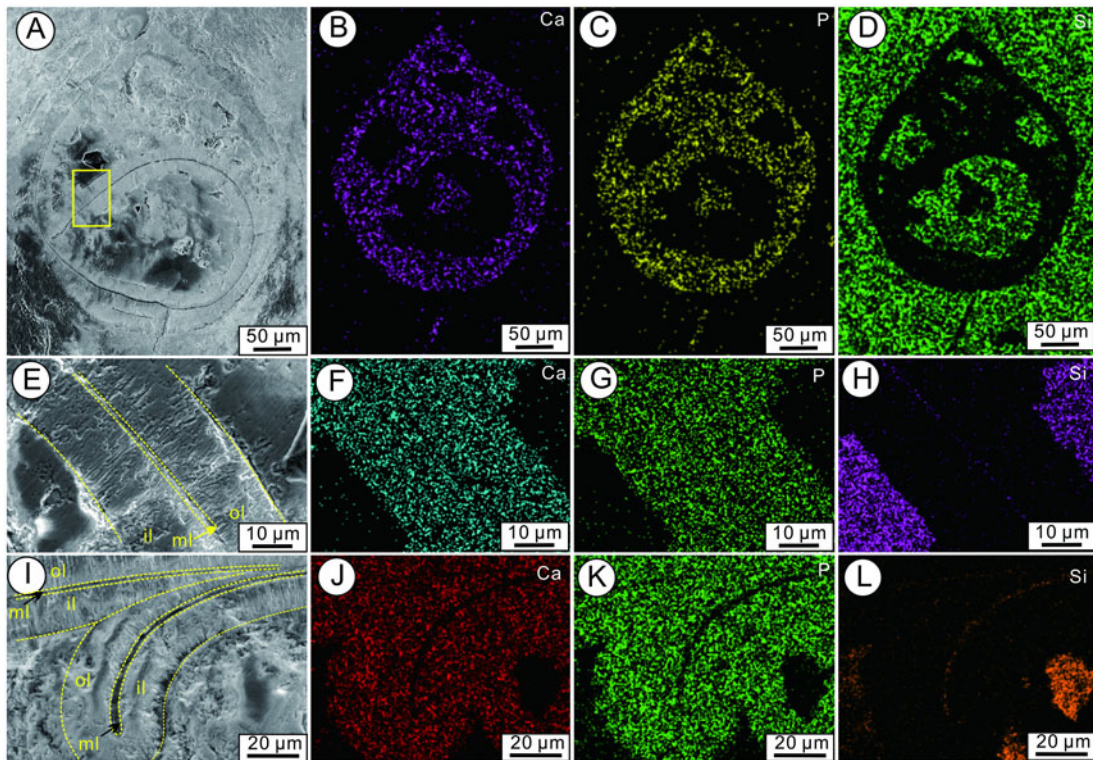
studied carapaces of caryocaridids may be composed of carbonate-fluorapatite (CFA). Miranda (2002) suggested that the presence of a small amount of silicon and aluminum may be due to clay pollution. However, we found that these elements are consistent with the composition of the middle lamella of the carapace. Therefore, the presence of these components indicates that the cuticle of caryocaridid carapaces contains not only a large amount of phosphorus and calcium, but also a small amount of other elements.

The cuticular structure of adult crustaceans typically consists of four layers: the epicuticle, exocuticle, endocuticle, and membranous layer (Dillaman *et al.* 2013). The epicuticle is the outermost and thinnest layer of the cuticle (Roer and Dillaman 1984), while the exocuticle, located beneath the epicuticle, is formed by



**Figure 6.** Elemental composition of the natural cross-section and thin-section carapaces by energy dispersive spectroscopy (EDS). **A**, The natural cross-section of a carapace showing the obvious three-layered ultrastructure. **B–E**, EDS maps of a natural cross-section of a carapace. **F**, EDS map of the carapace from the yellow line in **A** showing the variation in the abundance of different elements in the carapace. **G**, The thin section of a carapace showing the obvious three-layered ultrastructure. **H–J**, EDS maps of a thin section of a carapace. **K**, EDS map of the three-layered ultrastructure of the carapace along the yellow line in **G**, showing that the elemental composition of the inner and outer lamellae is obviously different from that of the middle lamella. Abbreviations: C., carapace; il, inner lamella; ml, middle lamella; ol, outer lamella; S.R., surrounding rock.





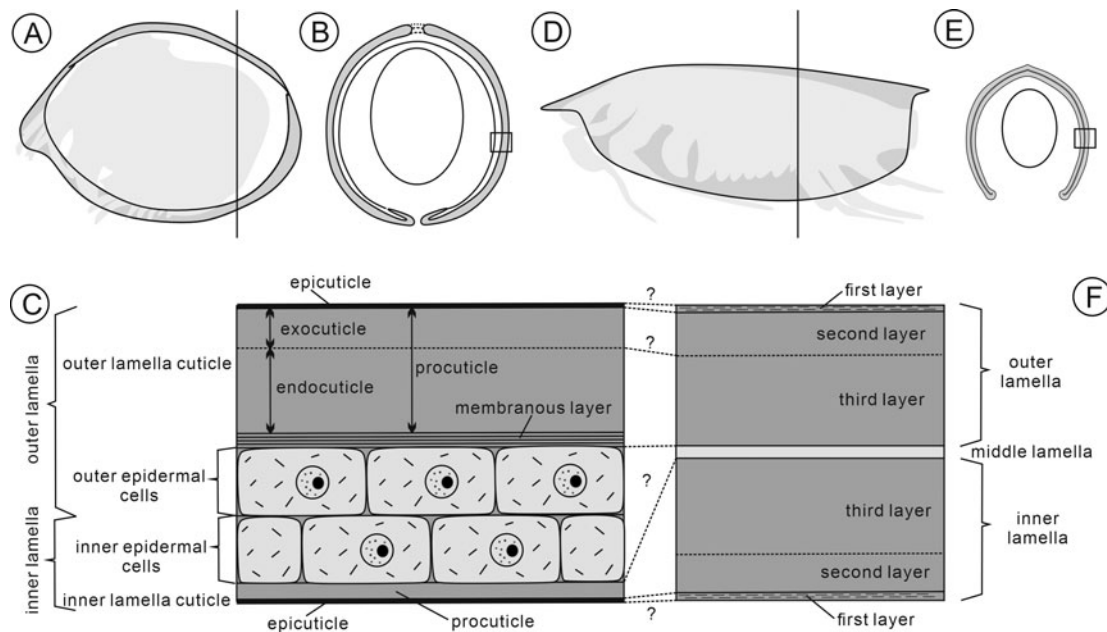
**Figure 7.** Elemental composition of the carapace by energy dispersive spectroscopy (EDS). **A–D**, EDS maps of a thin section through an enrolled carapace. **E–H**, EDS maps of the carapace showing the obvious three-layered ultrastructure of the carapace from the yellow box in **A**. **I–L**, EDS maps of the carapace showing the closed ventral margin of a carapace. Abbreviations: il, inner lamella; ml, middle lamella; ol, outer lamella.

stacking chitin–protein fibers (Green and Neff 1972). The endocuticle, the thickest and most calcified layer of the cuticle (Roer and Dillaman 1984), is located beneath the exocuticle. Vannier et al. (2003) suggested that the three mineralized lamellae of the carapace valves in caryocaridids may represent the typical three-lamella cuticular structure of living crustaceans. However, the thickness and ultrastructure of the three mineralized lamellae cannot be compared with the cuticles of living and fossil crustaceans (e.g., Pütz and Buchholz 1991; Haj and Feldmann 2002). In our study, the cuticular structure of the carapace valves in caryocaridids is mainly divided into three mineralized lamellae: inner, middle, and outer. The inner and outer lamellae can be further divided into three layers from the outside to the inside (Figs. 5J, 8D–E). Recent studies by Šlinger (2023) and Kovář et al. (2024) have also noted the two distinguishable layers of the caryocaridid cuticle: a white and translucent layer and a mostly opaque brown-black layer. However, the figures show two translucent layers with an opaque brown-black layer stuck on top of the second translucent layer, indicating a three-layer structure (Kovář et al. 2024: fig. 2Cb, plate 1, figs. 9, 10).

Wirkner and Richter (2013) presented the thin section of carapace of leptostracan phyllocarids for the first time in the study of crustacean circulation and respiration. However, they did not provide a detailed explanation or discussion of these findings. *Nebalia bipes* has a carapace cuticular structure consisting of inner and outer lamellae cuticles. The outer lamella cuticle is much thicker than the inner lamella cuticle. Epidermal cells and hemolymph sinuses are present between the two cuticles (Wirkner and Richter 2013: fig. 14.11C). However, our current understanding of the carapace cuticle in leptostracans is still limited. For instance, we have yet to study the ultrastructure and

function of the cuticle, as well as whether the inner cuticle will calcify in detail.

Extant ostracods have a complex carapace ultrastructure comprising the outer lamella (outer lamella cuticle and outer epidermal cells) and inner lamella (inner lamella cuticle and inner epidermal cells) (Bate and East 1972, 1975; Okada 1982a,b; Vannier and Abe 1995; Yamada et al. 2004; Yamada 2019; Fig. 8A–C). The outer lamella cuticle is significantly thicker and more calcified than the inner lamella cuticle. Calcification and thickening of the inner lamella cuticle have been observed in both living and fossil ostracod species (e.g., Olempska 2001, 2008; Yamada 2007; Yamada and Keyser 2010). However, Becker and Adamczak (1990) argued that the marginal thickenings found in fossil species are not true calcified inner lamella cuticle, but rather infold-like structures, similar to those of myodocopans. According to Horne et al. (2002), the infold structure is likely equivalent to the calcified inner lamella cuticle of ostracods. Yamada and Keyser (2010) also studied the calcification process of the inner lamella cuticle of living ostracod carapaces and found that the calcification ability of the outer epidermal cells extends to parts of the inner lamella cuticle in the adult stage, resulting in significant calcification of part of the inner lamella cuticle. Therefore, under certain conditions, the inner cuticle of some ostracod species will calcify, forming structures similar to the outer cuticle (Yamada and Keyser 2010). The ultrastructure of the carapace in caryocaridids appears to be comparable to that of ostracods (Fig. 8). The outer lamella cuticle of the ostracod carapace corresponds to the outer lamella of the caryocaridid carapace. Similarly, the inner lamella cuticle of ostracods is comparable to the inner lamella of caryocaridid carapace after the inner lamella cuticle has undergone obvious calcification. The



**Figure 8.** Schematic drawing of the carapace ultrastructures of ostracods and caryocaridids. **A**, Internal view of right carapace valve of ostracods. **B**, Transverse section of the black perpendicular lines in **A**. **C**, Carapace ultrastructure of the black box in **B** (modified after Yamada *et al.* 2004; Yamada 2019). **D**, Internal view of left carapace valve of caryocaridids. **E**, Transverse section of the black perpendicular lines in **D**. **F**, Carapace ultrastructure of the black box in **E**.

epidermal cells of ostracods appear to correspond to the middle lamella of the caryocaridid carapace, but there are significant differences in their thickness. Therefore, the cuticular ultrastructure of caryocaridid and ostracod carapaces is similar, although the inner lamella cuticle mineralization ability of caryocaridids is inferred to be much higher than that of ostracods. This high mineralization ability is necessary for caryocaridids to have their special cuticular ultrastructure (Fig. 8D,E).

Pütz and Buchholz (1991) examined the cuticle ultrastructure of some pelagic, nektobenthic, and benthic malacostracans. They suggested that certain cuticular characteristics correlate with the lifestyle. For instance, they claimed that the degree of cuticle mineralization and the thickness of the cuticle against body volume decrease from benthic to pelagic animals. Additionally, the thickness of laminae decreases, while the number of laminae in the cuticle increases from benthic to pelagic animals. Barrande (1872) and Rolfe (1962) recorded the cuticle thickness of other phyllocarid fossils. For example, the cuticle thickness of nektobenthic *Ceratiocaris papilio* and *Ceratiocaris scharyi* is 30–600  $\mu\text{m}$  and 250  $\mu\text{m}$ , respectively, and benthic *Aristozoe regina* is more than 1000  $\mu\text{m}$ . The cuticle thickness of caryocaridid carapaces is typically between 15 and 35  $\mu\text{m}$  (Churkin 1966; Miranda 2002; the present study; Figs. 5, 6). However, some species have carapaces as thin as 5–10  $\mu\text{m}$  (Vannier *et al.* 2003). Pelagic caryocaridids have a much thinner carapace cuticle compared with benthic *Aristozoe* and nektobenthic *Ceratiocaris*. This adaptation is reasonable, because caryocaridids with a light, thin carapace cuticle are better suited for a pelagic lifestyle. When the cuticle of the crustacean is deposited, the chitin–protein fibers change their direction in a helicoidal sequence, resulting in a lamellar appearance in cross-section (Bouligand 1972; Roer *et al.* 2015). However, caryocaridid carapaces lack laminae in their natural cuticle section (Fig. 6A); instead, a weak fibrous structure exists perpendicular to the carapace valves. Vannier *et al.* (2001: fig. 10.1) found similar patterns in the cuticle of the Devonian

ostracod carapace. They claimed that the original lamellar structure of the cuticle is often destroyed and micritized, and then a similar radial ultrastructure is produced. Although caryocaridids lack a clear lamellar structure in their cuticle, similar laminae have been found in the cuticles of their closely related nektobenthic ceratiocaridids (Rolfe 1962). In Paleozoic oceans, thylacoccephalans, a mysterious group of pelagic, bivalved arthropods, coexisted with phyllocarids (e.g., Zatoń *et al.* 2014). Their carapaces also have a laminated ultrastructure with elongated tubular structures piercing through and slightly disturbing the cuticle laminae (Broda *et al.* 2015; Broda and Zatoń 2017). It is inferred that the cuticle of caryocaridids develops similar laminae, but with thinner layer and a greater number of laminae than ceratiocaridids (Pütz and Buchholz 1991).

The primary function of the carapace in crustaceans is to protect the body and limbs, and this is also true for leptostracan phyllocarids (Olesen 2013). According to Vannier *et al.* (2003), the carapace of caryocaridids was initially chitinous and flexible, while Benson (1975) noted that the marginal infold of ostracods was a means of stabilizing the carapace. Therefore, thickening the endocoraeum may enhance the strength, hardness, and stability of the carapace of caryocaridids without adding too much weight, which is an obvious advantage for their pelagic lifestyle. After the carapace of leptostracans is fully developed, the inner and outer cuticles of the middle and posterior of the carapace are completely exposed to the ambient seawater (Manton 1934; Newman and Knight 1984; Olesen and Walossek 2000). The thickened inner cuticle is preferred to counteract the erosion or damage to the carapace caused by the external environment, in addition to stabilizing the carapace.

In the carapace cuticle of caryocaridids, we observe a unique polygonal reticulate ornamentation in the inner and outer lamellae near the middle lamella, with polygonal diameters typically ranging from 10 to 20  $\mu\text{m}$  (Fig. 4G,H). Such a polygonal fabric is also present in other caryocaridids (Tolmacheva *et al.* 2004;



Šilinger 2023; Kovář et al. 2024), such as *Ceratiocaris* (Rolfe 1962; Šilinger 2023; Kovář et al. 2024) and *Dictyocaris* (Salter 1860; Størmer 1935; Gensel et al. 1990). Kovář et al. (2024) interpreted this polygonal pattern as a common epicuticular structure of arthropods. However, the carapace surfaces of the caryocaridids in their figures are smooth (Kovář et al. 2024: plate 1, figs.14–18). The thylacocephalan carapaces also preserve this special polygonal ornamentation (Broda et al. 2015, 2020; Broda and Zatoň 2017). In most cases, it is distributed on the surface of the carapace to strengthen the thin carapace (Vannier et al. 2007; Broda and Zatoň 2017). Rolfe (1962) found a similar polygonal fabric in his study of the cuticle of *Ceratiocaris*, which he interpreted as being similar to the prism structure of cuticles in living crustaceans. Modern hexagonal prisms are thought to correspond to the underlying epidermal cell outlines (Dillaman et al. 2013) and are composed of the chitin–protein complex forming the cuticle (Rolfe 1962). The prismatic size of the carapace cuticle of crustaceans is typically between 4 and 15  $\mu\text{m}$ , and caryocaridids typically have individual polygonal diameters of 10–20  $\mu\text{m}$ . However, the individual polygon in *Ceratiocaris* is 52–270  $\mu\text{m}$ , and as great as 800–1000  $\mu\text{m}$  in *Dictyocaris* (Størmer 1935; Rolfe 1962; Šilinger 2023; Kovář et al. 2024). If such a polygonal reticulate structure corresponds to the cuticular prisms of living crustaceans, then the epidermal cells of phyllocarids will simply be too large. Vannier et al. (1997b: fig. 4C–G) reported that the carapaces of *Dahllella* and *Nebalia* have the obvious hemolymph sinuses consisting of small polygons connected to each other, forming a branching network, and the anterior and posterior margins of the carapace are not consistent. The anterior carapace of *Dictyocaris* has a less-developed reticular structure compared with the posterior (Salter 1860), suggesting that the development of this structure is not uniform throughout the carapace. The hemolymph sinuses are spaces between the inner and outer cuticle that guide hemolymph flow. Each space is connected to the others and blocked by the supporting epidermis formed by the interconnecting inner and outer epidermal cells (Talbot et al. 1972; Wirkner and Richter 2013). The polygonal reticulate structure of the archaeostracans may correspond to the hemolymph sinuses of the leptostracans in shape and size (Vannier et al. 1997b; Wirkner and Richter 2013). The uneven distribution of inner and outer epidermal cells is caused by the presence of hemolymph sinuses. The thickness of the middle lamella in the carapace cuticle of caryocaridids is irregular, and sometimes even disappears, which may be explained by its comparability to the epidermal cell layer (Fig. 4I). Pelagic arthropods require a higher oxygen supply to enhance and/or sustain their active metabolism for swimming, and the thin cuticle of their carapace allows for better diffusion of oxygen into the hemolymph (Perrier et al. 2015). Caryocaridids have larger hemolymph sinuses in their carapace, which can accommodate more hemolymph and transport more oxygen to sustain pelagic activities.

## Conclusion

This study examines the taphonomic types and ultrastructure of the caryocaridid carapace. The taphonomic types of the caryocaridid carapace can be classified into three categories (enrollment, wrinkles, and flattening) based on different compression deformations. Among these, the enrolled type is the rarest worldwide. The cuticle of the carapace in *Soomicaris cedarbergensis* from the Lower Ordovician at Guozigou, Xinjiang, NW China, consists of three mineralized lamellae (outer, middle, and inner). The outer

and inner lamella cuticles of caryocaridids are well calcified by CFA and have weak fibrous structures running perpendicular to the carapace valves. Similar to the exoskeleton of extant crustaceans, the outer and inner lamella cuticles of caryocaridids consist of three cuticular layers: epicuticle, exocuticle, and endocuticle. This special ultrastructure of the caryocaridid carapace appears to be comparable to that of ostracods. The carapace of archaeostracans has a polygonal reticulation structure that corresponds in shape and size to the hemolymph sinuses of leptostracans. The ultrastructure of the carapace of the Ordovician caryocaridids in this study, which includes a thin cuticle, thickened inner lamella cuticle, and large hemolymph sinuses, probably represents an adaptation to a pelagic lifestyle during the Ordovician plankton revolution (Servais et al. 2008, 2016).

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**Competing Interest.** The authors declare no competing interests.

## Literature Cited

- Abdullaev, R. N., and O. N. Khaletskaia. 1970. Nizhniy Paleozoy Chatkalskogo Khrebta: Trilobityi graptolity Ordovika Pskemskogo Khrebta [Lower Paleozoic of the Chatkalskiy Range: Trilobites and graptolites of the Ordovician of the Pskemskiy Range]. *Izdatel'stvo FAN Uzbekskoy SSR Tashkent*.
- Barrande, J. 1872. Système Silurien du centre de la Bohême, Supplément au vol. 1. Praha, Paris.
- Bate, R. H., and B. A. East. 1972. The structure of the ostracode carapace. *Lethaia* 5:177–194.
- Bate, R. H., and B. A. East. 1975. The ultrastructure of the ostracode (Crustacea) integument. *Bulletins of American Paleontology* 65:529–547.
- Becker, G., and F. J. Adamczak. 1990. *Aurikirkbya wordensis* (Hamilton): ein vermutlich filtrierender palaeocopider Ostracode. *Paläontologische Zeitschrift* 64:91–102.
- Benson, R. H. 1975. Morphologic stability in Ostracoda. *Bulletins of American Paleontology* 65:13–46.
- Bouligand, Y. 1972. Twisted fibrous arrangements in biological materials and cholesteric mesophases. *Tissue and Cell* 4:189–217.
- Broda, K., and M. Zatoň. 2017. A set of possible sensory system preserved in cuticle of Late Devonian thylacocephalan arthropods from Poland. *Historical Biology* 29:1045–1055.
- Broda, K., M. Wolny, and M. Zatoň. 2015. Palaeobiological significance of damaged and fragmented thylacocephalan carapaces from the Upper Devonian of Poland. *Proceedings of the Geologists' Association* 126:589–598.
- Broda, K., Š. Rak, and T. A. Hegna. 2020. Do the clothes make the thylacocephalan? A detailed study of Concavocarididae and Protozoecidae (? Crustacea, Thylacocephala) carapace micro-ornamentation. *Journal of Systematic Palaeontology* 18:911–930.
- Bulman, O. M. B. 1964. Lower Paleozoic plankton. *Quarterly Journal of the Geological Society* 120:455–476.

- [BGXUAR] Bureau of Geology of Xinjiang Uygur Autonomous Region. 1987a. *Regional Geological Survey Report of Baskan Pass-Huocheng*. Scale 1:200,000. Bureau of Geology of Xinjiang Uygur Autonomous Region, Urumqi.
- [BGXUAR] Bureau of Geology of Xinjiang Uygur Autonomous Region. 1987b. *Regional Geological Survey Report of Jinghe*. Scale 1:200,000. Bureau of Geology of Xinjiang Uygur Autonomous Region, Urumqi.
- [BGXUAR] Bureau of Geology of Xinjiang Uygur Autonomous Region. 1987c. *Regional Geological Survey Report of Sailimu*. Scale 1:200,000. Bureau of Geology of Xinjiang Uygur Autonomous Region, Urumqi.
- Chapman, F. 1902. XII.—New or little-known Victorian fossils in the National Museum, Melbourne. Part I.—Some Paleozoic species. *Royal Society of Victoria* 16:104–122.
- Chapman, F. 1934. On some phyllocarids from the Ordovician of Preservation Inlet and Cape Providence, New Zealand. *Transactions of the Royal Society of New Zealand* 64:105–111.
- Chen, X., H. L. Lin, H. K. Xu, and Y. X. Zhou. 1998. Early Palaeozoic strata from northwest Xinjiang. *Journal of Stratigraphy* 22:241–251.
- Chlupáč, I. 1970. Phyllocarid crustaceans of the Bohemian Ordovician. *Sborník geologických věd, Paleontologie* 12:41–77.
- Chlupáč, I. 2003. Phyllocarid crustaceans from the Middle Ordovician Šárka Formation at Praha–Vokovice. *Bulletin of Geosciences* 78:107–111.
- Churkin, M., Jr. 1966. Morphology and stratigraphic range of the phyllocarid crustacean *Caryocaris* from Alaska and the Great Basin. *Palaeontology* 9:371–380.
- Collette, J. H., and J. W. Hagadorn. 2010. Early evolution of phyllocarid arthropods: phylogeny and systematics of Cambrian–Devonian archaeostracans. *Journal of Paleontology* 84:795–820.
- Copeland, M. J. 1967. An occurrence of *Caryocaris* (Crustacea, Phyllocarida) from the Canadian Arctic. *Journal of Paleontology* 41:1193–1194.
- Dillaman, R., R. Roer, T. Shafer, and S. Modla. 2013. The crustacean integument: structure and function. Pp. 140–166 in M. Thiel and L. Watling, eds. *The natural history of Crustacea*. Vol. I, *Functional morphology and diversity*. Oxford University Press, New York.
- Freiberger, O. N. 1947. O nakhodke rakoobraznogo *Caryocaris curvikata* Gurley v ordovike yugo-vostochnogo Kapa-tay (On the discovery of the crustacean *Caryocaris curvikata* Gurley in the Ordovician of the southeastern Kapa-Tay). *Doklady Akademii Nauk SSSR* 58:1155.
- Gensel, P. G., N. G. Johnson, and P. K. Strother. 1990. Early land plant debris (Hooker's "waifs and strays"?). *Palaios* 5:520–547.
- Green, J. P., and M. R. Neff. 1972. A survey of the fine structure of the integument of the fiddler crab. *Tissue and Cell* 4:137–171.
- Haj, A. E., and R. M. Feldmann. 2002. Functional morphology and taxonomic significance of a novel cuticular structure in Cretaceous raninid crabs (Decapoda: Brachyura: Raninidae). *Journal of Paleontology* 76:472–485.
- Horne, D. J., A. Cohen, and K. Martens. 2002. Taxonomy, morphology and biology of Quaternary and living Ostracoda. *Geophysical Monograph* 131:5–35.
- Jell, P. A. 1980. Two arthropods from the Lancefieldian (La 1) of central Victoria. *Alcheringa* 4:37–46.
- Kovář, V., M. Šilinger, O. Fatka, and R. Brocke. 2024. Chemical processing of fossil phyllocarid cuticle: a comparison of micro- and macrofossil remains. *Palynology* 48:1–8.
- Li, Y. T. 1995. Early Palaeozoic stratigraphy and sedimentary facies in the Bolhinur Mountain region of western Middle Tianshan in Xinjiang. *Journal of the University of Petroleum* 19:18–24.
- Li, M., W. H. Wang, and H. Z. Feng. 2018. Tremadoc graptolite biozonation of South China and its global correlation. *Acta Geologica Sinica* 57:444–459.
- Liu, Y. L., R. Y. Fan, R. W. Zong, and Y. M. Gong. 2022. The evolution and initial rise of pelagic caryocaridids in the Ordovician. *Earth-Science Reviews* 231:104097.
- Liu, Y. L., R. Y. Fan, R. W. Zong, and Y. M. Gong. 2023. First occurrence of caryocaridids (Crustacea, Phyllocarida) in the Ordovician of North China. *Palaeogeography, Palaeoclimatology, Palaeoecology* 623:111638.
- Manton, S. M. 1934. V. On the embryology of the crustacean *Nebalia bipes*. *Philosophical Transactions of the Royal Society B* 223:163–238.
- Miranda, J. A. 2002. *Origin and formation of Ordovician sedimentary phosphates at Phosphate Hill, Mansfield*. B.S. thesis. University of Melbourne, Melbourne.
- Möberg, J. C., and O. Segerberg. 1906. Bidrag till kännedomen om Ceratopygeregionen med särskild hänsyn till dess utveckling i Fogelsångstrakten. *Meddelande från Lunds Geologiska Fältklubb B* 2:1–113.
- Newman, W. A., and M. D. Knight. 1984. The carapace and crustacean evolution—a rebuttal. *Journal of Crustacean Biology* 4:682–687.
- Nikitina, O. I., L. E. Popov, R. B. Neuman, M. G. Bassett, and L. E. Holmer. 2006. Mid Ordovician (Darriwilian) brachiopods of South Kazakhstan. Pp. 145–222 in M. G. Bassett and V. K. Deisler, eds. *Studies in Palaeozoic palaeontology*. National Museum of Wales Geological Series 25. National Museum of Wales, Cardiff.
- Obut, A. I., and E. N. Zuvtsov. 1965. Stratigrafiya i graptolity ordovika gornogo obramleniya Narynskoy vpadiny [Ordovician stratigraphy and graptolites from the mountains framing the Naryn Depression]. Pp. 13–32. in V. S. Sokolov ed. *Stratigrafiya i paleontologiya paleozoya Aziatskoy chasti SSSR [Stratigraphy and paleontology of the Paleozoic Asian part of the USSR]* Nauka, Moskva.
- Okada, Y. 1982a. Ultrastructure and pattern of the carapace of *Bicornucythere bisanensis* (Ostracoda, Crustacea). *Bulletin of the University Museum, University of Tokyo* 20:229–255.
- Okada, Y. 1982b. Structure and cuticle formation of the reticulated carapace of the ostracode *Bicornucythere bisanensis*. *Lethaia* 15:85–101.
- Olempska, E. 2001. Palaeozoic roots of the sigilliid ostracods. *Marine Micropaleontology* 41:109–123.
- Olempska, E. 2008. Soft body-related features of the carapace and the lifestyle of Paleozoic beyrichioidean ostracodes. *Journal of Paleontology* 82:717–736.
- Olesen, J. 2013. The crustacean carapace: morphology, function, development, and phylogenetic history. Pp. 103–139 in M. Thiel and L. Watling, eds. *The natural history of Crustacea*. Vol. I, *Functional morphology and diversity*. Oxford University Press, New York.
- Olesen, J., and D. Walossek. 2000. Limb ontogeny and trunk segmentation in *Nebalia* species (Crustacea, Malacostraca, Leptostraca). *Zoomorphology* 120:47–64.
- Perrier, V., M. Williams, and D. J. Siveter. 2015. The fossil record and palaeoenvironmental significance of marine arthropod zooplankton. *Earth-Science Reviews* 146:146–162.
- Pillola, G. L., S. Piras, and E. Serpagli. 2008. Upper Tremadoc–Lower Arenig? Anisograptid–Dichograptid fauna from the Cabitza Formation (Lower Ordovician, SW Sardinia, Italy). *Revue de micropaléontologie* 51:167–181.
- Pütz, K., and F. Buchholz. 1991. Comparative ultrastructure of the cuticle of some pelagic, nektobenthic and benthic malacostracan crustaceans. *Marine Biology* 110:49–58.
- Qiao, X. D. 1989. On graptolite-bearing strata and the graptolite fauna division of Ordovician in the western North Tianshan. *Xinjiang Geology* 7:81–90.
- Racheboeuf, P. R., and S. Crasquin. 2010. The Ordovician caryocaridid phyllocarids (Crustacea): diversity and evolutionary tendencies. *Neues Jahrbuch für Geologie und Paläontologie Abhandlungen* 257:237–248.
- Racheboeuf, P. R., J. Vannier, and G. Ortega. 2000. Ordovician phyllocarids (Arthropoda; Crustacea) from Argentina. *Paläontologische Zeitschrift* 74:317–333.
- Racheboeuf, P. R., S. Crasquin, and E. Brussa. 2009. South American Ordovician phyllocarids (Crustacea, Malacostraca). *Bulletin of Geosciences* 84:377–408.
- Roer, R., and R. Dillaman. 1984. The structure and calcification of the crustacean cuticle. *American Zoologist* 24:893–909.
- Roer, R., S. Abehsera, and A. Sagi. 2015. Exoskeletons across the Pancrustacea: comparative morphology, physiology, biochemistry and genetics. *Integrative and Comparative Biology* 55:771–791.
- Rolfé, W. D. I. 1962. The cuticle of some middle Silurian ceratiocaridid crustacea from Scotland. *Palaeontology* 5:30–51.
- Rolfé, W. D. I. 1969. Phyllocarida. Pp. 296–331 in H. K. Brooks, F. M. Carpenter, M. F. Glaessner, G. Hahn, R. R. Hessler, R. L. Hoffman, L. B. Holthuis, *et al.*, eds. *Treatise on Invertebrate Paleontology, Arthropoda*



- 4(1). Part R. Geological Society of America, New York, and University of Kansas Press, Lawrence.
- Rolfe, W. D. I.** 1981. Phyllocarida and the origin of Malacostraca. *Geobios* 14:17–27.
- Ruedemann, R.** 1934. Paleozoic plankton of North America. *Geological Society of America Memoir* 2:1–141.
- Rushton, A. W. A., and J. K. Ingham.** 2000. Scotland: the Highland Border Complex and Grampian terrane. In R. A. Fortey, D. A. T. Harper, J. K. Ingham, A. W. Owen, M.A. Parkes, A. W. A. Rushton, and N. H. Woodcock, eds. *A revised correlation of Ordovician rocks in the British Isles. Geological Society of London Special Report* 24:48–49.
- Rushton, A. W. A., and M. Williams.** 1996. The tail-piece of the crustacean *Caryocaris wrightii* from the Arenig rocks of England and Ireland. *Irish Journal of Earth Sciences* 15:107–111.
- Salter, J. W.** 1860. XV.—On new fossil crustacea from the Silurian rocks. *Annals and Magazine of Natural History* 5:153–162.
- Schram, F. R., and C. H. J. Hof.** 1998. Fossils and the interrelationships of major crustacean groups. Pp. 233–302 in G. D. Edgecombe, ed. *Arthropod fossils and phylogeny*. Columbia University Press, New York.
- Sepkoski, J. J.** 2000. Crustacean biodiversity through the marine fossil record. *Contributions to Zoology* 69:213–222.
- Servais, T., O. Lehnert, J. Li, G. L. Mullins, A. Munnecke, A. Nützel, M. Vecoli.** 2008. The Ordovician biodiversification: revolution in the oceanic trophic chain. *Lethaia* 41:99–109.
- Servais, T., V. Perrier, T. Danelian, C. Klug, R. Martin, A. Munnecke, H. Nowak, et al.** 2016. The onset of the “Ordovician Plankton Revolution” in the late Cambrian. *Palaeogeography, Palaeoclimatology, Palaeoecology* 458:12–28.
- Shen, Y. B.** 1986. *Caryocaris* from the Lower Ordovician of Jiangshan, Zhejiang, China. *Science Bulletin* 31:765–769.
- Šilinger, M.** 2023. *Stavba a složení exoskeletů vybraných skupin členovců pražské pánve* (Exoskeletal structure and composition in selected arthropod groups of the Prague Basin). M.S. thesis. Univerzita Karlova, Praha.
- Sohn, I. G., and L. S. Kornicker.** 1988. Ultrastructure of Myodocopid shells (Ostracoda). Pp. 243–258 in T. Hanai and K. Ishizaki, eds. *Evolutionary biology of Ostracoda, its fundamentals and applications*. Kodansha, Tokyo.
- Spears, T., and L. G. Abele.** 1999. Phylogenetic relationships of crustaceans with foliaceous limbs: an <sup>18</sup>S rDNA study of Branchiopoda, Cephalocarida and Phyllocarida. *Journal of Crustacean Biology* 19:825–843.
- Størmer, L.** 1937. Planktonic crustaceans from the Lower Didymograptus shale (3b) of Oslo. *Norsk geologisk tidsskrift* 16:267–278.
- Størmer, L. E. I. F.** 1935. *Dictyocaris*, Salter, a large crustacean from the Upper Silurian and Downtonian. *Norsk geologisk tidsskrift* 15:267–298.
- Talbot, P., W. H. Jr. Clark, and A. L. Lawrence.** 1972. Ultrastructural observations of the muscle insertion and modified branchiostegite epidermis in the larval brown shrimp, *Penaeus aztecus*. *Tissue and Cell* 4:613–628.
- Tolmacheva, T. Y., L. Holmer, L. Popov, and I. Gogin.** 2004. Conodont biostratigraphy and faunal assemblages in radiolarian ribbon-banded cherts of the Burubaital Formation, West Balkhash Region, Kazakhstan. *Geological Magazine* 141:699–715.
- Tolmacheva, T. J., K. E. Degtyarev, J. Samuelsson, and L. E. Holmer.** 2008. Middle Cambrian to Lower Ordovician faunas from the Chingiz Mountain Range, central Kazakhstan. *Alcheringa* 32:443–463.
- Tolmacheva, T. Y., K. E. Degtyarev, and A. V. Ryazantsev.** 2021. Ordovician conodont biostratigraphy, diversity and biogeography in deep-water radiolarian cherts from Kazakhstan. *Palaeogeography, Palaeoclimatology, Palaeoecology* 578:110–572.
- Vannier, J., and K. Abe.** 1995. Size, body plan and respiration in the Ostracoda. *Palaeontology* 38:843–873.
- Vannier, J., P. Boissy, and P. R. Racheboeuf.** 1997a. Locomotion in *Nebalia bipes*: a possible model for Palaeozoic phyllocarid crustaceans. *Lethaia* 30:89–104.
- Vannier, J., M. Williams, and D. J. Siveter.** 1997b. The Cambrian origin of the circulatory system of crustaceans. *Lethaia* 30:169–184.
- Vannier, J., S. Q. Wang, and M. Coen.** 2001. Leperditicopid arthropods (Ordovician–Late Devonian): functional morphology and ecological range. *Journal of Paleontology* 75:75–95.
- Vannier, J., P. R. Racheboeuf, E. D. Brussa, M. Williams, A. W. Rushton, T. Servais, and D.J. Siveter.** 2003. Cosmopolitan arthropod zooplankton in the Ordovician seas. *Palaeogeography, Palaeoclimatology, Palaeoecology* 195:173–191.
- Vannier, J., J. B. Caron, J. L. Yuan, D. E. Briggs, D. Collins, Y. L. Zhao, and M. Y. Zhu.** 2007. Tuzoia: morphology and lifestyle of a large bivalved arthropod of the Cambrian seas. *Journal of Paleontology* 81:445–471.
- Vega, F. J., V. M. Dávila-Alcocer, and H. F. Filkorn.** 2005. Characterization of cuticle structure in Late Cretaceous and Early Tertiary decapod crustacea from Mexico. *Bulletin of the Mizunami Fossil Museum* 32:37–43.
- Wang, J., H. Hua, X. Wang, P. Y. Gu, R. M. Chen, Y. J. Zhuang, S. P. He, and Y. H. Li.** 2019. Ziyang fauna: a slope facies fossil Lagerstätte of Early Ordovician in South China. *Chinese Science Bulletin* 64:2342–2350.
- Whittle, R. J., S. E. Gabbott, R. J. Aldridge, and J. N. Theron.** 2007. Taphonomy and palaeoecology of a Late Ordovician caryocaridid (Crustacea, Phyllocarida) from the Soom Shale Lagerstätte, South Africa. *Palaeogeography, Palaeoclimatology, Palaeoecology* 251:383–397.
- Wirkner, C. S., and S. Richter.** 2013. Circulatory system and respiration. Pp. 376–412 in M. Thiel and L. Watling, eds. *The natural history of Crustacea*. Vol. I, *Functional morphology and diversity*. Oxford University Press, New York.
- Woodward, H.** 1912. Note on a new species of *Caryocaris* (*C. kilbridensis*) from the Arenig rocks of the Kilbride Peninsula. *Quarterly Journal of the Geological Society of London* 68:99–101.
- [XBGMR] Xinjiang Bureau of Geology and Mineral Resources.** 1993. *Regional geology of Xinjiang Uygur Autonomy Region*. Geology Publishing House, Beijing.
- [XBGMR] Xinjiang Bureau of Geology and Mineral Resources.** 1999. *Lithostratigraphy of Xinjiang Uygur Autonomous Region*. China University of Geosciences Press, Wuhan.
- Xiang, W. L., and T. R. Zhang.** 1984. Tremadocian trilobites from western part of Northern Tianshan, Xinjiang. *Acta Palaeontologica Sinica* 23:399–410.
- Xu, J., and Z. G. Huang.** 1979. Lower Ordovician graptolite fauna of Guozigou area, Hocheng, Xinjiang. *Acta Geologica Sinica* 53:1–19.
- Yamada, S.** 2007. Ultrastructure of the carapace margin in the Ostracoda (Arthropoda: Crustacea). *Hydrobiologia* 585:201–211.
- Yamada, S.** 2019. Ultrastructure and cuticle formation of the carapace in the myodocopan ostracod exemplified by *Euphilomedes japonica* (Crustacea: Ostracoda). *Journal of Morphology* 280:809–826.
- Yamada, S., and D. Keyser.** 2010. Calcification of the marginal infold in podocopid ostracods. *Hydrobiologia* 638:213–222.
- Yamada, S., A. Tsukagoshi, and N. Ikeya.** 2004. Ultrastructure of the carapace in some *Semicytherura* species (Ostracoda: Crustacea). *Micropaleontology* 50:381–389.
- Zatoń, M., P. Filipiak, M. Rakociński, and W. Krawczyński.** 2014. Kowala Lagerstätte: Late Devonian arthropods and non-biomineralized algae from Poland. *Lethaia* 47:352–364.
- Zhang, T. R.** 1987. New finding of trilobites Tremadocian Sairam Formation from Bolhinur mountain, Xinjiang. *Xinjiang Geology* 5:62–70.
- Zhang, S. P.** 2010. *Tectonic evolution and lithofacies paleogeography of the early Paleozoic in the West Tianshan of Yili*. M.S. thesis. Northwestern University, Xi’an.
- Zhang, Y. D., B. D. Erdtmann, and H. Z. Feng.** 2004. Tremadocian (Early Ordovician) graptolite biostratigraphy of China. *Newsletters on Stratigraphy* 40:155–182.
- Zhang, Y. D., Z. H. Wang, H. Z. Feng, T. T. Luo, and B. D. Erdtmann.** 2005. Tremadocian (Ordovician) graptolite biostratigraphy of China: a review. *Journal of Stratigraphy* 29:215–135.