Hinge and ecomorphology of *Legumen* Conrad, 1858 (Bivalvia, Veneridae), and the contraction of venerid morphospace following the end-Cretaceous extinction

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**Abstract.**—The Veneridae are the most speciose modern family of bivalves, and one of the most morphologically conservative and homoplastic, making subfamily- and sometimes even genus-level classification difficult. The widespread Cretaceous genus *Legumen* Conrad, 1858 is currently placed in the subfamily Tapetinae of the Veneridae, although it more closely resembles the Solenoida (razor clams, Pharidae and Solenidae) in general shell form. Here we provide high-resolution images of the *Legumen* hinge for the first time. We confirm from hinge morphology that *Legumen* belongs in Veneridae, but it should be referred to incertae subfamiliae, rather than retained in the Tapetinae, particularly in light of the incomplete and unstable understanding of venerid systematics. *Legumen* represents a unique hinge dentity and a shell form—and associated life habit—that is absent in the modern Veneridae despite their taxonomic diversity. Veneridae are hyperdiverse in the modern fauna, but strikingly ‘under-disparate,’ having lost forms while gaining species in the long recovery from the end-Cretaceous extinction.

**Introduction**

The modern Veneridae are the most speciose modern family of the Bivalvia, with ∼750 species in 135 genera, outnumbering the next most diverse families, the Galeommatidae and Tellinidae, each with ∼500 species in 100 genera (Huber, 2015; Collins et al., 2018; Edie et al., 2018). Molecular analyses have established a backbone phylogeny for Veneridae (Mikkelsen et al., 2006; Chen et al., 2011), but species coverage within the group is still insufficient to confirm monophyly of some long-established taxonomic groupings, such as subfamilies. A further complication (or source of interest) is the high level of homoplasies in infaunal bivalve shell form owing to the functional requirements of burrowing method and life habit. Consequently, systematics within the Veneridae has been unstable, with species frequently moved between genera and with a much-revised subfamilial classification system (e.g., Keen, 1969; Harte, 1998; Huber, 2010; Alvarez, 2019). The classification of Bivalvia in the ongoing revision of the Treatise on Invertebrate Paleontology denotes many historically accepted vener subfamilies to tribes, some of which, including Tapetini (=Tapetinae), are thought to be paraphyletic (Carter et al., 2011), although subsequent studies continue to use subfamilies, and recent molecular work recovered a monophyletic Tapetinae (Kappner and Bieler, 2006; Mikkelsen et al., 2006; Chen et al., 2011).

Despite their taxonomic richness, the modern Veneridae are not as morphologically diverse as other species-rich bivalve families (e.g., the Pectinidae and Tellinidae), particularly if specialized rock-boring Petricolinae (subtribe Petricolini in Carter et al., 2011) are omitted from consideration. Soft-bottom infaunal venerids are equivalent, inequilateral, and prosogyrous. Most of their variation in form derives from posterior elongation, but even in this they are conservative compared to other heterodont bivalves such as the Tellinidae or the razor clams in the superfamilial Solenoidea, and Veneridae lack anteriorly elongate forms, such as occur in the fast-burrowing families Donacidae and Mesodesmatidae.

*Legumen* Conrad, 1858, a Cretaceous venerid genus, comprises ∼22 valid species, most of which are small, extremely posteriorly elongate, and very compressed, and thus a morphological outlier to the family. The genus has been placed in the Tapetinae since Stoliczka (1871); as “Tapesinae”), following the designation of *Venus* (*Tapes*) *fragilis* d’Orbigny, 1845 as the type of his new genus Baroda, subsequently synonymized with *Legumen* by Stephenson (1923). That synonymy was retained by Keen (1969, p. N682), who diagnosed Tapetinae as “Ovate to elongate, shell surface somewhat polished, inner margins smooth on at least posterior third; hinge plate narrow, with cardinals 3a entire, 3b normally entire, others frequently bifid; lateral teeth wanting.” The first appearance of *Legumen* is given as ‘lower Cretaceous’ in Keen (1969), which may have been a reference to “Tapes” *parallela* Coquand, 1865 (Aptian) or to an unnamed Albian Angolan *Legumen* sp. (Rennie, 1929; see Darteyelle and Freneix, 1957). Since the Treatise on Invertebrate Paleontology, a potential Albian species has been named as *L. iraniense* Collignon, 1981, but this is based on a single poor specimen with no hinge preserved. Because of the extremely thin, delicate nature of the shell of most *Legumen* species, this is not an uncommon situation: few specimens are available in museum collections and even fewer with interior
characters (i.e., hinge plate, teeth, muscle scars, pallial line, and sinus) preserved. Illustrations of internal characters in the literature are sparse for all known species. Our aim here is to provide, for the first time, high-resolution images of the hinge of this genus, and to place this unusual venerid in a broader context in terms of shell and hinge morphology. We find that the form of the dentition in *Legumen*, while it mostly conforms to Keen’s broad diagnosis of Tapetinae, is very distinctive.

We use 3D micro-CT scans to digitally ‘excavate’ the hinge of two specimens of *L. ellipticum* Conrad, 1858, one embedded in matrix and the other articulated. This technique, along with a well-preserved single valve of *L. ellipticum* and two of *L. carolinense* (Conrad, 1875), has allowed us to include *Legumen* in a 3D morphometric dataset. Using landmark/semilandmark morphometrics on the interior surface of the shell and on hingeplate configurations, we quantify and compare the form and hinge position in *Legumen* to an exemplar species from each extant genus in Veneridae (excluding the rock-boring Petricolinae) and to the related and superficially *Legumen*-like heterodont families Pharidae and Solenidae, colloquially known as ‘razor-clams.’ This analysis of form, plus close observation of hinge anatomy, suggests that *Legumen* fits poorly within the Subfamily Tapetinae (=Tribe Tapetini). This analysis also suggests that *Legumen* was a fast, vertically burrowing, deep-infaunal siphonate, silty-sand dweller similar to the smaller species of Phari-dae. This represents a distinct life-habit within the Veneridae that was vacated at the Cretaceous-Paleogene boundary when *Legumen* went extinct, and has not been re-occupied by other, surviving lineages within the family.

Materials and methods

Specimens.—The dataset consists of micro-CT scans of 151 Recent and Cretaceous bivalve specimens (138 non-petricoline venerids, 10 pharids, and three solenids), representing all extant sediment-burrowing genera in the three families, plus a small number of morphologically representative Cretaceous species from the southeastern USA. We exclude the Petricolinae because they are a monophyletic and functionally restricted group that occupies a hard substratum, requiring special modifications of the shell. We treat bivalve subgenera as operational genera (hereafter “genera”), following general paleobiological practice (e.g., Sepkoski, 2002). Genera are represented by one valve of a single adult specimen, most often of the type species (where available), and usually a left valve. Right valves were operationally mirrored about the plane of the commissure for inclusion in shape analyses when a left valve was unavailable for study.

Two species of *Legumen* are included in this dataset: *L. ellipticum* (N = 3: USNM 76669, Nacatoh Sand, Kaufman County, Texas; UFIP 180951, “Ripley” [presumably Owl Creek] Formation, Owl Creek, Tippah County, Mississippi; USNM 728210, Coon Creek Formation [or Coon Creek Member, Ripley Formation], Lee’s Old Mill Site, Union County, Mississippi) and *L. carolinense* (N = 2: USNM 31947 and 31802, Tar Heel Formation, Snow Hill, Greene County, North Carolina) (Fig. 1). Other species of *Legumen* and other Cretaceous venerids (N = 114) that were unavailable to us for scanning are included in our dataset using published shell heights and lengths, which we use to compare them to all other taxa using the commonly used metrics of aspect ratio and size, as geometric mean of length and height, a metric closely correlated to centroid size (Kosnik et al., 2006).

Shape analyses.—Shapes of the interior surface of the shell are characterized using an automated semi-landmarking procedure adapted from the ‘eigensurface’ procedure of Polly and MacLeod (2008), following the method detailed in Collins et al. (2019). The interior surface of the shell best represents the general shell form used to infer life modes within the Bivalvia, without adding noise related to sculptural variation (Collins et al., 2019). A grid of semilandmarks is placed over the digitized 3D surface of the shell, superimposed across specimens using a Generalized Procrustes Analysis, and then summarized into a morphospace using Principal Components Analysis (PCA).

Hinge configurations are captured using a minimal set of user-defined landmarks designed to capture the mechanical shape of the hingeplate without relying on specifics of tooth homology, which is violated by the comparison of species from two distantly related superfamilies (or even simply inclusion of both left and right valves in the dataset). The first three landmarks describe the beak and cardinal teeth, delineating the area of the hinge that acts to reduce shear during active burrowing. In order to capture variation in the position of this cardinal area relative to the rest of the hinge area, four more landmarks are included: the junctions of the adductor muscle scars with the hingeplate and the anterior-posterior-most points on the shell commissure parallel to the vector joining the adductor muscle landmarks (i.e., the vector commonly used to define the shell length). These anterior and posterior commissure points were computationally determined by finding the largest magnitude vector connecting two points along the shell commissure that is parallel to the vector defined by the two adductor muscle scar landmarks (Fig. 2).

Repositories and institutional abbreviations.—Specimens examined in this study are deposited in the following institutions: The National Museum of Natural History, Smithsonian Institution (USNM), the Field Museum of Natural History (Invertebrate Zoology) (FMNHIZ), the Florida Museum of Natural History (Invertebrate Zoology [UFIZ] and Paleontology [UFIP]), the Natural History Museum (NHM) London, and the California Academy of Sciences (CASIZ).

Results

The Appendix contains descriptions of the morphology and dentition of *Legumen ellipticum* and *L. carolinense*, and additional images of their dentition (Appendix Fig. 1).

Aspect ratio.—Figure 3 shows that *Legumen* species occupy a portion of the size/aspect ratio space intermediate between the Veneridae and the families in the Solenoida. Other species placed in the Tapetinae are also highlighted in this plot in the
Maastrichtian and the Recent (although the Veneridae first appear in the Jurassic and the Tapetinae in the Albian, these occurrences are omitted from the plot for clarity). In the Maastrichtian, these other species are encompassed wholly within the same area of the size/aspect ratio space as the rest of the Veneridae, and in the Recent, they are scattered, but still
generally lie in the same space as the other Veneridae. Of the species of *Legumen* scanned for this study, *L. carolinense* plots within the range of sizes and aspect ratios exhibited by the other Tapetinae, and *L. ellipticum* overlaps the sizes and aspect ratios of the Solenoida, closest to the smaller Pharidae. Confiirmed occurrences of *Legumen* begin in the Cenomanian, where specimens are already small and elongate relative to most of the modern venerids, and their morphospace expands to include a less elongate, more ‘veneriform’ extreme (Fig. 3). By the Campanian, the full range of aspect ratios known from this genus is in place, from the “veneriform” *Legumen ooides* (Gabb, 1864) (“L. o.” in plots) to the “phariform” *Legumen ellipticum*. By the Maastrichtian, diversification and/or improved sampling shows a wide range of *Legumen* forms between these extremes, clearly separable from the rest of the Veneridae.

**Shell form morphospace.**—Figure 4 illustrates the shell form morphospace (the shapes of the interior surface of the shell). Principal Component (PC) 1 describes 49% of the total variance in shell form, mostly anteroposterior elongation of the shell with a small component of inflation; equilateral species such as *Tivela tripla* (Linnaeus, 1771) plot at high positive values, and posteriorly elongate forms such as the Solenoida plot at extreme negative values. PC2 describes 16% of the total variance, dominantly shell height and inflation, with tall and inflated forms plotting at extreme negative values and low, compressed forms at high positive values. It is immediately apparent that the Veneridae and the Solenoida are well separated in this morphospace—except for *Legumen ellipticum* (Conrad, 1858), which plots close to *Siliqua* and *Neosiliqua* (Pharidae). Its congener, *L. carolinense* (Conrad, 1875), is much more like a “typical” venerid in outline, although notably compressed compared to other Tapetinae of similar outline. The callocardiine venerid *Macrocallista nimbosa* (Lightfoot, 1786) also plots in this space. *Macrocallista* is elongate, but considerably larger and more inflated than *Legumen*.

**Hinge morphospace.**—Figure 5 shows the morphospace of the cardinal area relative to the hingeplate and overall shell elongation (“hinge geometry”). PC1 describes 39% of the variance in shape, mostly posterior elongation (unsurprising, given that the vast majority of heterodont bivalve shape variation is related to this trait), but also relative size of cardinal area: shells with large cardinal areas have high PC1 scores and are more equilateral overall. Shells with small cardinal areas have low PC1 scores and tend to be posteriorly

![Figure 2. Landmarking scheme. The landmarks delineating the cardinal area and the line of the hingeplate are user-defined, the landmarks delineating the anteroposterior axis are placed automatically (see text). Figured specimen is Recent chionine venerid *Chione cancellata* (Linnaeus, 1767) (FMNH-IZ 184007, Guadalupe)](image)

![Figure 3. Size versus aspect ratio morphospace for specimens of *Legumen* and the modern venerid, solenid and pharid fauna (all lengths and heights from the published literature, for the species included in the 3D dataset, plus additional specimens of *Legumen* that were unavailable for scanning). Panels decrease in age left to right from the Cenomanian (base 97 Ma) to the Recent (today). Note the increase in the range of aspect ratios displayed by *Legumen* through time, without a concomitant increase in the range of sizes. Species of *Legumen* with more than one specimen represented are indicated by labels and lines: “L. m.” = *L. martinius* (Matheron, 1843), “L. v.” = *L. venetii* d’Archaic, 1854, “L. o.” = *L. ooides* (Gabb, 1864), “L. c.” = *L. carolinense* (Conrad, 1875), “L. e.” = *L. ellipticum* Conrad, 1858. Also indicated are venerids “I. tr.” = *Itris tris* (Linnaeus, 1758), and “M. nm” = *Macrocallista nimbosa* (Lightfoot, 1786).]
elongate. PC2, which explains 15% of the variance, includes a component of cardinal area size, but dominantly describes the rotation of the cardinal area relative to the hingeplate—the cardinal area being delineated by a triangle where the “apex” (relative to the shell) is the proximal end of whichever cardinal tooth is closest to the beak (in practice, almost always the anterior cardinal 2a), and the “base” of the triangle is formed by the distal end of the anterior cardinal and the distal end of the posterior cardinal. Low PC2 scores indicate a small cardinal area with a base subparallel to the line of the hingeplate, and high scores on PC2 indicate a larger cardinal area with a base at \( \sim 45° \) to the line of the hingeplate. In this hinge geometry space, a pattern similar to the shell morphospace (Fig. 4) emerges, where \( L. \) ellipticum groups with the Pharidae, whereas \( L. \) carolinense plots within the venerid point cloud. Point shapes and colors are as given in the legend to Figure 3.

**Discussion**

**Evaluation of the position of Legumen within the Tapetinae.**—These new images of the dentition of species assigned to *Legumen* suggest two genus-level groups of species have been included under this name. A revision of the group is accordingly now in progress—however, we accept monophyly faute de mieux in this study because the species included are undoubtedly each other’s closest relatives, regardless of how many genera are involved (Fig. 1). Hinge tooth differences strongly suggest that all species currently assigned to *Legumen* should be excluded from Tapetinae (Fig. 1). The narrow ligamental area and extreme elongation and perfect straightness of the posterior cardinal (teeth 4b [LV], 3b [RV]) even in the more “veneriform” *L. carolinense* contrasts with the hinge morphology of modern Tapetinae, in which a broad, curved ligamental area has a slightly raised edge and the posterior cardinal is confined to be short, and directed closer to subvertical. The striking, widely separated cardinals are described as bifid in both *Legumen ellipticum* and *L. carolinense* by Stephenson (1923, 1941), but despite examining and scanning some of the specimens he illustrated (Fig. 1.1–1.6, 1.10–1.12 *Legumen* versus Fig. 1.16–1.18, 1.22–1.24 other Tapetinae) where a single tooth is grooved or divided only part of the way down to the hingeplate (Fig. 1). A selection of
other Cretaceous Tapetinae (e.g., *Cyclorismina*, *Flaventia*, *Sinonia*, *Cyclorisma*; Fig. 1.20, 1.23) are also illustrated in Keen (1969), all of which fit the diagnosis of Tapetinae better. The grouping of Cretaceous tapetines *Cyclorismina*, *Cyclorisma*, and *Amakusatapes* with other Cretaceous venerids in the size/aspect ratio plot (Fig. 3) suggests that the group was well established by mid-Cretaceous time and highlights the unusual character of *Legumen*. Developmentally based tooth homologies and more densely sampled molecular analyses are needed to truly understand the systematics of modern Tapetinae and their precursors. We suggest that *Legumen* is certainly a venerid, but placement in the Tapetinae is difficult to defend morphologically. Given the currently uncertain view of subfamilies in the Veneridae and the ongoing molecular work attempting to build a more stable systematic understanding of the family (e.g., Lemer et al., 2019 and references therein), we refer *Legumen* to Veneridae incertae subfamiliae, rather than erect a monogeneric subfamily for it. We contend that *Legumen* represents a venerid lineage that was lost in the K/Pg extinction. The similarities of form between *Legumen* and the Solenoida are due to convergence, promoted by similar modes of life, as discussed in the next section.

**Figure 5.** The morphospace for the PCA of the hinge configuration. The upper panel shows the distribution of taxa within the space, with exemplar specimens at the margins. It should be noted that the PCA was performed on the landmark configuration shown in Figure 2, not on the full specimen shown (for the morphospace of the full internal shell-shape, see Fig. 4). The lower panel shows the simulated shapes for points across the space. The central configuration (shown at [0,0] and enclosed in a box) is the mean shape. Note that in this morphospace the Veneridae are much closer to the Solenoida, and that *Legumen* bridges the space between groups. Point shapes and colors are as given in the legend to Figure 3. *Macrocallista nimbosa* is indicated (“M. nm.”), though not illustrated (see Fig. 4 for an image) as a point of comparison.

**Ecomorphology of Legumen and the contraction of shell disparity within the Veneridae.**—The intermediate position of *Legumen* between the Solenoida and the rest of the Veneridae in all three morphospaces (Figs. 3–5) indicates that, whereas the genus contains members that are similar in outline to modern infaunal venerids, although distinctively compressed (e.g., *L. carolinense*, *L. ooides*), it also includes a number of species that represent a form-group that is entirely absent in the modern venerid fauna. We will devote most of this discussion to the ecomorphology of the scanned representative of this group, *Legumen ellipticum* Conrad, 1858.

Bivalve shell morphology is tightly linked to life habit (Stanley, 1970, 1975; Seilacher and Gishlick, 2014); anecdotal examples of convergence in form can be seen in even a glance at any large collection of shells. Shell anatomy can be broken down into a number of functional structures related to life habit, such as overall form and sculpture, which interact with the substratum, and the ligament, adductor muscles, and hinge teeth, which operate together to open and close the shell. The position of the hinge relative to the muscle scars and the overall longest axis of the shell controls both the orientation of the shell during the burrowing period, and the amount of “rocking” the animal is capable of during a burrowing sequence.
Legumen ellipticum is strikingly close to the smaller Pharidae (e.g., Siliqua) in shell form and hinge geometry, and given the strong relationship between shell form and life habit, we infer similar life modes for the two genera. The reduced cardinal area in L. ellipticum, which would have much diminished resistance to shear during burrowing compared to, for example, Dosiinia, which are fast burrowers, but have large cardinal areas, suggests that, like the pharids, its burrowing method did not have a significant rock component, so that it did not need to expend resources on structures for counteracting shear.

Pharids and the related solenids have an anterior pedal gape, which allows them to burrow almost vertically in semi-permanent tubes, and to make fast escapes from their burrows if disturbed (Stanley, 1970). Lack of a pedal gape indicates that Legumen were likely not tube dwellers, but their thin, smooth shell, small size, and long posterior all suggest adaptation for unconsolidated, muddy sediment (Stanley, 1970), and the small size and truncation of the posterior suggests a more vertical final life position than that occupied by Macrocallista nimbosa, which is otherwise the most comparable venerid. Limited literature exists on the life habits and ecology of the smaller Pharidae, but L. ellipticum plots most consistently with Siliqua, Neosiliqua, and Sinonovacula, all of which live buried sub-vertically in shallow-marine or estuarine, muddy or silty sand substrata, and L. ellipticum evidently lived in similar sediments (e.g., Sohl and Koch, 1983, 1984, 1987; Ebersole, 2016).

Modern Veneridae occupy a wide variety of habitats today across most available shell sediments. Many species of venerid co-occur, with co-existence perhaps mediated by predation pressure, by micro-partitioning their habitats, or simply because competition is low between filter-feeders (e.g., the striking diversity of Chioninae in western North America; Roopnarine 2001; Coan and Valentich-Scott, 2012). Despite this species richness, in the 66 Myr since the K-Pg extinction, no living venerids have recapitulated the shell form of Legumen to re-occupy this area of morphospace and exploit this life-habit, currently occupied by the smaller Pharidae. Cretaceous venerids were less diverse taxonomically, with 21 known genera in the Cretaceous venerids that represent both a lineage and a unique mode of life for venerids that was lost in the K/Pg extinction.

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Accessibility of supplemental data

All metadata, measurement data, and the PCA scores for both morphospaces are available from the Dryad Digital Repository: https://doi.org/10.5061/dryad.ncjsxksqz. CT scans of Legumen are available on Morphosource: http://www.morphosource.org/Detail/ProjectDetail/Show/project_id/773.

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Appendix: Additional descriptions of the morphology of Legumen Conrad, 1858

The original description of the genus *Legumen* Conrad, 1858 is as follows: “Shell equivale, very inequilateral, flattened; hinge with two very slender teeth in the right valve under the beak, and one posterior very oblique prominent lamelliform tooth. This genus is perhaps most nearly related to *Calidess*, Schum. It can readily be distinguished by external characters; its flat valves, straight and more produced anterior side, &c.” In the Treatise on Invertebrate Paleontology (Keen, 1969, p. N683), the genus is diagnosed as “elongate, slender, subelliptical, beaks at anterior fourth; sculpture concentric only; no lunule or escuteon; hinge plate narrow; pallial sinus moderate.” Both diagnoses rely on the markedly plain exterior, rather than the extremely distinctive denticulation. We provide an updated diagnosis, and descriptions for the two species imaged by micro-CT as part of this study.

Genus *Legumen*, Conrad 1858

**Diagnosis.**—Shell elongate, equivale, beaks at the anterior quarter to anterior fifth; sculpture weak concentric to absent with growth lines; cardinals thin, prominent, the anterior cardinal always vertical to subvertical; posterior cardinals thin and oblique, following the straight dorsal margin; no lunule or escuteon; lamelliform teeth; hinge plate narrow; pallial sinus moderate.

*Legumen ellipticum* Conrad, 1858

Appendix Figure 1.1–1.8

**Description.**—Equivale, inequilateral, with umbones at the anterior fifth. Highly compressed. Exterior smooth with fine growth lines. In the right valve, cardinals 3a (anterior) and 1 (median) are thin, subvertical, and very prominent; cardinal 3b (posterior) is thin, prominent, at least four times the length of 3a, and directed extremely obliquely, paralleling the posterior of the shell. In the left valve, cardinals 2a (anterior) and 2b...
Appendix Figure 1.  (1–8) *Legumen ellipticum* Conrad, 1858 (USNM 76669, Nacatoch Sand, Kaufman County). (1) Right valve, dorsal view; (2) right valve, internal view; (3) right valve, anterior view; 10 mm scale bar applies to (1–3); (4) closeup of right valve hingeplate, cardinal teeth annotated; (5) left valve, dorsal view; (6) left valve, internal view; (7) left valve, anterior view; (8) closeup of left valve hingeplate, cardinal teeth annotated. (9–12) *Legumen carolinense* (Conrad, 1875) (USNM 31802, Tar Heel Formation, Snow Hill, North Carolina). (9) Right valve, dorsal view; (10) right valve, internal view; (11) right valve, anterior view; (12) closeup of right valve hingeplate, cardinal teeth annotated. (13–16) *Legumen carolinense* (Conrad, 1875) (USNM 31947, Tar Heel Formation, Snow Hill, North Carolina). (13) Left valve, dorsal view; (14) left valve, internal view; (15) left valve, anterior view; (16) closeup of left valve hingeplate, cardinal teeth annotated. 10 mm scalebar applies to (1–3, 5–7, 9–11, 13–15); 5 mm scale bar applies to (4, 8, 12, 16).

*Legumen carolinense* (Conrad, 1875)

Appendix Figure 1.9–1.16

Description.—Equivalve, inequilateral, with umbones at the anterior quarter. Neither strongly compressed nor particularly inflated. Exterior smooth with occasional strong growth lines. In the right valve, cardinals 3a (anterior) and 1 (median) are thin, subvertical, and very prominent; cardinal 3b (posterior) is thin, prominent, unequally bifid, the anterior partition being shorter. In the left valve, cardinals 2a (anterior) and 2b (median) are narrowly triangular and divergent. Cardinal 4b is partly coalescent with the ligamental nymph. Lunule and escutcheon absent. Ligamental groove long, narrow, and well marked, with narrow nymph below. The adductor muscle scars are subequal with the posterior slightly larger, oval, and slightly impressed. The pallial line is wide and follows the curvature of the ventral margin. The pallial sinus is V-shaped, nearly horizontal, extending to about the midpoint of the internal disc. The inner margin is smooth.