Tarimspira from the Cambrian (Series 2, Stage 4) of Laurentia (Greenland): extending the skeletal record of paraconodontid vertebrates

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Abstract.—Phosphatic sclerites of the problematic *Tarimspira* Yue and Gao, 1992 (Cambrian Series 2) recovered by weak acid maceration of limestones display a unique range of mainly strongly coiled morphologies. They were likely organized into multielement scleritomes, but the nature of these is poorly known; some sclerites may have had a grasping function. *Tarimspira* sclerites grew by basal accretion in an analogous fashion to younger paraconodonts (Cambrian Series 3–4) but lack a basal cavity. Based on proposed homologies, *Tarimspira* may provide an extension of the early vertebrate paraconodont–euconodont clade back into the early Cambrian. *Tarimspira* is described for the first time from Laurentia (North Greenland), extending its known range from China and Siberia in Cambrian Series 2. In addition to the type species, *Tarimspira plana* Yue and Gao, 1992, the Greenland record of *Tarimspira* includes two morphotypes of a new species, *Tarimspira artemi*.

UUID: http://zoobank.org/c7c536c8-cdaf-49a9-ae1d-77c392f553fc.

Introduction

The discovery and description of the conodont animal from the Carboniferous of Scotland (Briggs et al., 1983) inevitably stimulated discussion concerning the relationship of conodonts to early vertebrates (Aldridge et al., 1986, 1993; Kemp and Nichol, 1995; Donoghue et al., 1998, 2000; Blieck et al., 2010; Turner et al., 2010; Donoghue and Keating, 2014). In these deliberations, the main focus of discussion was on euconodonts, the most familiar of the three descriptive groups (protoconodonts, paraconodonts, euconodonts) recognized on the basis of mainly Cambrian material by Bengtson (1983). The affinity of protoconodonts with chaetognaths rather than vertebrates was established by Szaniawski (1982, 1983, 1987, 2002; Vannier et al., 2007), but Bengtson’s (1983) suggestion that euconodonts were derived from paraconodonts ultimately formed the focus for the investigation of conodont element structure between these two groups by Murdock et al. (2013, 2014). Similarity between the crown tissue of euconodont elements and vertebrate enamel was attributed to convergence rather than homology (Murdock et al., 2013, 2014), although the vertebrate affinity of conodonts was not questioned. Donoghue and Keating (2014) accepted the conclusions of Murdock et al. (2013) while maintaining assignment of conodonts to the vertebrates on the basis of their soft part anatomy (Aldridge et al., 1993; Pridmore et al., 1997), despite the objections of Blieck et al. (2010) and Turner et al. (2010).

Microscopic, coiled, phosphatic early Cambrian sclerites described herein show little overall morphological similarity to the diverse and widely distributed conodont elements recovered from Cambrian–Triassic sediments. Their extraordinary mode of growth indicates, however, that they can be compared to some Cambrian (Series 3–4) paraconodonts, and this observation is developed to promote a tentative extension of the model of early vertebrate (conodont) evolution proposed by Murdock et al. (2013) back into the early Cambrian (Cambrian Series 2).

The Cambrian sclerites are assigned to *Tarimspira* Yue and Gao, 1992. The type species, *Tarimspira plana* Yue and Gao, 1992, was originally described from the lower Cambrian (Cambrian Series 2) of western China and is unusual because of its tightly coiled, laterally compressed, planispiral form (Fig. 1). Yue and Gao (1992) compared its shape with that of bellerophontiform mollusks, and there is a seductive resemblance to isostrophically coiled shells of macromollusk genera such as *Joleaudella* Patte, 1929. However, leaving aside the great difference in size, geological age, and phosphatic composition, examination of the method of shell accretion of *Tarimspira* quickly demonstrates that the similarity is superficial, as recognized by Yue and Gao (1992), merely reflecting their common logarithmic growth. *Tarimspira plana* is a relatively solid sclerite with externally deposited growth lamellae that extend from the base to envelop the lateral surfaces. While this method of accretion is also seen in the early growth stages of paraconodonts, the latter rapidly develop a conical form and a deep basal cavity during ontogeny that is not present in *Tarimspira* (Müller, 1959; Andres, 1988; Müller and Hinz, 1991; Murdock et al., 2013).

*Tarimspira* is documented herein from several horizons within the lower Cambrian (Cambrian Series 2, Stage 4) of North Greenland (Fig. 2), where it is represented by *Tarimspira artemi* n. sp. and a single specimen of the type species, *Tarimspira plana*. The occurrences represent its first description from Laurentia, consolidating a brief identification (as *Fengzuella*).
Kouchinsky et al. (2015, p. 481). Tarimspira has previously been described from Siberia (Kouchinsky et al., 2015) and several terranes in China (Yue and Gao, 1992; Steiner et al., 2003; Yang et al., 2015). Although few, these currently known occurrences display a clear equatorial distribution (Fig. 2.4).

Steiner et al. (2003) commented that Chinese specimens occurred in marginal shelf environments. This pattern is confirmed by the present records from shelf deposits of the Aftenstjernesø and Henson Gletscher formations of North Greenland (Ineson and Peel, 1997; Fig. 2.3).

Steiner et al. (2003) recognized that the co-occurrence of Tarimspira (as Fengzuella zhejiangensis) (He and Yu, 1992) with two other sclerite morphotypes in samples from South China indicated that it formed part of the scleritome of an undetermined organism. This deduction is confirmed herein with the description of a new species, Tarimspira artemi, from the Henson Gletscher Formation of North Greenland in which two sclerite morphotypes are known currently from the scleritome. Tarimspira artemi n. sp. is also recorded from the Emyaksin Formation (Cambrian Series 2, Stage 3) of northern Siberia, where it was described under the name Fengzuella zhejiangensis by Kouchinsky et al. (2015). Tarimspira plana, originally described from the Tarim terrane of western China (Yue and Gao, 1992), is now described from the basal Aftenstjernesø Formation of Peary Land, North Greenland (Figs. 1, 2); this species also appears to contain at least two sclerite morphotypes in its scleritome.

**Geological background and materials**

The geological evolution and lithostratigraphy of the Cambrian of North Greenland were described by Higgins et al. (1991) and Ineson and Peel (1997, 2011). Siliciclastic shelf sediments of the Buen Formation (Cambrian Series 2, Stages 3–4; Fig. 2.3) are overlain by carbonates (Aftenstjernesø Formation) that represent the initial stage in the establishment of a major carbonate platform that extended east–west across North Greenland during the Cambrian–Silurian. In the Freuchen Land and Peary Land region (Fig. 2.2), the Cambrian carbonates form a northward prograding complex in which recessive units of dark carbonates and shales (Henson Gletscher, Ekspedition Bræ, and Holm Dal formations; Fig. 2.3), representing periods of relative lowstand of sea level, are separated by carbonate turbidites and mass flow deposits (Aftenstjernesø, Sydpasset, and Fimbuldal formations; Fig. 2.3) deposited during periods of sea level highstand (Ineson and Peel, 1997).

Material of Tarimspira from North Greenland was collected from the Aftenstjernesø and Henson Gletscher formations (early Cambrian; Cambrian Series 2, Stage 4; Fig. 2.3). In Peary Land and Freuchen Land, the Aftenstjernesø Formation yields fossils only from its basal member, a widespread, sediment-starved, outer ramp sequence in which hardgrounds are conspicuous (Frykman, 1980; Ineson and Peel, 1997; Peel, 2017). The member is 3–7 m thick and is mainly composed of glauconitic and phosphatic dolostones.
and limestones that vary in texture from massive to laminated and burrowed.

The Henson Gletscher Formation is dominated by thinly bedded, sooty black limestones, dolostones, and shales that were deposited in an off-platform setting (Higgins et al., 1991; Ineson and Peel, 1997; Geyer and Peel, 2011). In most sections, a lower dark recessive member is overlain by a prominent median unit of pale sandstones and a recessive upper dark member. Diverse faunas (Cambrian Series 2, Stage 4) from the Henson Gletscher Formation were monographed by Blaker and Peel (1997), Geyer and Peel (2011), and Peel et al. (2016). Trilobites of Cambrian Series 3 age were described by Robison (1984) and Babcock (1994a, b).

Samples were collected during regional mapping campaigns (1978–1980; 1984–1985) of the Geological Survey of Greenland (Peel and Sønderholm, 1991). Specimens were etched from the limestone with 10% acetic acid, hand picked from sieved fractions (250 µm and coarser), and gold-coated prior to scanning electron microscopy. Images were assembled in Adobe Photoshop CS4.

**Locality information.**—GGU sample 255522 was collected by P. Frykman on 14 July 1979 from limestone within the lowest 1 m of the basal member of the Aftenstjernesø Formation in Løndal, Peary Land (82°17.5'N, 37°03'W; Fig. 2).

GGU sample 301354 was collected by J.S. Peel on 17 August 1985 from the lower member of the Henson Gletscher Formation on a nunatak in southern Freuchen Land (82°10.5'N, 42°09'W; Fig. 2). GGU sample 271748 was collected by J.S. Peel on 16 July 1978 from about 3 m below the median sandstone member of the Henson Gletscher Formation, Løndal, Peary Land (82°17.5'N, 37°03'W; Fig. 2). The sample is located in the stratigraphic section illustrated by Peel et al. (2016, fig. 2A). GGU sample 218584 was collected by J.R. Ineson on 28 July 1979 in southwest Peary Land, at the head of Henson Gletscher (82°10'N, 39°40'W; Geyer and Peel 2011, fig. 1D, E, locality 5; Blaker and Peel, 1997, fig. 8A, locality 4) from immediately below sandstones forming the upper 13 m of the formation (Fig. 2).

**Repositories and institutional abbreviations.**—GGU prefix indicates a sample collected during field work by Grønlands Geologiske Undersøgelse (Geological Survey of Greenland), now a part of the Geological Survey of Denmark and Greenland, Copenhagen. Type and figured specimens from Greenland (PMU prefix) are deposited in the paleontological type collection of the Museum of Evolution, Uppsala University. Other repositories are noted in the text.

**Systematic paleontology**

*Genus Tarinspira* Yue and Gao, 1992

1992 *Tarinspira* Yue and Gao, p. 153.

1992 *Fengzueella* He and Yu, p. 5, non-*Fengzueella* Li and Han, 1980.
1992 *Otoformilites* Qin and Li in Ding et al., p. 89.
2003 *Fengzuesta*: Steiner et al., p. 858.
2015 *Fengzuesta*: Kouchinsky et al., p. 479.
2015 *Tarimspira*: Yang et al. 2015, p. 1561.

**Type species.**—*Tarimspira plana* Yue and Gao, 1992 from the Aksu-Wushi region, Xinjiang, China.

**Diagnosis.**—Sclerites varying from bilaterally symmetrical, laterally compressed, planispirally coiled with up to three whorls, through shallowly curved and laterally compressed, to almost straight with circular cross section. Coiled morphotypes open coiled or with whorls in contact, often with a spiral keel or carina, or a shallow spiral depression, on periphero-lateral areas. Multilayered with growth lamellae overlapping lateral areas from the basal (abapical) surface. Lamellae compact externally but may be only loosely in contact internally, with cavities, and possibly a circumperipheral canal.

**Occurrence.**—Cambrian Series 2 of China, Siberia, and Laurentia (Fig. 2.4).

**Remarks.**—The diagnosis of Steiner et al. (2003), itself an emendation, is abbreviated and modified to take account of the greater number of whorls in sclerites of *Tarimspira artemi* n. sp. Furthermore, the central cavity is not present in all sclerites assigned to *Tarimspira* herein.

Steiner et al. (2003) synonymized *Tarimspira* Yue and Gao, 1992 and *Otoformilites* Qin and Li in Ding et al., 1992 with *Fengzuesta* He and Yu, 1992 following analysis of the dates of their publication. The latter genus, however, is a junior homonym of *Fengzuesta* Li and Han, 1980 that prompted Yang et al. (2015) to propose *Tarimspira* as a replacement name for *Fengzuesta* He and Yu, 1992. Ironically, both Li and Han (1980) and He and Yu (1992) also employed *zhejiangensis* as the species-group epithet for their respective genera. On account of this species-group homonymy, Yang et al. (2015) gave taxonomic priority to *Tarimspira plana* Yue and Gao, 1992, considering the entire combination *Fengzuesta zhejiangensis* He and Yu, 1992 to be invalid. However, application of International Code of Zoological Nomenclature (ICZN) article 57.8.1 states that “Homonymy between identical species-group names in combination (originally or subsequently) with homonymous generic names having the same spelling but established for different nominal genera is to be disregarded” (International Commission on Zoological Nomenclature, 1999, 2012). *Fengzuesta* Li and Han, 1980, an Ordovician brachiopod, and *Fengzuesta* He and Yu, 1992, a problematic Cambrian sclerite, are different nominal genera with the consequence that *Tarimspira zhejiangensis* (He and Yu, 1992) is a valid name. It is employed in the present context for specimens described by Steiner et al. (2003) from Zhejiang.

Contrary to the opinion of Steiner et al. (2003), *Tarimspira plana* Yue and Gao, 1992 is not considered to be a junior synonym of *Tarimspira zhejiangensis* (He and Yu, 1992). Furthermore, the collective synonymization under *Tarimspira plana* proposed by Yang et al. (2015) is not accepted.

The vitreous, translucent, phosphatic composition of specimens from Greenland suggests that *Tarimspira* sclerites...
were composed of primary calcium phosphate. Steiner et al. (2003) and Kouchinsky et al. (2015) suggested that *Tarimspira* phosphatized remains may have been originally unmineralized, as proposed for early Cambrian protoconodonts by Slater et al. (2018).

*Tarimspira plana* Yue and Gao, 1992

Figure 1

1992 *Tarimspira plana* Yue and Gao, 1992, p. 153, pl. 5, figs. 7–9.

2015 *Tarimspira plana*; Yang et al., p. 1561, fig. 100, P.

**Holotype.**—Specimen 33055/BH7-3a, illustrated by Yue and Gao (1992, pl. 5, fig. 9a, b), Xiaoeerbuk Formation (Cambrian Series 2, Stage 4), Aksu-Wushi Region, Tarim Basin, Xinjiang, northwest China (repository unknown).

**Diagnosis.**—Tightly coiled, planispiral, and bilaterally symmetrical, with a pair of peripheral keels.

**Description.**—The laterally compressed, planispiral, bilaterally symmetrical sclerite is tightly but uniformly coiled through about one and a quarter whorls. Shallowly convex lateral surfaces become concave prior to their transition into a pair of robust spiral keels, located at the periphery, one on each side of the plane of symmetry. The peripheral keels become reduced in relief at the latest growth stage. They delimit a median channel that is slightly V-shaped in transverse profile, with shallowly convex sides, in the early growth stages (Fig. 1.3) but becomes flattened at the latest growth stage (Fig. 1.4). Adaxially, each lateral surface curves abruptly into a deep narrow umbilicus formed as the growing sclerite overlaps earlier growth stages. At the latest growth stage, the sartorial area between the enveloping whorl and the earlier growth stage consists of a deep invagination, with the latest growth lamellae not in direct contact with the earlier whorl (Fig. 1.1).

Externally, the sclerite is compact and appears to be solid, formed by a series of lamellae that slope obliquely abapically in from the lateral areas toward the axial plane in the direction of growth. Thus, the latest growth lamella forms a continuous surface that curves around the growing margin of the sclerite from one lateral area to the other (Fig. 1.5, arrow). The lamellae produce regular transverse lines of growth that are concave toward the direction of growth of the sclerite.

**Materials.**—PMU 31847 from GGU sample 255522, Aftenstjernesø Formation, Cambrian Series 2, Stage 4, Løndal, Peary Land, North Greenland.

**Remarks.**—In following discussion, the morphology of this sclerite is referred to as morphotype A (Fig. 3.8).

The embracive synonymy of previously described taxa with *Tarimspira plana* Yue and Gao, 1992 proposed by Yang et al. (2015) is not accepted. Yue and Gao (1992) described *Tarimspira plana* from the lowermost beds of the Xiaoeerbuk Formation. Two of the specimens illustrated by Yue and Gao (1992, pl. 5, figs. 8, 9) show the bilaterally symmetrical form of the Greenland specimen. In a third specimen, the keels are displaced to one side and bilateral symmetry is lost, suggesting that this may be a second sclerite type within the scleritome.

While the outer surface of the only known specimen from Greenland is compact, the broken peripheral tip at the latest growth stage reveals an inner cavity within several widely spaced lamellae (Fig. 1.5). It is not known whether this cavity is partly original or a function of preservation. Steiner et al. (2003) described similar loose contact between internal lamellae in *Tamispira zhenjaingensis* and inferred the presence of a peripheral internal canal, discussed in the following.

The holotype illustrated by Yue and Gao (1992) from the Xiaoeerbuk Formation (Cambrian Series 2, Stage 4) of the Aksu-Wushi region of Xinjiang, northwest China, is twice the length of the single sclerite known from the Aftenstjernesø Formation. Two specimens illustrated by Yang et al. (2015) from the Shuijingtuo Formation (Cambrian Series 2, Stage 3) of South China are even larger. In one of these (Yang et al., 2015, fig. 10P), the whorl uncoils in its latest fraction. Coiling becomes tighter in the other specimen, occluding the umbilici (Yang et al., 2015, fig. 10O), but it seems likely that this may be a variable character.

*Tarimspira zhejiangensis* (He and Yu, 1992)

Figure 3.3–3.6

1992 Fengzuella zhejiangensis He and Yu, p. 5, pl. 2, fig. 17 non–Fengzuella zhejiangensis Li and Han, 1980.

2003 Fengzuella zhejiangensis; Steiner et al., p. 38, fig. 3a, b, p. q.

**Holotype.**—Specimen number 89-11-2-2452 in the Nanjing Institute of Geology and Palaeontology illustrated by He and Yu (1992) and by Steiner et al. (2003, fig. 3a, b) from the Hetang Formation, Jiangshan County, Zhejiang Province, China.

**Remarks.**—Steiner et al. (2003) described three types of sclerite (here referred to as morphotypes B–D; Fig. 3) within the scleritome of *Tarimspira zhejiangensis* (as Fengzuella zhejiangensis). The eponymous sclerite (morphotype B; Fig. 3.3, 3.6) is rare in the assemblages from the Hetang Formation. They are dominated numerically by sclerites of morphotype C (Fig. 3.4), which can be considered to characterize the scleritome.

*Tarimspira artemi* new species

Figures 4, 5

2015 Fengzuella zhejiangensis; Kouchinsky et al., p. 479, fig. 54.

**Holotype.**—PMU 31848 from GGU sample 301354, Henson Gletscher Formation, Cambrian Series 2, Stage 4, southern Freuchen Land, North Greenland.

**Diagnosis.**—Laterally compressed, planispiral, bilaterally symmetrical sclerite that is tightly but uniformly coiled through about one and a half to three whorls; lateral areas shallowly concave, without spiral keels or carinae (based on sclerite morphotype E).
Description.—The diagnostic sclerite (morphotype E) is laterally compressed, planispiral, bilaterally symmetrical and uniformly coiled through about one and a half to three whorls. The lateral surfaces are shallowly concave between the rounded periphery and the umbilical shoulders; spiral keels or carinae are absent. Adaxially, each lateral surface curves abruptly into a wide umbilicus formed as the growing sclerite overlaps earlier growth stages. The sutural area between the enveloping whorl and the earlier growth stage is deeply invaginated.

Externally, the sclerite is compact and appears to be solid. Growth lamellae slope obliquely abapically in from the lateral areas toward the axial plane in the direction of growth. Each growth lamella forms a continuous surface that curves from one lateral area to the other around the growing margin (base) of the sclerite (arrows in Figs. 4.2, 5.7). The growth lamellae produce regular transverse lines of growth that are concave toward the direction of growth (base) of the sclerite. Adjacent to the periphery, growth lamellae curve forward to form an elevation on the basal surface and appear to delimit a circumperipheral tubular canal (arrow in Fig. 4.3, 4.6, 4.7).

A single specimen similar to the holotype of Tarimspira zhejiangensis was found in the Henson Gletscher Formation of North Greenland. (1) PMU 31849; (2, 8) PMU 31848, holotype, with arrow in (2) indicating growth lamellae extending from the base onto the lateral areas; (3) PMU 31850, broken specimen with circumperipheral cavity on penultimate whorl (arrow); (4, 5) PMU 31851; (6, 7) PMU 31852. (1–6, 8) Scale bars = 50 µm; (7) scale bar = 25µm.

Figure 4. Tarimspira artemi n. sp., morphotype E, Henson Gletscher Formation, Cambrian Series 2, Stage 4; GGU sample 301354, southern Freuchen Land, North Greenland. (1) PMU 31849; (2, 8) PMU 31848, holotype, with arrow in (2) indicating growth lamellae extending from the base onto the lateral areas; (3) PMU 31850, broken specimen with circumperipheral cavity on penultimate whorl (arrow); (4, 5) PMU 31851; (6, 7) PMU 31852. (1–6, 8) Scale bars = 50 µm; (7) scale bar = 25µm.

The associated sclerite (morphotype B2; Fig. 5.1–5.7, 5.9) is laterally compressed, bilaterally symmetrical, and typically open coiled through about a half to a full whorl; it is characterized by a prominent spiral carina with concave sides on each lateral surface close to the acute periphery (Fig. 5.2).

Etymology.—For Artem Kouchinsky in recognition of his contribution to the study of Cambrian faunas.

Materials.—PMU 31849–PMU 31857 from GGU sample 301354, southern Freuchen Land; PMU 31858 from GGU sample 271748, Løndal; PMU 31859 from GGU sample 218584, Løndal. Henson Gletscher Formation, Cambrian Series 2, Stage 4, North Greenland.

Remarks.—The diagnostic morphotype (morphotype E) is distinguished from Tarimspira plana (morphotype A) by its wider umbilici and lack of spiral keels around the periphery. The associated sclerite (morphotype B2) resembles sclerite morphotype B of Tarimspira zhejiangensis in terms of the prominent spiral carinae on the lateral surfaces, but most specimens are more openly coiled through less than a whorl (Fig. 5.5, 5.6). However, a single specimen from GGU sample 271748 in Løndal (Fig. 5.8), tentatively placed here, is more strongly...
coiled and therefore morphologically close to the holotype of Tarimspira zhejiangensis (Fig. 3.3).

The nature of the possible circumperipheral canal in Tarimspira artemi is obscure, not least since it is developed at the apex of the subperipheral elevation (Fig. 4.6). It may be an artifact produced by fracturing, or imperfect stacking of growth lamellae that fail to maintain contact just at their apices in the cone-in-cone structure (Fig. 4.7). Gaps between lamellae may also result in part from diagenesis or etching during preparation, as visible in the B2 morphotype (Fig. 5.4). Well-preserved specimens of this latter morphotype show no evidence of a canal (Fig. 5.6, 5.7). Spine-like sclerites (Fig. 3.5, morphotype D) assigned to Tarimspira zhejiangensis by Steiner et al. (2003, fig. 3m-o, s) seem to have a hollow core, but the separation of the spiral lamella suggests a diagenetic origin. However, Steiner et al. (2003, fig. 3f) described a circumperipheral central cavity extending from the base to the apex in morphotype C (Fig. 3.4), although the relationship of this to the basal part of the growth lamellae is unclear.

Sclerite growth

Sclerites of problematic organisms are diverse and abundant in assemblages of shelly fossils from the lower Cambrian (Bengtson, 2005; Rozanov et al., 2010; Kouchinsky et al., 2011, 2015), but Tarimspira sclerites differ from most described examples in their mode of secretion; they are constructed of abapically sloping lamellae (Figs. 1.5, 1.6, 3.7–3.9). Thus, the latest (basal) growth lamella was deposited externally; it overlies the previously formed lamella as it passes continuously from one lateral surface to the other around the abapical (basal) extremity (Fig. 1.5, arrow). As noted by Steiner et al. (2003) and Kouchinsky et al. (2015), this growth pattern indicates that the developing sclerites were rooted in an invagination in underlying soft tissues. Contemporaneous Cambrian sclerites such as lapworthellids (Devaere and Skovsted, 2017) and protoconodonts (Szaniawski, 1982, 1983, 1987, 2002; Bengston, 1983; McIlroy and Szaniawski, 2000; Vannier et al., 2007) grew by the addition of lamellae on the inside of a hollow.
basal cavity, with deposition essentially upon papillae within the basal cavity. Both methods of growth increase the anchoring surface area of attachment relative to a simple, smaller, planar contact, which may enhance their defensive role or, in spinose grasping sclerites such as those of protoconodonts, even their maneuverability.

Comparable incremental growth around the lateral areas and across the base of sclerites occurs in paraconodonts such as Furnishina Müller, 1959, Prooneotodus Müller and Nogami, 1971, and Problematococonites Müller, 1959 (Murdock et al., 2013; Fig. 3.1, 3.2), although the acute spinose form and deep basal cavity of these paraconodonts find no morphological equivalence in Tarimspira. Instead of a basal cavity, the abapical termination of most sclerites of Tarimspira is rounded or wedge shaped (arrows in Fig. 1.5, 1.6; Fig. 5.1–5.7). The abapically concave shape of the growth lamellae as they cross the lateral areas of the sclerite in Tarimspira plana indicates that mineralizing tissue embraced the sclerite mainly on its lateral areas such that the earlier formed coil and umbilical areas in Tarimspira plana, together with most of the supra-apical surface, were probably exposed, at least periodically. While sclerite formation was within an invagination, the shape of the growth lamellae indicates that this pocket was likely formed as a cleft traversing either a papilla or a transverse ridge of secreting tissues.

The growing margin (abapical termination of the sclerite) is clearly visible in Tarimspira artemi n. sp. (arrow in Fig. 4.2) and in sclerites with pronounced lateral spiral carinae (Fig. 5.1, 5.6, 5.9). Lateral ridges in the latter form a prominent transverse element that both reinforces the narrow sclerite and stabilizes its attachment in the soft tissues perpendicularly to the otherwise narrow attachment surface along the plane of symmetry.

The prominent spiral lateral carinae of some Tarimspira sclerites (Fig. 5.1–5.9) are reminiscent of the lateral costae of distacodiform euconodont elements (Robison, 1981), but such structures are not typical of paraconodonts of the Furnishina–Prooneotodus–Problematococonites morphological group. In Coelocerodontus Ethington, 1959, however, the tall, laterally spiral keels on the supra-apical surface (Andres, 1988; Dong and Zhang, 2017). Specimens of Coelocerodontus from the early Ordovician of Sweden illustrated by Andres (1988, fig. 18) show a similar degree of curvature, rate of expansion, and ornamentation of spiral carinae to some sclerites of Tarimspira (Fig. 5.5, 5.6). Andres (1988) considered Coelocerodontus to be a paraconodont. Szaniawski (2015) suggested a separate group on the basis of its histology, but Dong and Zhang (2017) assigned it to the euconodonts. While the development of carinae perpendicular to the plane of symmetry in Tarimspira, Coelocerodontus, and distacodiform euconodont elements may suggest an evolutionary relationship, it is more likely that it reflects the common functions of strengthening the sclerite and increasing its anchorage in secreting soft tissues.

A basal cavity is not present in Tarimspira where the basal surface is usually protruding. There is evidence of the presence of a narrow spiral internal canal extending from the basal surface toward the apex in some sclerites (Fig. 4.6), as also discussed by Steiner et al. (2003), but the nature of the canal and, if present, its function are uncertain. A relationship to the basal cavity of members of the Furnishina–Prooneotodus–Problematococonites morphological group might be suggested, although the initial growth stages of their elements do not show a comparable structure (Andres, 1988; Murdock et al., 2013).

Scleritome

Reconstruction of isolated sclerites into their original scleritomes is a vital step in the determination of the affinity of Cambrian problematic organisms. The special preservation offered by Lagerstätten has proved invaluable with regard to groups such as the chancellorids (Bengtson and Collins, 2015) and halkieriids (Conway Morris and Peel, 1995). However, the reconstruction of many other organisms remains speculative and often a mental challenge, as demonstrated, for example, by the multiplied Trachyplax arctica Larsson, Peal, and Högstrom, 2009 that occurs in North Greenland in co-eval strata to those yielding the unrelated Tarimspira plana (Larsson et al., 2009).

Steiner et al. (2003) noted, but did not illustrate, clusters of similar sclerites of Tarimspira (as Fengzuela zhejiangiensis in samples from the early Cambrian of South China (Fig. 3.4–3.6) but no recognizable scleritomes. However, they recognized three co-occurring sclerite morphotypes, here referred to as morphotypes B–D (Fig. 3; the tightly coiled sclerites of Tarimspira plana are designated as morphotype A). Coiled sclerites of morphotype B (Fig. 3.6) are rare. They are usually curved through slightly more than one revolution and more open coiled than the holotype of Fengzuela zhejiangensis He and Yu, 1992, refigured by Steiner et al. (2003, fig. 3a; b; Fig. 3.3, 3.6), but otherwise similar. Morphotype B sclerites carry a prominent spiral fold on each lateral area.

More than 85% of the specimens available to Steiner et al. (2003) are laterally compressed curved sclerites (length 410–1,340 μm) with subparallel sides, a uniformly curved supra-apical surface, and a narrow base (morphotype C). These curved sclerites are coiled through less than half a whorl in lateral perspective (Fig. 3.4), with a median keel along the supra-apical surface, but lack spiral folds or ridges on the lateral areas (Steiner et al., 2003, fig. 3c). Narrow, straight or only slightly curved sclerites (morphotype D) are pointed and attain greater length than the curved sclerites. While they show the characteristic overlapping lamellae (Fig. 3.5), a cross section illustrated by Steiner et al. (2003, fig. 3s) indicates that this is a continuous spiral lamella rather than a series of discrete, stacked, cone-shaped lamellae. This difference in the mechanism of growth from morphotypes A–C makes it uncertain whether this sclerite belongs to the Tarimspira zhejiangiensis scleritome, although the two forms may represent end members in a morphological series. The sclerite in morphotypes A–C is planispirally coiled, without any translation along the axis (Fig. 3). This coiling is tight in morphotype A but widely open coiled in morphotype C. In the spine-like morphotype D, there is a high rate of translation of the growing lamella along the axis of coiling with the result that the direction of growth of the sclerite approximates to the orientation of the coiling axis. Growth increase in morphotypes A and B is perpendicular to the axis of coiling.
The increase in translation in morphotype D thus provides a mechanism for the development of a spinose sclerite with a possible grasping function. While its function may parallel the grasping function of sclerites of protoconodonts (Szaniawski, 1983) and panderodid euconodonts (Sansom et al., 1995), the growth of morphotype D is constrained by the underlying protruding base characteristic of *Tarimspira*.

Greenland material assigned to *Tarimspira artemi* n. sp. includes two sclerite morphotypes. Curved to open-coiled sclerites (Fig. 3.7, morphotype B2; Fig. 5.1–5.9) are similar to the morphotype B sclerites of *Tarimspira zhejiangensis* (Figs. 3.3, 4.6) and carry a prominent spiral ridge on either side of the acute supra-apical keel. They occur together with tightly coiled sclerites in which coiling varies between two and three whorls, but which lack spiral ridges on the lateral areas (Fig. 3.9, morphotype E; Fig. 4). Sclerites of this type are not described in *Tarimspira zhejiangensis*, but both sclerite morphotypes B2 and E are recognized in material from the Emyaksin Formation of northern Siberia described by Kouchinsky et al. (2015, fig. 54), which is here assigned to *Tarimspira artemi* n. sp. The single specimen of *Tarimspira plana* known from Greenland (Fig. 3.8, morphotype A; Fig. 1) is the same morphotype as the tightly coiled holotype (Yue and Gao, 1992) and specimens figured by Yang et al. (2015), but the former authors also illustrated an open-coiled morphotype with a lateral ridge similar to that noted here in morphotype B of *Tarimspira zhejiangensis* (Fig. 3.3, 3.6) and in *Tarimspira artemi* (morphotype B2; Fig. 5.1–5.9).

Thus, at least partial scleritomes with distinctive sclerites can be recognized for each of the three described species of *Tarimspira* (*T. plana*, *T. zhejiangensis*, and *T. artemi*), although it is likely that these scleritomes contain other as yet unknown sclerite morphotypes.

The arrangement of individual sclerites and their precise function within the scleritomes are not known. Steiner et al. (2003) speculated that sclerites were closely packed in *Tarimspira zhejiangensis* with morphotypes B and C located laterally to morphotype D. Interpretation of the sclerites as dorsal armor might promote comparison with well-known but unrelated forms such as the early–middle Cambrian *Wiwaxia* Walcott, 1911 (Slater et al., 2017, fig. 14) or *Halkeria* Poulsen, 1967 (Conway Morris and Peel, 1995), although the morphology, method of formation, and composition of individual sclerites in the latter are fundamentally different (Smith, 2014; Zhang et al., 2015). Furthermore, there is little direct evidence concerning the orientation of sclerites in *Tarimspira*.

The present comparison with paraconodonts in terms of sclerite growth promotes interpretation of the sclerites of *Tarimspira* as elements within an oropharyngeal feeding array (Donoghue et al., 2000), but there is little morphological similarity with known arrays. Andres (1988, fig. 17) presented a partial reconstruction of *Coelocerodontus* that might be applied to sclerites of morphotype B2, although these are more curved and less pointed than elements of *Coelocerodontus* and lack any obvious grasping function. However, this function may have been provided by the spine-like morphotype D. The position, orientation, and function of the tightly coiled sclerites of morphotypes A and E in such a reconstruction are unknown, but they may have had a crushing function deep in the pharynx in contrast to the grasping function of elements of the *Furnishina–Problematoconites* type (Murdock et al., 2014). Whereas the basal (proximal) surface of sclerites probably would be located anteriorly in an elongate dorsal sclerite, with the curvature toward the posterior, no preferred single orientation can be assumed in oropharyngeal conodont arrays (Sansom et al., 1995; Szaniawski, 2002; Murdock et al., 2013).

**Systematic position**

The style of basal accretion with lateral overlap promotes comparison of *Tarimspira* with paraconodonts, but there is little similarity in terms of overall morphology between *Tarimspira* and fully developed paraconodont elements (Müller, 1959; Müller and Nogami, 1971; Andres, 1988; Müller and Hinz, 1991; Murdock et al., 2013), suggesting that the arrangement and function of the respective hard parts were dissimilar. The morphological disparity may imply that the unusual method of accretion represents convergence in the method of growth, although a different but equally great range in morphology is seen when comparing other groups widely accepted as paraconodonts, namely the arcane westergaardodinids (Müller, 1959; Müller and Hinz, 1991) and the slender, curved, simple cones with a deep basal cavity extending almost to the tip seen in *Furnishina*, *Prooneotodus*, and *Problematoconites* (Fig. 3.1, 3.2).

Sclerites of *Tarimspira* lack a basal cavity, but the basal cavity in paraconodonts such as *Furnishina*, *Prooneotodus*, and *Problematoconites* is not developed until after elements attain a length of 100–300 µm (Andres, 1988, fig. 19; Murdock et al., 2013, fig. 2; Fig. 3.1). Prior to this stage, the characteristic basal overlap of the lateral areas by successive accreted layers closely resembles that seen in *Tarimspira* (Figs. 1.5, 5.4). Thus, *Tarimspira* sclerites could be interpreted as early ontogenetic stages of paraconodonts formed prior to the development of a basal cavity, although...
this interpretation is opposed by the large size of Tarimspira elements (1.5 mm) reported by Steiner et al. (2003) and the lack of direct evidence of later ontogenetic stages with a semblance of a basalt cavity in any of the samples.

It is proposed that Tarimspira sclerites may reflect a stage in paraconodont evolution prior to the development of a basalt cavity, suggesting a simple expansion of the model of paraconodont–euconodont evolution proposed by Murdock et al. (2013; Fig. 6). As concerns Tarimspira, however, the model is simplistic in that it is mainly based on the form of morphotypes A, B, and B2 (Fig. 3). Morphotype C is readily interpreted as a more open-coiled version of morphotype B2. Morphotype D of Tarimspira zhejiangensis is bizarre and may not belong to the scleritome, although greatly increased translation along the coiling axis is a feasible explanation of its growth form.

The interpretation of Tarimspira as a paraconodont is consistent with the geological record and pushes the known record of paraconodont vertebrates back into the early Cambrian. The oldest previously recognized paraconodonts, represented by Furnishina in the model of Murdock et al. (2013), are reported from the Psychagnostus gibbus Biozone (uppermost Stage 5) of Cambrian Series 3 (Dong and Bergström, 2001; Dong et al., 2001; Kouchinsky et al., 2011; Dong and Zhang, 2017) whereas Tarimspira occurs in Cambrian Series 2 (Stage 4).

Paraconodonts thus encompass at least three distinct morphological groups: the Furnishina–Prooneotodus–Problematoconites complex and Tarimspira. The inhomogeneity suggests that the relationships of these groups of early vertebrates are poorly understood, adding substance to the demonstration of paralogy by Murdock et al. (2013).

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