

Sympatric speciation driving evolution of Late Ordovician brachiopod *Zygospira* in eastern North America

Colin D. Sproat^{1*}  and Jessica S.A. McLeod¹

¹Department of Geological Sciences, University of Saskatchewan, Saskatoon, Saskatchewan, Canada. <c.sproat@usask.ca>, <j.mcleod@usask.ca>

Abstract.—The morphology of *Zygospira*, an early atrypide brachiopod, was analyzed using a multivariate approach. Principal component analysis and discriminant analysis clearly differentiated species as they are currently defined primarily based on differences in shell size and ornamentation but not in terms of overall shell shape. The older *Zygospira modesta* was able to persist into the late Katian (Richmondian) while smaller early species in other brachiopod lineages mostly went extinct. This may have been possible through niche partitioning because the smaller shells have been found attached to other filter feeders and no larger species have been found in these associations so far. This could represent a rare example of sympatric speciation preserved in the fossil record. In the future, detailed study of the spiralia and their associated structures may provide clues as to the ultimate evolutionary affinities of this group in relation to the other atrypide brachiopods evolving at this time.

Introduction

The brachiopod order Atrypida originated during the Ordovician Radiation (sensu Stigall et al., 2019) alongside most other brachiopod lineages (Harper et al., 2013) and other fossil groups that encompassed the Paleozoic Evolutionary Fauna. Unlike the more well-known atrypides from the Devonian that typically have large, strongly dorsibiconvex shells (such as the namesake of the order, *Atrypa*), these early atrypides had small ventribiconvex shells and less elaborate spiralia with fewer whorls supporting the lophophore. Because these early species have been understudied, the origin of this important brachiopod order remains uncertain.

One of the most common of these early atrypide genera in Laurentia is *Zygospira* Hall, 1862. The type area of the genus is the upper Katian of the Cincinnati tri-state area (Hall, 1862), but species have been reported from rocks of similar age in Ontario (Foerste, 1924), Hudson Bay Lowlands (Jin et al., 1997), Iowa (Wang, 1949), and Texas (Howe, 1965) (Fig. 1). A number of species reported from older rocks and localities from outside of Laurentia probably belong to other genera, such as the earlier *Anazyga* Davidson, 1883, or a variety of atrypide genera in the plates that now comprise China and Central Asia that are more distantly related to *Zygospira* (Copper, 1977; Rong et al., 2017). An exception to this is *Zygospira carinata* Percival, 1991, reported from New South Wales (Australia) that extends into the early Katian (although the spiralia in this species has

not yet been documented to confirm its assignment to the genus).

Many species of *Zygospira* have been defined based on generalized descriptions of differences in shell size and ornamentation but have not been critically re-assessed using contemporary methods. Devising a taxonomic scheme to classify these species is made difficult by the nature of early atrypide morphology and development. For example, unlike other brachiopod orders, such as Orthida or Strophomenida that commonly feature distinctive cardinalia and muscle scars that can be used as diagnostic characters, early atrypides generally have poorly impressed muscle scars and lack distinctive cardinalia and other diagnostic internal structures that have proven useful in distinguishing brachiopod species within other lineages.

The structure of the calcified supports for the lophophore that evolved in the spire bearers (spiralia) has proven to be a useful character for suprageneric classification of the group. Variations in the number of whorls and geometry of the spiralia, as well as variation in structures associated with the spiralia, have been used to differentiate lineages within these early Atrypida (Copper, 2002). Unfortunately, many of these features within the shell remain poorly documented due to the labor- and time-intensive process of serial sectioning necessary to produce meaningful cross-sections of these features.

Serial sections illustrate key characteristics of the most common species herein, but the present study focuses on the external morphology of shells to evaluate the existing taxonomic framework of this evolutionary lineage and investigate any functional or paleoecological implications of the morphological disparity present in *Zygospira*. A new collection of *Zygospira*

*Corresponding author.

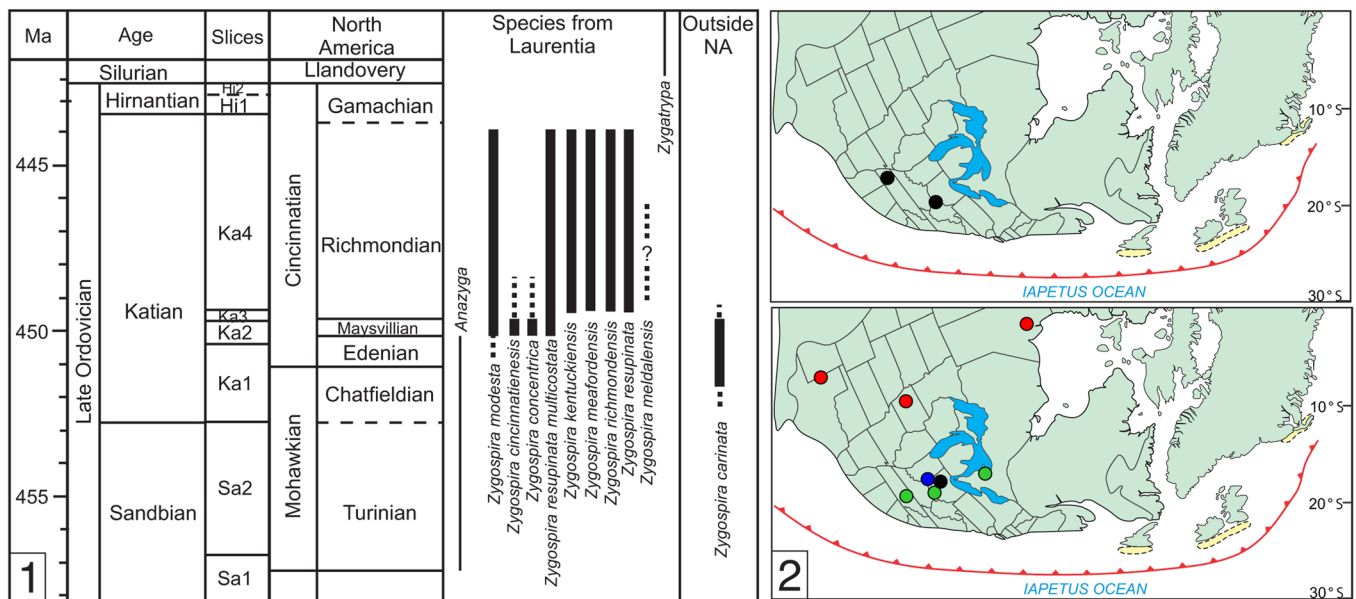


Figure 1. (1) Stratigraphic range of *Zygospira* and closely related genera *Anazyga* and *Zygotrypa* (left). (2) Maps showing the range of *Zygospira* in eastern North America in the middle (top) to late Katian (bottom). Note that this does not include the numerous species now classified under other atrypide genera or suspected to belong to *Anazyga* (see Systematic Paleontology). Colors: black = *Z. modesta*; green = *Z. kentuckiensis*; blue = *Z. cincinnatensis*; red = *Z. resupinata* and subspecies.

kentuckiensis James, 1878, from the upper Katian Queenston Shale near Owen Sound, Ontario, Canada, is described and illustrated, including traced serial sections showing the interior of the shell.

The results of this study will contribute to our understanding of the evolution of the shelly benthos during the Ordovician Radiation. Although the effects of this evolutionary radiation on brachiopod biodiversity are becoming increasingly clear, the underlying processes driving this event are not yet fully understood. Detailed specimen-based analyses such as this may hold the key to understanding the underlying dynamics and evolutionary innovations that drove the rise in biodiversity and morphological disparity during this event.

Previous studies

The genus *Zygospira* was initially erected by Hall (1862) to include all of the coarsely costate brachiopods with spiralia in the Upper Ordovician rocks of North America based on the earlier described *Atrypa modesta* Say in Hall, 1847. In the initial description of the genus, he rightly recognized the significance of the spiralia as a diagnostic character, noting the presence of a strong loop that differentiates these species from later Atrypida. The similarities in ventral position of the bases of the spiralia were used to hypothesize that these early forms were distant ancestors to the later atrypides of the Silurian and Devonian. Hall (1862) also noted the distinctive external form of these earlier atrypides in comparison to later species, but never elaborated on this point.

Hall and Clarke (1894) summarized much of the work done on the genus in the latter part of the nineteenth century. Notably, they synonymized the species assigned to *Anazyga* Davidson, 1883, with *Zygospira* based on the similar apparent variability in the spiralia and range in shell shapes of species assigned to both genera. Rather than two separate genera, they saw *Anazyga* and *Zygospira* as two endmembers on a continuous spectrum.

Since these earlier studies, few others have attempted a comprehensive study of the early atrypides in Laurentia. Copper (1977) proposed that *Zygospira* should be divided into two main groups, restoring *Zygospira* and *Anazyga* as separate genera. *Zygospira sensu stricto* included species similar to the type species *Zygospira modesta* that are strongly ventribiconvex and generally larger in size. These species have spiralia with spires that are dorso-medially oriented within the mantle cavity and have a jugum that connects the spires together high in the dorsal valve posterior to the apices of the spiralia.

Species restored to *Anazyga* were slightly older, common in the lower Katian (an interval referred to as the Trentonian in North America) of eastern North America, while *Zygospira* is largely confined to the middle to upper Katian. *Anazyga* species are united in having spiralia with medial-oriented spires in comparison to the dorso-medial spiralia of *Zygospira*. Although both genera possess a jugum, Copper (1977) recognized a consistent difference in its location within the shell. *Anazyga*, so far as is known, typically have a jugum closer to the anterior than in *Zygospira*, although this needs to be studied in detail to determine how stable this position is among species of the genus. Variability in the shape and configuration of this structure within this lineage remains poorly studied, however. Externally, *Anazyga* tends to be more biconvex than the later *Zygospira* species and has a more strongly carinate ventral valve with more prominently differentiated mid and lateral ribs.

Species of *Zygotrypa* are apparently separated from the genera of the Anazygidae by a stratigraphic gap in the Silurian (Copper, 1977). This may be an example of the Lazarus Effect, or this genus may be more closely related to one of the other atrypide lineages from this time. These species are only poorly documented and do not occur as widely as earlier forms.

Although *Zygospira* was once thought to be a cosmopolitan genus, reviews of atrypide brachiopods from Central Asia (Popov et al., 1999) and China (Rong et al., 2017) have revealed

that species from these plates and terranes mostly belong to the Atrypinidae rather than the Anazygidae. Late Ordovician Atrypinidae usually have a dorsal fold and ventral sulcus rather than the dorsal sulcus and ventral fold of Anazygidae and are so far only known to possess separated jugal processes rather than one solid jugum connecting the spiralia. The functional significance of these differences remains uncertain, but *Zygospira* sensu stricto are now almost entirely known from Laurentia. The only known occurrence of the genus as it is currently defined outside of Laurentia is *Zygospira carinata* Percival, 1991, from the Upper Ordovician of New South Wales, Australia, but no information on its spiralia is available to confidently assign the species to *Zygospira*.

Materials and methods

Several species of *Zygospira* were examined from collections made by Sproat and Dr. Jisuo Jin (Western University) now deposited at the Royal Ontario Museum (ROM). These were supplemented with specimens from the Geological Survey of Canada (GSC), Cincinnati Museum Center (CMC), Field Museum (UC, IP), American Museum of Natural History (AMNH), and the University of Iowa (SUI).

Diagnostic measurements of well-preserved specimens were recorded using a set of digital calipers and a protractor. These measurements included length, width, and depth of the shell in addition to length of the dorsal valve, depth of each valve measured from the posterior part of the commissure, depth of the deviation of the commissure at the anterior caused by the dorsal sulcus and ventral fold, apex angle formed by the ventral umbo, and number of ribs on the ventral valve (Fig. 2). Care was taken to avoid measuring shells with obvious signs of damage or deformation (e.g., cracks, parts of the shell missing, high degree of asymmetry that could indicate post-depositional deformation, etc.).

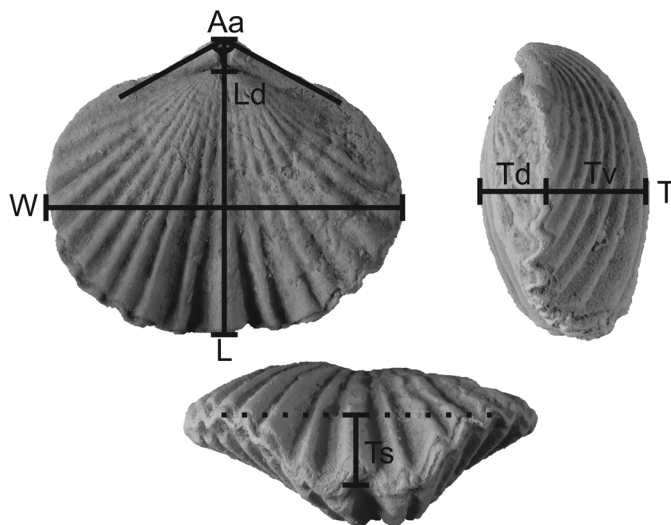


Figure 2. Measurements of *Zygospira* species in this study. L = length (equivalent to length of ventral valve); Ld = length of dorsal valve; W = width; T = thickness (depth) of specimens; Td = thickness of dorsal valve; Tv = thickness of ventral valve; Ts = depth of the sulcus at the anterior commissure; Aa = apical angle (angle formed by the ventral umbo as it projects across the hinge line). Number of ribs on the ventral valve also was counted.

Measurement data were analyzed using PAST v.4.09 (Hammer et al., 2001). Linear regressions were plotted using the reduced major axis method. Principal component analysis used a correlation matrix that standardizes data to account for differences in the dimensions measured. Discriminant analysis was carried out as a further test of the interspecific variation in morphology.

To examine the internal morphology of shells, brachiopods were serial sectioned using a Croft Parallel grinder. This process involves grinding the fossil down in set intervals and replicating each polished surface with cellulose acetate replication film after a brief rinse with dilute (2–3%) hydrochloric acid (HCl) to expose the shell from the micritic matrix material that fills the mantle cavity. Sections were made every 0.05–0.1 mm and digitally traced using the CorelDraw Graphics Suite.

Repositories and institutional abbreviations.—Specimens studied are deposited at the American Museum of Natural History (AMNH), Cincinnati Museum Center (CMC), Field Museum (FM or UC), Geological Survey of Canada in Ottawa (GSC), and Royal Ontario Museum (ROM).

Systematic paleontology

Class Rhynchonellata Williams et al., 1996
 Order Atrypida Rzhonsnitskaya, 1960
 Suborder Anazygidina Copper in Copper and Gourvenec, 1996
 Superfamily Anazygoidea Davidson, 1883
 Family Anazygidae Davidson, 1883
 Subfamily Anazyginae Davidson, 1883
Zygospira Hall, 1862

Type species.—*Producta modesta* Say in Hall, 1847.

Other species.—Copper (1977) briefly reviewed species assigned to *Zygospira* and restored *Anazyga* Davidson, 1883, to include species with smaller, medially directed spiralia. Using the shell morphology and stratigraphic ranges of the type species as a guide, he then assigned described species of *Zygospira* either to *Zygospira* or *Anazyga*. Some ambiguity remains regarding the spiralia in many species due to the time-intensive nature of serial sectioning necessary to examine the internal morphology of the shells and the lack of density contrast between the shell material and surrounding matrix that makes imaging using modern CT techniques difficult to impossible. That said, several species can confidently be assigned to *Zygospira* based on their ventribiconvex lateral profile, wider than long outline, strong simple ribs, and known configuration of the spiralia. These include:

Zygospira modesta (Say in Hall, 1847).—Say in Hall (1847, p. 141–142, pl. 33, fig. 15); type specimens initially identified as *Producta modesta* by Say (see Hall, 1847).

Zygospira kentuckiensis Nettelroth, 1889.—Nettelroth (1889, p. 138–139, pl. 34, 21–23; not pl. 24, 25).

Zygospira resupinata Wang, 1949.—Wang (1949, p. 18–19, pl. 10A, figs. 1–12); only a holotype and a single paratype remain in the SUI collections but Jin et al. (1997) illustrated a specimen of the subspecies *Zygospira resupinata multicostata* Howe, 1965, that clearly possesses dorso-medially directed

spiralia consistent with *Zygospira* (see Jin et al., 1997, pl. 30, fig. 21).

Several species can be synonymized with the above species based on strong similarities in shell morphology including:

Zygospira concentrica Ulrich, 1879.—Ulrich (1879, p. 14, pl. 7, figs. 10a, b); likely a synonym of *Z. modesta*.

Zygospira meafordensis Foerste, 1924.—Foerste (1924, p. 125, pl. 15, figs. 3a–c); herein considered a subspecies of *Z. kentuckiensis*.

Zygospira raymondi Foerste, 1924.—Foerste (1924, p. 127, 128); not figured, but based on specimens described as *Zygospira uphami* by Raymond (1921); likely a synonym of *Z. kentuckiensis* with a slightly flattened ventral valve in comparison to the type collection.

Zygospira richmondensis Caley, 1936.—Caley (1936, p. 60, 78, pl. 1, figs. 4, 6); likely a synonym of *Z. kentuckiensis*, although more evenly convex than typical *Z. kentuckiensis*.

Most of the early (early Katian, pre-Maysvillian) species previously assigned to *Zygospira* have been reassigned to *Anazyga* (see also Copper, 1977, for a list with minor differences of opinion). Their taxonomic assignment should be critically reassessed after their spiralia and associated structures have been documented. These are not examined in detail here, but include:

Atrypa recurvirostra Hall, 1847.—Type species of *Anazyga*, sometimes referred to in literature as *Zygospira recurvirostris*.

Zygospira calhounensis Fenton and Fenton, 1922.—Fenton and Fenton (1922, p. 76–77, pl. 2, figs. 4–6).

Zygospira circularis Cooper, 1956.—Cooper (1956, p. 670, pl. 141C, figs. 18–21, pl. 142B, figs. 6–10, pl. 142D, fig. 16).

Zygospira elongata Cooper, 1956.—Cooper (1956, p. 670–671, pl. 268G, figs. 29–32).

Zygospira gutta Oraspöld, 1956.—Oraspöld (1956, p. 64–65, pl. 4, figs. 14, 15).

Zygospira lebanonensis Cooper, 1956.—Cooper (1956, p. 671–672, pl. 142C, figs. 11–15).

Zygospira matutina Cooper, 1956.—Cooper (1956, p. 672, pl. 141B, figs. 13–17).

Zygospira maynei Roy, 1941.—Roy (p. 102–103, fig. 69).

Zygospira mediocostellata Cooper, 1956.—Cooper (1956, p. 672–673, pl. 143D, figs. 13–18).

Zygospira recurvirostris aequivalvis Twenhofel, 1928.—Twenhofel (1928, p. 214, pl. 19, figs. 10–12).

Zygospira recurvirostris noquettensis Hussey, 1926.—Hussey (1926, p. 162–163, pl. 11, figs. 1–3).

Zygospira recurvirostris turgida Foerste, 1917.—Foerste (1917, p. 103, pl. 5, fig. 15 a–c).

Zygospira variabilis Fenton and Fenton, 1922.—Fenton and Fenton (1922, p. 75–76, pl. 2, figs. 7–9).

Zygospira variabilis fountainensis Fenton and Fenton, 1922.—Fenton and Fenton (1922, p. 76, pl. 2, figs. 1–3).

A few species previously assigned to *Zygospira* share similarities with other atrypide genera or have already been assigned to other genera, including the species below.

Athyris headi Billings, 1862.—See Meek (1873, p. 127, pl. 11 a–d) for description of species as *Zygospira*; assigned to *Catazyga* as type species by Hall and Clarke (1894).

Zygospira uphami Winchell and Schuchert, 1893.—Convex profile suggests affinities with *Catazyga*, but spiralia shape and configuration remain unknown.

Zygospira maynei Roy, 1941.—See Bolton (2000); very convex profile suggests affinities with a different lineage. *Idiospira* was suggested by Bolton, but the illustrated specimen would be unusual for *Idiospira*. It seems likely that the shells described by Roy (1941) belong to a slightly older collection and may be *Anazyga* while the shells described and figured by Bolton (2000) are an unusual species of *Zygospira* or a different lineage altogether.

Zygospira putilla Hall and Clarke, 1894.—Hall and Clarke (1894, p. 157, fig. 150, pl. 54, figs. 35–37; not plate 55 as indicated in text); considered *Eospirigerina* by Amsden (1974, p. 72), the elongate shell form would be unusual for *Zygospira* and possesses a distinctive plate in the interior (see Amsden, 1974, text-figs. 49, 50) not known in other *Zygospira* species and more variation in rib bifurcation (Amsden, 1974, text-fig. 48).

The affinities of the following species remain uncertain because they are only known from a few specimens, sometimes are poorly preserved, and/or their external morphology is unusual for the genus.

Zygospira sulcata Howe, 1965.—Howe (1965, p. 655–656, pl. 81, figs. 9–12); fine ribs indicate affinities with *Anazyga*, but species is known only from poorly preserved fragmentary material.

Zygospira tantilla Bradley, 1921.—Also resembles *Anazyga* in shape, but if specimens are truly Richmondian in age, this is by far the youngest known species of *Anazyga*.

Two species of *Zygospira* from Scotland and Norway cannot be confidently assigned to *Zygospira* and require further material to make any definitive assignment.

Zygospira orbis Reed, 1917.—Reed (1917, p. 944, pl. 24, figs. 24–27); was assigned to *Zygospira* from Scotland (part of Laurentia during the Ordovician) alongside an unnamed questionably assigned specimen (*Zygospira?*, sp. Reed, 1917, pl. 24, figs. 28, 29). Described to have a broad dorsal depression on the ventral valve and a fold containing two ribs on the dorsal valve, suggesting affinities with *Zygospira*. The questionably assigned specimens are unusually elongate for *Zygospira*, however, and may belong to the Atrypinidae. Additional material will need to be examined before anything definitive can be said about either species.

Zygospira meldalensis Reed, 1932.—Reed (1932, p. 144, pl. 22, figs. 12, 12a); from the Upper Hovin Group of Norway, only known from a single poorly preserved specimen that cannot definitively be located, although a single specimen within a limestone fragment bearing the same specimen number contains an external mold of the specimen consisting of only the anterior was described by Neuman et al. (1997); however, the configuration of the brachidial structures remains unknown; resembles *Z. kentuckiensis* in size and shape.

Several species from the Kazakh terranes and North and South China previously have been assigned to *Zygospira*, but these have almost all since been reassigned to other genera. These all differ from *Zygospira* in lacking a complete jugum, although the spiralia of many species are still inadequately known. Most also possess a ventral sulcus and dorsal fold rather

than the ventral fold and dorsal sulcus that is typical of *Zygospira*. These species include:

Zygospira parva Rukavishnikova, 1956.—Rukavishnikova (1956, p. 162–163, pl. 5, figs. 14–16); assigned to *Schachriomonia* by Popov et al., 1999.

Zygospira qinghaiensis Xu in Jin et al., 1979.—Xu in Jin et al. (1979, p. 108, pl. 21, figs. 10–15, 19–21, text fig. 51; see also Rong et al., 2017, p. 188).

Zygospira shaanxiensis Fu, 1982.—Fu (1982, p. 145, pl. 39, fig. 9a–c; see also Rong et al., 2017, p. 192).

Zygospira (Kuzgunia) bankanasensis Klenina, Nikitin, and Popov, 1984.—Klenina et al. (1984, p. 116); assigned to *Sulcatospira* by Popov et al., 1999.

Zygospira (Sulcatospira) plicata Xu in Jin et al., 1979.—Xu in Jin et al. (1979; see also Rong et al., 2017, p. 190).

Zygospira carinata Percival, 1991 (p. 165, 167, 169; figs. 20, 27) is one of the few probable *Zygospira* species from outside Laurentia. It is unusual for the genus in that it has a prominent ventral medial rib rather than an interspace, and generally has fewer and coarser ribs. The shell of this species is more elongate than typical *Zygospira*, which are usually shorter in length than width. The spiralia and associated structures remain unknown for this species, so it can only be assigned to the genus provisionally until additional specimens with intact shell interiors are found. The unusual morphology may reflect the relative isolation of Australia (part of Gondwana in the Ordovician) from Laurentia and lends the species considerable paleobiogeographic significance.

Zygospira modesta kagawongensis Caley, 1936 (Caley, 1936, p. 58) was published without description or figures and thus should be considered a nomen nudum (see also Copper, 1977). A search of the database of the Royal Ontario Museum where the other specimens described by Caley (1936) were deposited revealed no results.

Diagnosis.—See Copper, 2002.

Occurrence.—Common in middle to upper Katian (Maysvillian–Richmondian) rocks of eastern North America, but its range outside of Laurentia is limited. Species from lower Katian strata are almost certainly *Anazyga* or another early atrypide genus, but need re-examination.

Remarks.—*Zygospira* has referred to a number of different species in literature. Here, the Treatise on Invertebrate Paleontology (Copper, 2002) is followed in referring to only the coarsely ribbed species with mediodorsally directed spiralia. Other authors have included species more recently considered to belong to *Anazyga* (see above). These species generally have spiralia directed medially rather than dorso-medially (although the internal shell morphology has not yet been documented for all species) and can be recognized externally by a generally more convex dorsal valve, weaker ribs, and often is significantly smaller. *Zygospira* also characteristically has one or two very prominent ribs on each side of the ventral fold, a feature that is usually absent from species that tentatively have been assigned to *Anazyga*.

Anazyga has been reported only from rocks that are older than *Zygospira*, being particularly abundant in the lower Katian

Trenton Limestone and equivalent strata in eastern North America (Chatfieldian and Maysvillian in North American terminology). The species referred to *Zygospira* above are all middle to late Katian (Maysvillian–Richmondian in North American terminology) in age. This led Copper (1977) to suggest that *Anazyga* was a likely ancestor to *Zygospira*, but this hypothesis has yet to be tested in a broader cladistic analysis of the lineage.

The paleoecology of *Zygospira* and other early atrypides has not yet been examined in detail (although see Copper, 1977, for a brief discussion), but the genus is known to occur in dense clusters (Fig. 3). These dense accumulations could reflect an opportunistic life habit where *Zygospira* may have been able to multiply rapidly and take advantage of regular disruptions in the environment, such as regular storm events in eastern North America during the Ordovician (e.g., Brookfield and Brett, 1988; Kerr and Eyles, 1991; Jennette and Pryor, 1993). Alternatively, these dense shell beds could represent periods of quiescence with reduced sedimentation that enabled *Zygospira* to proliferate (Dattilo et al., 2008, 2012). McFarland et al. (1999) analyzed shell beds in the lower Katian Verulam Formation of Ontario where the older atrypide *Anazyga* is known to be locally abundant and concluded that most of the shell beds must have been formed from allogenic processes such as winnowing by storms. They mentioned that the dense accumulations of atrypide shells may have been at least partially derived from autogenic processes given their common preservation in apparent life position.

These dense clusters are also sometimes associated with filter feeders, such as bryozoans and crinoids (Fig. 3). The smaller *Z. modesta* are preserved in positions that suggest that they may have been attached to these filter feeders when alive via their pedicle (Fig. 3.2, 3.3). There is no evidence of such associations with the larger *Zygospira* species thus far, however, and the extent of this phenomenon across the Anazygidae has not yet been investigated.

Zygospira modesta Say in Hall, 1847
Figures 4, 5

1847 *Atrypa modesta* Say in Hall, p. 141, pl. 33, fig. 15a, b; not pl. 33, fig. 15c.

1862 *Zygospira modesta*; Hall, p. 155, text-figs. 1, 2.

1879 *Zygospira concentrica*; Ulrich, p. 14, pl. 7, figs. 10, 10a, b.

1894 *Zygospira modesta*; Hall and Clarke, pl. 54, figs. 7–10, 12.

1910 *Zygospira modesta*; Foerste, p. 29, pl. 2, fig. 15a, b.

1924 *Zygospira modesta*; Foerste, p. 127, pl. 10, fig. 21a, b.

1977 *Zygospira modesta*; Copper, p. 303, pl. 37, figs. 1–8, text-figs. 3, 4.

1988 *Zygospira modesta*; Howe, p. 205, fig. 2.1–2.7.

Types.—Lectotype AMNH 29835 (formerly AMNH 1356A) selected by Foerste (1910) from Hall's collection at the AMNH (Fig. 3, AMNH 1356a–d). The specimen tags associated with the type collection lack specific information on the location of the original type locality and strata from which the specimens were collected. The older, typed tags indicate that specimens were collected from the Hudson River Group at Cincinnati, Ohio. These are unlikely to be

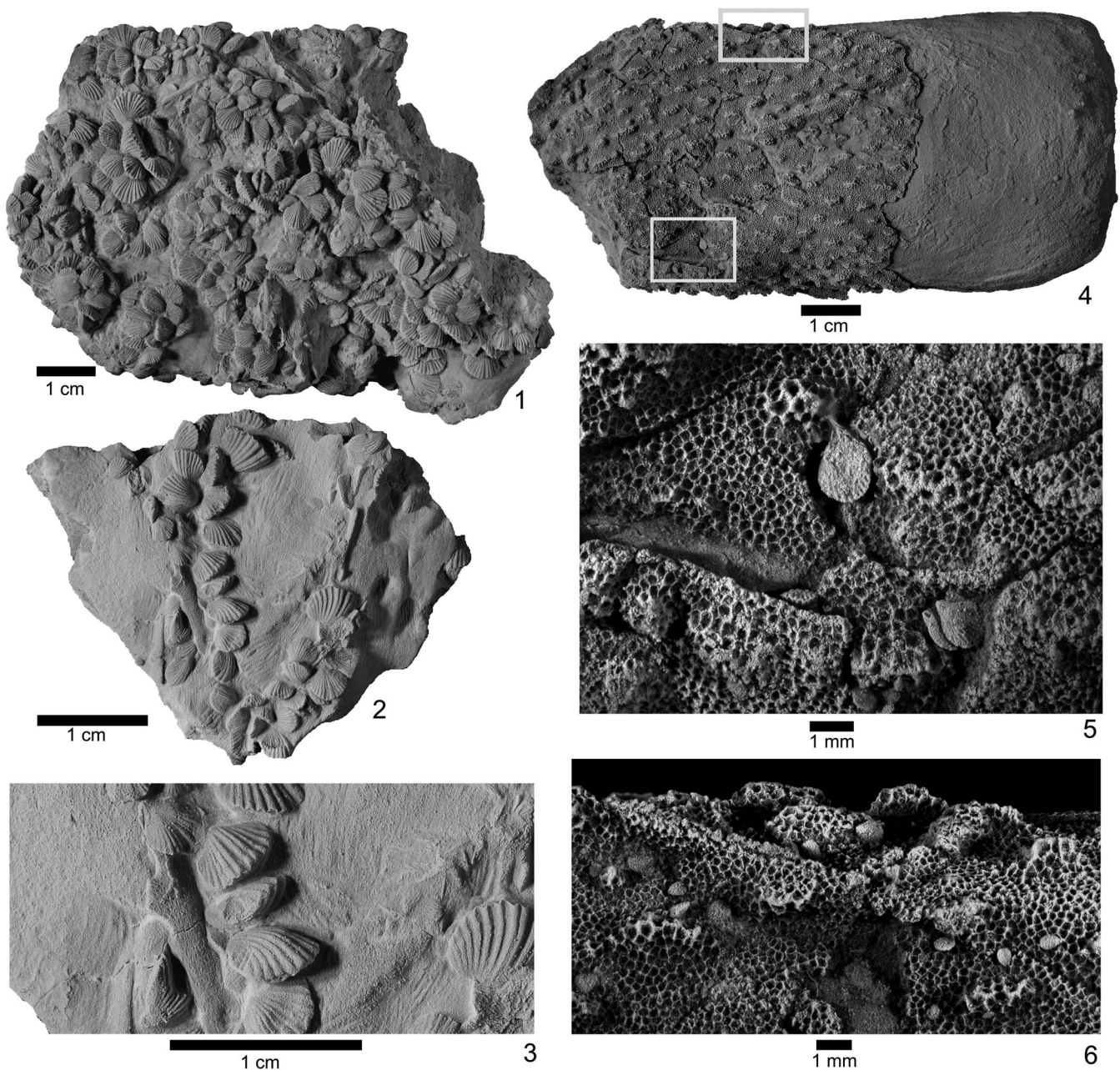


Figure 3. *Zygospira modesta* attached to bryozoans from Tanner's Creek Formation in Indiana. (1) PE 16630, dense cluster of *Zygospira modesta* from near Brookville; (2, 3) FM PE16583 *Zygospira modesta* preserved attached to bryozoan; (4–6) FM PE16631 *Zygospira modesta* attached to *Spatiopora corticans* (Nicholson, 1874) from near Clifton; (5) and (6) magnified views of (4) in blue and green, respectively. Scale bars for (1–4) = 1 cm; (5, 6) = 1 mm.

the original tags, however, because they refer to the specimens as *Zygospira* while Hall (1847) described the specimens as a species of *Atrypa*. The newer tags suggest that the specimens are from the Trenton Limestone, but this is highly unlikely. Foerste (1910) suggested that similar specimens are known from the Fairmount beds near Hamilton, Ohio, which should be considered the type locality. Although this is a reasonable inference, in the absence of any further information regarding their origin, all that can be said is that the types were likely, but not definitively, collected from Cincinnati-aged strata in the area around Cincinnati.

One of the types, now labeled AMNH 29837 but formerly labeled AMNH 1356C, is characteristic of *Z. cincinnatiensis* (see below). We tentatively assign it to that species, but a more extensive review is needed to determine the full range of morphological variability within *Z. cincinnatiensis*, along with a more extensive examination of the internal morphology of both species.

There are 12 unfigured paratypes deposited in the collection along with 2 casts under the collection number AMNH 1356d–1356o (probably following the original numbering scheme—no updated numbers are associated with the specimens).

Diagnosis.—See Copper, 2002.

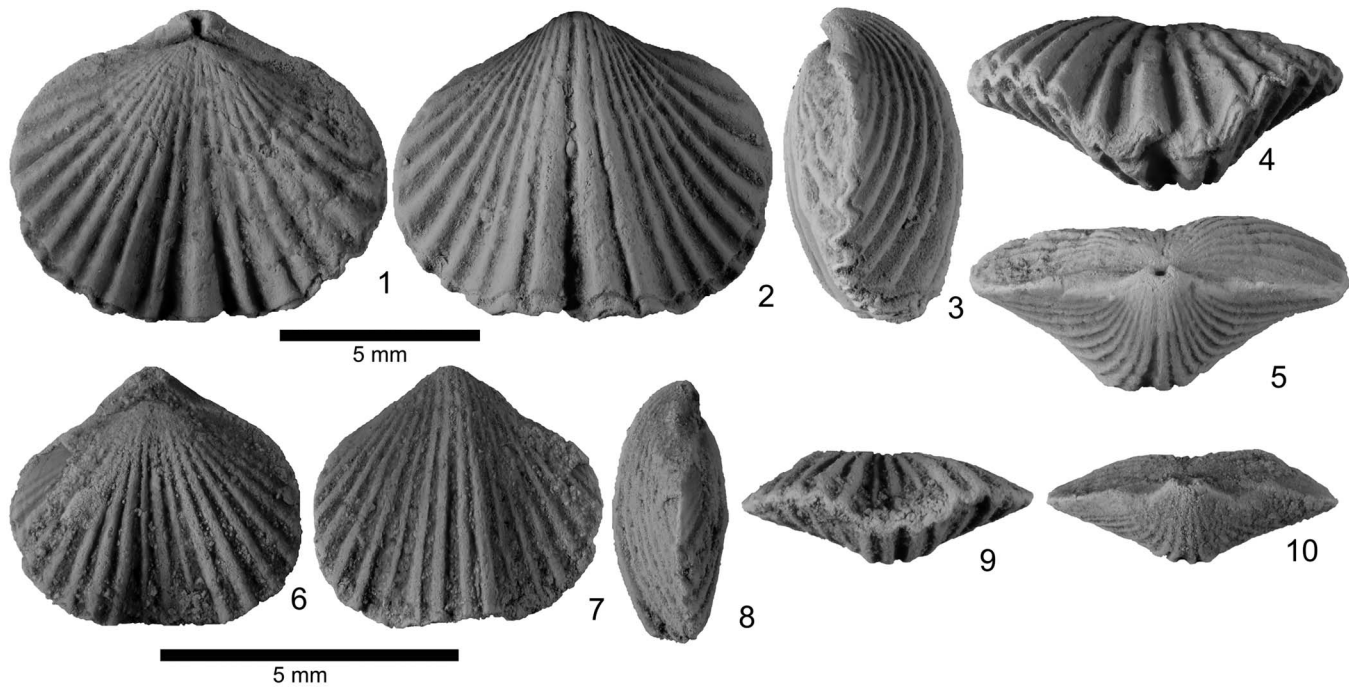


Figure 4. (1–10) *Zygospira modesta* types from the Hudson River Group (Katian) near Cincinnati, Ohio. (1–5) Lectotype AMNH 29835, dorsal, ventral, lateral, anterior, and posterior views; (6–10) paratype AMNH 29836, dorsal, ventral, lateral, anterior, and posterior views. Scale bars = 5 mm. This is part of Hall's type collection at the AMNH along with several unfigured paratypes in the collection (labeled AMNH 1356d–o).

Occurrence.—*Zygospira modesta* is most common in middle Katian (Maysvillian) strata of eastern North America near Cincinnati in Ohio, Kentucky, Indiana, and Illinois, but has been reported from upper Katian (Richmondian) strata as well. It is considered the most widespread species of *Zygospira*, but other species are commonly misidentified as *Z. modesta* in historical collections.

Description.—Shell small, measuring on average 6.4 mm in length, 7.4 mm in width, and 3.7 mm in depth (Fig. 6, Table 1); outline subpentagonal and always significantly wider than long (mean of 87% as long as wide) with ventribiconvex lateral profile with ventral valve ~1.5 times as deep as dorsal valve. Astrophic hingeline with rounded lateral flanks. Anterior margin strongly unisulcate forming prominent flat-topped tongue. Ribs strong and rounded, simple with rare bifurcations and expanding towards anterior, numbering between 14–20, but usually 16–18. Fine growth lamellae on well-preserved specimens cover the shell.

Ventral umbo strongly curved featuring prominent apical foramen. Strongly anacline interarea that becomes nearly perpendicular to commissural plane towards tip, projecting over hingeline. Delthyrium mostly open, flanked by minute deltidial plates visible on well-preserved specimens. Carinate, with flat-topped ventral fold with medial interspace flanked by a pair of prominent ribs on each side.

Dorsal umbo minute and incurved, obscuring open dorsal interarea. Broad dorsal sulcus usually containing single prominent medial rib flanked by two smaller ribs beginning near umbo with a single rib forming each flank of the sulcus. Medial rib larger than other ribs on shell, but similar in prominence to ribs flanking ventral fold.

Dental plates thin, thickening slightly towards anterior, especially at bases, with small dental cavities infilled with secondary shell material towards posterior. Small teeth pointed dorso-medially with weak medial grooves forming accessory lobes.

Minute myophragm near posterior of dorsal valve. Sockets small with weak medial ridge forming tight interlocking hinge with teeth. Hinge plates fuse to form cruralium, rising off the base of the ventral valve towards anterior and supported by a medial septum. Crural bases flat with crura becoming rod-like towards anterior. Dorsomedially directed spiralia consisting of up to four whorls connected by ventro-anteriorly pointed jugum high in dorsal valve near posterior of spiralia.

Remarks.—*Zygospira modesta* appears to be the most common and long ranging species of *Zygospira* in the eastern United States. Middle Katian (Maysvillian) specimens of *Z. modesta* have been widely reported as occurring alongside *Z. cincinnatiensis* James in Meek, 1873, and *Z. concentrica* Ulrich, 1879 (the latter species herein synonymized with *Z. modesta*) in this region while late Katian (Richmondian) specimens occur in strata of similar age to *Z. kentuckiensis*. The younger late Katian specimens are notably longer than wide in comparison to the middle Katian specimens based on the collections measured herein (Fig. 6).

The late Katian (Richmondian) *Z. modesta* primarily differ from *Z. kentuckiensis* based on their much smaller size, a more prominent and angular dorsal sulcus and ventral fold, and fewer ribs on the flanks of the shell than *Z. kentuckiensis*. Given that ribs generally do not increase in number as the shell grows (other than a few uncommon examples of bifurcation in some

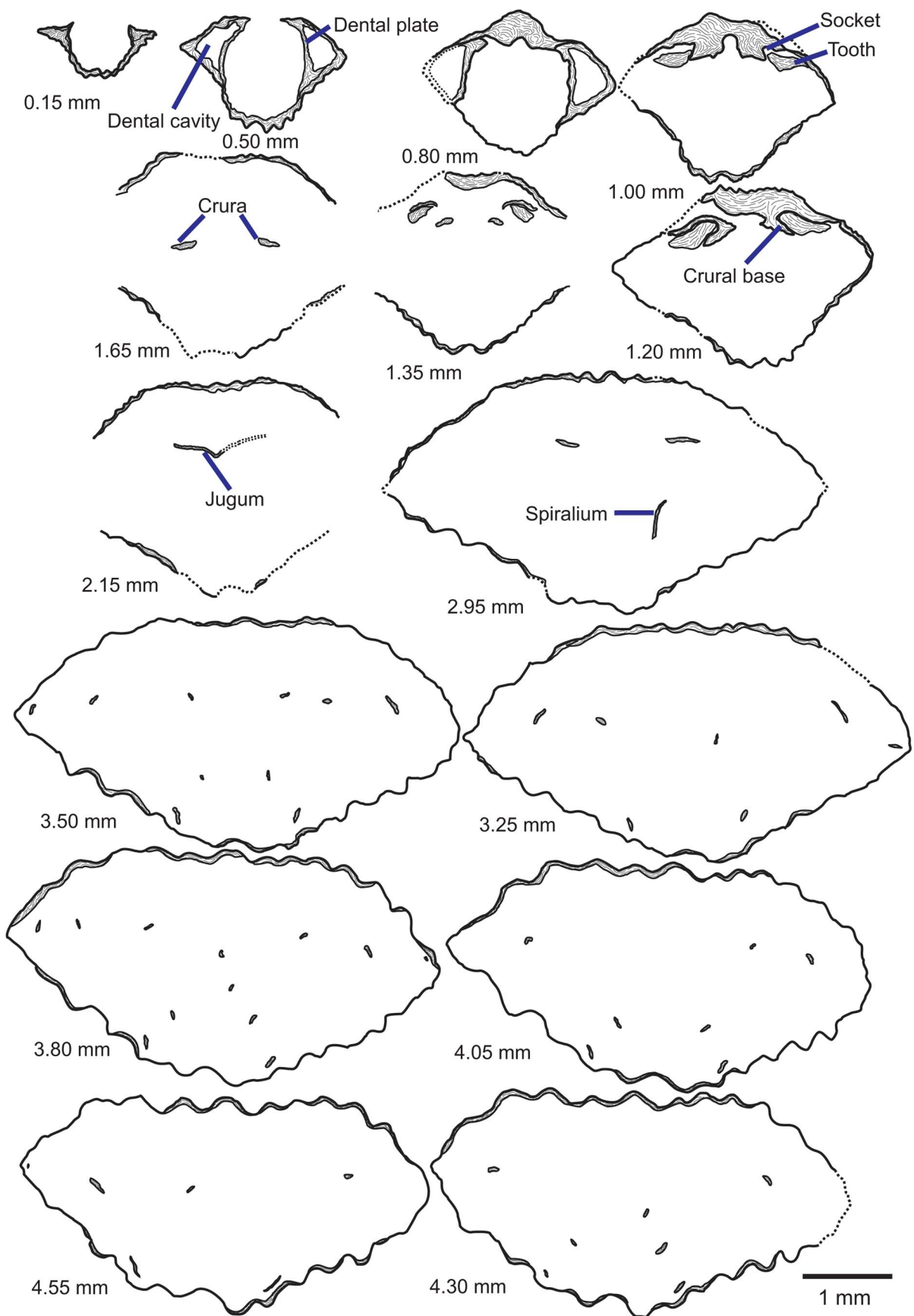


Figure 5. Tracings of serial sections of a *Zygospira modesta* shell from the Grant Lake Formation near Sharonville in the region around Cincinnati, Ohio (CMC IP96903). Numbers represent distance from posterior of ventral umbo in mm at which the section was ground. Scale bar = 1 mm.

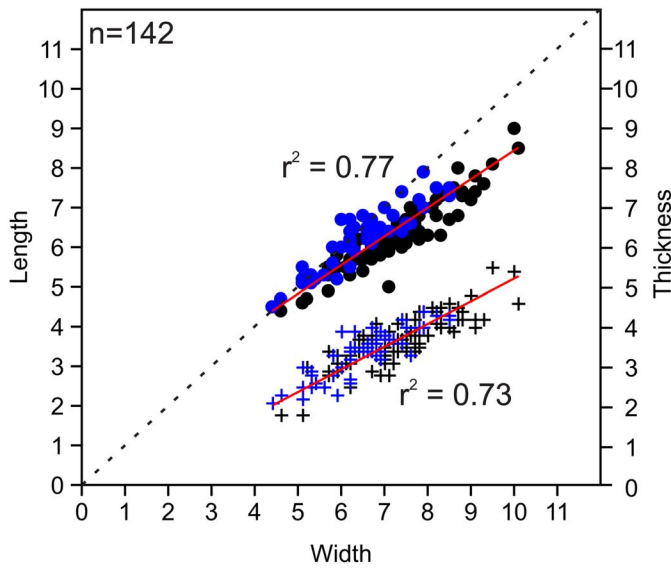


Figure 6. Bivariate plots of length vs. width and thickness vs. width for measured *Zygospira modesta* specimens. Black = middle Katian (Maysvillian) specimens (n=92); blue = late Katian (Richmondian) specimens (n=50); circles indicate length vs. width; plus signs represent thickness vs. width. The r^2 values shown (red lines) represent values for combined ages; r^2 values for individual datasets are as follows: length/width Maysvillian=0.85759; thickness/width Maysvillian=0.70637; length/width Richmondian=0.82215; thickness/width Richmondian=0.75109.

specimens), this seems a reliable diagnostic character to differentiate the species.

The earlier species are more problematic to differentiate. *Zygospira cincinnatiensis* was differentiated from *Z. modesta* in having fewer ribs on the lateral flanks of the shell (typically five) and generally larger size (Foerste, 1910). It would be difficult to differentiate *Z. cincinnatiensis* and *Z. modesta* of similar sizes, however, perhaps indicating that *Z. cincinnatiensis* is more common than reports would suggest. *Zygospira cincinnatiensis* is tentatively retained here, but the division between these specimens remains somewhat indistinct.

Zygospira concentrica was differentiated from *Z. modesta* based on having stronger concentric striae, no radiating plications (ribs), straighter posterior margins, and a more pointed ventral beak (Foerste, 1910). The apparent lack of ribs in specimens of *Z. concentrica* seems to reflect preservation rather than any real diagnostic difference because examination of collections of species from the Cincinnati region revealed weak ribs that appear to have been abraded in most specimens identified as *Z. concentrica*. Furthermore, the strong concentric striae probably represent a prominent interruption in the growth of

Table 1. Summary statistics for *Zygospira modesta* specimens measured for this study (n = 92). L = length; W = width; T = thickness (depth); Ad = deflection at anterior of the commissure (all in mm); Aa = apical angle in degrees; R = number of ribs counted on the ventral valve; L/W = length/width ratio; T/W = thickness/width ratio.

	L	W	T	Ad	Aa	R	L/W	T/W
Mean	6.4	7.4	3.7	1.0	116	17	0.87	0.50
Median	6.2	7.3	3.7	0.9	115	18	0.87	0.50
SD	0.8	1.1	0.6	0.4	7.14	1.6	0.050	0.049
Min	4.4	4.6	1.8	0.5	101	14	0.70	0.35
Max	9	10.1	5.5	2.8	131	20	0.99	0.61

the shell and should not be considered a reliable diagnostic character. *Zygospira concentrica* is thus herein synonymized with *Z. modesta*.

An in-depth analysis of the configuration of the spiralia is outside the scope of this study, but it is perhaps notable that the jugum appears slightly closer to the posterior in our sectioned specimen than in the sections illustrated by Copper (1977). It is still located postero-dorsally, however.

Zygospira cincinnatiensis James in Meek, 1873
Figure 7

- 1847 *Atrypa modesta* Say in Hall, p. 141, pl. 33, fig. 15c; not pl. 33, fig. 15a, b.
- 1873 *Zygospira cincinnatiensis* James in Meek, p. 126, pl. 11, fig. 5a–c.
- 1894 *Zygospira cincinnatiensis*; Hall and Clarke, pl. 54, figs. 13, 14.
- 1910 *Zygospira cincinnatiensis*; Foerste, p. 30, pl. 6, figs. 16a, b.

Types.—Foerste (1910) explained that the original types (collected by James and described by Meek, but credited to James by Meek) were not deposited in a known institution and suggested that the collection numbered UC164 from the Fairmount Beds near Cincinnati, Ohio, be regarded as “typical” (Foerste, 1910, p. 30). He reasoned that Meek would have received the original type specimens from James whose collection was then housed at the University of Chicago but is now located at the Field Museum in Chicago. One of the specimens from the original UC164 collection has been selected here as a more typical neotype (FM UC164-a).

A single specimen in the Field Museum collection is marked as a plesiotype (FM UC12391) and was apparently figured by Hall and Clarke (1894), but this specimen has far more ribs than is typical for *Z. cincinnatiensis*. It is unlikely that this specimen was collected from the same locality, although this is not clear from the label associated with the specimen. In museum records, this is noted as belonging to the Hall collection while the other specimens are part of the James collection. This may represent a single, unusual example of *Z. cincinnatiensis* or could represent a subspecies, but it is difficult to draw any conclusions based on a single shell. Although it has a similar number of ribs to *Z. kentuckiensis*, it more closely resembles *Z. cincinnatiensis* in having a more prominent fold and sulcus.

Occurrence.—*Zygospira cincinnatiensis* seems to be restricted to the middle Katian (Maysvillian) in the Cincinnati area, although further detailed study of larger collections of *Zygospira* may extend its range into the late Katian (Richmondian). Foerste (1910) mentioned that ancestral forms of the species are found in the Edenian (lower to middle Katian) that have less conspicuous bifurcation of ribs on the fold. Given the doubtful utility of rib bifurcation as a diagnostic feature in this genus, these specimens should likely be considered *Z. cincinnatiensis* as well.

Remarks.—*Zygospira cincinnatiensis* was distinguished from *Z. modesta* by Meek based on the presence of bifurcating ribs

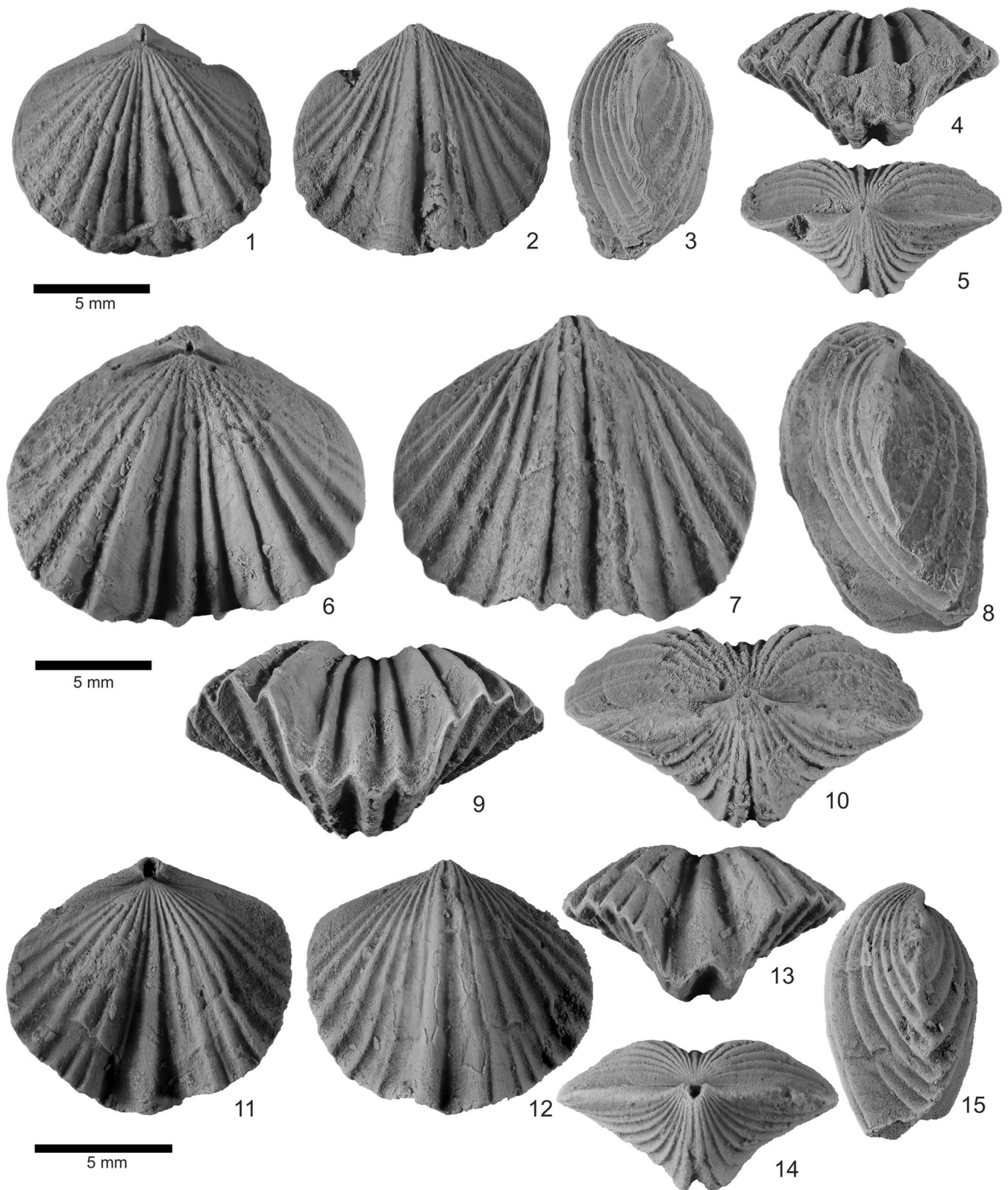


Figure 7. *Zygospira cincinnatiensis* types from the Fairmount Beds (Cincinnatian = Katian) near Cincinnati, Ohio. (1–5) Lectotype FM UC16-a: dorsal, ventral, lateral, anterior, posterior views; (6–10) figured paratype FM UC164-b: dorsal, ventral, lateral, anterior, posterior views. *Zygospira cincinnatiensis* from the Hudson River Group near Cincinnati, Ohio (from Hall's type collection for *Atrypa modesta*). (11–15) AMNH 29837, dorsal, ventral, anterior, posterior, and lateral views. Scale bars = 5 mm.

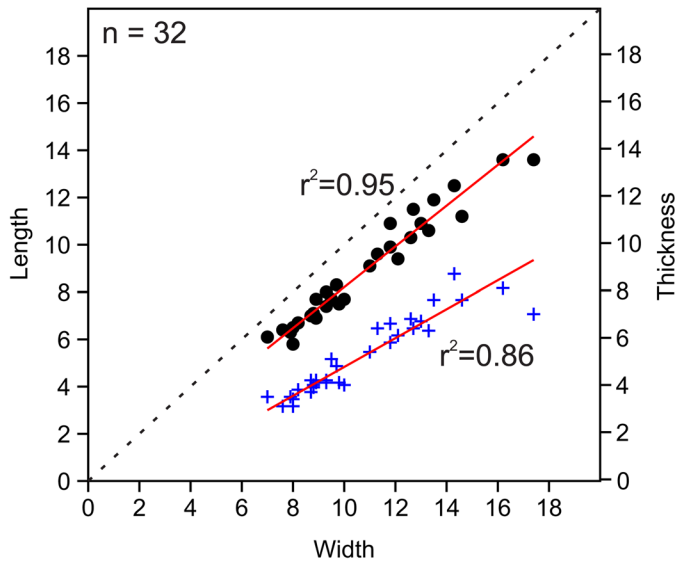


Figure 8. Bivariate plots of length vs. width and thickness vs. width for measured *Zygospira cincinnatiensis* specimens. Circles indicate length vs. width; plus signs represent thickness vs. width.

on the ventral fold, but bifurcating ribs are sometimes present in large collections of *Z. modesta* as well and may not be a reliable diagnostic character (also noted by Foerste, 1910). The elevated ventral fold that was mentioned, however, is more diagnostic, and *Z. cincinnatiensis* is, on average, larger than *Z. modesta* (Fig. 8). Both differ from the similar-sized *Z. kentuckiensis* in having a significantly more prominent fold with a matching prominent medial rib in the dorsal sulcus. *Zygospira cincinnatiensis* also has consistently fewer ribs than both *Z. modesta* and *Z. kentuckiensis* when large collections are compared.

It is difficult to determine the precise abundance and stratigraphic range of this species. Although apparently easily differentiated from *Z. modesta* in isolated collections (especially when the *Z. cincinnatiensis* are larger), it can be difficult to differentiate these species when specimens of similar sizes are compared. When *Z. cincinnatiensis* is small, the shells very closely resemble large *Z. modesta* in every way other than having fewer ribs on average (14–16 in *Z. cincinnatiensis* rather than 16–18 in *Z. modesta*). These smaller shells are nearly identical in terms of length, width, and depth to *Z. modesta* and lack the distinctive raised fold of larger shells (Table 2). Larger shells are easily differentiated from *Z. kentuckiensis* shells, which have more ribs and much less prominent dorsal sulcus and ventral fold.

Table 2. Summary statistics for *Zygospira cincinnatiensis* specimens measured for this study (n = 32). L = length; W = width; T = thickness (depth); Ad = deflection at anterior of the commissure (all in mm); Aa = apical angle in degrees; R = number of ribs counted on the ventral valve; L/W = length/width ratio; T/W = thickness/width ratio.

	L	W	T	Ad	Aa	R	L/W	T/W
Mean	8.8	10.7	5.2	2.9	114	15	0.82	0.48
Median	7.9	9.8	4.5	2.5	115	15	0.81	0.48
SD	2.28	2.65	1.62	1.03	7.19	1.77	0.04	0.05
Min	5.8	7	3.1	1.5	100	12	0.73	0.39
Max	13.6	17.4	8.7	4.9	129	22	0.92	0.61

One of the paratypes of *Z. modesta* from the American Museum of Natural History (AMNH 29837) is herein considered *Z. cincinnatiensis*. It is unclear whether all the *Z. modesta* type specimens were collected from the same locality, but if that was the case, *Z. modesta* co-occurs with *Z. cincinnatiensis* as the species are currently defined.

Zygospira kentuckiensis James, 1878
 Figures 9, 10.1–10.10, 11

- 1878 *Zygospira modesta* var. *kentuckyensis* James, p. 7.
- 1889 *Zygospira kentuckiensis* James in Nettelroth, p. 138, pl. 34, figs. 21–25.
- 1894 *Zygospira kentuckiensis*; Hall and Clarke, pl. 54, figs. 11, 15, 16.
- 1924 *Zygospira kentuckiensis*; Foerste, p. 127, pl. 10, fig. 20a–c, pl. 15, figs. 1a–p, 2a, b, 4a–c.

Types.—The species was first mentioned by James (1878) in brief, describing fossils from Nettelroth from the upper part of the Cincinnati Group, Jefferson County, Kentucky, 18 miles east of Louisville. Nettelroth (1889) later described the species in more detail himself, explaining that the species is found in different places in Oldham County, Kentucky, and that it is found in great abundance at Taylor’s Station but also on the Shelby Railroad in the shales of the Hudson River Group. Unfortunately, there is no indication of where Nettelroth may have deposited his collection, although a large collection of Nettelroth’s fossils was apparently donated to the US National Museum (now Smithsonian Institution) after his death.

A small collection of six shells within James’ collection at the Field Museum labeled UC96 apparently were collected from the exact locality mentioned in James’ (1878) initial description of the species (Jefferson County, 18 miles east of Louisville, Kentucky) and is thus likely the original type collection. A single specimen from UC96 (FM UC96-a) is herein selected as a lectotype for *Z. kentuckiensis*.

Another collection of *Z. kentuckiensis* from the Field Museum labeled IP 12390 consists of a single well-preserved and carefully cleaned specimen. This shell apparently was illustrated by Hall and Clarke (1894), but the specimens from the James collection (FM UC96) have priority as types.

Diagnosis.—Shells large for the genus, usually wider than long, and ventribiconvex in profile; simple ribs numbering >18 but less prominent than other species; relatively weak ventral fold with medial interspace accentuated by two ribs on each flank and broad and shallow dorsal sulcus with weak medial ridge; thick dental plates and hinge plates for the genus with simple dorso-posteriorly located jugum; prominent teeth with hook-like transverse profile towards anterior.

Occurrence.—Upper Katian (Richmondian) of eastern North America. Common in the upper Fairmount (and correlated units) in the Cincinnati area and the Queenston Shale in Ontario.

Description.—Shell medium-sized, with a mean length of 9.8 mm, width of 10.9 mm, and a thickness of 5.5 mm

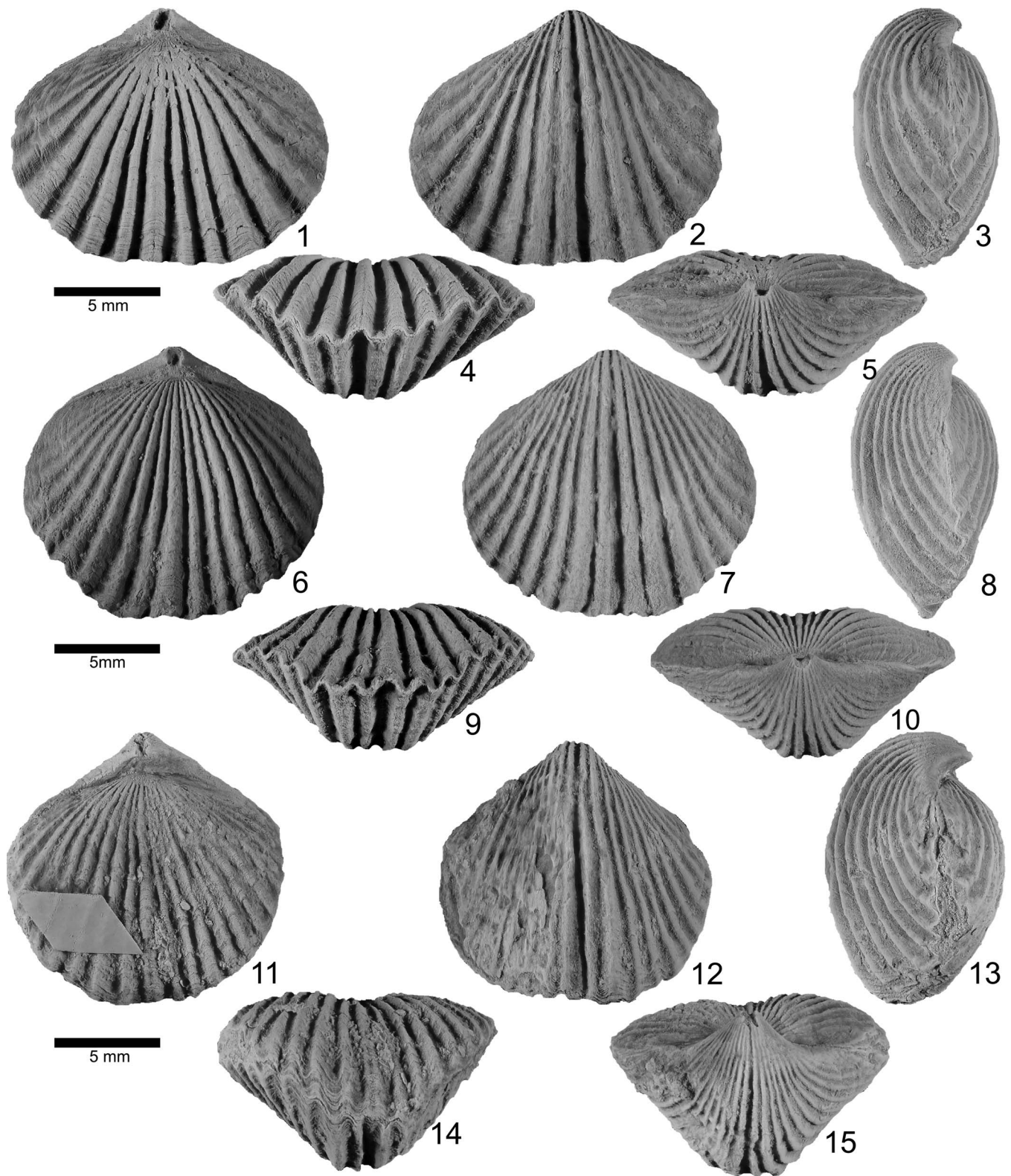


Figure 9. *Zygospira kentuckiensis*. (1–10) Types from Cincinnatian (= Katian) of Jefferson County, 18 miles east of Louisville, Kentucky; (1–5) neotype FM UC96-a dorsal, ventral, lateral, anterior, posterior views; (6–10) figured paratype FM UC96-b, dorsal, ventral, lateral, anterior, posterior views; (11–15) illustrated plesiotype FM UC 12390 in Hall and Clarke (1894) from Cincinnatian (=Katian) of Oldham County, Kentucky, dorsal, ventral, lateral, anterior, posterior views. Scale bars = 5 mm.

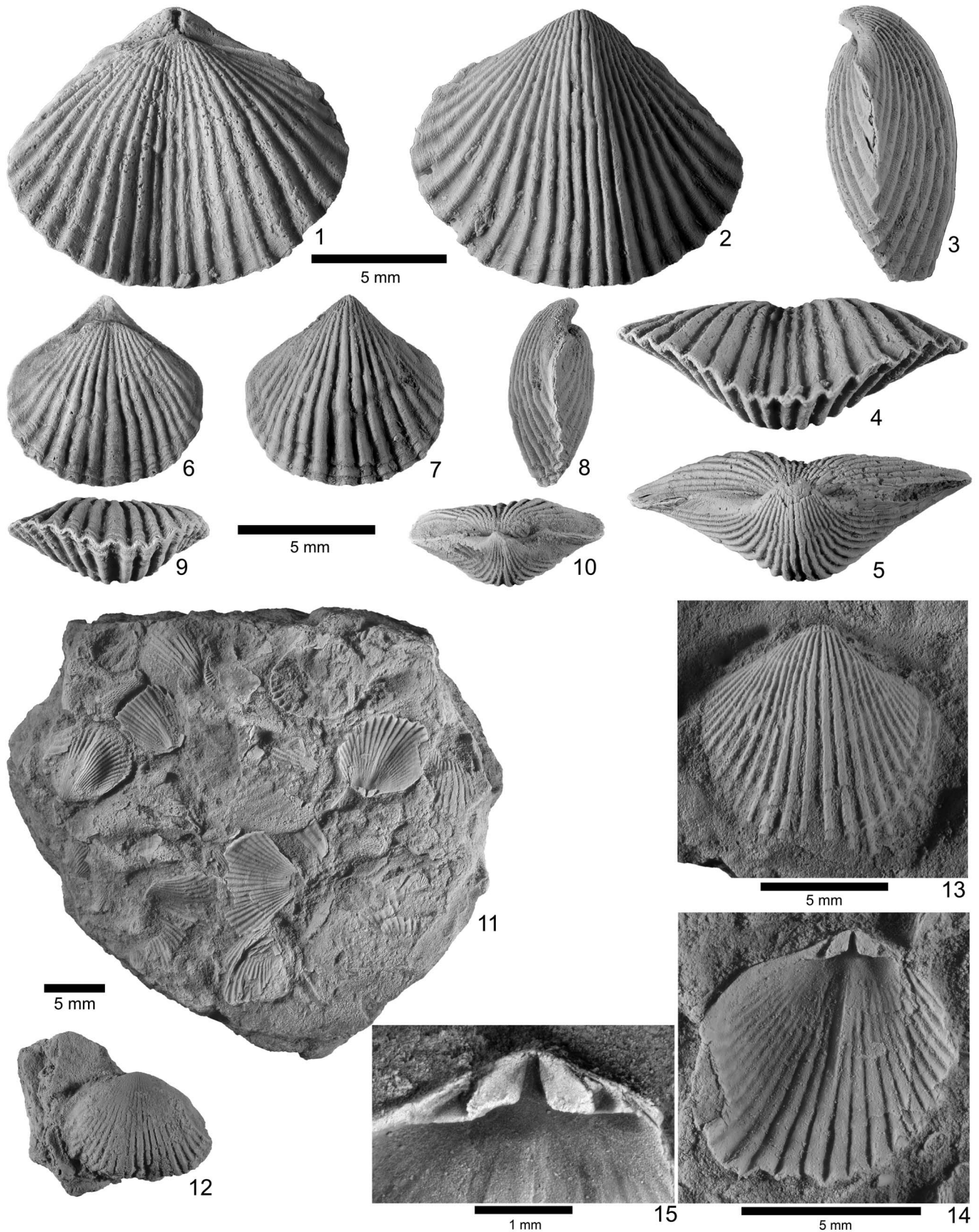


Figure 10. (1–10) *Zygospira kentuckiensis* from the Queenston Shale (late Katian) at Big Bay near Owen Sound, Ontario. (1–5) ROMIP 66852 dorsal, ventral, lateral, anterior, posterior views; (6–10) ROMIP 66853 dorsal, ventral, lateral, anterior, posterior views. (11–15) *Zygospira kentuckiensis meafordensis* from the Queenston Shale near Meaford, Ontario (GSC 8514); (11) small slab with several shells on surface; (12) dorsal valve attached to piece of rock; (13) exterior of isolated dorsal valve attached to slab in (11); (14) interior of dorsal valve attached to slab in (11); (15) magnified view of (14). Scale bars = 5 mm (1–14); = 1 mm (15).

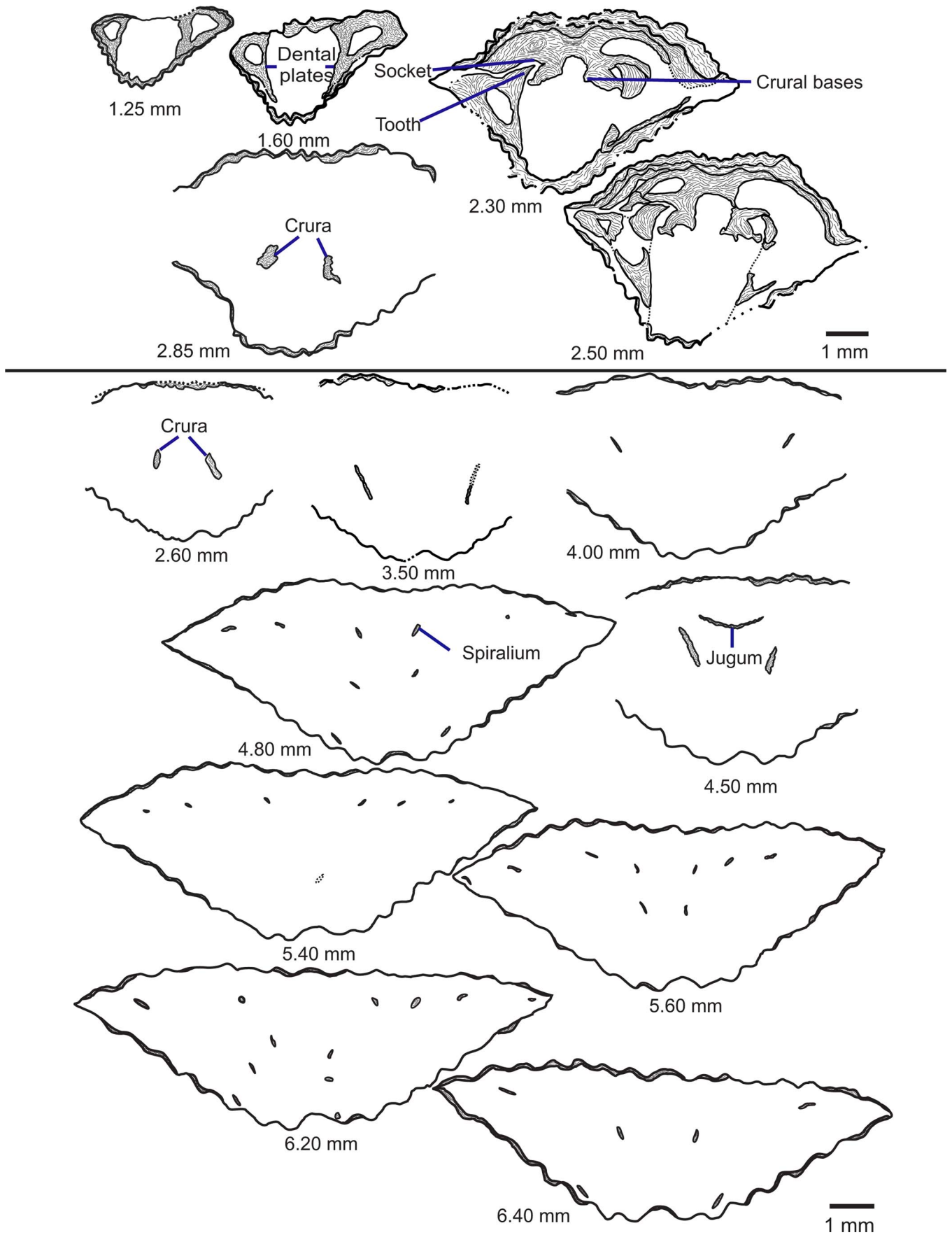


Figure 11. Tracings of serial sections of *Zygospira kentuckiensis* from the Queenston Formation at Big Bay near Owen Sound, Ontario (ROMIP 66854). Numbers represent distance from posterior of ventral umbo in mm at which the section was ground. Figure represents two sectioned shells to show the complete internal morphology of species (above and below line). Scale bar = 1 mm.

Table 3. Summary statistics for *Zygospira kentuckiensis* specimens measured for this study (n = 110). L = length; W = width; T = thickness (depth); Ad = deflection at anterior of the commissure (all in mm); Aa = apical angle in degrees; R = number of ribs counted on the ventral valve; L/W = length/ width ratio; T/W = thickness/width ratio.

	L	W	T	Ad	Aa	R	L/W	T/W
Mean	9.8	10.9	5.5	2.1	107	21	0.91	0.51
Median	9.8	11.0	5.6	1.9	106	20	0.90	0.51
SD	2.0	2.4	1.3	0.7	10	2	0.06	0.07
Min	5.5	5.4	2.4	0.8	82	15	0.78	0.39
Max	14.3	16.4	9.4	4.3	129	30	1.17	0.70

(Table 3); outline subpentagonal with outline wider than long (mean of 91% as long as wide) and ventribiconvex in lateral profile with ventral valve ~1.5 times as deep as dorsal valve. Astrophic hingeline with rounded lateral flanks. Anterior margin unisulcate forming prominent tongue. Ribs strong and rounded, simple with rare bifurcations and becoming larger towards anterior, typically numbering from 18–22 but can be more numerous (up to 28 in collections examined herein). Fine growth lamellae visible especially near anterior margin on well-preserved shells.

Ventral umbo strongly curved, featuring prominent apical foramen. Strongly anacline interarea that becomes nearly perpendicular to commissural plane towards tip, projecting over hinge-line. Delthyrium open, flanked by minute deltidial plates visible on some well-preserved specimens. Carinate, with less prominent flat-topped ventral fold with medial interspace flanked by a pair of prominent ribs on each side beginning near umbo.

Dorsal umbo minute and incurved, obscuring dorsal interarea. Broad, weak dorsal sulcus containing single prominent medial rib flanked by two smaller ribs beginning near umbo with no obvious prominent flanking ribs. Medial rib generally most prominent.

Dental plates moderately thick, divergent near posterior of ventral valve and convergent and thinner towards anterior. Dental cavities small. Teeth prominent, thickened, and curved, becoming increasingly curved towards anterior with variable weak accessory lobes separated from main lobe by weak medial groove. Ventral muscle scars poorly impressed but serial sections show evidence of small divergent adductor scars separated by slightly raised diductor platform.

Minute myophragm variably present near posterior of dorsal valve. Sockets curved with weak medial ridges, forming tight interlocking fit with teeth in dorsal valve. Median septum forms cruralium with hinge plates. Crural bases flat but crura become rod-like and laterally directed towards the anterior. Spiralia with dorso-medially directed apices consist of up to five whorls. Jugum located high in dorsal valve posterior to spiralia and curved towards anterior.

Remarks.—*Zygospira kentuckiensis* is similar in shape to the type species *Z. modesta*, but *Z. kentuckiensis* is significantly larger (Fig. 12) and has more numerous ribs (18–22 in *Z. kentuckiensis* in comparison to the 16–18 in *Z. modesta*). The two ribs that flank each side of the ventral medial fold are less prominent in this species in comparison to *Z. modesta* and form a shallower medial interspace on the ventral valve. The corresponding medial rib in the dorsal sulcus is also less prominent in *Z. kentuckiensis*.

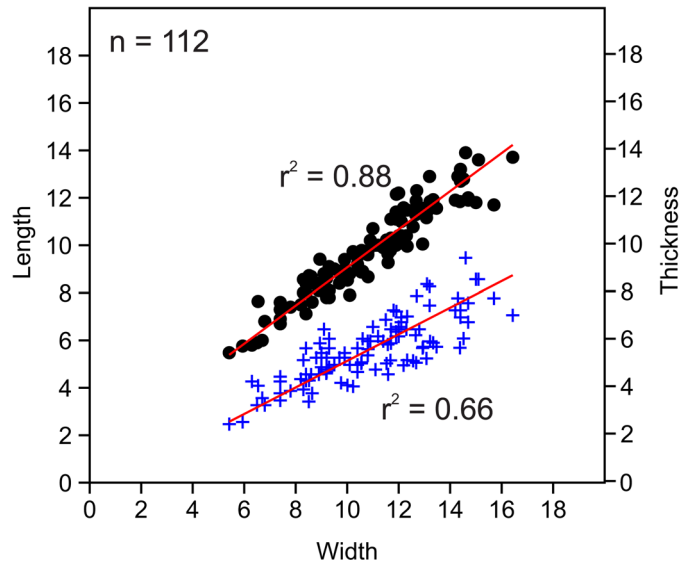


Figure 12. Bivariate plots of length vs. width and thickness vs. width for measure *Zygospira kentuckiensis* specimens. Circles indicate length vs. width; plus signs represent thickness vs. width.

There are slight differences in the interior of the shell as well. *Zygospira kentuckiensis* has significantly thicker dental plates and hinge structures in comparison to *Z. modesta*, as well as a more thickened posterior shell. This thickening may simply reflect the growth of a larger shell in the case of *Z. kentuckiensis*, but *Z. modesta* appears to have consistently thinner shells (based on the 5 specimens of each species sectioned here).

New material (Figs. 10.1–10.10, 11) was collected from the Queenston Formation at Big Bay near Owen Sound, Ontario (44.796537°N, 80.924807°W). These shells are similar to typical shells from the Cincinnati area but are slightly less convex. *Zygospira kentuckiensis* shells are common and numerous at this locality, forming low-diversity shell beds that form packstones comprised of this one species. Some of the exposed bedding planes show wave ripples, implying that *Zygospira* may have been able to tolerate relatively high energy environments above fair-weather wave base.

Zygospira richmondensis Caley, 1936, which was described from the Kagawong Formation (Richmondian, latest Katian) on Manitoulin Island, is represented by a single shell in the collections of the Royal Ontario Museum in Toronto (ROMIP 12448). This specimen resembles *Z. kentuckiensis* in size but seems to have a shallower fold and sulcus than is typical from the species. Given that all other characteristics are similar, however, it is here synonymized with *Z. kentuckiensis*.

Zygospira kentuckiensis meafordensis (Foerste, 1924)
Figure 10.11–10.14

1924 *Zygospira meafordensis* Foerste, p. 128, pl. 15, figs. 3a–c.

Types.—Lectotype GSC 8514A selected from GSC 8514 collected from Concession VIII, lot 24, ~4 miles northwest of Meaford (Foerste, 1924). Although these shells are illustrated as isolated shells, all three illustrated specimens are embedded in matrix. GSC 8514A is a ventral valve, GSC 8514B is a dorsal interior, and GSC 8514C is wider than a typical ventral valve.

Occurrence.—Specimens in the GSC collection were collected from only a single locality near Meaford, Ontario, from the Queenston Formation, which was described by Foerste (1924) as being at Concession VIII, lot 24, 4 miles northwest of Meaford. Two other localities are noted from the same region.

Remarks.—This species very closely resembles *Z. kentuckiensis* differing only in the much more numerous ribs (26–30 in this subspecies versus 18–22 in *Z. kentuckiensis*) and in its slightly smaller size, based on the limited number of specimens available for study. The fold and sulcus are broad and shallow, as is typical of *Z. kentuckiensis*, but Foerste noted that the slope between the fold and sulcus and the shell flanks is more gradual in these specimens in comparison to *Z. kentuckiensis*. Foerste (1924) described a cardinal process, but no cardinal process is visible in the type material (GSC 8514, Fig. 10). Given the extremely limited range of this species and overall similarity in shell size and shape, *Z. meafordensis* is herein regarded as a subspecies of *Zygospira kentuckiensis* with which it is found in Ontario.

Zygospira resupinata Wang, 1949
Figure 13

1949 *Zygospira resupinata* Wang, p. 18, pl. 10, fig. A (1–12).

Types.—SUI 1874 (holotype), SUI 1873 (paratype) from the *Cornulites* zone of the Brainard member; south corner of section 29, Fairfield township, Jackson County, Iowa (Wang, 1949).

Occurrence.—Upper Katian (Richmondian) of central North America.

Description.—See Wang (1949).

Remarks.—Wang only deposited two specimens in the State University of Iowa collection. Based on these two specimens (Table 4) and his descriptions, this species differs from all other species of *Zygospira* in being longer than wide and having far fewer costae (12 versus the typical 18–22 of *Z. modesta*). It also differs from other species in having a

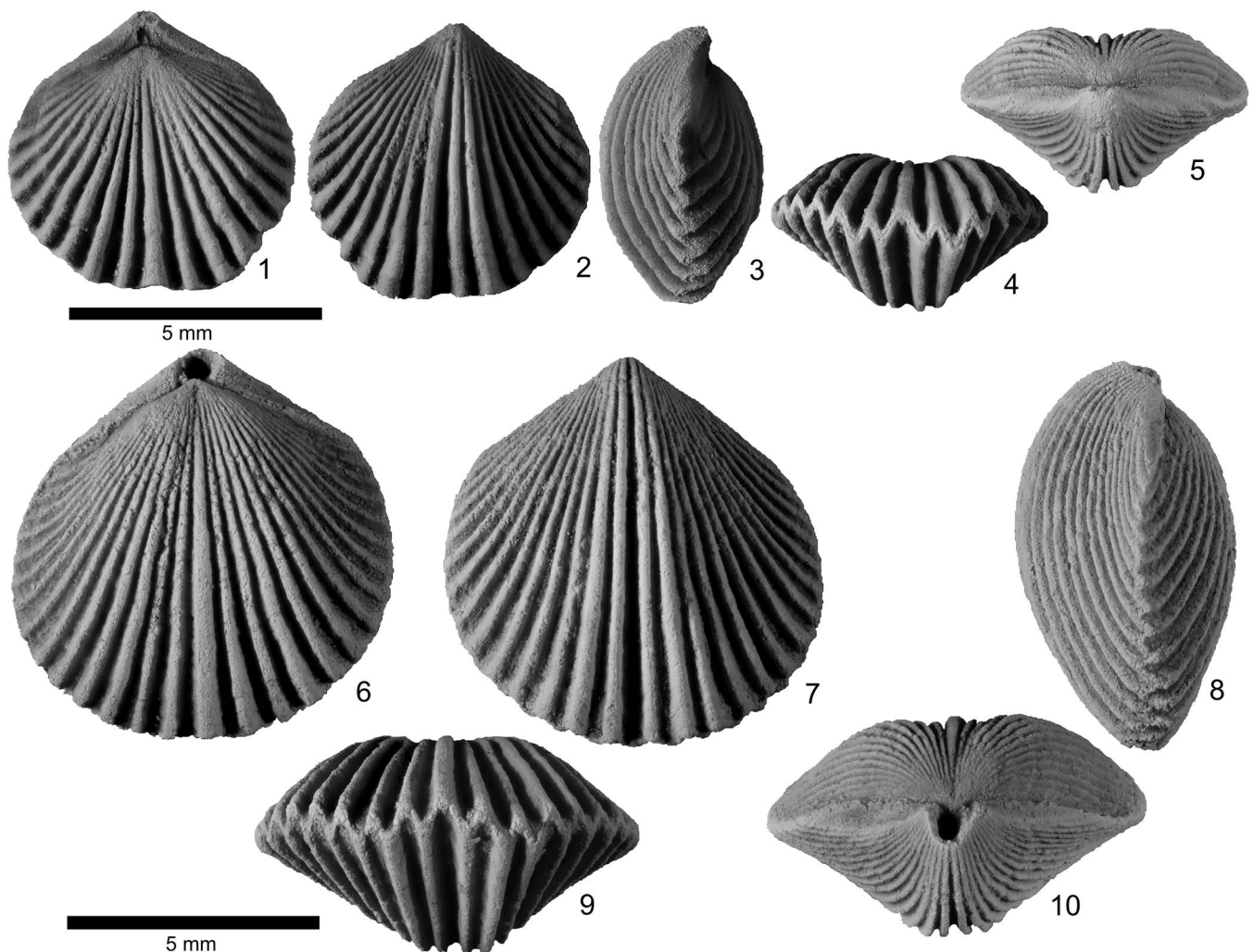


Figure 13. (1–10) *Zygospira resupinata* types from Brainard Shale, Maquoketa Formation in Jackson County, Iowa. (1–5) Holotype SUI 1874, dorsal, ventral, lateral, anterior, posterior views; (6–10) paratype SUI 1873, dorsal, ventral, lateral, anterior, posterior views. Scale bars = 5 mm.

Table 4. Measurements of *Zygospira resupinata* types. L = length; W = width; T = thickness (depth); Ad = deflection at anterior of the commissure (all in mm); Aa = apical angle in degrees; R = number of ribs counted on the ventral valve; L/W = length/width ratio; T/W = thickness/width ratio.

Specimen	L	W	T	Ad	Aa	R	L/W	T/W
SUI 1873	5.3	5.4	3	1	100	18	0.98	0.56
SUI 1874	7.3	7.2	4.1	0.6	99	26	1.01	0.57

smaller and narrower ventral fold and dorsal sulcus, creating a less prominent deflection of the commissure near the anterior. The interior of this species remains unknown, but the external characteristics suggest affinities with *Zygospira*, although this species shares the more elongate shell shape and narrower fold and sulcus of the earlier *Anazyga*. A single damaged *Zygospira resupinata multicostata* Howe, 1965, shell from the Hudson Bay Lowlands (Jin et al., 1997, pl. 30, fig. 21) exhibits typical *Zygospira*-style spiralia, but this has not been described in the material from Iowa. If this feature is confirmed in *Zygospira resupinata sensu stricto*, the species may represent an earlier divergence from earlier atrypide stocks (such as *Anazyga*), although the lack of early occurrences of this species at a time when species such as *Z. modesta* were becoming widespread makes this unlikely.

Large collections of this species are currently unavailable for study, but *Z. resupinata* and *Z. resupinata multicostata* Howe, 1965, from the Montoya Group in Texas and the Surprise Creek and Caution Creek formations in the Hudson Bay Lowlands represent the farthest that *Zygospira* was able to infiltrate into the inland seas of Laurentia. No atrypides are yet known from the paleoequatorial seas of southern Manitoba despite a relatively diverse brachiopod fauna being reported there (Jin and Zhan, 2001).

Results

Two principal component analysis scattergrams were produced to illustrate the differences in morphology as represented shell measurements between *Zygospira* species in eastern North America (Figs. 14, 15). Specimens were primarily grouped by species (color) and region (symbols).

Two groups of species can be recognized in the PC1 (principal component one) versus PC2 scattergram (Fig. 14): one comprised primarily of *Z. modesta* shells and the other representing both *Z. kentuckiensis* and *Z. cincinnatiensis*. The two type specimens of *Z. resupinata* plot within the *Z. modesta* cluster, but given the small sample size here, the significance of this is debatable. The shells of *Z. concentrica* plot between these two main groups, although much higher on the PC2 axis. *Zygospira modesta* and *Z. kentuckiensis* overlap considerably, reflecting broad similarities in their shell morphology.

The biplots at the center of the axes provide some indication of the covariance of the measurements and proxies for shape. Most of the linear measurement vectors (length, width, thickness) approximately align with the horizontal axis (PC1) while the proxies for shape (ratios of measurements) align with the vertical axis (PC2). This indicates a strong size influence on PC1 while PC2 approximately represents most of the variance in shape within the dataset. These relationships also are clearly

illustrated in the loadings plots below the figure for PC1 (bottom left) and PC2 (bottom center). The scree plot (Fig. 14, bottom right) illustrates that most of the variance is captured by PC1 (52%) while PC2 represents 15% of the variance.

Some intraspecific morphological variation is revealed in the *Z. modesta* cluster. Younger specimens plot higher on axis 2 (Fig. 13) than older specimens. This is mainly due to a more elliptical shell (smaller length/width value), slightly larger apical angle, and slightly less prominent anterior fold and sulcus (anterior deflection/width) in the older, primarily Maysvillian shells in comparison to the younger Richmondian sample.

A second PCA scattergram (Fig. 15) plots PC2 and PC3. This shows a different pattern, with more-significant overlap between the groups represented by species. As in the previous figure, PC2 is positively correlated with the proxies for shape in the dataset while PC3 is influenced mainly by the number of ribs. The significant overlap between species in the scattergram reflects that size (represented mostly in PC1 rather than PC2 or PC3) is the most diagnostic character separating the species while shell shape is broadly similar among species of *Zygospira*. There is more significant interspecific variation in the number of ribs in this plot, however. PC3 represents a relatively small proportion of the variance, but the effects of shell size and shape are represented by a greater number of variables, and thus are perhaps over-represented in the variance of the dataset.

Note that in both cases, variables were standardized using a correlation matrix within PAST before plotting the scattergrams. This ensures that the variance in the dataset is more reasonably represented rather than working with the large ranges in values within the raw dataset. It also ensures that variables with large absolute ranges do not dominate the variance. In this study, the relatively large values for apical angle and ribs would overprint any trend in small values such as the depth of the sulcus.

Discriminant analysis (DA) also was run on the dataset to test whether the existing classification scheme can adequately discriminate between the species as they are currently defined, producing a scattergram that represents the maximum variance between samples (Fig. 16). Most of the data points are near zero on axis 1, with the *Z. concentrica* samples clustering together far to the left (top). This supports the hypothesis that *Z. concentrica* should be synonymized and likely represents deformed and abraded shells of *Z. modesta*, given that they plot so far out of the normal morphological range of *Zygospira*.

When *Z. concentrica* is excluded, the specimens form a more even distribution across the scattergram with very little overlap between clusters (Fig. 16, bottom). The confusion matrix produced by the analysis shows relatively stable classifications (Tables 5, 6), seemingly indicating that size is a good diagnostic character to differentiate species as they are currently defined. Whether size should be considered a diagnostic character requires some further consideration.

Discussion

Speciation or ontogeny?—If species have similar shapes but different sizes, perhaps they represent different stages of growth (ontogeny) rather than different trajectories of

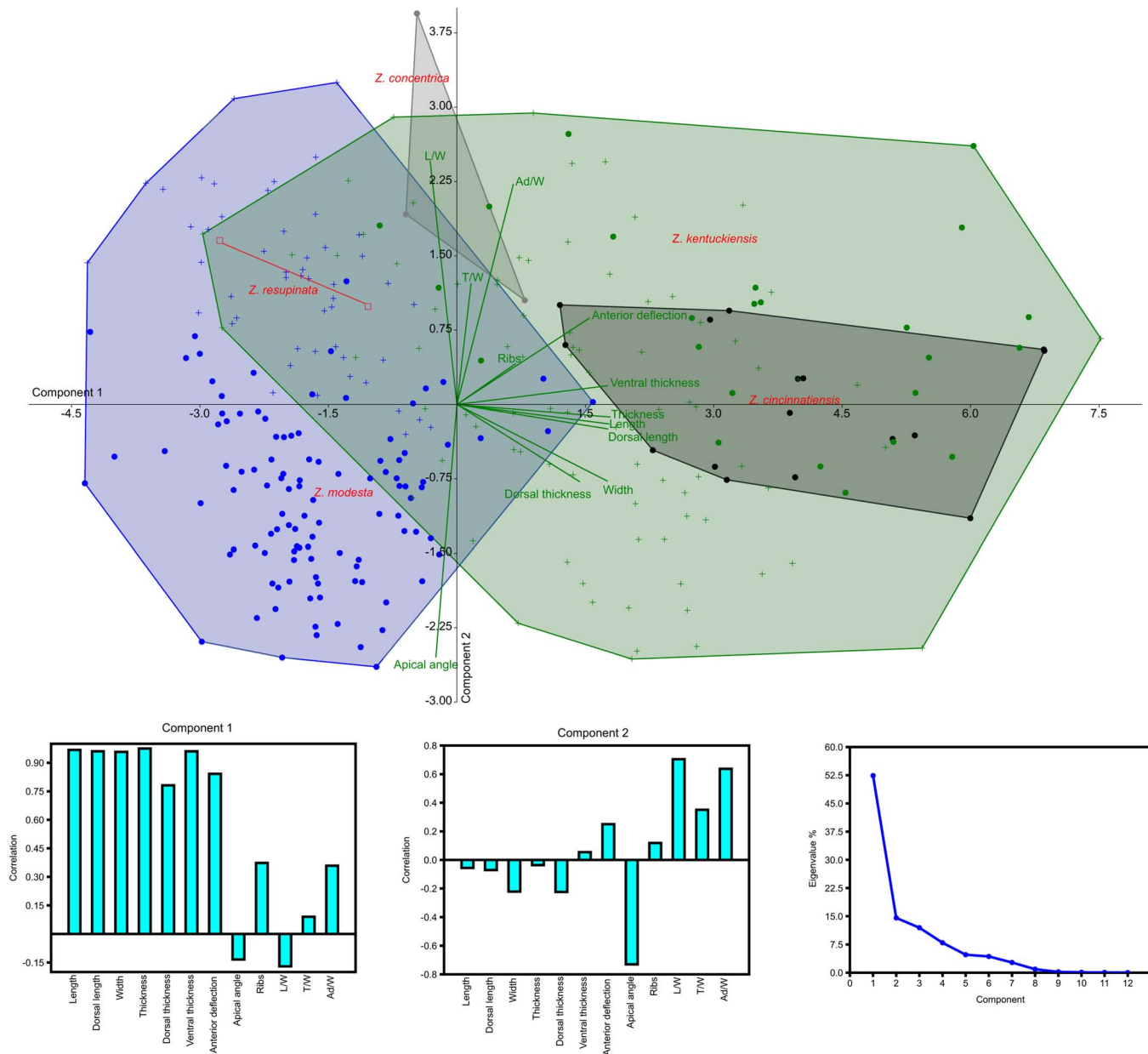


Figure 14. Principal component analysis (PCA) scattergram showing PC1 plotted against PC2. Loadings for PC1 (bottom left) and PC2 (bottom center) shown along with scree plot (bottom right). Blue = *Z. modesta* from localities near Cincinnati, Ohio (dots = specimens from the middle Katian, plus signs = specimens from upper Katian); green = *Z. kentuckiensis* (dots = specimens from the Cincinnati region, plus signs = specimens from Ontario); black = *Z. cincinnatiensis* from area near Cincinnati; red = *Z. resupinata* types from upper Katian of Iowa; gray = *Z. concentrica* from area near Cincinnati.

evolution (speciation). The two most common species of *Zygospira* are found in rocks of the same age in the late Katian (Richmondian) with only minor differences in the shell shape (as shown by slightly different L/W and T/W values for *Z. modesta* and *Z. kentuckiensis*; Tables 2, 3, see also Howe, 1965, text-fig. 4). These differences are much less significant than the difference between the eastern species and the western species, such as *Z. resupinata* from Iowa and the subspecies *Z. resupinata multicostata* from Texas and the Hudson Bay Lowlands.

There are morphological differences between these species that are more difficult to explain through ontogeny, however. The number of ribs differ between these two species (mean of

17 on *Z. modesta* specimens compared to 21 on *Z. kentuckiensis*). Although it is not unusual for a larger brachiopod shell to have more numerous ribs, either through bifurcation of ribs towards the anterior or insertion of additional ribs in interspaces through intercalation as the shell grows, *Zygospira* show little to no multiplication of ribs as shells become larger within species (except for some populations of *Z. resupinata multicostata*—see Jin et al., 1997, pl. 30). If *Z. kentuckiensis* were simply a form of *Z. modesta* that was able to biomineralize a larger shell in a more favorable environment, for example, *Z. kentuckiensis* should have larger ribs rather than more numerous ribs based on the limited variability in the number of ribs within a species, regardless of size.

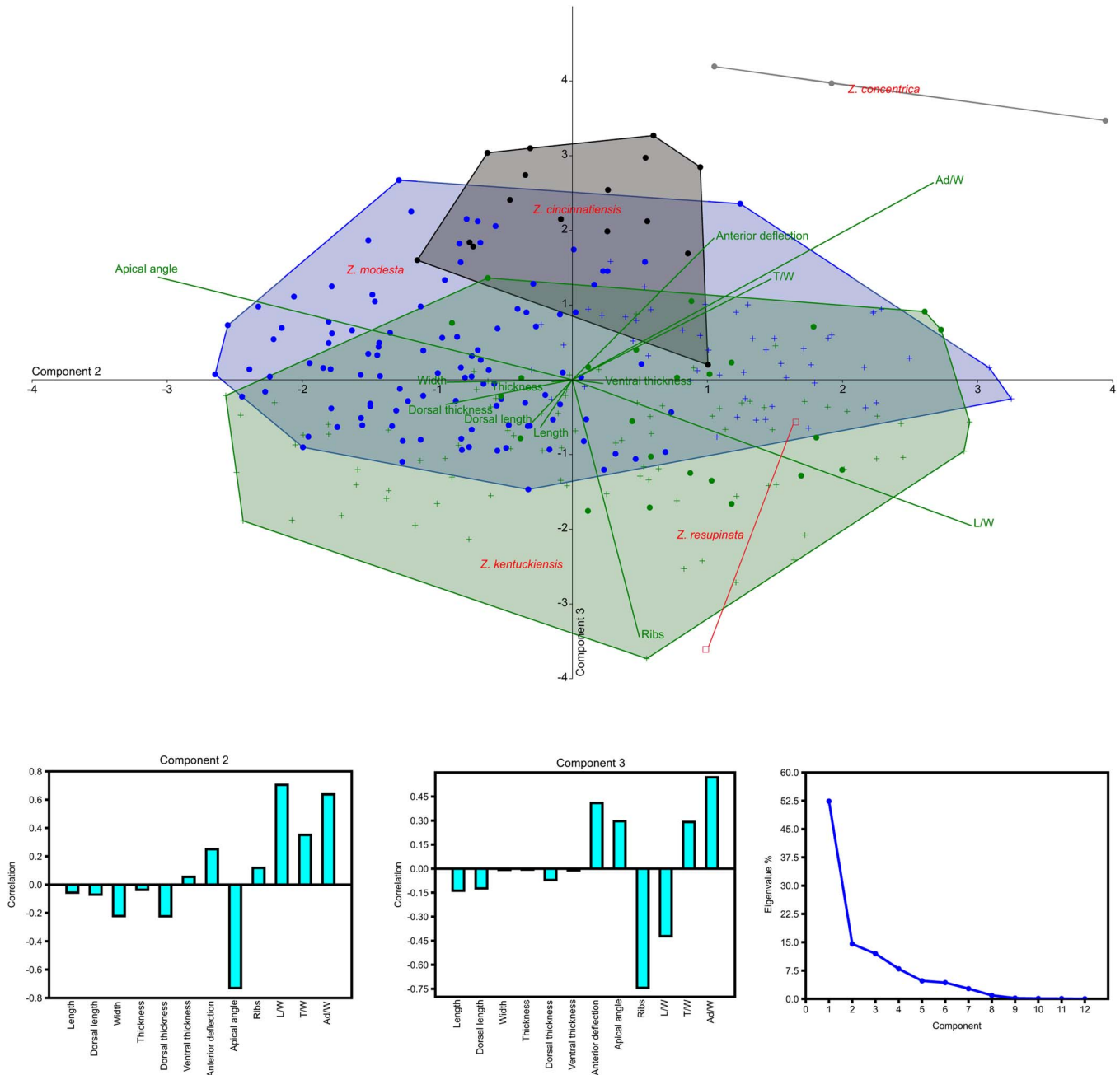


Figure 15. Principal component analysis (PCA) scattergram showing PC2 plotted against PC3. Loadings for PC2 (bottom left) and PC3 (bottom center) shown along with scree plot (bottom right). Blue = *Z. modesta* from localities near Cincinnati, Ohio (dots = specimens from the middle Katian, plus signs = specimens from upper Katian); green = *Z. kentuckiensis* (dots = specimens from the Cincinnati region, plus signs = specimens from Ontario); black = *Z. cincinnatiensis* from area near Cincinnati; red = *Z. resupinata* types from upper Katian of Iowa; gray = *Z. concentrica* from area near Cincinnati.

Furthermore, the shapes of the ventral fold and dorsal sulcus differ considerably between smaller and larger species, with *Z. modesta* exhibiting a distinctive raised fold and depressed sulcus in comparison to the much less prominent fold and sulcus of *Z. kentuckiensis*. This significant change in shell shape, which would be difficult to explain as an effect of shell growth, is more reasonably explained as interspecific variation. The larger *Z. cincinnatiensis* measured herein maintain their more distinctive fold and sulcus in larger shells, similar to the smaller *Z. modesta*, indicating that this feature is a useful diagnostic tool, although it is difficult to recognize in deformed shells. The profile of the

fold and sulcus is known to vary with shell growth among species of other atrypide lineages (e.g., *Thulatrypa* and *Meifodia* from the early Silurian—see Huang et al., 2016; Baarli, 2022).

There are also subtle differences in the size and shape of the internal characteristics of *Z. modesta* and *Z. kentuckiensis*. The larger, more robust dental plates of *Z. kentuckiensis* could be a result of shell thickening with growth of the animal, but this pattern of growth is known in some early Silurian pentameride lineages that are thought to have increased the thickness of the posterior of their shells to stabilize their living position (Ziegler

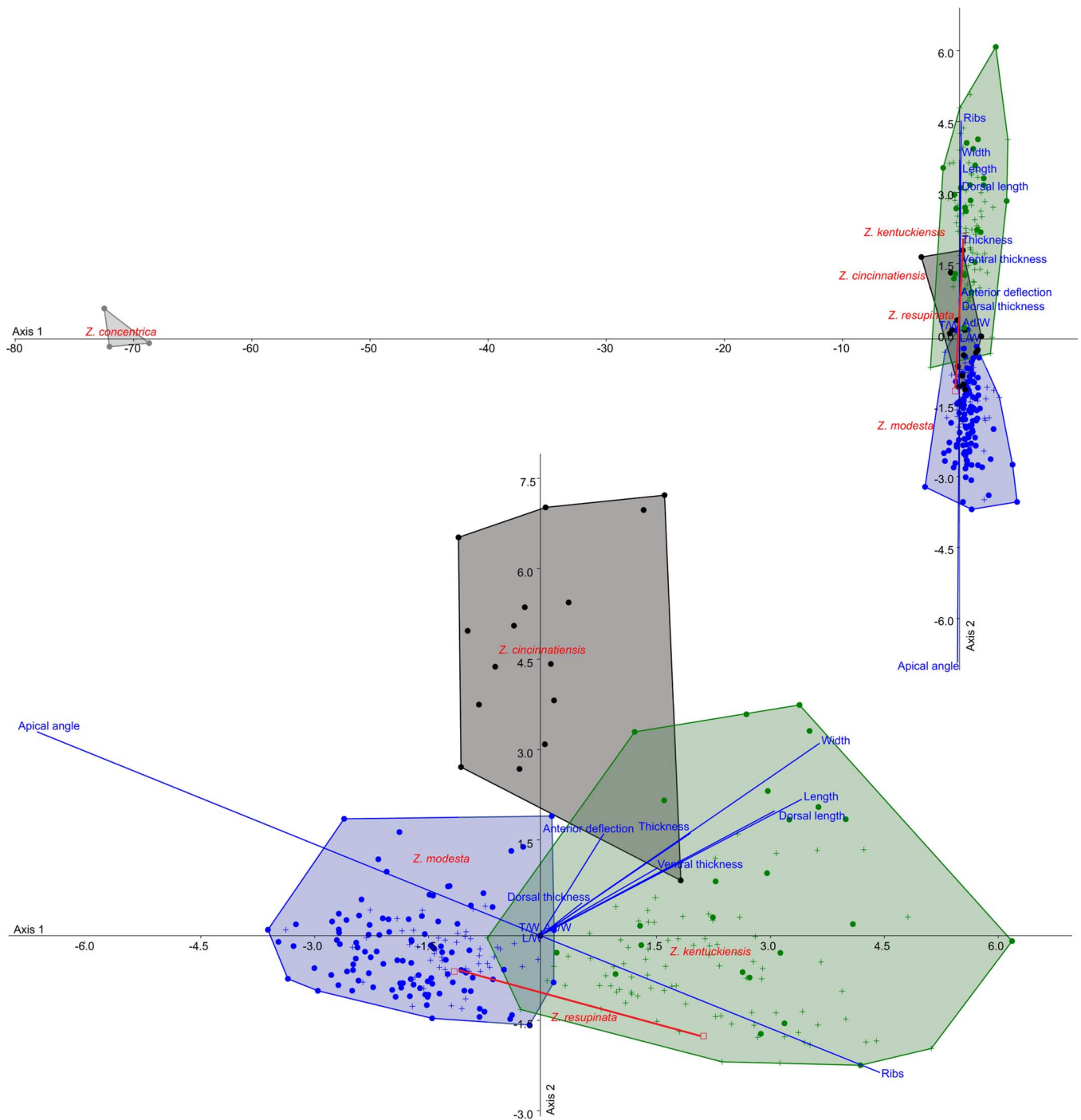


Figure 16. Discriminant analysis (DA) scattergram showing the morphological difference between all species (top) and all species excluding *Z. concentrica* (bottom). Blue = *Z. modesta* from localities near Cincinnati, Ohio (dots = specimens from the middle Katian, plus signs = specimens from upper Katian); green = *Z. kentuckiensis* (dots = specimens from the Cincinnati region, plus signs = specimens from Ontario); black = *Z. cincinnatiensis* from area near Cincinnati; red = *Z. resupinata* types from upper Katian of Iowa; gray = *Z. concentrica* from area near Cincinnati.

et al., 1966). This may have been an adaptation in the larger *Z. kentuckiensis* to stabilize their shell in position the seafloor with the commissure oriented upward. The shape of the jugum associated with the spiralia differs between these species as well (Figs. 5, 11), but given how fragile this structure is, this may be a taphonomical artefact rather than a real morphological difference. Further study of the internal morphology of the brachidia in these and other early atrypide species is needed to

determine the degree of intra- and interspecific variation of this structure in early members of the lineage.

Besides morphological differences that are difficult to reconcile, the geographic range of species differ. *Zygospira modesta* is almost entirely restricted to the area around Cincinnati throughout its stratigraphic range, while *Z. kentuckiensis* has a wider distribution across eastern North America during the late Katian (Richmondian), extending on to the Trenton Shelf in Ontario. If

Table 5. Confusion matrix for *Zygospira* species in Figure 15 (top). Rows are given groups while columns are predicted groups. Note that 93.4% of specimens were correctly identified.

	<i>Z. modesta</i>	<i>Z. concentrica</i>	<i>Z. cincinnatiensis</i>	<i>Z. kentuckiensis</i>	<i>Z. resupinata</i>	Total
<i>Z. modesta</i>	153	0	0	0	5	158
<i>Z. concentrica</i>	0	3	0	0	0	3
<i>Z. cincinnatiensis</i>	0	0	15	1	0	16
<i>Z. kentuckiensis</i>	5	0	4	96	4	109
<i>Z. resupinata</i>	0	0	0	0	2	2
Total	158	3	19	97	11	288

Z. modesta truly were an earlier ontogenetic stage of *Z. kentuckiensis*, then younger, smaller shells should be found across the same geographic range as the larger shells where both occur in the same stratigraphic interval. Even in the Cincinnati region where both species are known to occur, they are never found in the same beds or at the same locality despite both species being widespread throughout upper Katian (Richmondian) strata in the area, which perhaps indicates that the two species were adapted to different environmental conditions.

The external morphology, internal morphology, and differences in the geographic range of these shells seem to suggest that these shells do, in fact, represent different species. It is relatively uncommon to have multiple fossil brachiopod species of the same genus in the same area, so atypical processes could have driven speciation in *Zygospira*.

Niche partitioning driving evolution.—A variety of processes can contribute to the evolution of new species, but allopatric speciation where populations become isolated by a barrier causing the populations to evolve along different trajectories (Fig. 17) is thought to have produced most species in the fossil record (Eldredge, 1971; Johnson, 1975) because of the divergence of evolution caused by major environmental shifts in a region (e.g., sea level change, tectonic activity) that would isolate populations of species from one another eventually leading to speciation. This can happen either through vicariance (i.e., a barrier separates two groups within their range) or dispersal (i.e., when a subgroup of a species moves into a new geographic area and subsequently becomes geographically isolated) (Stigall, 2013; Stigall et al., 2017). *Zygospira resupinata* and *Z. resupinata multicostata* may have evolved through dispersal, spreading to the midcontinent region during a time of lower sea level before becoming isolated from the Cincinnati region by the Sebree Trough (Kolata et al., 2001).

The lack of apparent geographic barriers between most *Zygospira* species in the Cincinnati region rules out allopatric speciation. Other types of speciation, such as peripatric and parapatric speciation, are rarely reported in the fossil record

due to the complexities of interpreting niche partitioning in the fossil record, although these processes likely played important roles in planktonic groups (e.g., Lazarus, 1983; Wei, 1994; Jackson and Cheetham, 1999). The evolution of late Katian species of *Zygospira* may be an uncommon example of sympatric speciation in the fossil record where a single ancestral species gives rise to new species within the same geographic range commonly through the exploitation of slightly different niches or sexual selection that then drives evolution along two different trajectories. Factors influencing sympatric speciation are often difficult to determine in modern ecosystems and are particularly problematic to interpret based on the fossil record where only a small part of the ecosystem is ever preserved.

The larger size of *Z. kentuckiensis* could have been an advantage in competing with other filter feeders for space on the seafloor where the posterior weighting of the shell, keeping the commissure upright, would have provided an advantage in dense shell beds. *Zygospira* sometimes occurs in dense accumulations, although whether these are always in situ or reflect taphonomic processes that form condensed beds is not always clear. Other factors could have favored the establishment of one species over the other (e.g., seasonal shifts in nutrient supply, changes in sediment supply, or even changes in the marine ecosystem such as predator-prey cycling), which could have affected whether larger or smaller shells were more successful. The effects of these factors on a single brachiopod lineages from one geographic area are difficult to quantify based on fossil material and would require a more comprehensive study at a regional scale to be meaningfully investigated.

The smaller size of species such as *Z. modesta* may have been a beneficial adaptation under certain circumstances as well. Shells are commonly found in dense shell beds associated with other filter feeders such as bryozoans, sponges, and crinoids. Often these are only loose associations, but there are examples of *Zygospira* preserved in place attached to other filter feeders (Fig. 3; see also Sandy, 1996, for an example of *Zygospira* attached to articulated crinoid columnals). These shells are oriented with the pedicle opening facing the host filter feeders, so these associations are unlikely to be the result of brachiopods

Table 6. Confusion matrix for *Zygospira* species in Figure 15 without *Z. concentrica* (bottom). Rows are given groups while columns are predicted groups. Note that 93.31% of specimens were correctly identified.

	<i>Z. modesta</i>	<i>Z. cincinnatiensis</i>	<i>Z. kentuckiensis</i>	<i>Z. resupinata</i>	Total
<i>Z. modesta</i>	151	1	0	5	157
<i>Z. cincinnatiensis</i>	0	15	1	0	16
<i>Z. kentuckiensis</i>	5	4	97	3	109
<i>Z. resupinata</i>	0	0	0	2	2
Total	156	20	98	10	284

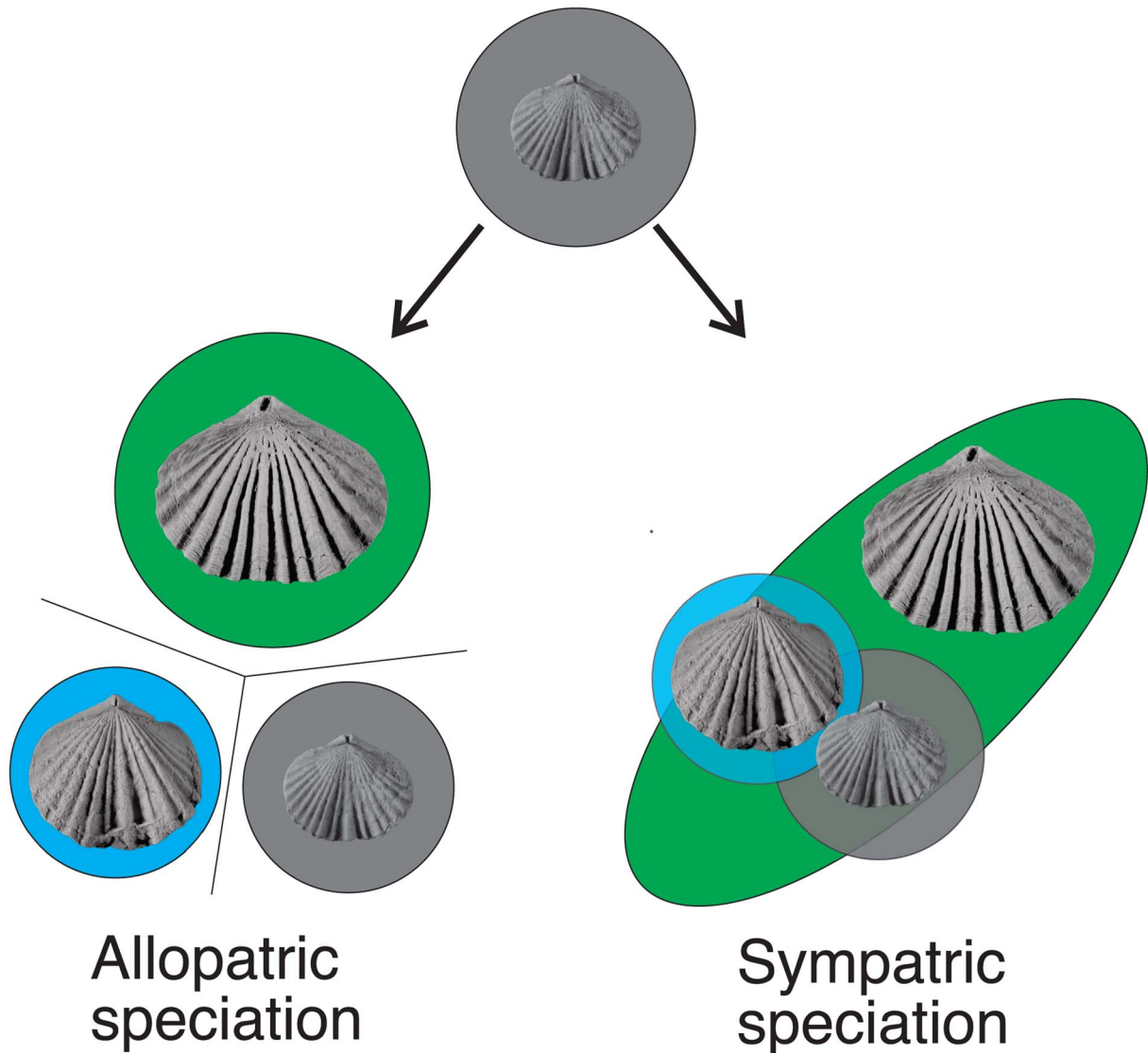


Figure 17. Allopatric and sympatric speciation. Species represented by different background colors (gray = *Z. modesta*; blue = *Z. cincinnatiensis*; green = *Z. kentuckiensis*). The late Katian species of *Zygospira* in eastern North America likely arose through sympatric speciation, although it isn't yet clear from which lineage *Z. resupinata* in the interior of North America arose.

randomly settling on the filter feeder after their death. Only the smaller *Z. modesta* specimens have been preserved attached in place, however. Their smaller size may have been an advantage in comparison to the larger *Z. kentuckiensis*, if they were living as epibionts on other filter feeders. These attachments would have enabled the brachiopods to feed at a much higher level in the water column than would otherwise be possible, thus avoiding competition for space and food with other brachiopods and filter feeders closer to the seafloor below. This could explain why *Z. modesta* was able to thrive through the late Katian when many of the other smaller brachiopod species in other lineages mostly disappeared as larger species took over across Laurentia (e.g., as in orthide and rhynchonellide lineages: Sohrabi and Jin, 2013; Sproat and Jin, 2013; Sproat et al., 2014).

Epibiotic life strategies have been exploited by other brachiopod lineages. Carboniferous chonetids are thought to have used their posterior spines to attach to crinoid stems (e.g., Grant, 1963). Other lineages, such as the craniiformean

brachiopods, are commonly found cemented to filter feeders (including other brachiopods) and were already widespread by the Ordovician (e.g., Chen and Rong, 2019; Bruthansová and Van Iten, 2020), although these brachiopods are most commonly cemented to flat surfaces. Most rhynchonelliform brachiopods are not widely known as epibiota, but this life strategy may have been more common than the fossils would suggest given how unlikely this arrangement is to survive taphonomic processes and be preserved in the fossil record.

These types of associations may have played important roles in the evolution of shelly benthos through the Ordovician Radiation and the greater radiation of filter-feeding benthos that occurred through the early Paleozoic. Although the increase in biodiversity associated with this event and some of the contributing extrinsic environmental factors are becoming increasingly clear, we are only beginning to understand some of the intrinsic factors driving radiations (Stigall et al., 2019, and references therein). We know particularly little about the complex interplay between filter

feeders that led to the opening of new niches and evolution of a wide variety of body plans (although see Vinn and Wilson, 2015). Brachiopods, which were among the most common components of the benthic ecosystem at this time, must have played a key role in the evolution of these relationships, but further study is needed to clarify how they may have affected the evolution of other elements of the benthic ecosystem at this time.

Evolution of larger shells with increased sediment influx.—Few studies have confidently linked changes in brachiopod shell morphology to environmental factors, but Laurin and García-Joral (1990) concluded that an increase in clay sedimentation in the Jurassic of the Iberian Range in Spain caused miniaturization of rhynchonellid shells (i.e., evolution of smaller, mature shells). They proposed that these smaller shells were better adapted to an r-mode life strategy (rapid development and maturation to overcome high juvenile mortality), which would have allowed the brachiopods to colonize a difficult living environment quickly. Larger shells would otherwise be an advantage in competing for space in a more favorable living environment where the larger size would be beneficial in gathering food and nutrients from the water column more quickly than smaller shells and could have excluded smaller shells from becoming established.

The carbonate platforms and basins of eastern North America (e.g., the Trenton Shelf across southeastern Canada and New York) were subjected to an increase in clastic sedimentation during the Late Ordovician from the nearby Taconic Mountains, which were uplifted and eroding by this time (Waldron and van Staal, 2001; van Staal et al., 2007; Ganis and Wise, 2008). Rather than miniaturization, however, larger species of *Zygospira* evolved and become more widespread than the earlier smaller species. Detailed sedimentological study of all collection localities has not been attempted here, but in general terms, sediments on the St. Lawrence Platform in Ontario become notably argillaceous in the middle to late Katian, especially in the Queenston Formation (Armstrong et al., 2010) where *Z. kentuckiensis* is found in thick successions of dense shells beds. An influx of clay into the region does not seem to have excluded the larger *Zygospira*, but rather seems to have favored their establishment. Perhaps a larger shell was less likely to have been buried by clastic sediment entering the basin, but further morphometric and paleoecological studies on this and other brachiopod lineages in different regions across the geological time scale are needed to determine whether this is a consistent ecophenotypic trend.

Variation in the spiralia.—From the interior of the five shells of *Z. modesta* and five shells of *Z. kentuckiensis* examined here, it appears that there is very little intraspecific variation in the structure of the brachidia of *Zygospira*. Both large and small shells have similar spiralia with almost identical shapes and numbers of whorls. The only notable differences are the shape of the jugum, but the position of the structure is relatively stable and not as variable as once thought (Hall and Clarke, 1894), the posterior thickening in the larger *Z. kentuckiensis*, and thicker, more hook-shaped teeth in *Z. kentuckiensis*. The internal morphology of other species, especially those now assigned to *Anazyga*, remains to be studied.

The shape and size of the calcified brachial support structures (spiralia) in *Zygospira* and the other Anazygidae could reveal the evolutionary affinities of this group to other early atrypides, however. Unfortunately, it is currently difficult to study the internal morphology of large samples of shells unless a researcher comes upon a damaged shell that reveals the spiralia intact. The Atrypidae (and the Rhynchonellidae, along with the other spire-bearing brachiopod lineages) are only rarely found disarticulated due to their tight-fitting cyrtomatodont hinge that holds the shells together even after death, and even then the spiralia is almost always missing. To study the internal morphology of atrypides, researchers must serial section shells—a destructive process that is slow and laborious and requires the use of a parallel grinder, which has not been manufactured in years (although see Zhang et al., 2019, for open-source design diagrams to create new machines). Sectioning a single atrypide shell may take several days to capture the entire spiralia and associated structures. The destructive nature of the process makes it difficult to examine historical collections at museums and in other collections where destructive techniques are usually prohibited. Contemporary computed tomography (CT) methods allow for more rapid and less destructive 3-D imaging of brachiopod shells, but have had little success thus far in imaging calcite marine invertebrates preserved in limestones because of the minimal contrast that is common between shell material and surrounding sediment. These techniques hold promise in allowing paleontologists to rapidly image the interiors of a large number of shells to efficiently study variation in structures such as the spiralia and better understand the degree of variation in these structures in early atrypides.

Although not the focus of this study, the shape of the jugum could have some diagnostic utility given the apparent differences between species. The function of this distinctive section of the brachidium that connected the spiralia remains unknown, perhaps playing a role in supporting the two spires of the spiralia that it connects (Rudwick, 1970) or providing support for the mouth of the brachiopod in life, providing a balance to keep the spiralia level, or playing a role in raising or lowering the spiralia as the brachiopod opened and closed its shell (Copper, 1967). Regardless of its function, the jugum may have some utility in the taxonomy of the spire-bearing brachiopods given the conservative nature of the evolution of internal shell structures, but it has been poorly studied in many spire-bearing lineages. Despite this variation in shape of the jugum noted here in *Zygospira*, there is no indication of the variation in position of the jugum noted by previous workers (e.g., Hall and Clarke, 1894). This apparent variation was likely a result of lumping both *Anazyga* and *Zygospira* under a single genus because a lower position for the jugum near the anterior of the shell is more characteristic of *Anazyga* (Copper, 1977). Again, the significance of these differences remains uncertain until a better understanding of the jugum has been developed.

Conclusions

The early atrypide brachiopod *Zygospira* spread across much of Laurentia during the late Katian. *Zygospira* species are difficult to differentiate because of the lack of typical diagnostic features useful in other brachiopod lineages, but they differ from one

another mostly based on size and the number of ribs on the shell rather than by shape. Although this may indicate that species simply represent different stages in growth or growing conditions, this does not seem to be the case. Although the large *Z. kentuckiensis* do consistently possess more ribs, large examples of *Z. cincinnatiensis* show fewer ribs despite reaching similar shell sizes. Key differences between species in the shape of the ventral fold and dorsal sulcus and structures in the interior such as the jugum would be difficult to produce through ontogeny.

Despite the difficulty in classifying species, the taxonomy of the genus can be refined although the key to classifying these species may rely on a more detailed examination of the morphology of the spiralia and its associated structures. *Zygospira concentrica* can be synonymized with *Z. modesta* because the concentric growth lamellae are more likely to represent local environmental disturbances and are unlikely to be diagnostic. The newly reported specimens from the area near Owen Sound, Ontario are typical *Z. kentuckiensis* that differ only in their slightly less convex shells from the type specimens from the Cincinnati area. *Zygospira meafordensis* is herein considered a subspecies of *Z. kentuckiensis* based on its similar morphology and its extremely limited distribution.

While many other brachiopod lineages seem to have evolved in a stepwise fashion through the Late Ordovician with older, smaller species being replaced by younger, larger, and more globose species, the older and smaller *Zygospira modesta* co-existed with younger species that evolved throughout the Katian (e.g., *Z. cincinnatiensis*, *Z. kentuckiensis*, and *Z. resupinata*). Although the species vary significantly in size, their shell shapes are notably similar, except for *Z. resupinata*, which differs in ornamentation, shell shape, and shape of the dorsal sulcus and ventral fold. That several species evolved in the Cincinnati area at the same time makes this a rare example of sympatric speciation in the fossil record. The ability of *Z. modesta* to live as an epibiont on other filter feeders may have contributed to the longevity of species. This niche partitioning likely played a key role in the Ordovician Radiation, although the role of brachiopods as epibionts remains understudied and the fossil record of these direct associations remains poor.

Further research on the internal morphology of the early spire-bearing brachiopods may reveal how the evolution of their distinctive spiralia may have initially occurred. This seems to have been a key evolutionary innovation because the atrypides and spiriferides dominated shallow marine seas of the later Silurian and Devonian, but it remains unclear how and why the characteristic calcified lophophore supports of these lineages evolved.

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