

Paleontological Contributions

Number 8

Comprehensive database on Induan (Lower Triassic) to Sinemurian (Lower Jurassic) marine bivalve genera and their paleobiogeographic record

Sonia Ros-Franch, Ana Márquez-Aliaga, and Susana E. Damborenea





April 10, 2014 Lawrence, Kansas, USA ISSN 1946-0279 (online) paleo.ku.edu/contributions http://hdl.handle.net/1808/13433



Paleontological Contributions

Number 8

COMPREHENSIVE DATABASE ON INDUAN (LOWER TRIASSIC) TO SINEMURIAN (LOWER JURASSIC) MARINE BIVALVE GENERA AND THEIR PALEOBIOGEOGRAPHIC RECORD

Sonia Ros-Franch,^{1,2,3*} Ana Márquez-Aliaga,¹ and Susana E. Damborenea^{2,3}

¹Geology Department and Cavanilles Institute of Biodiversity and Evolutionary Biology, University of Valencia, Av. Doctor Moliner 50, Burjassot, Valencia 46100, Spain, soniaros@fcnym.unlp.edu.ar, Ana.Marquez@uv.es, ²Invertebrate Paleozoology Division, La Plata Museum, Paseo del Bosque s/n, La Plata 1900, Argentina, sdambore@fcnym.unlp.edu.ar; ³Consejo Nacional de Investigaciones Científicas y Técnicas (CONICET), Rivadavia 1917, Buenos Aires, Argentina

ABSTRACT

Marine bivalve genera that were described or mentioned for Triassic and Lower Jurassic deposits worldwide are reviewed in terms of their validity, stratigraphic range, paleogeographic distribution, paleoautecology, and shell mineralogy. Data were originally compiled at species level and are arranged systematically. A brief discussion for each genus includes synonymy, taxonomic status, and included subgenera, as well as current uncertainties about their validity and range. The distribution of each genus is also shown on paleogeographic maps. Type species and first and last appearances of each genus are also mentioned. We recognize as valid 281 genera and their included subgenera, and we further discuss 148 genera (arranged alphabetically) that were mentioned for the study interval but are not included for different reasons. The purpose of this paper is to provide an updated critical assessment of all available basic information for each genus, in order to obtain a sound database to study the generic paleodiversity of marine bivalves in the time interval from the Induan (Early Triassic) to the Sinemurian (Early Jurassic). This was a critical time for bivalve evolution and diversification, which began with the recovery from the Permian–Triassic extinction and ended with the recovery from the Triassic–Jurassic extinction.

Keywords: Bivalvia, paleoecology, paleogeography, Triassic, Early Jurassic

INTRODUCTION

In order to study the generic paleodiversity of marine bivalves in the time interval from Induan (Early Triassic) to Sinemurian (Early Jurassic) (In-Sin), which begins with the recovery from the Permian– Triassic extinction and ends with the recovery from the Triassic–Jurassic extinction, a significant critical review of each genus was badly needed. The purpose of this paper is not merely to offer such a compendium of data extracted from the published literature, but to provide an updated critical assessment of all available information for each genus as well.

*Corresponding author.

This paper is thus a review of all marine bivalve genera that were described for Triassic and Lower Jurassic deposits worldwide, in terms of their validity, stratigraphic range, paleogeographic distribution, paleoautecology, and shell mineralogy. We include marine bivalves only, and thus the families Pachycardiidae, Unionidae, and Neomiodontidae are not considered. Data were compiled at species level, with the purpose being to contrast the assignment of each species to the genera. Only published monographs and papers that include images and descriptions of the taxa were considered. This paper is a revised version of part of a Ph.D. thesis (Ros, 2009).

Methodology

We based our study mostly on bibliographic data, but the different records attributed to each genus were revised, checked, and critically updated as far as possible from systematic, stratigraphic, and geographic points of view.

First, we looked for all genera described in our study interval. The *Treatise on Invertebrate Paleontology* (Cox & others, 1969; Stenzel, 1971) was taken as reference for the period previous to 1965, accepting in most cases the proposed synonyms, although in some genera, subsequent literature that changed the *Treatise* views was also taken into account. However, we frequently used literature prior to the *Treatise* to try to elucidate some pending questions. For the time span between 1965 and the present, an exhaustive literature search was performed. To carry out this task, we referred to *Zoological Record* from 1965 to 2005 and other sources (Diener, 1923; Kutassy, 1931; Neave, 1939; Cox & others, 1969; Sepkoski, 2002; and several electronic sources, such as Paleobiology Database [PBDB] and *Nomenclator Zoologicus*).

Then we compiled all references into a bibliography about these genera and reviewed it. Since the amount of literature and data discussed here is quite large, we first proceeded to develop a bibliographic database that allowed us to handle them rapidly and effectively. This database was made using the computer program FileMaker Pro 8.5. The selection of this program was based on the simplicity of its construction, handling, and relation of information. The introduction of the species with author and year in the database is very important, instead of compiling the genera directly, since it allows us to control the different scope given to them over time. For example, the species decidens Bittner, 1899, was described in the genus Pseudomonotis, assigned later to Streblochondria and Claraia, in the end to be assigned by Newell and Boyd (1995) to Crittendenia and also as type species of *Claraia* (Bittnericlaraia) by Gavrilova (1996). If we had not introduced the species in the database, it would have been impossible to continue the history of this species, and we would have had the occurence of the same species in four different genera.

Through a careful revision of the literature, we eliminated all genera that, for one reason or other, should not be considered, including those with only doubtful occurrence in our study interval. These are treated in the section Genera not Included (see p. 156), listed alphabetically, with a brief explanation of the reason for their exclusion. Specifically, we do not include: (1) genera convincingly placed in synonymy; (2) subgenera for which we did not find any publication elevating them to genus level; (3) genera with no solid presence during the temporal study range, although they could have been mentioned for the study range; and (4) generic homonyms. Finally, there are some taxa that were listed in the *Compendium of Fossil Marine Genera* published by Sepkoski (2002), but which were not included in this analysis because they are regarded as subgenera.

Taxonomic data were reviewed as far as possible in order to assign all species to genera. We compared the generic diagnosis with the species descriptions and figures offered in the literature, and, in some instances, we consulted different specialists. We are aware that, even considering these meticulous analyses, it is impossible to eliminate all mistakes, and taxonomic decisions are always subjective, so the opinion of different authors is frequently indicated in the genus discussion. For the assignment of genera to different families, we followed Cox and others (1969) for genera described earlier than the *Treatise*, but more recent bibliographies that proposed changes in assignation are discussed and listed for each case. For genera described after 1965, we follow critically the assignment published in the literature. The survey includes only papers published before 2011.

Data Organization

Stratigraphic Ranges.—We follow the stratigraphic chart of Gradstein and Ogg (2004), using their stage names, except for the Permian, which is subdivided into three epochs: Cisuralian, Guadalupian, and Lopingian, here listed as early, middle, and late Permian, respectively.

For the equivalence between the different charts used in the reviewed bibliography, we used the conversion tables provided by the Paleobiology Database (http://www.paleodb.org/) and GeoWhen Database (http://www.stratigraphy.org/geowhen/index.html).

We had some problems distinguishing between Rhaetian and Norian in papers older than the redefinition of Rhaetian by Dagys and Dagys (1994). At least the Kössen Formation in Austria and the Gabbs Formation in the United States can be regarded as Rhaetian in age (Dagys & Dagys, 1994; Hallam, 2002).

We used H. J. Campbell and Raine (in Cooper, 2004) and H. J. Campbell, Raine, and Wilson (in Cooper, 2004) for the correlation of New Zealand stages with the Global Geochronological Scale.

For each genus, we indicate the entire stratigraphic range observed after reviewing the literature, and the two records we regard as the first and last appearances. Stratigraphic ranges are compared with those in Cox and others (1969), which is the most recent published review with a thorough taxonomic revision and stratigraphic data. In most cases, those ranges are changed with the new information here considered. We also compared our data with Sepkoski's (2002) compilation.

It should be pointed out that the stratigraphic ranges offered are observed, i.e., the first and last occurrences of a taxon are the limits of its stratigraphic range, which is only an approximation to the real range, and therefore also to the moment of origination and extinction. The effects of sampling, stratigraphic hiatuses, transgressions and regressions, Signor-Lipps effect, and other factors may greatly influence or distort the actual ranges (Holland, 1995). We must also bear in mind that the distribution of fossil bivalves is particularly dependent on facies.

Paleogeographic distribution.—For each genus, we provide the paleogeographic distribution during the time interval considered in this review. For those genera present in the Paleozoic, the paleogeographic distribution during the late Permian is also given. This section is not intended as a paleobiogeographic study, although the domains considered were established both in a paleogeographic and a paleobiogeographic sense (Westermann, 2000). We merely review the distribution of each genus and represent it on paleogeographic maps. Consequently, the term domain as used here has no paleobiogeographic implication.

The distribution of each genus in space and time is shown in three maps, for Permian–Triassic, Middle Triassic, and Triassic–Jurassic intervals. These representations sketch the position of continents at the three time moments selected, but they are not strictly faithful to all recent knowledge in every detail. The maps are based on and adapted from various sources. The first map was compiled mainly following Ziegler, Hulver, and Rowley (1997) and Christopher Scotese's maps available on his website Paleomap Project (http:// www.scotese.com); the following two maps were based on Golonka and Ford (2000) and Golonka (2004, 2007). We introduced some changes, especially in the configuration of Cimmerian block, Lhasa block, and southern part of the Tethys following Dèzes (1999), Nicoll (2002), J. Yin and Grant-Mackie (2005), and J. Yin and McRoberts (2006).

The paleogeographic distribution for each genus was recorded by using contemporary country names, which were then grouped into the following paleogeographic informal domains: Tethys, Circumpacific, Boreal, and Austral. As here understood, the Tethys domain covers the entire length of the Tethys Sea during the time interval here considered, without differentiating between Neotethys and Paleotethys. The Boreal domain includes mainly the northern part of Russia, Greenland, northern Canada, and Alaska. The Austral domain comprises the southern part of South America (part of Argentina and Chile), New Zealand, New Guinea, and Antarctica. Finally, the Circumpacific domain covers the Paleopacific, being limited to the north and south by the Boreal and Austral domains, respectively. For the location of countries in the different domains, we mostly follow Nakazawa (1991), Dercourt, Ricou, and Vrielynck (1993), Metcalfe (1998, 1999), Gaetani and others (2000a, 2000b, 2000c), Acharyya (2000), Stampfli and others (2001), Stampfli and Borel (2002, 2004), Chumakov and Zharkov (2003), and Klets (2005), in addition to those cited above for construction of the maps. Every genus, even those with localized occurrences, was referred to one or more of these domains.

Although we indicate the distribution of each genus only during the time interval considered in this review, in some cases, we also discuss their paleogeographic range before or after this interval, if we find this relevant for any reason. The distribution of each genus is listed according to the domains just mentioned, and each indicates the countries in which the genus was found with the relevant bibliographic data sources. If a distribution for a genus in the literature is uncertain, we use a question mark (?) herein for that record. We occasionally discuss data included in papers with no illustrations of the specimens, but these records were not taken into account for the distribution of genera. These are mostly related to Russia and China, especially to some pre-1980 literature that we could not see for this study.

Paleoautecology.—The modes of life of the genera included are assigned according to the original bibliographic source, and, when this was not possible, they were inferred by functional morphology or analogy with related Recent species. The categories recognized here are based mainly on S. M. Stanley (1968, 1969, 1970, 1972), Kauffman (1969), Bambach (1977, 1983), Bambach, Bush, and Erwin (2007), and others. We are aware that there are many exceptions to the general guidelines given for the recognition of modes of life, so they will necessarily be tentative and always referred to adult specimens (the different modes of life that a bivalve can display along its ontogeny were not taken into account). Sometimes it was not possible to assign a unique mode of life to one genus, because the included species may differ in this aspect. When there was not enough information about the genus morphology or about the environment in which it is recorded, we refer to the predominant mode of life within the family.

The following aspects were taken into account for establishing the different modes of life: life position in relation to water column and substrate, trophic group, mobility, and fixation. The mode of life assigned to each genus is coded by letters, as follows:

Position on the water column: benthic [B] or pseudoplanktonic [Ps].

Trophic group: suspensivorous [S] and detritivorous [D]. Carnivorous bivalves are mostly beyond our study interval, because septibranchs appeared in the Jurassic (later than Sinemurian). In addition, we indicate possible photosymbiotic [Ph] and chemosymbiotic [Ch] relationships with microorganisms.

Life position in relation to the substrate: epifaunal [E], shallow infaunal [Is], deep infaunal [Id] or semi-infaunal [Se].

Mobility: sedentary [Sed], facultative mobile [FaM], slow mobile [SM], and fast mobile [FM]. Regarding mobility, SM and FM are categories only considered for burrower bivalves, while FaM refers to swimmer and pseudoplanktonic bivalves.

Fixation: we consider if they lived attached to the substrate or unattached [Un]; attached bivalves can be byssate (endobyssate [Endo] or epibyssate [Epi]) or cemented [C].

Several modes of life are then defined by the intersection of the categories just mentioned: shallow burrower in soft substrate [Sb], deep burrower [Db], borer [Bo], byssate [By], cemented [C], recliner [R], swimmer [Sw], nestler [N].

Mineralogy.—Shell mineralogy data provided here are taken mostly from J. D. Taylor, Kennedy, and Hall (1969, 1973), Carter (1990a, 1990b, 1990c), Carter, Lawrence, and Sanders (1990), and Carter, Barrera, and Tevesz (1998). In specific cases, we used other sources that are indicated in the discussion of each genus.

Mineralogy of shell layers is given for each genus, when this information is available; alternatively, we assign the predominant mineralogy for the family. Three types of mineralogy are considered: aragonitic, when all shell layers are fully formed by aragonite; bimineralic: when at least one of the shell layers is calcitic and the others aragonitic; and calcitic, when all shell layers are formed by calcite.

INCLUDED GENERA

The systematic arrangement used here follows Amler (1999), Amler, Fischer, and Rogalla (2000), and Bouchet and Rocroi (2010) with some modifications. Those are the most complete general Bivalve mollusk classifications that include fossil families introduced after the *Treatise on Invertebrate Paleontology*. The changes introduced here are: Lipodonta is recognized as a subclass following Cope (1995) and including the Solemyoidea; family Pichleriidae is included with the Limopsoidea; superfamilies Dimyoidea and Plicatuloidea are included within the Ostreida rather than the Pectinida; the name Terquemiidae Cox, 1964, is replaced by Prospondylidae Ptchelincev, 1960 (Hautmann 2001a), the classification of the superfamily Kalenteroidea Marwick, 1953, has been emended, according to Z. Fang and Morris (1997) and Damborenea (2004).

The genera included in this review are listed in systematic order below (Table 1). Each of them is then briefly discussed separately, with indication of type species, possible synonym names, and details of stratigraphic and paleogeographic distribution, mode of life, and shell structure. Table 1. Summary of various data for included genera, arranged in the order in which they are discussed herein. Abbreviations: (1) Stratigraphic range: O, Ordovician; Tre, Tremadocian; Dev, Devonian; Llan, Llanvirnian; Fam, Famennian; Car, Carboniferous; Miss, Mississippian; Vis, Visean; Penn, Pennsylvanian; Pe, Permian; Sak, Sakmarian; Art, Artinskian; Guad, Guadalupian; Wuch, Wuchiapingian; Chang, Changhsingian; Tr, Triassic; In, Induan; Ol, Olenekian; Ani, Anisian; Lad, Ladinian; Car, Carnian; Nor, Norian; Rha, Rhaetian; J, Jurassic; Hett, Hettangian; Sin, Sinemurian; Plie, Pliensbachian; Toa, Toarcian; Aal, Aalenian; Baj, Bajocian; Call, Callovian; Oxf, Oxfordian; Kim, Kimmeridgian; Tit, Tithonian; Cret, Cretaceous; Berr, Berriasian; Val, Valanginian; Haut, Hauterivian; Apt, Aptian; Alb, Albian; Cen, Cenomanian; Tur, Turonian; Cam, Campanian; Maa, Maastrichtian; P. Paleocene; Dan, Danian; L., Lower; M., Middle; U., Upper; (2) Paleoautoecology: B; benthic, Ps, pseudoplanktonic, D, detritivorous, S, suspensivorous, Ph, photo-symbiotic; Ch, chemosymbiotic, E, epifaunal, Is, shallow infaunal, Id, deep infaunal, SI, semi-infaunal, FM, fast mobile, SM, slow mobile, FaM, facultative mobile, Sed, sedentary, Un, unattached, Endo, endobyssate, Epi, epibyssate, C, cemented, Sb, burrower in soft substrate, Db, deep burrower, Bo, borer, By, byssate, R, recliner, Su, swimmer, N, nestler.

Genus	Stratigraphic range	Paleogeographic distribution	Paleoautoecology	Mineralogy
Palaeonucula	M. Tr (l. Ani)–L. Cret (Apt)	Tethys, Circumpacific, and Boreal	B, D, Is, FM; Sb	Aragonitic
Trigonucula	U. Tr (Car–u. Rha)	Tethys and Circumpacific	B, D, Is, FM; Sb	Aragonitic
Nuculoma	U. Tr (Rha)–L. Cret (Val)	Tethys and Circumpacific	B, D, Is, FM; Sb	Aragonitic
Nuculana	M. Tr (Ani)–Recent	Cosmopolitan	B, D, Is, FM; Sb	Aragonitic
Phestia	M. O (l. Llan)–U. Tr (Car)	Tethys, Circumpacific and Boreal	B, D, Is, FM; Sb	Aragonitic
Veteranella	L. Pe (Art)–U. Tr (Nor)	Tethys	B, D, Is, FM; Sb	Aragonitic
Eleganuculana	U. Tr (Nor)	E Tethys	B, D, Is, FM; Sb	Aragonitic
Xiaoschuiculana	U. Tr	Tethys and Circumpacific	B, D, Is, FM; Sb	Aragonitic
Phaenodesmia	M. Tr (Ani)–U. Tr (Rha)	Tethys and Circumpacific	B, D, Is, FM; Sb	Aragonitic
Prosoleptus	M. Tr (Ani)–U. Tr (Car)	Tethys and Boreal?	B, D, Is, FM; Sb	Aragonitic
Palaeoneilo	L. O (Tre)–L. J (Toa)	Cosmopolitan	B, D, Is, FM; Sb	Aragonitic
Lapteviella	M. Tr (Ani)	Boreal	B, D, Is, FM; Sb	Aragonitic
Dianucula	U. Tr (Nor)	E Tethys	B, D, Is, FM; Sb	Aragonitic
Ningliconcha	U. Tr (Nor)	E Tethys E Tethys	B, D, Is, FM; Sb	Aragonitic
0		E Tethys		
Yongshengia	U. Tr (Nor)		B, D, Is, FM; Sb	Aragonitic
Rollieria Budania	L. J (Hett)–L. Cret	Tethys and Circumpacific	B, D, Is, FM; Sb	Aragonitic
Ryderia	U. Tr (Rha)–L. J (Toa)	Tethys and Circumpacific	B, D, Is, FM; Sb	Aragonitic
Dacryomya	U. Tr (Nor)–U. J (Kim)	Tethys, Circumpacific and Boreal	B, D, Is, FM; Sb	Aragonitic
Mesoneilo	U. Tr (Nor–Rha)	Tethys and Boreal?	B, D, Is, FM; Sb	Aragonitic
Nucinella	L. J (Hett)–Recent	E Tethys	B, D/Ch, Is, FM; Sb	Aragonitic
Solemya	U. Car (u. Penn)–Recent	Tethys and Circumpacific	B, Id, S/Ch, FM; Db	Aragonitic
Modiolus	U. Dev (Fam)–Recent	Cosmopolitan	B, Se, S, Endo, Sed; By	Bimineralic
Promytilus	Car (Miss)–L. Tr (In)	Boreal	B, E, S, Epi, Sed; By	Bimineralic
Inoperna	U. Tr (Rha)–U. Cret (Maa)	Tethys and Austral?	B, Se, S, Endo, Sed; Bo?	Bimineralic
Falcimytilus	U. Tr (Car)–U. J (Tit)	Tethys, Circumpacific and Boreal	B, E, S, Epi, Sed; By	Bimineralic?
Lycettia	L. J (Sin)–U. Cret (Maa)	Austral	B, E, S, Epi, Sed; By	Aragonitic
Lithophaga	U. Tr (Nor)–Recent	Tethys	B, I, S, By, Sed; Bo	Bimineralic
Mysidiella	U Tr (Car–Rha)	Tethys and Circumpacific	B, E, S, Epi, Sed; By	Bimineralic
Botulopsis	M. Tr (Lad)–U. Tr (Car)	W Tethys	B, E, S, Epi, Sed; By	Bimineralic
Promysidiella	M. Tr (l. Ani)–U. Tr (l. Nor)	Tethys, Circumpacific and Boreal?	B, E, S, Epi, Sed; By	Bimineralic
Protopis	M. Tr (Ani)–U. Tr (Car)	Tethys	B, E, S, Epi, Sed; By	Bimineralic ?
Joannina	L. Tr (In)?–U. Tr (Car)	Tethys	B, Se, S, Endo, Sed; By	Bimineralic ?
Leidapoconcha	M. Tr (Ani)	E Tethys	B, Se, S, Endo, Sed; By	Bimineralic ?
Waijiaonella	M. Tr (Ani)	E Tethys	B, Se, S, Endo, Sed; By	Bimineralic ?
Qingyaniola	M. Tr (Ani)	E Tethys	B, Se, S, Endo, Sed; By	Bimineralic ?
Macrodontella	M. Tr (Ani)	Tethys	B, E/Se, S, Epi/Endo, Sed; By	Aragonitic
Catella	U. Tr (Car)–L. P (Dan)	E Tethys	B, Se, S, Êndo, Sed; By	Aragonitic
Parallelodon	M. Dev–U. Cret	Cosmopolitan	B, E, S, Epi, Sed; By	Aragonitic
Grammatodon	L. Pe (Art)–U. Cret (Maa)	Cosmopolitan	B, E, S, Epi, Sed; By	Aragonitic
Bapristodia	U. Tr (Nor)	E Tethys	B, E, S, Epi, Sed; By	Aragonitic
Cucullaea	L. J (Hett)–Recent	Circumpacific	B, Is/Se, S, SM; Sb	Aragonitic
Eophilobryoidella	M. Tr (u. Ani)	E Tethys	B, E, S, Epi, Sed; By	Aragonitic
Hoferia	U. Tr (Car)	Tethys	B, Is, S, Endo, SM; Sb	Aragonitic
Pichleria	U. Tr (Car)	Tethys	B, Is/Se, S, Sed; ?	Aragonitic
Elegantarca	M. Tr (Ani)–U. Tr (Car)	Tethys	B, Se, S, Endo, Sed; By	Aragonitic
Myalina	Car (l. Miss)–U. Pe, ;L. Tr?	Circumpacific	B, E, S, Epi, Sed; By	Bimineralic
Myalinella	Car (Vis)–L. Tr (u. Ol)	Tethys, Circumpacific and Boreal	B, Se, S, Endo, Sed; By	Bimineralic
Promyalina	U. Pe (u. Chang)–L. Tr (u. Ol)	Tethys, Circumpacific and Boreal	B, Se, S, Endo, Sed; By	Bimineralic
Aviculomyalina	M. Tr (Ani)	W Tethys	B, E, S, Epi, Sed; By	Bimineralic
Parainoceramus	L. J (Hett)–U. J (Tit)	Tethys, Circumpacific and Boreal	B, E, S, Epi, Sed; By	Bimineralic
Pseudomytiloides	L. J (Hett) $-M$. J (Aal)	Tethys	B/Ps, E, S, Epi, Sed/FaM; By	Bimineralic
Arctomytiloides	L. J (Sin-Toa?)	Boreal and Circumpacific	B, E, S, Epi, Sed; By	Bimineralic
Pteria	L. Tr (Ol)–Recent	Tethys, Circumpacific and Boreal	B, E, S, Epi, Sed; By B, E, S, Epi, Sed; By	Bimineralic
Arcavicula	L. Tr (l. Ol)–U. Tr (Rha?)	Tethys and Circumpacific	B, E, S, Epi, Sed, By B, E, S, Epi, Sed; By	Bimineralic
Rhaetavicula	U. Tr (Rha)	Tethys and Circumpacific	B, E, S, Epi/Un, Sed; By/R	Bimineralic
Stefaninia	M. Tr (u. Lad)–U. Tr (Car)	W Tethys	B, E, S, Epi/Oil, Sed; By/R B, E, S, Epi, Sed; By	Bimineralic
Bakevellia	U. Pe–U. Cret	Cosmopolitan	B, Se, S, Endo, Sed; By B, Se, S, Endo, Sed; By	Bimineralic
Gervillella	L. J (Hett)–U. Cret (?)	W Tethys, Circumpacific and Austral		
			B, Se, S, Endo, Sed; By B/De Se/E S Endo/Eni Sed; By	Bimineralic
Gervillia	M. Tr (Lad)–U. Cret (Maa)	Tethys and Circumpacific	B/Ps, Se/E, S, Endo/Epi, Sed; By	Bimineralic
Hoernesia	L. $Tr(Ol)-U$. $Tr(Rha)$	Tethys and Boreal	B, Se, S, Endo, Sed; By	Bimineralic
Langsonella	M. Tr (Ani) $-$ U. Tr (Car)	E Tethys	B, Se, S, Endo, Sed; By	Bimineralic
Cuneigervillia	L. J (Hett)–L. Cret $(?)$	W Tethys	B, Se, S, Endo, Sed; By	Bimineralic
			K Ne N Hindo Nedi By	Bimineralic
Gervillaria Gervillancea	M. Tr (Ani)–U. Cret (Tur) U. Tr (Car–Nor)	Tethys, Circumpacific and Austral S Tethys	B, Se, S, Endo, Sed; By B, Se, S, Endo, Sed; By	Bimineralic

Table 1 (continued). See facing page for heading.

Genus	Stratigraphic range	Paleogeographic distribution	Paleoautoecology	Mineralogy
Songdaella	U. Tr (Nor)	E Tethys	B, E, S, Epi, Sed; By	Bimineralic
Aguilerella	U. Tr (Rha)–L. Cret (Haut)	E Tethys and Circumpacific	B, E, S, Epi, Sed; By	Bimineralic
Towapteria	L. Pe (Sak)–L. Tr (In)	Tethys and Circumpacific?	B, E, S, Epi, Sed; By	Bimineralic
1		S Tethys	B, Se, S, Endo, Sed; By	Bimineralic
<i>Virgellia</i>	U. Tr (Car) $-$ U. J (Kim)	,		
Gervilleioperna	L. J (Sin)–M. J (Aal)	Circumpacific	B, Se, S, Endo, R, Sed; By	Aragonitic
Cassianella	Pe?, M. Tr (Ani)–U. Tr (Rha)	Cosmopolitan	B, E, S, Un, Sed; R	Aragonitic
Burckhardtia	U. Tr (Car)	Circumpacific	B, E, S, Un, Sed; R	Aragonitic
Ioernesiella	U. Tr (Car)	W Tethys	B, E, S, Un, Sed; R	Aragonitic
ilangina	M. Tr (Ani) - U. Tr (Car)	E Tethys	B, E, S, Un, Sed; R	Aragonitic
Peubenia	U. Tr (Car)	W Tethys	B, E, S, Un, Sed; R	Aragonitic
eptihoernesia	M. Tr (Lad)–U. Tr (Car)	Tethys and Circumpacific	B, E, S, Un, Sed; R	Aragonitic
1	. , , , ,			
Datta	U. Tr (Rha)	S Tethys	B, E, S, ?, ?	Bimineralic
ognomon	U. Tr (Car)–Recent	Tethys	B, Se, S, Endo, Sed; By	Bimineralic
eproconcha	M. Tr	W Tethys	B, E, S, ?, ?	Unknown
Vaagenoperna	L. Pe (Sak)–U. Tr (u. Nor)	Tethys and Circumpacific	B, Se, S, Endo, Sed; By	Bimineralic
ositra	L. Tr (l. Ol)–M. J (l. Oxf)	Tethys, Circumpacific and Boreal	B, E, S, Un, Sed; R	Bimineralic
monotis	U. Tr (Car)	Tethys	B, E, S, Epi, Sed; By	Bimineralic
eldidenella				Bimineralic
	M. Tr (u. Ani–u. Lad)	W Tethys	B, E, S, Epi, Sed; By	
aenodiotis	L. J (Sin–Plie)	W Tethys	B, E, S, Epi, Sed; By	Bimineralic
einmannia	L. J (u. Sin–l. Toa)	Boreal	B, E, S, Un, Sed; R	Bimineralic
llesmerella	L. Tr (u. Ol)	Boreal	B, E, S, Un, Sed; R	Bimineralic
inna	L. Tr (?)–Recent	Tethys, Circumpacific and Austral	B, Se, S, Endo, Sed; By	Bimineralic
trina	M. Tr (Ani)–Recent	Tethys and Circumpacific	B, Se, S, Endo, Sed; By	Bimineralic
alaeolima	U. Dev (Fam?)–U. Tr (Nor)	Tethys and Circumpacific	B, E, S, Epi, Sed; By	Bimineralic
viculolima	M. Tr (Ani)	W Tethys	B, E, S, Epi, Sed; By	Bimineralic
adiotella	M. Tr (Lad)–U. Tr (Car)	Tethys	B, E, S, Epi/Un, Sed; By/R	Bimineralic
imatula	M. Tr (Lad) - Recent	Tethys, Circumpacific and Austral	B, E, S, Épi, FaM; By/Św	Bimineralic
imea	M. Tr (Ani)–Recent	Cosmopolitan	B, E, S, Epi, Sed; By	Bimineralic
lysidioptera	L. Tr (Ol)–U. Tr (Rha)	Cosmopolitan	B, E, S, Epi, Sed; By	Bimineralic
		*		
lagiostoma	M. Tr (Ani)–U. Cret (Maa)	Cosmopolitan	B, E, S, Epi, Sed; By	Bimineralic
rrania	U. Tr (Nor - Rha)	E Tethys	B, E, S, Epi, Sed; By	Bimineralic
irolidia	M. Tr (Lad)–U. Tr (Car)	W Tethys	B, E, S, Epi, Sed; By	Bimineralic
ntiquilima	M. Tr (Lad)-L. Cret (Apt)	Cosmopolitan	B, E, S, Epi, Sed; By	Bimineralic
tenostreon	U. Tr (u. Rha)–L. Cret (Val?)	Tethys, Circumpacific and Austral	B, E, S, Epi, Sed; By	Bimineralic
Fryphaea	U. Tr (Car)–U. Cret (Cam)	Cosmopolitan	B, E, S, Un, Sed; R	Bimineralic?
21				
Imbrostrea	M. Tr (Ani)–U. Tr (Rha)	Tethys	B, E, S, C, Sed; C	Bimineralic
ctinostreon	U. Tr (Rha)–U. Cret (Maa)	Tethys, Circumpacific and Austral	B, E, S, C, Sed; C	Calcitic
iostrea	U. Tr (Car)–U. Cret (Cen)	Cosmopolitan	B/Ps, E, S, C, Sed/FaM; C	Calcitic?
treta	U. Tr (Car)–U. Cret (Maa)	Tethys	B, E, S, C, Sed; C	Bimineralic
rotostrea	M. Tr (Ani)	E Tethys	B, E, S, C, Sed; C	Bimineralic
Iarpax	U. Tr (Nor)–L. J (Toa)	Austral and Boreal, Tethys?	B, E, S, C, Sed; C	Bimineralic
Toplicatula	U. Tr (Car–Rha)	Tethys	B, E, S, C, Sed; C	Bimineralic
seudoplacunopsis	M. Tr (Ani)–U. Tr (Rha)	Tethys and Circumpacific	B, E, S, C, Sed; C	Bimineralic
laraia	U. Pe (Wuch)–L. Tr (m. Ol)	Cosmopolitan	B/Ps, E, S, Epi, Se/FM	Bimineralic
umorphotis	L. Tr (In–Ol)	Tethys and Circumpacific	B, E, S, Epi, Sed; By	Bimineralic
rnithopecten	M. Tr (Ani)–U. Tr (Car)	Tethys	B, E, S, Epi, Sed; By	Bimineralic
xypteria	U. Tr (Car)	W Tethys	B, E, S, Epi, Sed; By	Bimineralic
· · · ·	M. Tr (Ani)–U. Tr (Car)	W Tethys	B, E, S, Epi, Sed; By B, E, S, Epi, Sed; By	Bimineralic
ntıjanıra				
mphijanira	M. Tr (Ani)–U. Tr (Car)	W Tethys	B, E, S, Epi, Sed; By	Bimineralic
rimahinnites	U. Tr (Nor–Rha)	Tethys	B, E, S, Epi, Sed; By	Bimineralic
eomorphotis	M. Tr (Ani)	Tethys	B, E, S, Epi, Sed; By	Bimineralic
rittendenia	L. Tr	Tethys and Circumpacific	B/Ps, E, S, Epi, Sed/FaM; By	Bimineralic
reblopteria	Car (Miss)–M. Tr (Ani)	Boreal	B, E, S, Epi, Sed; By	Bimineralic
eptochondria	M. Pe (Guad) $-$ U. Tr (Nor)	Tethys and Circumpacific	B, E, S, Epi, Sed; By	Bimineralic
1	. , . ,			
ittneria	U. Tr (Car)	W Tethys	B, E, S, Epi, Sed; By	Bimineralic?
lokonuia	U. Tr (u. Car– Nor)	Austral	B, E, S, Epi, Sed; By	Bimineralic
chuania	U. Tr (Nor)	E Tethys	B, E, S, Epi, Sed; By	Bimineralic?
larwickiella	M. Tr (Ani)	Austral	B, E, S, Epi, Sed; By	Bimineralic
nningella	L. J (Hett - Sin)	W Tethys	B, E, S, Epi, Sed; By	Bimineralic
lonotis	U. Tr (Nor–I. Rha)	Cosmopolitan	B/Ps, E, S, Epi, Sed/FaM; By	Calcitic?
		*		
tapiria	L. Tr (Ol)–U. J (Kim)	Cosmopolitan	B, E, S, Epi, Sed; By	Calcitic?
xytoma	L. Tr?, M. Tr (Lad)–U. Cret (Maa)	Cosmopolitan	B/Ps, E, S, Epi, Sed/FaM; By	Calcitic
vicularca	M. Tr (Lad)–U. Tr (Car)	W Tethys	B, E, S, Epi, Sed; By	Unknown
leleagrinella	U. Tr (Nor)–L. Cret (Alb)	Tethys, Circumpacific and Boreal	B/Ps, E, S, Epi, Sed/FaM; By	Bimineralic
anchuania	U. Tr (Rha?)	E Tethys	B, E, S, Epi, Sed; By	Bimineralic
almoxytoma	L. J (Hett–Plie)	Cosmopolitan	B, E, S, Epi, Sed; By	Calcitic
soella	M. Tr (Ani)–L. J (Plie)	Tethys, Circumpacific and Austral	B, E, S, Epi/Un, Sed; By/R	Calcitic?
talia	M. Tr (Ani)	Austral	B, E, S, Epi, Sed; By	Calcitic?
erquemia	L. J (Sin)–U. J?	Tethys	B, E, S, Ĉ, Sed; C	Bimineralic
ewaagia	M. Tr (Ani)–U. Tr (Rha)	E Tethys and Boreal	B, E, S, C, Sed; C	Bimineralic
ersia	U. Tr (Nor) $-L$. J (l. Hett)	Tethys	B, E, S, C, Sed; C	Bimineralic
		Circumpacific	B, E, S, C, Sed; C B, E, S, C, Sed; C	Bimineralic
Pegmavalvula	L. Pe (Art)–L. Tr (Ol)			

 Table 1 (continued). See p. 4 for heading.

 Paleogeographic distribution

Genus
Pergamidia
Krumbeckiella Manticula
Semuridia
Parapergamidia
Oretia Halobia
Daonella
Enteropleura
Aparimella Kolymonectes
Parvamussium
Chlamys
Weyla Indopecten
Camptonectes
Crenamussium
Avichlamys Pseudopecten
Agerchlamys
Canadonectites
Eopecten Janopecten
Ochotochlamys
Tosapecten
Nevadapecten Loxochlamys
Pleuronectites
Periclaraia Radulonectites
Entolium
Scythentolium
Filopecten Posidonotis
Entolioides
Permophorus
Curionia Triaphorus
Somareoides
Kalentera
Ouamoia Weixiella
Myoconcha
Pseudomyoconcha
Healeya Hippopodium
Trigonia
Praegonia
Prorotrigonia Prosogyrotrigonia
Geratrigonia
Vaugonia Vuushutuigania
Kyushutrigonia Acanomyophoria
Jaworskiella
Guineana Frenguelliella
Costatoria
Myophoria
Neoschizodus Agonisca
Gruenewaldia
Elegantinia
Caledogonia Atalantia
Liotrigonia
Erugonia
Guizhoumyophoria Minetrigonia
Myophorigonia
Perugonia
Maoritrigonia

Stratigraphic range U. Tr (Nor) U. Tr (Car-u. Rha) U. Tr (Car? - Nor), L. Cret (Berr) L. J (Sin) U. Tr (u. Car?-l. Nor) U. Tr (l. Nor) U. Tr (l. Car-m. Nor) M. Tr (Ani-Lad) M. Tr (m. Ani) M. Tr (u. Ani)–U. Tr (l. Car) U. Tr (Nor?)-L. J (l. Toa) M. Tr (Ani)-Recent M. Tr (Ani)-Recent L. J (Hett-Toa) U. Tr (Nor-Rha) M. Tr (Ani)–U. Cret (Maa) U. Tr (Car-Nor) L. Tr (Ol) L. J (Hett)-M. J (Baj) U. Tr (Car)-L. J (Toa) L. J (Sin-Plie) U. Tr (Car)-L. Cret (Alb) M. Tr (Ani)–U. Tr (l. Car) U. Tr (Nor) - L. J (Toa) U. Tr (Car-Rha) M. Tr (u. Lad) M. Tr (Lad)–U. Tr (Car) M. Tr (Ani)–U. Tr (Car) L. Tr (u. Ol) - M. Tr (l. Ani) L. J (Hett - Plie) L. Tr-U. Cret (Maa) L. Tr U. Tr (Car-Rha) L. J (Sin-Toa) L. Tr–U. Tr (Car) Car (Miss)-L. Tr (Ol) L. Tr (?)-U. Tr (Rha) U. Tr (Car-Nor) U. Tr (Car) U. Tr (Nor)-L. J (Toa) U. Tr (Nor - Rha) U. Tr (Nor - Rha) U. Tr (Rha)–U. Cret (Maa) M. Tr (Lad)-U. Tr (Rha) U. Tr (Nor-Rha) U. Tr (Rha?)-U. J (Tit) M. Tr (Ani)–U. Cret (Cen) M. Tr (Lad) U. Tr (Nor) U. Tr (Nor)-L. J (Sin) L. J (Hett-Toa) L. J (Hett)–U. J (Oxf) U. Tr (Car-Nor) U. Tr (Car) L. J (Hett-Plie) U. Tr (Car-Nor) U. Tr (Car)-M. J (Baj) U. Pe–U. Tr (Rha) M. Tr (Ani)–U. Tr (Rha) M. Pe (Guad)-U. Tr (Rha) M. Tr (Lad) U. Tr (Car - Rha) L. Tr (Ol)-U. Tr (Rha) U. Tr (u. Car-l. Nor) U. Tr (l. Car) L. J (Hett?-Plie) U. Tr (Nor) M. Tr (Ani) U. Tr (Car–Rha) U. Tr (Car-Rha) U. Tr (Car?-Rha?) U. Tr (Car-Rha)

Tethys Tethys Austral W Tethys E Tethys Austral Cosmopolitan Cosmopolitan Tethys and Circumpacific E Tethys, Austral and Boreal Circumpacific, Austral and Boreal Tethys, Circumpacific and Boreal Cosmopolitan Circumpacific and Austral Tethys Cosmopolitan Tethys and Circumpacific E Tethys W Tethys Circumpacific, Austral and Boreal Circumpacific Tethys, Circumpacific and Austral Boreal Boreal Circumpacific and Boreal Circumpacific Circumpacific Tethys and Circumpacific E Tethys E Tethys Cosmopolitan Tethys Tethys and Boreal Circumpacific Tethys and Circumpacific Circumpacific Tethys Circumpacific, Austral and Boreal S Tethys Circumpacific and Austral Austral Tethys Tethys and Austral Tethys, Circumpacific and Boreal Tethys W Tethys and Boreal Tethys, Circumpacific and Austral Austral Tethys Tethys and Circumpacific Circumpacific Circumpacific Circumpacific E Tethys Circumpacific and Austral Austral Circumpacific y Austral Tethys, Circumpacific and Austral Tethys Cosmopolitan Austral Tethys and Circumpacific Tethys and Circumpacific Austral W Tethys Tethys and Circumpacific Circumpacific E Tethys Circumpacific, Austral and Boreal Circumpacific Circumpacific Circumpacific and Austral

Paleoautoecology	Mineralogy
B, E, S, Epi, Sed; By	Bimineralic
B, E, S, Epi, Sed; By	Bimineralic
B, E, S, Epi, Sed; By	Bimineralic
B, E, S, Epi, Sed; By	Bimineralic
B, E, S, Epi, Sed; By	Bimineralic
B, E, S, Epi, Sed; By	Unknown Binnin anglia
B/Ps, E, S/Ch, Epi/Un, Sed/FaM; By/R B/Ps, E, S/Ch, Epi/Un, Sed/FaM; By/R	Bimineralic Bimineralic
B/Ps, E, S/Ch, Epi/Un, Sed/FaM; By/R	Bimineralic
B/Ps, E, S/Ch, Epi/Un, Sed/FaM; By/R	Bimineralic
B, E, S, Un, FaM; R/Sw	Bimineralic
B, E, S, Un, FaM; R/Sw	Bimineralic
B, E, S, Epi/Un, FaM; By/R/Sw	Bimineralic
B, Se, S, Un, Sed; R	Bimineralic
B, E, S, Epi/Un, FaM; By/R/Sw	Aragonitic
B, E, S, Epi, FaM; By/Sw	Bimineralic
B, E, S, Epi, Sed; By	Bimineralic
B, E, S, Epi, Sed; By B, E, S, Un, FaM; R/Sw	Bimineralic Bimineralic
B, E, S, Epi, Sed; By	Bimineralic
B, E, S, Epi, Se; By	Bimineralic
B, E, S, Epi/C, Sed; By/C	Bimineralic
B, E, S, Epi, FaM; By/Sw?	Bimineralic
B, E, S, Epi, Sed; By	Bimineralic
B, E, S, Epi, Sed; By	Bimineralic
B, E, S, Epi, Sed; By	Bimineralic
B, E, S, Epi, Sed; By	Bimineralic
B, E, S, Epi, Sed; By B, E, S, Epi, Sed; By	Bimineralic Bimineralic
B, E, S, Epi, Sed; By B, E, S, Epi, Sed; By	Bimineralic
B, E, S, Un, FaM/Sed; R/Sw	Bimineralic
B, E, S, Epi, Sed; By	Bimineralic
B, E, S, Epi, Sed; By	Bimineralic
B, E, S, Un, Sed; R	Bimineralic
B, E, S, Epi, FaM; By/Sw	Bimineralic
B, Is, S, Endo/Un, Sed/SM; By/Sb B, Is/Se, S, Endo/Un, Sed/SM; By/Sb	Aragonitic Aragonitic
B, Is/Se, S, Elido/Oli, Scarsivi, By/55 B, Is/Se, S, SM; Sb	Aragonitic
B, Se, S, Endo/Un, Sed/SM; Sb	Aragonitic
B, Se/Is, S, SM; Sb	Aragonitic
B, Is, S, Endo/Un, Sed/SM; By/Sb	Aragonitic
B, Is, S, Endo/Un, Sed/SM; By/Sb	Aragonitic
B, Se, S, Endo, Sed; By B, Se, S, Endo, Sed; By	Aragonitic Aragonitic
B, Se, S, Endo, Sed; By B, Se, S, Endo, Sed; By	Aragonitic
B, Se, S, Endo, Sed; By	Aragonitic
B, Is, S, FM; Sb	Aragonitic
B, Is, S, FM; Sb	Aragonitic
B, Is, S, FM; Sb	Aragonitic
B, Is, S, FM; Sb	Aragonitic
B, Is, S, FM; Sb B, Is, S, FM; Sb	Aragonitic Aragonitic
B, Is, S, FM; Sb	Aragonitic
B, Is, S, FM; Sb	Aragonitic
B, Is, S, FM; Sb	Aragonitic
B, Is, S, FM; Sb	Aragonitic
B, Is, S, FM; Sb	Aragonitic
B, Is, S, SM; Sb	Aragonitic
B, Is, S, FM; Sb	Aragonitic
B, Is, S, FM; Sb B, Is, S, FM; Sb	Aragonitic Aragonitic
B, Is, S, FM; Sb	Aragonitic
B, Is, S, FM; Sb	Aragonitic
B, Is, S, FM; Sb	Aragonitic
B, Is, S, FM; Sb	Aragonitic
B, Is, S, FM; Sb	Aragonitic
B, Is, S, FM; Sb B, Is, S, FM; Sb	Aragonitic
B, Is, S, FM; Sb B, Is, S, FM; Sb	Aragonitic Aragonitic
B, Is, S, FM; Sb	Aragonitic
B, Is, S, FM; Sb	Aragonitic
B, Is, S, FM; Sb	Aragonitic

Genus	Stratigraphic range	Paleogeographic distribution	Paleoautoecology	Mineralog
Groeberella	L. J (Sin)–M. J (Baj)	Circumpacific and Austral	B, Is, S, FM; Sb	Aragonitic
Antiquicorbula	U. Tr (Nor–Rha)	Tethys	B, Is, S, SM; Sb	Aragonitic
Pseudosaxicava	L. J (Hett)–U. J (Tit)	E Tethys	B, E, S, Epi, Sed; By/N	Aragonitic
Sonchodon	U. Tr (u. Nor? - Rha)	W Tethys	B, E/Se, S, Un, Sed; R	Aragonitic
		,		
Pachyrisma	L. J (Sin)–U. J (Tit)	Tethys	B, E/Se, S, Un, Sed; R	Aragonitic
Paramegalodus	U. Tr (Nor)	Tethys and Circumpacific	B, E/Se, S, Un, Sed; R	Aragonitic
Pomarangina	U. Tr (Car)	E Tethys	B, E/Se, S, Un, Sed; R	Aragonitic
Ampezzania	U. Tr (Nor)	W Tethys	B, E/Se, S, Un, Sed; R	Aragonitic
Friadomegalodon	U. Tr (Car–Rha)	Tethys, Circumpacific?	B, Se, S/Ph, Un, Sed; R	Aragonitic
Quemocuomegalodon	U. Tr (Nor)	E Tethys	B, E/Se, S, Un, Sed; R	Aragonitic
Neomegalodon	U. Tr (Car - Rha)	Tethys and Boreal	B, E/Se, S, Un, Sed; R	Aragonitic
Rhaetomegalodon	U. Tr (Nor–Rha)	W Tethys	B, E/Se, S, Un, Sed; R	Aragonitic
Dicerocardium		Tethys and Circumpacific	B, E/Se, S/Ph, Un, Sed; R	
	U. Tr (u. Car–Rha)	1		Aragonitic
Cornucardia	U. Tr (Car)	Tethys	B, E/Se, S, Un, Sed; R	Aragonitic
Physocardia	M. Tr (Ani)–U. Tr (Car)	W Tethys	B, E/Se S, Un, Sed; R	Aragonitic
Carinocardia	U. Tr (l. Car)	W Tethys	B, E/Se, S, Un, Sed; R	Aragonitic
aubeia	U. Tr (Car)	W Tethys	B, E/Se, S, Un, Sed; R	Aragonitic
Vallowaconcha	U. Tr (Nor)	Tethys and Circumpacific	B, E/Se, S/Ch/Ph, Un, Sed; R	Aragonitic
inbadiella	L. Tr (Ol)	Circumpacific	B, Is, S/Ch?, SM; Sb	Aragonitic
uciniola	U. Tr (Rha) $-L$. J (Plie)	Tethys and Circumpacific	B, Is, S/Ch?, SM; Sb	Aragonitic
		W Tethys		
Storthodon	M. Tr (Lad)		Unknown	Unknown
Schafhaeutlia	M. Tr (Ani)–U. Tr (Rha)	Tethys, Circumpacific and Boreal	B, Is, S, SM; Sb	Aragonitic
phaeriola	L. J (Sin)–M. J (Call)	Tethys, Circumpacific and Austral	B, Is, S/Ch, SM; Sb	Aragonitic
Futcheria	U. Tr (Car)–M. J (Aal)	Tethys, Circumpacific and Austral	B, Is, S, SM; Sb	Aragonitic
Palaeocardita	U. Tr (Car–Rha)	Tethys, Circumpacific and Austral	B, Is, S, Endo/Un, Sed/SM; By/Sb	Aragonitic
eptocardia	U. Tr (Car–Rha)	Tethys, Circumpacific and Austral	B, Is, S, Endo/Un, Sed/SM; By/Sb	Aragonitic
Schizocardita	U. Tr (Nor)	Circumpacific	B, Is/Se, S, SM; Sb	Aragonitic
Coelopis	M. Tr (Lad)–L. Cret (?)	Tethys	B, E, S, Epi, Sed; By	Aragonitic
lstarte	M. Tr (Ani)–Recent	Tethys, Circumpacific and Austral?	B, Is, S, SM; Sb	Aragonitic
Coelastarte	L. J (Hett)–U. Cret (Tur)	Circumpacific	B, Is, S, SM; Sb	Aragonitic
lstartopsis	L. J (Sin), U. J (Oxf)	E Tethys	B, Is, S, SM; Sb	Aragonitic
Vicaniella	L. J (Hett)–Pa	E Tethys	B, Is, S, SM; Sb	Aragonitic
Radiastarte	U. Tr (Car)	E Tethys	B, Is, S, SM; Sb	Aragonitic
Veocrassina	L. J (Hett)-L. Cret (Berr)	Circumpacific	B, Is, S, SM; Sb	Aragonitic
Praeconia	U. Tr (Nor)–U. J (Tit)	Tethys and Circumpacific	B, Is, S, SM; Sb	Aragonitic
	· · · · · · · · · · · · · · · · · · ·			
Opis .	U. Tr (Nor)–U. Cret (Maa)	E Tethys	B, E, S, Un, Sed; R	Aragonitic
Cardinia	U. Tr (Car)–L. J (Toa)	Cosmopolitan	B, Is, S, SM; Sb	Aragonitic
Balantioselena	M. Tr (Lad)	Austral	B, Is, S, SM; Sb	Aragonitic
sopristes	U. Tr (Nor)	Circumpacific	B, Is, S, SM; Sb	Aragonitic
Minepharus	U. Tr (Car)	Circumpacific	B, Se, S, Endo, Sed; By	Aragonitic
Forastarte	U. Tr (Rha)–L. J (Toa)	Tethys and Austral	B, Is, S, MS; Sb	Aragonitic
Pseudastarte	L. J (Hett–Sin)	Tethys	B, Is, S, SM; Sb	Aragonitic
Myophoricardium	U. Tr (Car–Rha)	Tethys	B, Is, S, FM; Sb	Aragonitic
Myophoriopis	M. Tr (Ani)–U. Tr (Rha)	Tethys and Circumpacific	B, Is, S, FM; Sb	Aragonitic
Pseudocorbula	M. Tr (Ani)–U. Tr (Rha)	Tethys	B, Is, S, FM; Sb	Aragonitic
Protocardia	U. Tr (Nor)–U. Cret (Maa)	Tethys and Circumpacific	B, Is, S, SM; Sb	Aragonitic
Fulongocardium	U. Tr (Nor–Rha)	Tethys	B, Is, S, SM; Sb	Aragonitic
ntegricardium	L. J (Hett)-P (Dan)	Circumpacific	B, Is, S, MS; Sb	Aragonitic
Rhaetidia	M. Tr (Lad)–U. Tr (Nor)	W Tethys	B, Is, S?, SM; Sb	Aragonitic
Fancredia	L. J (Sin) –P (Dan)	Tethys	B, Id, S, FM; Db	Aragonitic
akawanella	U. Tr (Car)	Circumpacific	B, Id, S, FM; Db	Aragonitic
socyprina	U. Tr (Rha)–L. Cret (Apt)	Tethys	B, Is, S, FM; Sb	Aragonitic
Pseudotrapezium	L. J (Hett)–U. J (Oxf)	Tethys	B, Is, S, FM; Sb	Aragonitic
angvopĥorus	U. Tr (Nor–Rha)	E Tethys	Unknown	Aragonitic
Pholadomya	U. Tr (Rha)–Recent	Tethys, Circumpacific and Austral	B, Id, S, SM; Db	Aragonitic
Inomalopleuroides	U. Tr (Car)	W Tethys	B, Id?, S, SM; Db?	Aragonitic
Cortinia	U. Tr (Car)	W Tethys	B, Is?, S, SM; Sb?	Aragonitic
	L. J (Hett)–U. Cret (Maa)	Circumpacific		Aragonitic
Goniomya			B, Id, S, SM; Db	
Нототуа	M. Tr (Ani)–L. Cret (Val)	Tethys and Circumpacific	B, Id, S, SM; Db	Aragonitic
Pachymya	M. Tr (Ani)–U. Cret (Tur)	Tethys and Circumpacific	B, Id, S, SM; Db	Aragonitic
Dianomya	M. Tr (Ani)	E Tethys	B, Id, S, SM; Db	Aragonitic
Dsteomya	L. J (Sin)-M. J (Call)	Circumpacific	B, Id, S, SM; Db	Aragonitic
Pleuromya	L. Tr (Ol)–L. Cret (Val)	Tethys, Circumpacific and Austral	B, IP, S, SM; Db	Aragonitic
Burmesia		E Tethys and Circumpacific	B, Id, S, SM; Db	Aragonitic
	U. Tr (Car)–L. J (Hett)			
Prolaria	U. Tr (Nor–Rha)	E Tethys	B, Id, S, SM; Db	Aragonitic
Pseudoburmesia	U. Tr (Car)	Tethys	B, Id, S, SM; Db	Aragonitic
Pteromya	U. Tr (Rha)–L. J (Hett)	W Tethys	B, Is, S, SM; Sb	Aragonitic
Dchotomya	U. Tr (Nor - Rha)	Austral and Boreal	B, Is, S, SM; Sb	Aragonitic
Gresslya	L. J (Hett) - U. J (Tit)	E Tethys and Circumpacific	B, Id, S, SM; Db	Aragonitic
Thracia		Tethys and Circumpacific		
	U. Tr (Nor) - Recent		B, Id, S, SM; Db	Aragonitic
Cercomya	U. Tr (Car)–U. Cret (Maa)	Tethys	B, Id, S, SM; Db	Aragonitic
Fulongella	M. Tr (Ani) - U. Tr (Rha)	E Tethys	B, Id, S, SM; Db	Aragonitic
wonge w				

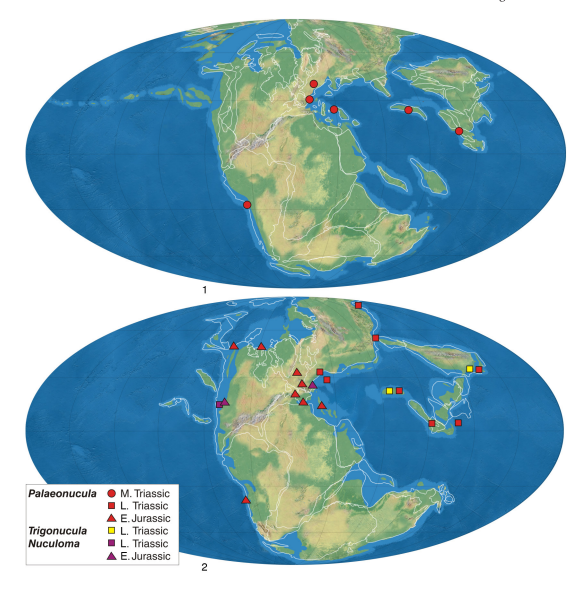


Figure 1. Paleogeographical distribution of Nuculidae (Palaeonucula, Trigonucula, Nuculoma). 1, Middle Triassic; 2, Late Triassic-Early Jurassic.

Superfamily NUCULOIDEA Gray, 1824 Family NUCULIDAE Gray, 1824 Genus PALAEONUCULA Quenstedt, 1930, p. 110

Type species.—Nucula hammeri Defrance, 1825b, p. 217.

Remarks.—Palaeonucula was regarded as subgenus of *Nuculoma*, *Nucula*, and *Nuculopsis* (see Hodges, 2000, p. 13) and even as a synonym of the last (Nakazawa & Newell, 1968); it is here regarded as a valid genus, following Carter (1990a) and Hodges (2000).

Stratigraphic range.—Middle Triassic (lower Anisian)–Lower Cretaceous (Aptian) (Komatsu, Chen, & others, 2004; Gang, 2001). The stratigraphic range was here extended, both with respect to the Triassic–Jurassic range in Cox and others (1969), and also to Sepkoski (2002), who assigned a Triassic (Ladinian)–Jurassic (Tithonian) age. The oldest records we accept are Anisian (Tamura & others, 1975; Wen & others, 1976; Komatsu, Chen, & others, 2004). Although Bailey (1978) mentioned the species *Palaeonucula strigilata* from the Mississippian of Arkansas, this will not be taken into account because the source is an abstract in a conference proceedings volume, and we did not find any reference where the author figured or described the species. The genus was widely mentioned throughout the interval Middle to Late Jurassic (Pugaczewska, 1986; Sha & Fürsich, 1994; Holzapfel, 1998; Harries & Little, 1999; Gahr, 2002; Delvene, 2003; J. Yin & Grant-Mackie, 2005).

Paleogeographic distribution.—Tethys, Circumpacific, and Boreal (Fig. 1).

Tethys domain: Middle Triassic: northern Vietnam (Komatsu, Huyen, & Huu, 2010); Anisian of China (Wen & others, 1976; Komatsu, Chen, & others, 2004), Spain (Márquez-Aliaga, 1985), Malaysia (Tamura & others, 1975); Ladinian of Germany (Ürlichs, 1992), Spain (Márquez-Aliaga, 1985; Niemeyer, 2002), Malaysia (Tamura & others, 1975); Late Triassic: of China (Gou, 1993); Carnian of China (Wen & others, 1976), Italy (Fürsich & Wendt, 1977), Malaysia (Tamura & others, 1975); Norian of China (Lu, 1981), Iran (Repin, 2001; Hautmann, 2001b); Rhaetian of Iran (Hautmann, 2001b), Hungary (Vörös, 1981); Early Jurassic: Hettangian–Sinemurian of England (Liu, 1995; Hodges, 2000); Sinemurian of Turkey (M. A. Conti & Monari, 1991); Sinemurian of southwestern France, Spain, & Portugal (Liu, 1995).

Circumpacific domain: Middle Triassic: Anisian of Chile (Barthel, 1958); Late Triassic: Carnian of Japan (Hayami, 1975); Early Jurassic: Hettangian–Sinemurian of Chile (Aberhan, 1994a; Damborenea, 1996a); Sinemurian of Canada (Aberhan, 1998a).

Boreal domain: Late Triassic: northeastern Siberia (Yakutia Region) (Kurushin, 1987); Triassic–Jurassic: northeastern Asia (Kurushin, 1990).

Paleoautoecology.—B, D, Is, FM; Sb. Holocene nuculids dig into the surface layers of the sediment and remain very close to its surface. They actively use the foot to dig and move around and use the palp proboscis to feed on detritus (Reid, 1998). A similar mode of life is suggested for *Palaeonucula*. Its external form would facilitate a fairly quick movement through the sediment. Pallial sinus is not observed, and thus it probably did not have siphons. Living nuculids are commonly found in shallow waters and fine-grained sandy sediments, and this is consistent with the associated lithology of fossil species (Hodges, 2000). All previous authors considered *Palaeonucula* as an infaunal, mobile, and shallow burrowing detritivorous bivalve (see e.g., Pugaczewska, 1986; Damborenea, 1987a; M. A. Conti & Monari, 1991; Holzapfel, 1998; Delvene, 2003).

Mineralogy.—Aragonitic (Carter, 1990a, p. 150). Outer shell layer: aragonite (irregular prismatic). Middle and inner shell layers: aragonite (homogeneous).

Genus TRIGONUCULA Ichikawa, 1949, p. 267

Type species.—Trigonucula sakawana Ichikawa, 1949, p. 268.

Stratigraphic range.—Upper Triassic (Carnian-upper Rhaetian) (Hayami, 1975; Hautmann, 2001b). Both Cox and others (1969) and Sepkoski (2002) assigned a Upper Triassic range to this genus. This is here maintained, despite references from the Jurassic and Cretaceous: Trigonucula yunshanensis Yu & Li (in Z. Li & Yu, 1982, p. 94, fig. 13-14) and Trigonucula? yunshanensis (Gu, Li, & Yu, 1997, p. 15, pl. 2,8-9). Nevertheless, both the hinge and shell outline of these species are totally different from those described for Trigonucula sakawana and therefore probably do not belong to the genus (Jingeng Sha, personal communication, 2008). Dickins and McTavish (1963) mention the genus (Trigonucula sp.) in the Lower Triassic (Scythian), but this will not be taken into account for two reasons: (1) it was not included in Cox and others (1969); and (2) the figures in Dickins and McTavish (1963) are impossible to compare due to their poor quality; also the authors pointed out in their discussion (p. 129): "In shape Trigonucula sp. is not unlike Nucula sp. juv. ind. Spath (1930, p. 53, pl. 12,12) from the Lower Triassic (Otoceratan) of Greenland."

Paleogeographic distribution.—Tethys and Circumpacific (Fig. 1). Tethys domain: Late Triassic: Norian of Iran (Hautmann, Aghababalou, & Krystyn, 2011); Rhaetian of Iran (Hautmann, 2001b).

Circumpacific domain: Late Triassic: Carnian of Japan (Ichikawa, 1949; Hayami, 1975; Kobayashi & Tamura, 1983b).

Paleoautecology.—B, D, Is, FM; Sb. A relatively quick shallow burrower and detritivorous mode of life is attributed to this genus, similar to the living nuculids (Reid, 1998). Hautmann (2001b, p. 26) described divaricate ornamentation of the shell in his emended diagnosis of the genus. This type of ornamentation has been studied from a functional point of view by several authors (e.g., S. M. Stanley, 1969; Seilacher, 1972), and it is now known that it facilitates excavation (Checa & Jiménez-Jiménez, 2003b).

Mineralogy.—Aragonitic. No specific data about the *Trigonucula* mineralogy and shell microstructure is known, but a totally aragonitic mineralogy is assumed, according to the diagnosis of subclass Paleaeotaxodonta given by Allen and Hannah (1986).

Genus NUCULOMA Cossmann in Cossmann & Thièry, 1907, p. 124

Type species.—Nucula castor d'Orbigny, 1850, p. 339.

Stratigraphic range.—Upper Triassic (Rhaetian)–Lower Cretaceous (Valanginian) (Laws, 1982; Kaim, 2001). Although *Nuculoma* was regarded as a typical Jurassic genus (Cox & others, 1969; Sepkoski, 2002), it was found in the Rhaetian of New York Canyon (Laws, 1982; Guex & others, 2003, 2004; Lucas & Tanner, 2004; Lucas & others, 2007) and in the Lower Cretaceous of several localities (Kaim, 2001; Marinov & others, 2006; X. Li, 1990). Accordingly, its stratigraphic range is here extended.

Although originally used for Jurassic forms, several Recent species were also included in *Nuculoma*. There is no revision of these living species to test which ones are really consistent with the diagnosis, but they will not be taken into account here, because, according to Hansson (1998, p. 93), living species should be included in *Ennucula* Iredale, 1931 ["*Ennucula* Iredale, 1931 = *Nuculoma*: auct., *non* Cossmann in Cossman & Thièry, 1907 (*Nucula castor* d'Orbigny, 1850 - Jurassic fossil)"].

Paleogeographic distribution.—Tethys and Circumpacific (Fig. 1). Available information suggests that the genus appeared in the Circumpacific domain, specifically in the Rhaetian of Nevada (Guex & others, 2004), and, during the Early Jurassic, it spread to the Tethys (Hallam, 1972, 1977) and Boreal domains (Zakharov & others, 2006). Later, during the Jurassic, its distribution expanded not only to the European Tethys (X. Li & Grant-Mackie, 1994; Holzapfel, 1998), but also to the Proto-Atlantic (Liu, 1995).

Circumpacific domain: Late Triassic: Rhaetian of Nevada (Hallam & Wignall, 2000); Early Jurassic: Hettangian of Nevada (Hallam & Wignall, 2000).

Tethys domain: Early Jurassic: Sinemurian of Europe (Hallam, 1977).

Paleoautoecology.—B, D, Is, FM; Sb. Fürsich (1982) compared *Nuculoma* with Recent *Nucula*, and he assigned it a similar mode of life, moving just below the sediment surface, feeding on the detritus taken with palp proboscis, like other nuculids.

Mineralogy.—Aragonitic (Carter, 1990b, p. 307). Outer shell layer: aragonite (prismatic). Middle and inner shell layers: aragonite (nacreous).

Superfamily NUCULANOIDEA Adams & Adams, 1858 in 1854–1858 Family NUCULANIDAE Adams & Adams, 1858 in 1854–1858 Genus NUCULANA Link, 1807, p. 155

Type species.—Arca rostrata Chemnitz, 1784, pl. 55, fig. 550–551. *Remarks.—Nuculana* is a genus especially difficult to identify since the external characters alone, in absence of inner views, are

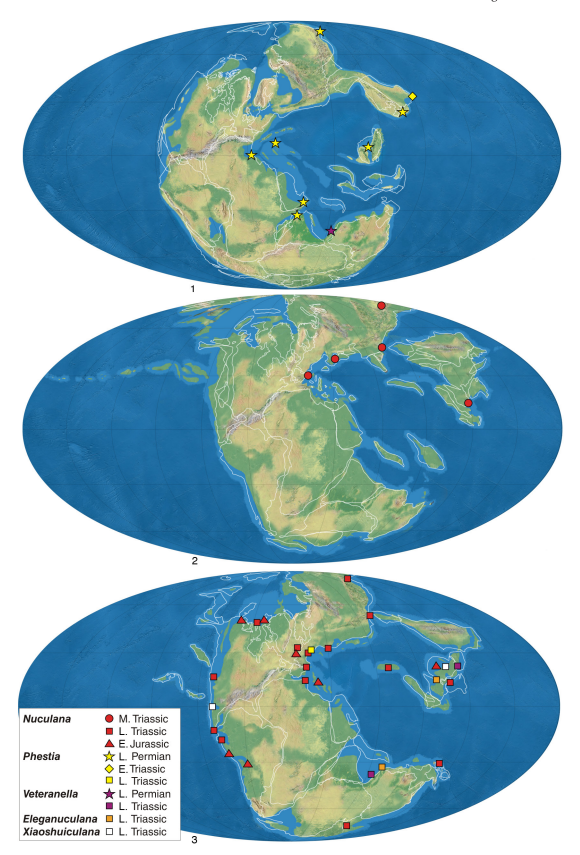


Figure 2. Paleogeographical distribution of Nuculanidae (Nuculana, Phestia, Veteranella, Eleganuculana, Xiaoschuiculana). 1, Early Triassic; 2, Middle Triassic; 3, Late Triassic-Early Jurassic.

undistinguishable from those of other genera, such as *Phaenodesmia* Bittner, 1894, *Phestia* Chernyshev, 1951, *Veteranella (Veteranella)* Patte, 1926, or *V. (Glyptoleda)* Fletcher, 1945; hence many Paleozoic and Mesozoic specimens attributed to *Nuculana* probably belong to other genera (Boyd & Newell, 1979; Damborenea, 1987a; Carter, 1990a; Z. Fang & Cope, 2004).

Leda Schumacher, 1817, is a junior objective synonym of *Nuculana* (both genera have the same type species). Although this was pointed out by Cox and others (1969), some authors still use the name *Leda* (e.g., Fürsich & Wendt, 1977; Ruban, 2006a).

Stratigraphic range.---Middle Triassic (Anisian)-Holocene (Tamura & others, 1975). Although there are references for this genus from the Paleozoic, we follow McAlester (in Cox & others, 1969), who considered its range to be from the Triassic to the present (see Nakazawa & Newell, 1968, p. 37-38 for discussion of this genus). Sepkoski (2002) extended it to the lower Induan, but the only mention from the Lower Triassic is Nuculana (Dacryomya) sp. from Japan in Nakazawa (1961); however, this was referred to as *Phestia* sp. by Nakazawa and Newell (1968). Therefore, the first appearance is regarded as Anisian (Tamura & others, 1975). Furthermore, the record is fairly continuous throughout the entire study interval (see Paleogeographic Distribution, below). However, not all authors agree about the presence of *Nuculana* in this time interval. Carter (1990a, p. 151) stated that Nuculana did not appear until the Cretaceous. In our opinion, a thorough review of this genus is needed, but it is beyond the scope of this paper, and we will tentatively use the proposed range.

Paleogeographic distribution.—Cosmopolitan (Fig. 2). Although the genus is also present in the Early Jurassic from both the Circumpacific and the Austral domains, these records are Pliensbachian in age (Damborenea, 1987a; Aberhan, 1994a, 1998a).

Tethys domain: Middle Triassic: Anisian of Malaysia (Tamura & others, 1975), northern Vietnam (Komatsu, Huyen, & Huu, 2010); Ladinian of Afghanistan (Farsan, 1975), Spain (Niemeyer, 2002), western Caucasus (Ruban, 2006a); Late Triassic: China (Cowper-Reed, 1927); Carnian of Italy (S. Conti, 1954), Slovenia (Jurkovsek, 1978), China (Wen & others, 1976; Lu, 1981; Gou, 1993; J. Yin, Enay, & Wan, 1999); Norian of Iran (Hautmann, 2001b; Repin, 2001); Norian–Rhaetian of England (Hallam & El Shaarawy, 1982; Hallam, 2002); Rhaetian of Italy (Sirna, 1968), Iran (Hautmann, 2001b; Repin, 2001), Spain (Márquez-Aliaga, Plasencia, & Ros, 2005), China (J. Yin, Enay, & Wan, 1999), Austria (Tomašových, 2006a, 2006b); Early Jurassic: Hettangian of China (J. Yin, Enay, & Wan, 1999); Sinemurian of Turkey (M. A. Conti & Monari, 1991).

Circumpacific domain: Late Triassic: Carnian of Mexico (Alencaster de Cserna, 1961), Peru (Jaworski, 1922; Cox, 1949); Norian of Nevada (Laws, 1982).

Boreal domain: Middle Triassic: northern Siberia (Dagys & Kurushin, 1985), Primorie (Kiparisova, 1972); Late Triassic: Carnian of Primorie (Kiparisova, 1972); Triassic–Jurassic: northeastern Asia (Kurushin, 1990). Holocene species have a wide distribution in the boreal domain and in cold-temperate regions.

Austral domain: Late Triassic: Carnian of New Zealand (Marwick, 1953), Carnian–Norian of New Guinea (Skwarko, 1967); Rhaetian of New Zealand (Grant-Mackie, 1960).

Paleoautoecology.—B, D, Is, FM; Sb. Holocene species of this genus are very fast burrowers (Gordillo & Aitken, 2000), moving in the surface of the sediment, with a detritivorous trophic regime (Damborenea, 1987a; M. A. Conti & Monari, 1991; Holzapfel, 1998; Hautmann, 2001b).

Mineralogy.—Aragonitic (Carter, 1990b, p. 311–312, for Recent species). All shell layers: aragonite (homogeneous).

Genus PHESTIA Chernyshev, 1951, p. 9

Type species.—Leda inflatiformis Chernyshev, 1939, p. 116. Remarks.—We regard Polidevcia Chernyshev, 1951, p. 25, as a subgenus of *Phestia* (see discusion for *Polidevcia* in Genera not Included, p. 168).

The generic name was first proposed in Chernyshev, 1943, p. 35, but no type species was designated, and it remained a nomen nudem until the type was designated by this author in 1951.

Stratigraphic range.—Middle Ordovician (lower Llanvirn)–Upper Triassic (Carnian) (Carter, 1990a; Z. Fang & Cope, 2004). Cox and others (1969) assigned this genus a Devonian–Lower Triassic range, and Sepkoski (2002), considered it was present from the Devonian (Givetian) to Lower Triassic (data taken from Skelton & Benton, 1993), but the range is extended in this paper. The first occurrence of the genus was Middle Ordovician (Llanvirnian [=overlaps with Darriwilian stage]), according to Z. Fang and Cope (2004), although doubtfully because the figured specimen is an external mold, and the internal features are unknown. Thus, although reference to Phestia seems justified, the same authors note that it externally resembles Glyptoleda (regarded in this paper as a subgenus of Veteranella). Carnian is used here as the upper limit, based on data provided by Carter (1990a, p. 153). This author considered that "Nuculana" sulcellata, from the Italian Carnian, would be better located within Phestia, since its form, ligament structure, and nacreous interior are typical of this genus. Nakazawa and Newell (1968) included Nuculana (Dacryomya) sp., figured by Nakazawa (1961, p. 270, pl. 14,5-7), in Phestia, and thus extended the range of this genus to the Triassic of southwestern Japan. Hautmann and others (2005) mentioned Phestia? cf. perlonga (Mansuy) from the upper Rhaetian of southern Tibet, but this is the type species of *Mesoneilo* Vu Khuc, 1977a, p. 676, to which the authors did not refer in their paper. This very doubtful record will not be taken into account here.

Paleogeographic distribution.—Tethys, Circumpacific, and Boreal (Fig. 2). Cox and others (1969) regarded *Phestia* as a cosmopolitan genus, but, although it was very widespread during the Paleozoic, during the studied time interval, it was only present in the Tethys, the Boreal, and the Circumpacific domains.

Tethys domain: late Permian: Iran (Teichert, Kummel, & Sweet, 1973), Tunisia and India (Boyd & Newell, 1979), southern China (L. Li, 1995; Clapham & Bottjer, 2007), Oman (Dickins, 1999); Late Triassic: Carnian of Italy (Carter, 1990a).

Circumpacific domain: late Permian: Japan (Nakazawa & Newell, 1968; Murata & Bando, 1975; Hayami & Kase, 1977); Early Triassic: Japan (Nakazawa & Newell, 1968).

Boreal domain: late Permian: northeastern Russia (Biakov, 1998, 2002, 2006, 2007; Klets & others, 2006).

Paleoautoecology.—B, D, Is, FM; Sb. Phestia most probably had a mode of life similar to the living Nuculana, but it lacked a pallial sinus, so it possibly did not have true siphons (as *Nuculana* has). Instead, it may have had pseudosiphons created by ciliary connections between the undulations of the mantle (see Bradshaw, 1999, p. 75–76). The genus was regarded as a superficial burrowing detritivore that used the palp proboscis to collect food particles (Hoare, Heaney, & Mapes, 1989; Bradshaw, 1999). R. Zhang and Yan (1993) agreed with this and provided a reconstruction of its mode of life, showing the similarity to *Palaeoneilo*. The elongated anterior part and the anterior pedal muscle scars, which can be observed in some specimens, suggest it had a large foot that would allow it to burrow effectively (Mángano & others, 1998). These authors associated the ichnofossil *Lockeia ornata* with *Phestia*, from which they concluded that *Phestia* was a vagrant detritivorous capable of moving subhorizontally in the sediment.

Mineralogy.—Aragonitic (Carter, 1990a, p. 154–155). Outer shell layer: aragonite (prismatic). Middle and inner shell layers: aragonite (nacreous).

Genus VETERANELLA Patte, 1926, p. 158

Type species.—Nuculana (Veteranella) strenua Patte, 1926, p. 158. Stratigraphic range.—lower Permian (Artinskian)–Upper Triassic (Norian) (Kutassy, 1931; Waterhouse, 1964). The stratigraphic range is here extended with respect to Sepkoski (2002), because we include *Glyptoleda* and *Nucundata* as subgenera of *Veteranella*, according to Cox and others (1969) (see discussion for *Glyptoleda* and *Nucundata* in Genera not Included, p. 161, 166).

Cox and others (1969) assigned a Permian–Triassic range to *Veteranella*; Sepkoski (2002) considered *Nucundata* to be present in the Permian following Cox and others (1969), *Glyptoleda* in the late Guadalupian following Waterhouse (1987), and a Norian range for *Veteranella* following Hallam (1981). J. Chen, Liu, and Lan (1983) mentioned *Glyptoleda* and other genera attributed to their new subfamily Veteranellinae from Devonian to Permian, but none of these genera was listed as being present before the Carboniferous (Z. Fang & Cope, 2004).

The oldest record of the genus (referred to *Nucundata* and *Glyp-toleda*) is lower Permian (Artinskian–Kunguarian) of New Zealand (Waterhouse, 1964). The genus was also mentioned from Lower Jurassic age; e.g., Kurushin (1990) quoted *Veteranella* from the Triassic–Jurassic boundary, confirming its presence in lower Het-tangian beds, and Zhakarov and others (2006) mentioned *Glyptoleda* from the Pliensbachian, but none of them justified the presence of this genus in the Lower Jurassic, because they neither figured the specimens nor included the original source of their data. The youngest record is Upper Triassic: *Veteranella* (*Ledoides*) Chen, Wen, & Lan in Wen & others, 1976, from Carnian–Norian of Tibet (Kobayashi & Tamura, 1983a), from Carnian of China (Wen & others, 1976), and from Norian of eastern Tethys (Hallam, 1981).

Paleogeographic distribution.—Tethys (Fig. 2). *Veteranella* had a wide distribution in Boreal and Austral domains during the early and middle Permian (Waterhouse, 1964, 1983; Biakov, 1998, 2006), but it was not found there during the late Permian.

Tethys domain: late Permian: Changhsingian of Nepal (Waterhouse & Chen, 2006); Late Triassic: China (Kutassy, 1931); Carnian of China (Wen & others, 1976); Carnian and Norian of southern Tibet (Kobayashi & Tamura, 1983a); Norian of Xizang (Tibet) (Z. Fang & others 2009).

Paleontological Contributions, number 8

Paleoautoecology.— B, D, Is, FM; Sb. *Veteranella reidi* (Fletcher, 1945) (Permian) is the oldest species with oblique chevron-type ornamentation, which became common among bivalves during the Cenozoic, and is interpreted as an adaptation to rapid escape from potential predators and for minimizing shell damage during burrowing (Checa & Jiménez-Jiménez, 2003a). Numerous studies demonstrated that this type of ornamentation facilitates excavation (S. M. Stanley, 1969, 1970; Seilacher, 1972), so this genus is regarded as a fast burrower.

Mineralogy.—Aragonitic (Carter, Barrera, & Tevesz, 1998). Data provided for subclass Protobranchia. All shell layers: aragonite. Inner shell layer: usually nacreous.

Genus ELEGANUCULANA J. Chen & Yang, 1983, p. 355 [358]

Type species.—Eleganuculana nyeruensis J. Chen & Yang, 1983, p. 356.

Stratigraphic range.—Upper Triassic (Norian) (J. Chen & Yang, 1983). J. Chen and Yang (1983) described *Eleganuculana* including only the type species from Norian of Knagmar region in Xizang province (southern China). J. Chen, Liu, and Lan (1983) mentioned the same species from the Norian of Tibet.

Paleogeographic distribution.—Eastern Tethys (Fig. 2).

Tethys domain: Late Triassic: Norian of southern China (J. Chen & Yang, 1983), Tibet (J. Chen, Liu, & Lan, 1983).

Paleoautoecology.-B, D, Is, FM; Sb. Similar to Nuculana.

Mineralogy.—Aragonitic (Carter, Barrera, & Tevesz, 1998). No data are available for *Eleganuculana*. Protobranchia shell mineralogy is fully aragonitic (Carter, Barrera, & Tevesz, 1998).

Genus XIAOSHUICULANA J. Chen in J. Chen, Liu, & Lan, 1983, p. 622, 626

Type species.—Reticulana elegansa Li & Li in R. Zhang, Wang, & Zhou, 1977, p. 9.

Stratigraphic range.—Upper Triassic. Xiaoshuiculana was described by J. Chen (in J. Chen, Liu, & Lan, 1983) from Upper Triassic of China (Guangdong province), including only the type species. McRoberts (1997a) described a new species: X. tozeri McRoberts, 1997a, from the lower Rhaetian Antimonio Formation of Sonora (Mexico).

Paleogeographic distribution.—Tethys and Circumpacific (Fig. 2).

Tethys domain: Late Triassic: China (J. Chen, Liu, & Lan, 1983).

Circumpacific domain: Late Triassic: Rhaetian of Mexico (McRoberts, 1997a).

Paleoautoecology.—B, D, Is, FM; Sb. *Xiaoshuiculana* is externally similar to *Nuculana*, but its rostrum is more elongated, and the shell bears oblique ribs (McRoberts, 1997a). These ribs would primarily strengthen the shell and probably also favored efficient excavation (Checa & Jiménez-Jiménez, 2003a).

Mineralogy.—Aragonitic (Carter, Barrera, & Tevesz, 1998). Data provided for subclass Protobranchia.

Family MALLETIIDAE Adams & Adams, 1858 in 1854–1858 Genus PHAENODESMIA Bittner, 1894, p. 188

Type species.—Phaenodesmia klipsteiniana Bittner, 1894, p. 188.

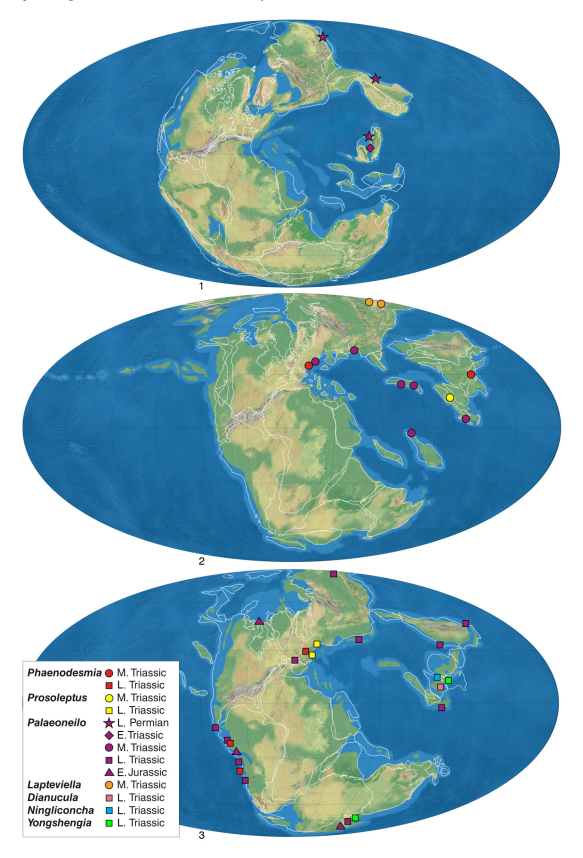


Figure 3. Paleogeographical distribution of Malletiidae (*Phaenodesmia, Prosoleptus, Palaeoneilo, Lapteviella, Dianucula, Ningliconcha, Yongshengia*). 1, Early Triassic; 2, Middle Triassic; 3, Late Triassic–Early Jurassic.

Stratigraphic range.—Middle Triassic (Anisian)–Upper Triassic (Rhaetian) (Hallam, 1981; F. Stiller, personal communication, 2008). Cox and others (1969) assigned a Triassic range in Europe to this genus. Sepkoski (2002), allegedly based on data provided by Hallam (1981), assigned it an Anisian–Norian range. However, Hallam (1981) considered that *Phaenodesmia* was present in Carnian and Norian (including Rhaetian) deposits. The oldest record is from Anisian beds of the Alps (Diener, 1923) and southwestern China (F. Stiller, personal communication, 2008).

Paleogeographic distribution.—Tethys and Circumpacific (Fig. 3).
Tethys domain: Middle Triassic: Anisian of southwestern China
(F. Stiller, personal communication, 2008), southern Alps (Diener, 1923); Late Triassic: Carnian of southern Alps (Diener, 1923).

Circumpacific domain: Late Triassic: South America (Hallam, 1981), Peru (Jaworski, 1922; Körner, 1937); Carnian of Chile (Nielsen, 2005).

Paleoautoecology.—B, D, Is, FM; Sb. We assign to this genus the same mode of life as other nuculids.

Mineralogy.—Aragonitic (Carter, Barrera, & Tevesz, 1998). Data provided for subclass Protobranchia.

Genus PROSOLEPTUS Beushausen, 1895, p. 95

Type species.—Nucula lineata Goldfuss, 1837 in 1833–1841, p. 153.

Stratigraphic range.—Middle Triassic (Anisian)–Upper Triassic (Carnian) (Fürsich & Wendt, 1977; Komatsu, Huyen, & Huu, 2010). Cox and others (1969) considered that *Prosoleptus* was present in the European Triassic. Sepkoski (2002) assigned it a Middle Triassic(?)–Carnian range following Hallam (1981), who mentioned *Prosoleptus* from Ladinian and Carnian deposits of western Tethys.

Fürsich and Wendt (1977) found *P. lineata* (Goldfuss, 1837 in 1833–1841) in the Cassian Formation of the southern Alps. This formation is regarded as upper Ladinian–Carnian in age, and probably for this reason, Hallam (1981) mentioned it from the Ladinian. Although Fürsich (in PBDB, 2005) confirmed that *P. lineata* occurs only in Carnian beds, it was recently reported from Anisian beds of northern Vietnam (Komatsu, Huyen, & Huu, 2010).

Paleogeographic distribution.—Tethys and ?Boreal (Fig. 3). *Prosoleptus* was present in the Tethys domain and probably also in northern Siberia (Carnian) (Kurushin, 1984).

Tethys domain: Middle Triassic: Anisian–Ladinian of northern Vietnam (Komatsu, Huyen, & Huu, 2010); Late Triassic: Carnian of Italy (south of the Alps) (Fürsich & Wendt, 1977), Germany (Goldfuss, 1863).

Paleoautoecology.—B, D, Is, FM; Sb. We assign the same mode of life as other nuculids.

Mineralogy.—Aragonitic (Carter, 1990a, p. 159–160; Carter, Lawrence, & Sanders, 1990, p. 315–316). All layers: aragonite (homogeneous).

Genus PALAEONEILO Hall & Whitfield, 1869, p. 6

The original spelling was *Palaeaneilo* Hall & Whitfield, 1869, p. 6, see McAlester, 1968, p. 41. For the authorship of the paper where this genus was named, see McAlester, 1968, p. 62.

Type species.—Nuculites constricta Conrad, 1842, p. 249.

Remarks.—There is some confusion in the literature among the genera *Palaeoneilo*, *Praesaccella* Cox, 1940, and *Mesosaccella* Chavan, 1947, p. 197 (see discussion in Damborenea, 1987a, p. 54). The problem stems from the fact that there are both Paleozoic and Mesozoic records of this genus and in the presence or absence of resilifer in different species referred to it. Some authors (cf. Damborenea, 1987a, p. 54; Aberhan, 1998a, p. 67) referred Paleozoic species to *Palaeoneilo* and Mesozoic species to *Mesosaccella*, but Cox (1937a) stated that there is no reason to separate the Paleozoic and Mesozoic species in different genera; this last criterion is followed here (but see Duff, 1978). Hodges (2000) regarded *Palaeoneilo* as a morphologically conservative genus that changed very little in general external shape through time.

Stratigraphic range.—Lower Ordovician (Tremadocian)–Lower Jurassic (Toarcian) (Gahr, 2002; Sánchez, 2002). Cox and others (1969) mentioned the range of this genus as Ordovician to the end of the Mesozoic, with a cosmopolitan distribution. Later, Sepkoski (2002) assigned it an Ordovician (upper Arenigian)–Jurassic (?upper Pliensbachian) range, following Pojeta (1971).

The first appearance is from the Lower Ordovician of Argentina (Sánchez, 2002). However, there are some problems with its last appearance. We accept Gahr's youngest record (2002) from the Toarcian; other younger records will not be taken into account, since almost all have some descriptive problems. For instance, Sha and Fürsich (1993) mentioned *Palaeoneilo* sp. from the Upper Jurassic and Lower Cretaceous of China, but they did not figure or systematically describe it. Later (Sha & Fürsich, 1994; Sha & others, 1998), they studied specimens from the same area and referred them to several species of *Nuculana (Praesaccella)* and *Mesosaccella*, but none to *Palaeoneilo*, although they discussed the problems of differentiating *Palaeoneilo* and *Mesosaccella*, and they even gave a series of guidelines to distinguish them. Hu, Jansa, and Wang (2008) also mentioned the genus from the Upper Jurassic–Lower Cretaceous interval, but not only did they not figure it, but they listed it as an ammonoid.

Paleogeographic distribution.—Cosmopolitan (Fig. 3). Palaeoneilo was a cosmopolitan genus during part of Paleozoic and Mesozoic, at least during the study interval considered.

Tethys domain: late Permian: southern China (L. Li, 1995; Y. Wang & others, 2006; Y. Yin & others, 2006; He, Feng, & others, 2007); Early Triassic: China (Z. Yang & Yin, 1979; L. Li, 1995; Sha & Grant-Mackie, 1996); Middle Triassic: Tethys (Hallam, 1981); Muschelkalk and Buntsandstein of Poland (Senkowiczowa, 1985); Anisian of southern China (Komatsu, Chen, & others, 2004); Anisian-Norian of Malaysia and Thailand (Tamura & others, 1975); Ladinian of Afghanistan (Farsan, 1975); Late Triassic: Tethys (Hallam, 1981); Carnian of China (Sha & Grant-Mackie, 1996), southern Alps (Diener, 1923; Kutassy, 1931; Fürsich & Wendt, 1977); Carnian-Norian of China (Wen & others, 1976; Lu & Chen, 1986; Gou, 1993); Norian of southwestern China (Lu, 1981), Singapore (Kobayashi & Tamura, 1968a); Rhaetian of Burma (Diener, 1923); Early Jurassic: Hettangian-Sinemurian of southwestern England (Hodges, 2000); Sinemurian-Pliensbachian of Europe (Hallam, 1987).

Circumpacific domain: late Permian: Japan (Nakazawa & Newell, 1968; Hayami & Kase, 1977); Late Triassic: Mexico, Chile, and Peru

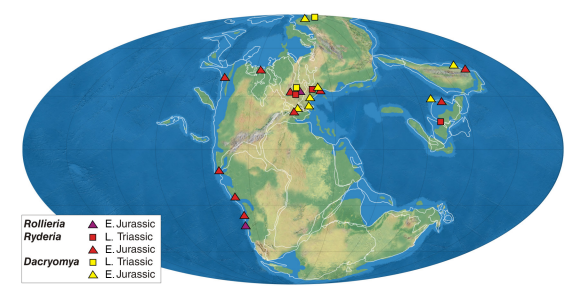


Figure 4. Paleogeographical distribution of Yoldiidae (Rollieria) and Polidevciidae (Ryderia, Dacryomya). Late Triassic-Early Jurassic.

(see references in Damborenea, 1987a); Carnian of Mexico (Diener, 1923), Japan (Hayami, 1975); Carnian–Norian of Japan (Onoue & Tanaka, 2005); Early Jurassic: Sinemurian of Chile (Covacevich, Pérez, & Escobar, 1991).

Boreal domain: late Permian: northeastern Russia (Biakov, 1998, 2007); Late Triassic: northeastern Russia (Polubotko & Repin, 1990).

Austral domain: Late Triassic: New Zealand (see references in Damborenea, 1987a), Rhaetian of New Zealand (MacFarlan, 1998) and Argentina (Damborenea & Manceñido, 2012); Early Jurassic: Pliensbachian of New Zealand (MacFarlan, 1998).

Paleoautoecology.—B, D, Is, FM; Sb. *Palaeoneilo* is regarded as a detritivorous bivalve, a superficial burrower living completely buried, very close to the sediment surface, like some living species of *Yoldia* (Damborenea, 1987a). It used a palp proboscis to feed on organic particles dispersed in the sediment (Hodges, 2000). Since in some species (e.g., *P. elliptica*) there is a shallow pallial sinus, presumably it had short siphons. Life position within the substrate most probably was with the posterior end up near the surface of the sediment, as interpreted for *Phestia* (see R. Zhang & Yan, 1993, p. 854, fig. 2). According to Hodges (2000), shell morphology indicates that *Palaeoneilo* was a quick burrower, and the foot could help to increase excavation speed.

Mineralogy.—Aragonitic (Carter & Tevesz, 1978; Carter, 1990a, p. 159–161; Carter, Lawrence, & Sanders, 1990, p. 315; Zhu & others, 1990). Outer shell layer: aragonite (homogeneous + fibrous prismatic). Middle and inner shell layers: aragonite (homogeneous).

Genus LAPTEVIELLA Kurushin in Dagys & Kurushin, 1985, p. 47

Type species.—Lapteviella prontchistshevi Kurushin in Dagys & Kurushin, 1985, p. 47.

Stratigraphic range.—Middle Triassic (Anisian) (Dagys & Kurushin, 1985). Lapteviella is a monospecific genus described

by Kurushin (in Dagys & Kurushin, 1985) from the Anisian of northern Central Siberia. It is similar to *Mesoneilo* Vu Khuc, 1977a, but it differs by having a pallial sinus, prosogyrous beaks, and the anterior part of hinge shorter than the posterior. It is also comparable to *Palaeoneilo* (both have prosogyrous beaks, a similar arrangement of hinge teeth, shallow pallial sinus, concentric ornamentation, adductor muscle scars of similar shape, size, and position) (see fig. 6 and 10 in Dagys & Kurushin, 1985, and fig. 31 in Hodges, 2000). On the other hand, *Lapteviella* figures in Dagys and Kurushin (1985) show neither an internal septum nor the typical radial groove of *Palaeoneilo*.

Paleogeographic distribution.—Boreal (Fig. 3).

Boreal domain: Middle Triassic: Anisian of Siberia (Dagys & Kurushin, 1985; Klets, 2006) and northeastern Russia (Konstantinov, Sobolev, & Yadernkin, 2007).

Paleoautoecology.—B, D, Is, FM; Sb. Dagys and Kurushin (1985) indicated the presence of a smooth pallial sinus, from which the presence of short siphons could be inferred. We assign a mode of life similar to other members of the family Malletiidae, i.e., detritivorous, feeding from the substrate surface, and constantly moving to find new food sources.

Mineralogy.—Aragonitic (Carter, Barrera, & Tevesz, 1998), data provided for the subclass Protobranchia.

Genus DIANUCULA Guo, 1988, p. 113

Type species.—Dianucula sulcata Guo, 1988, p. 113.

Remarks.—Guo (1988) included *Dianucula* in the family Malletiidae and we follow him, although Z. Fang and others (2009) included it in the family Afghanodesmatidae Scarlato & Starobogatov, 1979.

Stratigraphic range.—Upper Triassic (Norian) (Guo, 1988). Guo (1988) proposed *Dianucula* from upper Upper Triassic beds, and included two new species, the type and *Dianucula ovata* Guo, 1988. The genus was recorded from Dapingzhang formation, which is Norian in age (Feng & others, 2005). Sepkoski (2002) did not consider it in his compendium.

Paleogeographic distribution.—Eastern Tethys (Fig. 3).

Tethys domain: Late Triassic: Norian of southwestern China (Yunan province) (Guo, 1988).

Paleoautoecology.—B, D, Is, FM; Sb. We assign a mode of life similar to other members of the family Malletiidae, i.e., detritivorous, feeding from the substrate surface, and constantly moving to find new food sources.

Mineralogy.—Aragonitic (Carter, Barrera, & Tevesz, 1998), data provided for subclass Protobranchia.

Genus NINGLICONCHA J. Chen & Stiller, 2008, p. 364

Type species.—Ningliconcha ningliensis J. Chen & Stiller, 2008, p. 365.

Stratigraphic range.—Upper Triassic (lower Norian) (J. Chen & Stiller, 2008). J. Chen and Stiller (2008) proposed the monospecific genus *Ningliconcha* and reported it from upper lower Norian of China (Yunnan province).

Paleogeographic distribution.—Eastern Tethys (Fig. 3).

Tethys domain: Late Triassic: Norian of southwestern China (Yunnan province) (J. Chen & Stiller, 2008).

Paleoautoecology.—B, D, Is, FM; Sb. We assign this genus a mode of life similar to the rest of nuculids. The cancellate shell sculpture, characteristic of *Ningliconcha* (J. Chen & Stiller, 2008), probably aided in burrowing, as it does in other nuculids. The pallial line is integripalliate, and it either did not have siphons or they were short.

Mineralogy.—Aragonitic (Carter, Barrera, & Tevesz, 1998), data provided for subclass Protobranchia.

Genus YONGSHENGIA J. Chen & Stiller, 2008, p. 360

Type species.—Palaeoneilo cuneata Chen in Ma & others, 1976, p. 195.

Stratigraphic range.—Upper Triassic (middle Norian) (J. Chen & Stiller, 2008). J. Chen and Stiller (2008) proposed Yongshengia based on *P. cuneata* Chen in Ma & others, 1976 (lower middle Norian of Yunnan) and tentatively included *Palaeoneilo mundeni* Fleming (in Fleming, Munden, & Suggate, 1954) from the Norian of New Zealand.

Paleogeographic distribution.—Eastern Tethys (Fig. 3).

Tethys domain: Late Triassic: Norian of southwestern China (Yunan province) (Ma & others, 1976; Guo, 1985; Zhu & others, 1990; J. Chen & Stiller, 2008).

Austral domain: Late Triassic: Norian of New Zealand (Fleming, Munden, & Suggate, 1954).

Paleoautoecology.—B, D, Is, FM; Sb. Similar to *Palaeoneilo. Yong-shengia* is one of the largest nuculanoids and had a large posterior pedal muscle scar (J. Chen & Stiller, 2008). Its large size was probably inconvenient to fast burrowing, but this could be compensated for by a large foot (inferred by the pedal muscle scar).

Mineralogy.—Aragonitic. Zhu and others (1990) studied the shell microstructure of "*Palaeoneilo*" *cuneata*, and they described a probable aragonitic outer shell layer of homogeneous microstructure and aragonitic inner layers.

Family YOLDIIDAE Dall, 1908b Genus ROLLIERIA Cossmann, 1920, p. 82

Type species.—Nucula palmae J. de C. Sowerby, 1824, p. 117.

Remarks.—Cox and others (1969) and other workers (Liu, 1995; Gahr, 2002) considered *Rollieria* as a subgenus of *Nuculana* Link, 1807, but here we regard it as a separate genus, following Hodges (2000, p. 36), who remarked that *Rollieria* lacks some of the characters of *Nuculana*: "... it does not possess the characteristic elongated posterior, lacks an escutcheon and is suboval in outline." The name *Rollieria* also was used for an ammonoid genus, *Rollieria* Jeannet, 1951, p. 98, but *Rollieria* Cossman, 1920, has priority.

Stratigraphic range.—Lower Jurassic (Hettangian)–Lower Cretaceous (Hodges, 2000; Jingeng Sha, personal communication, 2008). Cox and others (1969) assigned it a Jurassic range, as did Sepkoski (2002), based on data from Hallam (1977). The oldest record is Hettangian (Hallam, 1972, 1976, 1977, 1987, 1990; Hodges, 2000). Sowerby described the type species from sediments of Carboniferous age, but Hodges (2000, p. 35–36) doubted the presence of *Rollieria* at that time, based on the fact that the genus was never found again in sediments older than Jurassic.

Paleogeographic distribution.—Tethys and Circumpacific (Fig. 4). Tethys domain: Early Jurassic: Hettangian–Sinemurian of southwestern Britain (Hodges, 2000; Mander, Twitchett, & Benton, 2008; Hallam & Wignall, 2000), Europe (Hallam, 1976, 1977, 1987).

Circumpacific domain: Early Jurassic: South America (Damborenea, 2002b).

Paleoautoecology.—B, D, Is, FM; Sb. We assign a mode of life similar to other nuculanoids. General morphology of the species of this genus suggests that it was a quick burrower (Hodges, 2000). *Rollieria* had a shallow pallial sinus, and thus probably had short siphons. We assume that the *Rollieria* mode of life was similar to *Palaeoneilo*.

Mineralogy.—Aragonitic (Carter, Barrera, & Tevesz, 1998), data provided for subclass Protobranchia.

Family POLIDEVCIIDAE Kumpera, Prantl, & Růžička, 1960 Genus RYDERIA Wilton, 1830, p. 72

Type species.—Leda renevieri Oppel, 1856 in 1856–1858, p. 215.

Remarks.—Although several authors include *Ryderia* as a subgenus of *Nuculana*, it is considered here as an independent genus, following Cox and others (1969). However, we regard *Teinonuculana* Zhang in Zhang, Wang, & Zhou, 1977, p. 9, as a synonym of *Ryderia* (see discussion of *Teinonuculana* in Genera not Included, p. 171).

Stratigraphic range.—Upper Triassic (Rhaetian)–Lower Jurassic (Toarcian) (Liu, 1995; J. Yin & McRoberts, 2006). Cox and others (1969) indicated that this genus was present in Europe during the Jurassic. Sepkoski (2002) restricted its range to the Lower Jurassic (lower Hettangian–upper Pliensbachian), following Hallam (1977, 1987). The oldest record we accept here is Rhaetian, according to Ivimey-Cook and others (1999) and J. Yin and McRoberts (2006). Hodges (2000) considered that the genus was present from Carboniferous to Early Jurassic times, but we did not find any mention older than Rhaetian nor any paper quoting this genus before the Late Triassic. Hodges did not provide references to his statement, and

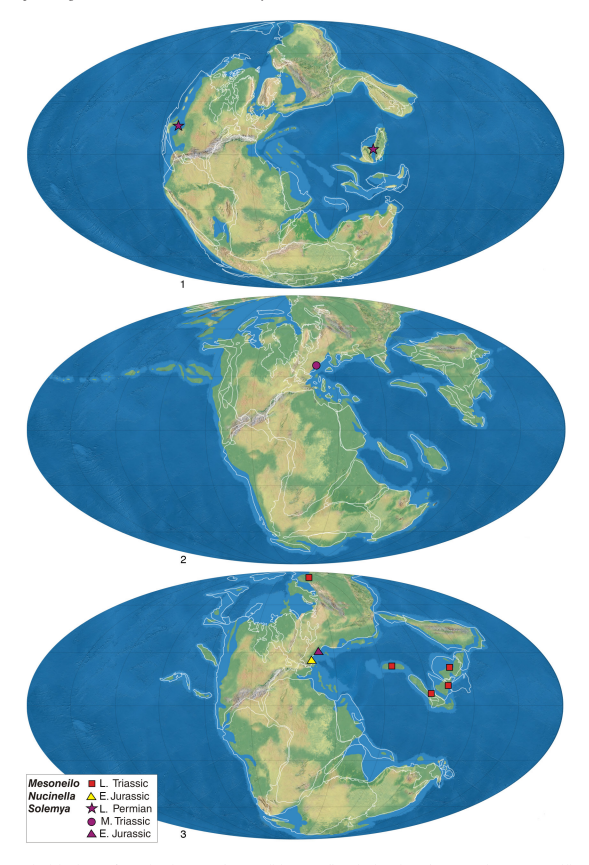


Figure 5. Paleogeographical distribution of Ctenodontidae (*Mesoneilo*), Nucinellidae (*Nucinella*) and Solemyidae (*Solemya*). 1, Early Triassic; 2, Middle Triassic; 3, Late Triassic–Early Jurassic.

thus it will not be taken into account here. The youngest record is from Toarcian beds (Liu, 1995; Fürsich & others, 2001).

Paleogeographic distribution.—Tethys and Circumpacific (Fig. 4). Tethys domain: Late Triassic: Rhaetian of Tibet (China) (J. Yin & McRoberts, 2006), northeastern England (Ivimey-Cook & others, 1999), southwestern United Kingdom (Mander, Twitchett, & Benton, 2008), southern Germany (Hodges, 2000); Early Jurassic: Germany and China (Hodges, 2000); Hettangian and Sinemurian of Europe (Hallam, 1976, 1977, 1987), eastern Asia and Australasia– Indonesia (Hallam, 1977), southwestern England (Hodges, 2000), southern England (Liu, 1995); Sinemurian of Portugal (Liu, 1995).

Circumpacific domain: Early Jurassic: Canada (Aberhan, 1998a), South America (Hodges, 2000), western Japan (Hodges, 2000).

Paleoautoecology.—B, D, Is, FM; Sb. According to Hodges (2000), *Ryderia* was a fast burrower. Puri in Cox and others (1969) mentioned a wide and shallow pallial sinus, but Hodges remarked that he did not observe such a feature in any of the specimens he studied, after making a thorough revision of the genus. Moreover, Hodges (2000, p. 45) indicated that "The lack of a pallial sinus and the extremely elongated rostrum suggest that the exposed siphons were short and that the tip of the rostrum lay just below the sediment surface." Like other nuculanids, it was probably detritivorous.

Mineralogy.—Aragonitic (Carter, Lawrence, & Sanders, 1990, p. 313; Carter, Barrera, & Tevesz, 1998). Outer and middle shell layers: aragonite (?). Middle shell layer: aragonite (?).

Genus DACRYOMYA Agassiz, 1842–1844, p. 500

Type species.—Nucula lacryma J. de C. Sowerby, 1824, p. 119. Remarks.—Dacryomya is a particularly problematical genus for several reasons. For a long time this genus has been (and still is) regarded as a subgenus of Nuculana. Externally, it is very similar to other nuculanoids, such as Ryderia, Nuculana, or Phestia, and the differences between these genera and Dacryomya are often subjective when internal structures are not seen. For example, the four genera have a rostrate posterior, but in different degree: Ryderia has the longest rostrum, while Dacryomya has the shortest. Hodges (2000, p. 21) pointed out that Dacryomya is very similar to Ryderia and Nuculana and they can be confused, but Dacryomya "is distinguishable from Ryderia by its much greater inflation and shorter rostrum and from Nuculana by its much shorter rostrum and lack of marked ridges bordering the escutcheon." Moreover, in Cox and others (1969, p. 239), Phestia was described as "Like Nuculana, but with prominent internal ridges," but Dacryomya and Ryderia also have internal ridges. A thorough review of this genus is needed.

Furthermore, there is no consensus about the family affiliation of this genus: it was referred to Nuculanidae (Cox & others, 1969; Hayami, 1975), Nuculidae (Ivimey-Cook & others, 1999; Hodges, 2000) or Polidevciidae (Carter, 1990a; Delvene, 2000), where we provisionally include it.

Stratigraphic range.—Upper Triassic (Norian)–Upper Jurassic (Kimmeridgian) (Okuneva, 1985; Delvene, 2000). There are some Lower Triassic records of this genus (Nakazawa, 1961; Hayami, 1975), but they will not be taken into account here because Puri in Cox and others (1969) ignored them and assigned it a Middle Jurassic range. Carter (1990a) and Hodges (2000) regarded the first appearance as Lower Jurassic, but Ivimey-Cook and others

(1999) reported it from the Rhaetian and Okuneva (1985) from Norian beds. Sepkoski (2002) assigned it a Lower Triassic–Lower Jurassic range, following Hayami (1975), but these data will not be taken into account due to the descriptive issues already mentioned. The youngest age accepted is Upper Jurassic, according to Delvene (2000).

Paleogeographic distribution.—Tethys, Circumpacific, and Boreal (Fig. 4).

Tethys domain: Late Triassic: Rhaetian of England (Ivimey-Cook & others, 1999). Early Jurassic: England (Watson, 1982); Sinemurian of Europe (Hallam, 1976, 1977, 1987), England and Portugal (Liu, 1995), England, Germany, Switzerland, France, and Portugal (Hodges, 2000); Hettangian of China (Hodges, 2000).

Circumpacific domain: Early Jurassic: Japan (Goto, 1983).

Boreal domain: Late Triassic: Norian of Transbaykal region (Siberia) (Okuneva, 1985). Early Jurassic: northern Siberia and Arctic region (Zakharov & others, 2006).

Paleoautoecology.-B, D, Is, FM; Sb. Similar to Palaeonucula.

Mineralogy.—Aragonitic (Carter, 1990a, p. 153–156). Outer shell layer: aragonite (prismatic). Middle and inner shell layers: aragonite (nacreous).

Superfamily CTENODONTOIDEA Wöhrmann, 1894 Family CTENODONTIDAE Wöhrmann, 1894 Genus MESONEILO Vu-Khuc, 1977a, p. 676

Type species.—Leda perlonga Mansuy, 1914, p. 82.

Remarks.—Vu Khuc (1977a) included this new genus in the family Ctenodontidae because it possesses a continuous hinge; in other words, the hinge is not interrupted below the umbo. He distinguished *Mesoneilo* from *Phaenodesmia* or *Palaeoneilo* because the first has opisthogyrous beaks and more teeth in the anterior part of the hinge. However, other authors included the type species of *Mesoneilo* in *Nuculana* (Gou, 1993; Hautmann, 2001b, p. 30) or in *Phestia* (Hautmann and others, 2005), but none of them mentioned *Mesoneilo*. A review of this species is needed to solve this question. Furthermore, some authors suggested that the family Ctenodontidae was exclusively Paleozoic and it is not well defined (Carter, 1990a); all these provide strong arguments to revise the familial affiliation of this genus.

Stratigraphic range.—Upper Triassic (Norian–Rhaetian) (Vu Khuc, 1977a). The genus was reported from Norian of northern Vietnam but a Norian to Rhaetian (Upper Triassic) range was given (Vu Khuc, 1977a). Later Okuneva (1985) referred a specimen from Norian beds of Siberia (Transbaykal region) to *Mesoneilo perlonga*, but she did not see its hinge and assigned it on the basis of its external shape alone. We think this assignation is very uncertain due to the external similarity with other nuculanoid genera.

Paleogeographic distribution.—Tethys and ?Boreal (Fig. 5).

Tethys domain: Late Triassic: China (Gou, 1993); Norian–Rhaetian of northern Vietnam, Laos, Burma, and southern China (Vu Khuc, 1977a; Vu Khuc & Huyen, 1998), Iran (Hautmann, 2001b).

?Boreal domain: Late Triassic: Norian of Transbaykal region (northern Siberia) (Okuneva, 1985).

Paleoautoecology.—B, D, Is, FM; Sb. Similar to Nuculana.

Mineralogy.—Aragonitic (Carter, Barrera, & Tevesz, 1998). Data about the shell mineralogy of the genus and the family are not avail-

able. We assign an aragonitic mineralogy following Carter, Barrera, and Tevesz (1998) for subclass Protobranchia.

Superfamily MANZANELLOIDEA Chronic, 1952 Family NUCINELLIDAE Vokes, 1956 Genus NUCINELLA Wood, 1851 in 1851–1882, p. 72

Type species.—Pleurodon ovalis Wood, 1840, p. 230.

Stratigraphic range.—Lower Jurassic (Hettangian)–Holocene (Vokes, 1956; McRoberts, Newton, & Allasinaz, 1995). The oldest record of *Nucinella* is *N. liasina* (Bistram, 1903) from Hettangian beds of the border between Switzerland and Italy (near lake Lucano) (Vokes, 1956). McRoberts, Newton, and Allasinaz (1995) mentioned the same species from Hettangian beds of the Lombardian basin (Alps). Cox and others (1969), as well as Skelton and Benton (1993) and Sepkoski (2002), agreed with the same time range.

Paleogeographic distribution.—Eastern Tethys (Fig. 5). The genus is now widely distributed geographically (see Cox & others, 1969, p. 269), as it was during older times (e.g., Cretaceous of Georgia [Pojeta, 1988] or Japan [Amano, Jenkins, & Hikida, 2007]). Nevertheless, during the study interval, the genus was only reported from Italy. During the Toarcian, it was also reported from Germany and England (Aberhan, 1993; Harries & Little, 1999).

Tethys domain: Early Jurassic: Hettangian of Italy (Vokes, 1956; McRoberts, Newton, & Allasinaz, 1995).

Paleoautoecology.—B, D-Ch, Is, FM; Sb. *Nucinella* was a nonsiphonate, active burrower, as indicated by the lack of pallial sinus. It was most probably detritivorous, and it possibly had chemosymbiotic bacteria. According to Allen and Sanders (1969), at least the living species possesses large gills and tiny palps, similar to *Solemya*, which is not considered as detritivorous (S. M. Stanley, 1970). This suggests that *Nucinella* may have had a similar feeding habit, but the wide bathymetric range of the living species (between 9 and 900 m, though most live at approximately 400 m) challenges this assumption. The possibility that they have symbiotic relations with chemosynthetic bacteria is strongly supported by the fact that some living species do not even posses palps or intestine (e.g., *N. viridula* Kuznetzov & Schileyko, 1984, *N. maxima* (Thiele & Jaeckel, 1931) (Beesley, Ross, & Wells, 1998).

Mineralogy.—Aragonitic (Carter, 1990a, p. 178). Outer shell layer: aragonite (prismatic). Middle and shell layers: aragonite (homogeneous).

Superfamily SOLEMYOIDEA Gray, 1840 Family SOLEMYIDAE Gray, 1840 Genus SOLEMYA Lamarck, 1818, p. 488

Type species.—Solemya mediterranea Lamarck, 1818, p. 488.

Remarks.—Cox and others (1969) regarded *Janeia* King, 1850, p. 177, as a subgenus of *Solemya*, but Pojeta (1988, p. 214–215) advised not to use that name since the generic concept lacks meaning. The genus *Acharax* Dall, 1908a, p. 2, is not included in the study interval, because no records from Triassic or Lower Jurassic deposits were found (see discussion in Genera not Included, p. 156).

Stratigraphic range.—Carboniferous (Upper Pennsylvanian)–Holocene (Pojeta, 1988). Cox and others (1969) assigned *Solemya* a Devonian–Holocene range, but Pojeta (1988) made an exhaustive revision of Paleozoic solemyoids and concluded that this genus ex-

tended from Pennsylvanian to Recent. We follow Pojeta, although other authors (Cope, 1997) extended its range back to the Devonian (Cope did not discuss this matter further), while others considered that the Paleozoic records are doubtful and regard the Jurassic as the first certain appearance (Imhoff & others, 2003; Little & Vrijenhoek, 2003). Ciriacks (1963) mentioned Solemya sp. from upper Permian deposits, but his assignation was only based on external shape. Seilacher (1990) described a new ichnofossil, Solemyatuba, that might be produced by Solemya or other related genera that live in a similar way. This ichnogenus has a wide distribution from Ordovician to Holocene (see Seilacher, 1990, p. 306-309). The only solemyoid genera known during Permian and Triassic times are Solemya and supposedly Acharax, and both have Recent representatives that are able to build Y-shaped tubes (S. M. Stanley, 1970; K. A. Campbell, Nesbitt, & Bourgeois, 2006); therefore, both are good candidates for Solemyatuba builders.

Paleogeographic distribution.—Tethys and Circumpacific (Fig. 5). During the time range of this review, *Solemya* was only reported from the Tethys, whereas it is mentioned from a wider geographic range at other moments of geologic time. Nevertheless, there is certain evidence (ichnogenus *Solemyatuba*) of its possible presence in Rhaetian beds of Germany and the Permian of Russia (Seilacher, 1990). During the middle Permian, it was reported from Russia and Siberia (Biakov, 2006; Ganelin & Biakov, 2006; Klets & others, 2006) and also from Middle and Late Jurassic from the Tethys (Fürsich, 1982; Komatsu, Saito, & Fürsich, 1993; Sha & Fürsich, 1994) and Austral domains (N. Hudson, 2003). There is a doubtful reference from the Late Triassic of Argentina (Damborenea & Manceñido, 2012).

Tethys domain: late Permian: Changhsingian of China (Teichert, 1990; M. Lin & Yin, 1991); Middle Triassic: Anisian of Hungary (Vörös & Pálfy, 2002); Early Jurassic: Germany (Seilacher, 1990).

Circumpacific domain: late Permian: Wyoming (United States) (Ciriacks, 1963).

Paleoautoecology.—B, Id, S-Ch, FM; Db. Solemya species have elongated and cylindrical shells with which they burrow deep Yshaped tunnels (S. M. Stanley, 1970). Although their feeding behavior is not fully understood, and many strategies have been proposed: detritivorous (Cope, 1997), filter feeder (S. M. Stanley, 1970; Fürsich, 1982), or the use of both strategies (Liljedahl, 1984), it is clear that most of their food requirements are provided by chemosynthetic symbiotic bacteria (Cavanaugh, 1983). Many Holocene species have very small gut and palp proboscides (Reid, 1998) but have disproportionately large gills where they lodge the chemosymbiotic bacteria (Stewart & Cavanaugh, 2006). They live most of their life inside the Y-shaped tunnels and have a well-developed foot used for burrowing and swimming (Reid, 1998). In addition to the typical Y- or U-shaped galleries (described for Solemya), other types of burrows, such as I- and J-shaped, were attributed to Acharax (Campbell, Nesbitt, & Bourgeois, 2006). Even though they can swim, this is not their main mode of life; and the foot may also be functional to move inside their galleries (S. M. Stanley, 1970). The known species usually live in shallow water areas and are almost always associated with low-oxygen environments rich in sulfur and organic matter (S. M. Stanley, 1970; Pojeta, 1988; Seilacher, 1990). This habitat provides a barrier against oxygen-dependent predators (A. G. Fischer & Bottjer, 1995).

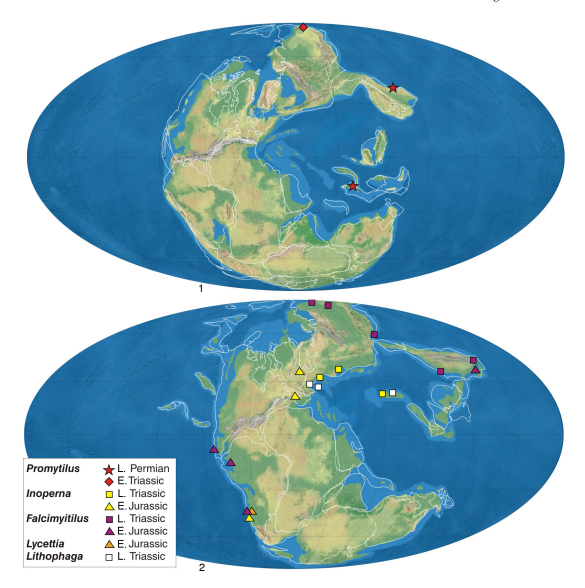


Figure 6. Paleogeographical distribution of Mytilidae (Promytilus, Inoperna, Falcimytilus, Lycettia, Lithophaga). 1, late Permian-Early Triassic; 2, Late Triassic-Early Jurassic.

Mineralogy.—Aragonitic (Carter, 1990a, p. 174). Outer shell layer: aragonite (prismatic). Middle and inner shell layers: aragonite (homogeneous).

Superfamily MYTILOIDEA Rafinesque, 1815 Family MYTILIDAE Rafinesque, 1815 Genus MODIOLUS Lamarck, 1799, p. 87

Type species.—Mytilus modiolus Linnaeus, 1758, p. 706.

Stratigraphic range.—Upper Devonian (Famennian)–Holocene (Cox & others, 1969). *Modiolus* is a long-ranging genus: having its origin in the Devonian, it is one of the oldest mussel genera with a good living representation (Cox & others, 1969). However, some authors believe that only Cenozoic to Holocene species should be referred to *Modiolus* (see Hodges, 2000).

Paleogeographic Distribution.—Cosmopolitan.

Paleoautoecology.—B, Se, S, Endo, Sed; By. As suggested by comparison with living *Modiolus* species, most fossil species are thought to have been semi-infaunal and endobyssate (S. M. Stanley, 1970, 1972). There are many examples of fossil *Modiolus* found in life position that confirm that they were gregarious and lived semiburied and fixed by their byssus to pebbles and other hard objects buried in the sediment (Fürsich, 1980, 1982). The byssus emerges from the anterior part of the shell. They tend to inhabit intertidal and subtidal, high-energy environments (S. M. Stanley, 1970; Hodges, 2000). Seilacher (1984, p. 228–229), in his own terminology, qualified this genus as a "mud-sticker."

Mineralogy.—Bimineralic (Hayami, Maeda, & Ruiz-Fuller, 1977; Carter, 1990a, p. 283). Outer shell layer: calcite (prismatic). Middle and inner shell layers: aragonite (nacreous).

Genus PROMYTILUS Newell, 1942, p. 37

Type species.—Promytilus annosus Newell, 1942, p. 38.

Stratigraphic range.—Carboniferous (Mississippian)-Lower Triassic (Induan) (Newell, 1942; Waller & Stanley, 2005). Cox and others (1969) assigned Promytilus a Carboniferous (Mississippian)-Permian range; Sepkoski (2002) assigned it to the Carboniferous (Mississippian)-Permian (upper Guadalupian), following Hayami and Kase (1977). But these last authors mentioned several Promytilus species reported by Nakazawa and Newell (1968) from middle and upper Permian (Changhsingian) of Japan (Tenjinnoki and Gujo formations), although Hayami and Kase (1977, p. 86) only indicate upper Permian (stage unknown). Boyd and Newell (1997) considered Gujo Formation to be of upper Permian age. The presence of Promytilus in Lower Triassic deposits was mentioned by Waller (in Waller & Stanley, 2005) based on P. borealis Kurushin, the original reference is Kurushin (in Dagys & others, 1989). In his original proposal of the genus, Newell (1942) noted that some Triassic and Jurassic specimens attributed to Modiolus could belong to Promytilus instead.

Paleogeographic distribution.—Boreal (Fig. 6). Promytilus had a cosmopolitan distribution during Carboniferous and early Permian times, and lived in the Tethys domain in the late Permian (Wuchi-apingian of China: Clapham, & Bottjer, 2007; Malaysia: Nakazawa, 1973), but during the study interval, we only found references from the Boreal domain.

Boreal domain: Early Triassic: Taimyr Peninsula (Russia) (Kurushin in Dagys & others, 1989).

Paleoautoecology.—B, E, S, Epi, Sed; By. According to S. M. Stanley (1972), *Promytilus* represents an intermediate stage between *Mytilus* and *Modiolus* and most probably lived attached by its byssus to hard substrates in the intertidal zone, by analogy with the extant species *"Modiolus" pulex* (Lamarck, 1819), with which it has a great similarity (see fig. 8 and 9 in S. M. Stanley, 1972). As described by Newell (1942), *Promytilus* had a well-defined byssal sinus. According to Waller (in Waller & Stanley, 2005), *Promytilus* is mytiliform in most shell features but possesses an anterior lobe that is smaller and less developed than in *Modiolus* species.

Mineralogy.—Bimineralic (Newell, 1942; Nakazawa & Newell, 1968). Newell (1942) and Nakazawa and Newell (1968) indicated the presence of a calcitic outer shell layer with prismatic microstructure, but Carter (1990a) doubts that this was the original mineralogy. Outer shell layer: ?calcite (prismatic). Middle and inner shell layers: aragonite (nacreous).

Genus INOPERNA Conrad in Kerr, 1875, p. 5

Type species.—Modiolus (Inoperna) carolinensis Conrad in Kerr, 1875, p. 5.

Stratigraphic range.—Upper Triassic (Rhaetian)–Upper Cretaceous (Maastrichtian) (Abdel-Gawad, 1986; Repin, 1996). Cox and others (1969) assigned it a Lower Jurassic (upper Liassic)–Upper Cretaceous range. However, findings referred to the subgenus *Triasoperna* Repin, 1996, p. 367 [7], indicate that the stratigraphic range of this genus should be extended back to the Upper Triassic (Repin, 1996; Hautmann, 2001b).

Paleogeographic distribution.—Tethys and ?Austral (Fig. 6). Although Inoperna was not widely distributed during our study interval, from Pliensbachian times and throughout all the Jurassic and Cretaceous, it had a cosmopolitan distribution (Freneix, 1965; Vörös, 1971; Hayami, 1975; Wen, 1982; Abdel-Gawad, 1986; Damborenea, 1987a; Liu, 1995; Holzapfel, 1998; Sha & others, 1998; Fürsich & others, 2001; Gahr, 2002; Delvene, 2003; Valls, Comas-Rengifo, & Goy, 2004).

Tethys domain: Late Triassic: Rhaetian of Iran (Repin, 1996; Hautmann, 2001b), northern Caucasus (Repin, 1996), Austria (Tomašových, 2006a, 2006b; Siblík & others, 2010); Early Jurassic: Hettangian of southwestern Great Britain (Hodges, 2000); Sinemurian of Portugal (Liu, 1995).

Austral domain: Early Jurassic: Hettangian–Sinemurian of Argentina (Damborenea, 1996a; Damborenea & Manceñido, 2005b).

Paleoautoecology.-B, Se, S, Endo, Sed; ?Bo. Because Inoperna was referred to the subfamily Lithophaginae, several authors suggested the possibility that it was a borer (Damborenea, 1987a; Hodges, 2000; Hautmann, 2001b). However, Pojeta and Palmer (1976) warned that although its morphology is similar to the current Lithophaga, which has a borer mode of life, this particular life habit cannot be certain unless we find the specimens within their holes. In addition, one of those authors found Inoperna plicata J. Sowerby (Middle Jurassic of England) in life position that indicates a semi-infaunal habit. Hodges (2000, p. 64) compared Inoperna with members of the living genus Adula H. Adams & A. Adams, 1857 in 1854-1858, and he suggested that, like them, Inoperna could be a mechanical borer, boring into the substrate with its anterior part and then fixed inside it by the byssus. Most authors regard Inoperna as semi-infaunal endobyssate (Fürsich & others, 1995, 2001; Hautmann, 2001b; Gahr, 2002; Delvene, 2003).

Mineralogy.—Bimineralic (Carter, 1990a, p. 185). There is no information about *Inoperna* shell mineralogy. We use the data provided for the subfamily Lithophaginae.

Outer shell layer: calcite (homogeneous-prismatic). Middle shell layer: aragonite (nacreous). Inner shell layer: aragonite (nacreous-prismatic).

Genus FALCIMYTILUS Cox, 1937c, p. 343

Type species.—Mytilus suprajurensis Cox, 1925, p. 142.

Stratigraphic range.—Upper Triassic (Carnian)–Upper Jurassic (Tithonian) (Kelly, 1984). Cox and others (1969) assigned it a Jurassic range, regardless of previous records that reported the genus from the Upper Triassic of Japan (Kobayashi & Ichikawa, 1950; Nakazawa, 1956; Hayami, 1958a). The latest undoubted record accepted here dates from Tithonian times (Kelly, 1984). Other authors, such as J. D. Taylor, Cleevely, and Morris (1983), mentioned *Falcimytilus lanceolatus* Sowerby from Lower Cretaceous (Albian), but they neither figured nor described the specimens, they just included them in a list of bivalve shells perforated by gastropods.

Paleogeographic distribution.—Tethys, Circumpacific, and Boreal (Fig. 6). During the study interval, the genus was known from the eastern Tethys, and later in the Jurassic, it was reported also from western Tethys (Hallam, 1976, 1996). The genus is often recorded from this area from Middle and Upper Jurassic beds (Fürsich, 1982; Kelly, 1984; Jaitly, 1988; Liu, 1995, 1999; Holzapfel, 1998; Sha & others, 1998).

Circumpacific domain: Late Triassic: Carnian of Japan (Kobayashi & Ichikawa, 1950; Nakazawa, 1956; Hayami, 1975); Early Jurassic: Mexico (Damborenea in Damborenea & González-León, 1997); Hettangian of Japan (Hayami, 1958a); Hettangian–Sinemurian of ?South America (Damborenea, 1996a).

Boreal domain: Late Triassic: northern Siberia (Dagys & Kurushin, 1985) and eastern Siberia (Kobayashi & Tamura, 1983b); Carnian of Primorie (Kiparisova, 1972); Norian of Russia (Zabaykal region) (Okuneva, 1985).

Paleoautoecology.—B, E, S, Epi, Sed; By. As indicated by its mytiliform outline and its triangular cross section, *Falcimytilus* probably lived as epibyssate on hard substrates, as living *Mytilus* species do (Fürsich, 1982; Damborenea, 1987a).

Mineralogy.—Unknown (Carter, 1990a, p. 283; 1990b, p. 400). Outer shell layer: calcite and/or aragonite (?). Middle shell layer: aragonite (nacreous). Inner shell layer: aragonite (nacreous-prismatic).

Genus LYCETTIA Cox, 1937c, p. 345

Type species.—Mytilus lunularis Lycett, 1857, p. 128.

Stratigraphic range.—Lower Jurassic (Sinemurian)–Upper Cretaceous (Maastrichtian) (Cox & others, 1969; Damborenea, 1996a). Cox (1937c) proposed Lycettia and reported it from the Jurassic. Cox and others (1969) considered it was present in the Jurassic and Upper Cretaceous. This discontinuous range is due to the fact that *Cuneolus* Stephenson, 1941, p. 156, is regarded as a synonym of Lycettia in Cox and others (1969). The type species of *Cuneolus* (Dreissena tippana Conrad, 1858, p. 328) is typical from Upper Cretaceous beds (Carter, 1990a). Since then, the genus was also reported from Lower Cretaceous deposits (Hayami, 1975; Villamil, Kauffman, & Leanza, 1998; Komatsu & Maeda, 2005).

Paleogeographic distribution.—Austral (Fig. 6). During the earliest Jurassic, *Lycettia* was only present in the Austral Domain, but during the rest of the Jurassic, it was also reported from the Tethys (Damborenea, 1987a; Liu, 1995; Fürsich & others, 2001; Gahr, 2002).

Austral domain: Early Jurassic: Sinemurian of Argentina (Damborenea, 1996a; Damborenea & Manceñido, 2005b; Damborenea & Lanés, 2007).

Paleoautoecology.—B, E, S, Epi, Sed; By. *Lycettia* presents some characters that indicate an epibyssate mode of life: external shape triangular in lateral view, the absence of anterior lobe and triangular cross section (S. M. Stanley, 1972). These features are taken to the extreme in this genus, which most probably lived fixed by the byssus to hard substrates in high-energy environments (Damborenea, 1987a). These hard substrates may be other bivalve shells, as *Myoconcha* from Middle Jurassic (Damborenea, 1987a), or *Steinmanella quintucoensis* (Weaver) from the Lower Cretaceous (Villamil, Kauffman, & Leanza, 1998).

Mineralogy.—Aragonitic (Carter, 1990b, p. 395–396). Outer shell layer: aragonite (prismatic). Middle shell layer: aragonite (nacreous). Inner shell layer: aragonite (prismatic-nacreous).

Genus LITHOPHAGA Röding in Bolten, 1798, p. 156

Type species.—Mytilus lithophagus Linnaeus, 1758, p. 705.

Stratigraphic range.---Upper Triassic (Norian)-Holocene (Kleemann, 1994). Cox and others (1969) considered the reports of this genus from Carboniferous age to be doubtful, and thus they attributed it a continuous range from Miocene to Recent. These doubtful data are surely from Newell (1942), who pointed out that, although Lithophaga was not very common in the Paleozoic, several species were described from Carboniferous and Permian beds around the world. However, Kleemann (1990) argued that the specimens attributed to Lithophaga from Carboniferous and Triassic age are indeed more than doubtful: they are externally similar to Lithophaga, but they probably did not have an endolithic mode of life. Kleemann (1994) and Carter and Stanley (2004) found specimens of Lithophaga in life position in holes bored inside Upper Triassic corals (as happens in many living species). Since the presence of Lithophaga in pre-Upper Triassic sediments cannot be assured, we will consider that the genus ranges from Upper Triassic to the present. Linck (1972) reported Lithophaga sp. cf. vermiculata Linck from Carnian beds, but he only assigned his specimen to Lithophaga on the basis of its external features, and he did not find it in life position, so we cannot really be sure that it was Lithophaga. Ivimey-Cook and others (1999) found borings that could belong to Lithophaga in Rhaetian deposits from England.

Paleogeographic distribution.—Tethys (Fig. 6). If finding specimens within their borings is a prerequisite to refer them to *Lithophaga*, then it is very difficult to know what the actual distribution of the genus was. In our study interval, it was reported from Tethys, but at other times (and also Recent), it had a cosmopolitan distribution (Cox & others, 1969).

Tethys domain: Late Triassic: Norian of Germany (Carter & Stanley, 2004); Rhaetian of Austria (Kleemann, 1994; Carter & Stanley, 2004), Iran (Hautmann, 2001b).

Paleoautoecology.—B, Is, S, By, Sed; Bo. Living Lithophaga species are borers in hard substrates, especially in dead and live coral skeletons (Kleemann, 1994; Scott, 1988). They are regarded as chemical borers, because they disaggregate the substrate with the aid of special chemical substances, although it appears that in some species, like Lithophaga nigra (d'Orbigny, 1853), this is supplemented by mechanical boring (L. Fang & Shen, 1988). Regarding fossil species, Lithophaga shells are reported in coral boreholes (Kleemann, 1994; Waller & Stanley, 2005), so the same mode of life is assumed for them. Savazzi (2001) mentioned a possible macrosymbiotic relationship between Lithophaga species burrowing into live corals and the corals themselves, as the coral provides protection and reduces competition with other borers that only bore on nonliving substrates. There is no evidence that the bivalve uses the coral as a food source.

Mineralogy.—Bimineralic (Carter, 1990a, p. 285). Outer shell layer: calcite (homogeneous-prismatic). Middle shell layer: aragonite (nacreous). Inner shell layer: aragonite (nacreous-prismatic).

Family MYSIDIELLIDAE Cox, 1964

Although Hautmann (2008) proposed that this family should be included in the superfamily Ambonychioidea, we follow Cox and others (1969), Carter (1990a), Amler (1999), Amler, Fischer, and Rogalla (2000), Waller and Stanley (2005), and Bouchet and Rocroi (2010) to refer it to the Mytiloidea (see discussions in Hautmann, 2001b, 2008; Waller & Stanley, 2005). Some disagreement has arisen in recent years about which genera should be included in this family, and this is summarized below.

Cox (1964) proposed the family and included within it Protopis Kittl, 1904, p. 718 (=Joannina Waagen, 1907, p. 94) and two new genera, Mysidiella Cox, 1964, p. 44 (pro Mysidia Bittner, 1891, p. 113, non Westwood, 1840) and Tommasina Cox, 1964, p. 44 (nom. nov. for Mytiliconcha Tommasi, 1911, non Mytiloconcha Conrad, 1862). Waller (in Waller & Stanley, 2005) proposed the synonymy Protopis (= Tommasina) and advised the exclusion of Protopis and its synonyms (Joannina and Tommasina) from the family Mysidiellidae, but he did not suggest a new allocation. Waller (in Waller & Stanley, 2005) proposed the inclusion of Botulopsis Reis, 1926 (emended by him) and his new genus Promysidiella Waller in Waller & Stanley, 2005. Stiller and Chen (2006), following Cox but without mentioning Waller's paper (in Waller & Stanley, 2005), added to the genera in the family (Protopis, Mytiliconcha, and *Mysidiella*), their three new genera from the Anisian of China: Leidapoconcha Stiller & Chen, Waijiaoella Stiller & Chen, and Qingyaniola Stiller & Chen. These authors regarded the name Mytiliconcha as valid, and, since Conrad's and Tommasi's names differ in one letter, we agree with Vokes (1980) in regarding Tommasina as an unnecessary name.

On the other hand, Hautmann (2008) proposed the new family Healeyidae, which includes Healeya Hautmann, 2001b, Joannina (which he regarded as valid for substantial differences with Protopis), and, with some hesitation, Protopis and the three genera created by Stiller and Chen (2006): Leidapoconcha, Waijiaoella, and Qingyaniola. In turn, he suggested that Mysidiella, Promysidiella, and Botulopsis should remain in Mysidiellidae, following Waller (in Waller & Stanley, 2005), but including this family within the Ambonychiodea. Hautmann (2008) based these conclusions on the study of the microstructure of one of his specimens of *Mysidiella imago* Hautmann, 2001b, where he found that the outer shell layer was subdivided into several sublayers (from outside to inside): prismatic, foliar, and coarsely prismatic. He argued that the foliar sublayer and the size of the prisms of the outer sublayer support an origin from myalinids rather than from mytilids. Nevertheless, although no microstructure studies are known for Promysidiella cordillerana (Newton), Newton (in Newton & others, 1987, fig. 13, p. 16) proposed that "abraded specimens exhibit an inner, very fine radial structure, representing silica replacement of primary fibrous prismatic microstructure, homologous to that occurring in outer ostracum of modern Mytilus, as well as Permian mytiloids (Newell, 1942)."

Waller (in Waller & Stanley, 2005) differentiated Mysidiellidae from Myalinidae, because they had different microstructure in the outer shell layers (fibrous prismatic and columnar prismatic, respectively) and different types of ligament (opisthodetic and duplivincular, respectively). But he did not study the microstructural shell details under the electronic microscope, and based his assumption on: "it superficially appears very similar to the microstructure of the outer shell layer of many modern mytilids" (Waller in Waller & Stanley, 2005, p. 8). Hautmann's (2008) proposal seems more solidly supported because it is based on microstructural studies and also on the revision of holotypes and several collections but, in our view, a lot of work is still needed to clarify this question. It is evident that the problems of this family are far from settled, and we therefore include all related genera in this family without further systematic discussions.

We follow Waller and Stanley (2005) and Stiller and Chen (2006) in the allocation of their genera into Mysidiellidae, and we also maintain the inclusion of *Protopis* and its synonyms. Meanwhile, in the absence of new studies, we believe it is still risky to accept the new family proposed by Hautmann (2008) with its original generic composition. We regard *Joannina* as a valid genus, following Hautmann (2008).

Genus MYSIDIELLA Cox, 1964, p. 44

nom. nov. pro Mysidia Bittner, 1891, non Westwood, 1840, p. 83 Type species.—Mysidia orientalis Bittner, 1891, p. 113.

Stratigraphic range.-Upper Triassic (Carnian-Rhaetian). Cox and others (1969) assigned this genus a Ladinian-Rhaetian range, as did also Sepkoski (2002) and Stiller and Chen (2006). The Ladinian record surely refers to Mysidia taramellii De Toni, 1913, because it is the only one from that stage (data from Diener, 1923); but Waller (in Waller & Stanley, 2005), in his revision of family Mysidiellidae and based on the illustrations provided by De Toni, assigned this species to Botulopsis Reis, 1926, p. 124. Moreover, they also considered that Mysidiella cordillerana Newton in Newton and others, 1987, p. 16, and Mysidiella americana (Körner, 1937) should belong to Promysidiella Waller in Waller & Stanley, 2005, p. 10. They renamed the specimens described by Newton (in Newton & others, 1987) as Krumbeckiella sp. cf. timorensis (Krumbeck, 1924) as their new species Mysidiella newtonae Waller & Stanley, 2005. Szente and Vörös in Budai and others (2003) reported ? Mysidiella sp. from Anisian beds, but this will not be taken into account because the material was not figured.

Paleogeographic distribution.—Tethys and Circumpacific (Fig. 7). Tethys domain: Late Triassic: Carnian of Greece (Diener, 1923), China (Wen & others, 1976); Norian of Turkey (Diener, 1923), China (Kobayashi & Tamura, 1983a); Norian–Rhaetian of Iran

(Hautmann, 2001b).
Circumpacific domain: Late Triassic: Carnian of British Columbia
(Waller & Stanley, 2005), of Japan (Nakazawa, 1994); Norian of
Oregon, United States (Wallowa Terrane) (Waller & Stanley, 2005).

Paleoautoecology.—B, E, S, Epi, Sed; By. The deep byssal notch (see diagnosis in Waller & Stanley, 2005) present in species attributed to this genus indicates an epibyssate mode of life. Their external morphology is similar to mytilid shells, and thus they probably lived in high-energy environments (Newton in Newton & others, 1987). The anterior part of the valves is flat, suggesting the animal lived orthothetically rested, i.e., with the commissure at a nearly right angle to the substrate surface (Hautmann, 2001b, 2008).

Mineralogy.—Bimineralic (Hautmann, 2008). According to Hautmann (2001b), the outer shell layer consists of foliated calcite, but Waller (in Waller & Stanley, 2005) did not find evidence of this type of microstructure in their specimens referred to *Mysidiella* species. Hautmann (2008), following a suggestion by Waller (in

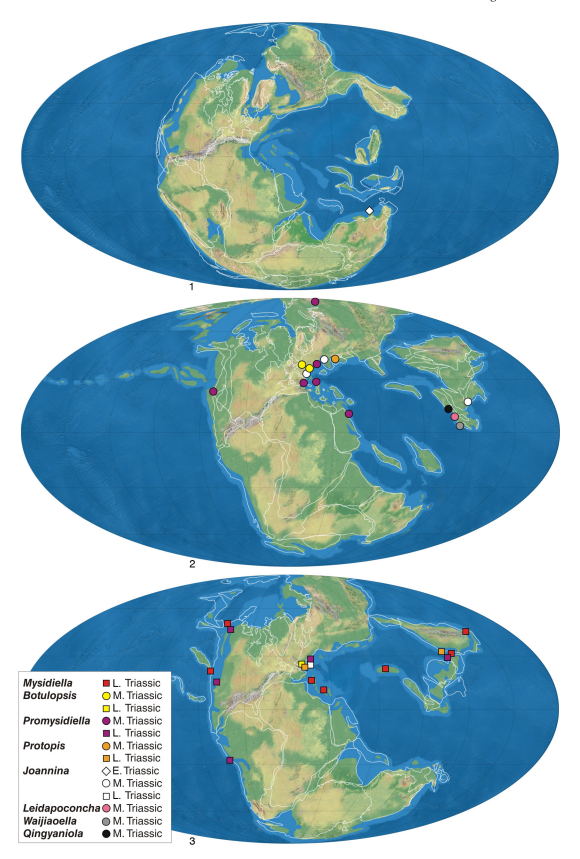


Figure 7. Paleogeographical distribution of Mysidiellidae (*Mysidiella, Botulopsis, Promysidiella, Protopis, Joannina, Leidapoconcha, Waijiaoella, Qingyaniola*). 1, Early Triassic; 2, Middle Triassic; 3, Late Triassic–Early Jurassic.

Waller & Stanley, 2005), studied the microstructure in tangential section (rather than in radial section, as he had done previously: Hautmann, 2001b) and found that the outer shell layer of *Mysidiella imago* Hautmann, 2001b, had several sublayers with prismatic, foliar, and coarsely prismatic microstructure (from outer to inner sublayers). Outer shell layer: calcite (prismatic). Middle and inner shell layers: aragonite (?).

Genus BOTULOPSIS Reis, 1926, p. 124

Type species.—Botulopsis reisi Waller in Waller & Stanley, 2005, p. 13 [*=Botulopsis cassiana* Reis, 1926, *non Botulopsis cassiana* (Bittner, 1895)].

Remarks.—We include *Botulopsis* in Mysidiellidae following Waller (in Waller & Stanley, 2005).

Stratigraphic range.—Middle Triassic (Ladinian)–Upper Triassic (Carnian) (Waller & Stanley, 2005). Cox and others (1969) assigned this genus an Upper Triassic range; Sepkoski (2002) considered *Botulopsis* present in Lanidian and Carnian times, based on data provided by Hallam (1981). Waller (in Waller & Stanley, 2005) emended the generic diagnosis and renamed the type species *Botulopsis cassiana* Reis, 1926, as *Botulopsis reisi* Waller in Waller & Stanley, 2005, present in Ladinian beds. Furthermore, Waller (in Waller & Stanley, 2005) included other two species within the genus: *Botula? cassiana* Bittner, 1895 (Carnian) and *Mysidia taramellii* De Toni, 1913 (Ladinian). Hautmann (2008) pointed out that *Botulopsis* was reported from Rhaetian deposits of Germany, and if that reference is correct, the range for this genus should be extended. Stiller (2001) mentioned *Botulopsis cassiana* from the Anisian of China, but this assignation is wrong (Stiller, personal comnunication, 2008).

Paleogeographic distribution.—western Tethys (Fig. 7).

Tethys domain: Middle Triassic: Ladinian of Austria (Kutassy, 1931), Italy (Reis, 1926; Waller & Stanley, 2005); Late Triassic: Carnian of the Alps (Bittner, 1895; Diener, 1923; Waller & Stanley, 2005).

Paleoautoecology.—B, E, S, Epi, Sed; By. *Botulopsis* was probably an epibyssate bivalve, on account of its external shell shape, which is quite inflated, and its shallow byssal gape, but possibly it did not inhabit high-energy environments, as *Mysidiella* did.

Mineralogy.—Bimineralic (Waller & Stanley, 2005). *Botulopsis* shell microstructure is not known; data used here are taken from the diagnosis of the family Mysidiellidae in Waller and Stanley (2005). Outer shell layer: calcite (fibrous prismatic). Middle and inner shell layers: aragonite (?).

Genus PROMYSIDIELLA Waller in Waller & Stanley, 2005, p. 10

Type species.—Mysidiella cordillerana Newton in Newton & others, 1987, p. 16.

Stratigraphic range.—Middle Triassic (lower Anisian)–Upper Triassic (lower Norian) (Waller in Waller & Stanley, 2005; Newton in Newton & others, 1987). The type species comes from lower Norian beds, but Waller (in Waller & Stanley, 2005) included also the following species: *Mysidia americana* Körner, 1937, *Mytilus eduliformis* Schlotheim, 1820, *Mytilus otiosus* McLearn, 1947, and two new species: *Promysidiella planirecta* Waller in Waller & Stanley, 2005, and *P. desatoyensis* Waller in Waller & Stanley, 2005. Waller (in Waller & Stanley, 2005) also pointed out that some species attributed to *Mytilus* Linnaeus, 1758, from the European Muschelkalk could be included in *Promysidiella*. In the same paper (Waller in Waller & Stanley, 2005, p. 10), he assigned it a Lower Triassic (Spathian)– Upper Triassic (Norian) range, but this is probably an error, because in the discussion he said, ". . . the oldest known *Promysidiella, P. eduliformis* (Schlotheim, 1820) from the lower Middle Triassic." Hautmann (2008) stated that *eduliformis* did not appear until early Anisian. There is, however, a Lower Triassic record of this species, although its systematic affiliation needs confirmation: Z. Yang & Yin (1979) mentioned *Mytilus eduliformis* from the upper Scythian Shihchienfeng Group of Shaanxi province (northern China), and Hautmann and others (2011) mentioned *Promysidiella?* sp. from southern China.

Paleogeographic distribution.—Tethys, Circumpacific, and ?Boreal (Fig. 7).

Tethys domain: Middle Triassic: Anisian–Ladinian of Spain (Mallada, 1880; Schmidt, 1935; Virgilli, 1958; Márquez-Aliaga, 1983, 1985; Budurov & others, 1991), Italy (Posenato, 2002; Posenato & others, 2002), Germany (Ürlichs, 1992), Jordan (Hautmann, 2008); Late Triassic: China (Gou, 1993), Germany (Warth, 1990).

Circumpacific domain: Middle Triassic: Ladinian of Nevada (United States) (Waller & Stanley, 2005); Late Triassic: Carnian of Peru (Körner, 1937), British Columbia (Canada) (Waller & Stanley, 2005); early Norian of Oregon (United States) (Newton, 1986; Newton & others, 1987).

?Boreal domain: Middle Triassic: Anisian of northern Siberia (Dagys & Kurushin, 1985), although we should check if this is taxonomically correct.

Paleoautoecology.—B, E, S, Epi, Sed; By. Newton (in Newton & others, 1987) suggested that the type species was epibyssate with a life style similar to Recent mytilids; based on its external shape, she inferred that it probably lived on hard substrates in high-energy open environments. Some species, such as *P. desatoyensis*, probably lived gregariously as the living *Mytilus edulis* Linnaeus does, according to taphonomic analysis of several individuals found in proximity to each other (Waller in Waller & Stanley, 2005). Waller (in Waller & Stanley, 2005) also argued that this species could have had a pendent mode of life, because some features (broad and flat anterior part, anterior margin concave, deep byssal invagination, and byssal notch) indicate that it was strongly attached by the byssus. However, he also suggested an epibyssate mode of life on hard substrates, but solitary (i.e., not forming clusters) for another species, *P. planirecta*.

Mineralogy.—Bimineralic (Newton in Newton & others, 1987, p. 16; Carter, 1990a, p. 286; Waller & Stanley, 2005, p. 10; but see Hautmann, 2008, p. 556). Outer shell layer: calcite (fibrous prismatic). Middle and shell layers: aragonite (?).

Genus PROTOPIS Kittl, 1904, p. 718

Type species.—Opis (Protopis) triptycha Kittl, 1904, p. 718.

Remarks.—We regard *Mytiliconcha* Tommasi, 1911 (=*Tommasina* Cox, 1964) as a synonym of *Protopis* following Waller (in Waller & Stanley, 2005) (see discussion for *Tommasina* in Genera not Included, p. 171). Waller (in Waller & Stanley, 2005, p. 9) removed *Protopis* from the Mysidiellidae and this was further discussed by Hautmann (2008, p. 559), who reillustrated the

original specimens of the type species and placed *Protopis* within the Modiomorphoidea (p. 562).

Stratigraphic range.---Middle Triassic (Anisian)--Upper Triassic (Carnian) (Waller & Stanley, 2005; Hautmann, 2008). Hautmann (2008) reviewed the genus Protopis and included there only the type species. Some species traditionally placed within this genus were transferred to Joannina by the author (see discussion for Joannina below). Therefore he assigned the genus to the Anisian. However, he did not refer to other species, as Protopis ginghaiensis Wen, from Carnian-Norian of Qinghai (data provided by Stiller & Chen, 2006). These authors suggested that this species might be included in Waijiaoella Stiller & Chen, 2006, based on its overall shape, but they noted a revision was needed. Waller (in Waller & Stanley, 2005) included Tommasina Cox, 1964 (see discussion in Genera not Included, p. 171) as a synonym of Protopis. Tommasina, or more correctly Mytiliconcha Tommasi, 1911, p. 35 (see Vokes, 1980), is also monospecific, including only the type, Mytiliconcha orobica Tommasi, 1911, from Carnian beds (Cox, 1964; Cox & others, 1969; Stiller & Chen, 2006). Waller (in Waller & Stanley, 2005) indicated its presence in Ladinian times, but this is most probably an error, because the only data source is Tommasi, 1911. Skelton and Benton (1993, p. 243) mentioned as first appearance of the family Mysidiellidae Protopis triptycha Kittl, 1904, from Scythian of the Werfen layers in the Austrian Alps, but it is not possible to verify this record because the authors did not indicate the original source.

Paleogeographic distribution.—Tethys (Fig. 7).

Tethys domain: Middle Triassic: Anisian of the Balkans (Hautmann, 2008); Late Triassic: Carnian of the Alps (Italy) (Cox, 1964; Cox & others, 1969; Stiller & Chen, 2006), China (Stiller & Chen, 2006).

Paleoautoecology.—B, E, S, Epi, Sed; By. We assign this genus an epibyssate mode of life, like most mysidiellids. The species here recognized within *Protopis* are *Opis (Protopis) triptycha* Kittl, 1904 (type species of *Protopis*) and *Mytiliconcha orobica* Tommasi, 1911 (type species of *Tommasina*, considered synonym of *Protopis*). These species lack the typical anterior lobe of *Joannina* species and show morphological features similar to other Mysidiellidae, so we suggest they had the same mode of life.

Mineralogy.—Bimineralic(?). There are no data about shell mineralogy and microstructure of this genus. Provisionally, we assign it bimineralic mineralogy.

Genus JOANNINA Waagen, 1907, p. 94

Type species.—Joannina joannae Waagen, 1907, p. 94.

Remarks.—Krumbeck (1924) included *Joannina* as a synonym of *Protopis* and Cox (1964) and Cox and others (1969), among others, accepted this situation, which Waagen (1907) already suspected (Hautmann, 2008). However, Hautmann (2008, p. 559–560) revised the holotypes of the type species of *Joannina* and *Protopis*, and he found differences that justify the separation of both genera. This author included within *Joannina* its type species and tentatively *Protopis timorensis* Krumbeck, 1924, from the Lower Triassic of Timor, *Joannina waageni* Schnetzer, 1934, and *Joannina aberrans* Schnetzer, 1934, both from the Anisian of Austria.

Stratigraphic range.—Lower Triassic (?Induan)–Upper Triassic (Carnian) (Krumbeck, 1924; Stiller & Chen, 2006). The oldest re-

cord is from the Lower Triassic (Krumbeck, 1924), and the youngest, *Protopis joannae*, from Carnian beds (Waagen, 1907; Stiller & Chen, 2006). This species was also mentioned by Hautmann (2008) from Ladinian [data provided by Waagen, 1907], and although Kochanová, Mello, and Siblík (1975, pl. 8,5) mentioned *Protopis* sp. cf. *joannae* from the Carnian of the Carpathians, the figured material is only a very poorly preserved fragment, not enough to ascertain if it really belongs to this species. Sha, Chen, and Qi (1990) also mentioned *Protopis*? sp. cf. *P. timorensis* Krumbeck, but according to Stiller and Chen (2006), this specimen is badly preserved and thus of doubtful relationship.

Paleogeographic distribution.—Tethys (Fig. 7).

Tethys domain: Early Triassic: ?Induan of Timor (Krumbeck, 1924); Middle Triassic: Anisian of Austrian Alps (Hautmann, 2008), China (Komatsu, Chen, & others, 2004), Hungary (Szente & Vörös in Budai & others, 2003); Ladinian of the Alps (Hautmann, 2008); Late Triassic: Carnian of the Alps (Stiller & Chen, 2006).

Paleoautoecology.—B, Se, S, Endo, Sed; By. According to Hautmann (2008, fig. 5), one of the main differences between *Joannina* and *Protopis* is that the former has a distinct anterior lobe. Furthermore, he pointed out that *Joannina* is modioliform, and thus he suggested an endobyssate mode of life, with the byssus emerging "between this anterior shell lobe and the main body of the shell, a faint radial shell fold creates a gape between both valves for the passage of the byssus" (Hautmann, 2008, p. 559, and see his fig. 5.1). *Joannina* is externally similar to *Leidapoconcha*, which probably had also an endobyssate mode of life.

Mineralogy.—Bimineralic(?). *Joannina* shell mineralogy or microstructure has not been studied. Due to the taxonomic problems already discussed, we cannot refer to the predominant mineralogy in the family (see explanation in Mineralogy of *Leidapoconcha* below). We provisionally assign it bimineralic mineralogy.

Genus LEIDAPOCONCHA Stiller & Chen, 2006, p. 215

Type species.—Leidapoconcha gigantea Stiller & Chen, 2006, p. 216. *Stratigraphic range.*—Middle Triassic (Anisian) (Stiller & Chen, 2006). *Leidapoconcha* has only been reported from sediments dated as lower upper Anisian (Stiller & Chen, 2006). It is a monotypic genus.

Paleogeographic distribution.—Eastern Tethys (Fig. 7).

Tethys domain: Middle Triassic: Anisian of southwestern China (Guizhou) (Stiller & Chen, 2006).

Paleoautoecology.—B, Se, S, Endo, Sed; By. According to the environment suggested for the deposits where *Leidapoconcha*, *Waijiaoella*, and *Qingyaniola* were found, they lived in fully marine, shallow water, and low-energy settings (Stiller & Chen, 2006). The authors also suggested an endobyssate or epibyssate mode of life for these genera, on the basis of their external morphology, since all of them have byssal gapes. Nevertheless, we think that a semi-infaunal, endobyssate, and sedentary mode of life is more feasible, similar to that proposed for *Healeya* by Hautmann (2001b).

Mineralogy.—Bimineralic(?). No data about the shell mineralogy and microstructure of this genus are available. Both Waller (in Waller & Stanley, 2005) and Hautmann (2008) agree that the family Mysidiellidae probably had a bimineralic shell, with calcitic outer shell layer and aragonitic middle and inner layers,

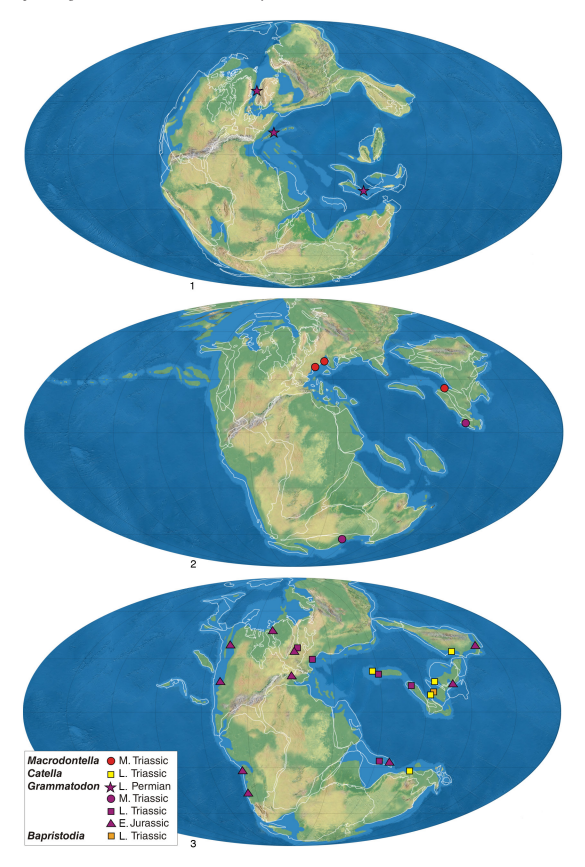


Figure 8. Paleogeographical distribution of Parallelodontidae (*Macrodontella, Catella, Grammatodon, Bapristodia*). 1, late Permian; 2, Middle Triassic; 3, Late Triassic–Early Jurassic.

but they disagree about the outer shell layer microstructure. However, Hautmann (2008) included *Leidapoconcha*, *Waijiaoella*, and *Qingyaniola* in his new family Healeyidae. No microstructural studies have been done on its type genus (*Healeya*), but an original aragonitic mineralogy is suggested by the fact that the shell is often found completely recrystallized (Hautmann, 2008). Stiller and Chen (2006) indicated the presence of recrystallized calcite in the shell of the specimens they studied.

Genus WAIJIAOELLA Stiller & Chen, 2006, p. 218

Type species.—Waijiaoella elegans Stiller & Chen, 2006, p. 219. Stratigraphic range.—Middle Triassic (Anisian) (Stiller & Chen, 2006). Waijiaoella was only reported from sediments dated as lower upper Anisian (Stiller & Chen, 2006). The genus includes two species, the type and Waijiaoella speciosa Stiller & Chen, 2006.

Paleogeographic distribution.—Eastern Tethys (Fig. 7). See *Leidapo-concha* (p. 26).

Paleoautoecology.—B, Se, S, Endo, Se. See *Leidapoconcha* (p. 26). *Mineralogy.*—Bimineralic(?). See *Leidapoconcha* (p. 26).

Genus QINGYANIOLA Stiller & Chen, 2006, p. 222

Type species.—*Qingyaniola mirabilis* Stiller & Chen, 2006, p. 223. *Stratigraphic range.*—Middle Triassic (Anisian) (Stiller & Chen, 2006). See *Leidapoconcha* (p. 26).

Paleogeographic distribution.—Eastern Tethys (Fig. 7). See *Leidapo-concha* (p. 26).

Paleoautoecology.—B, Se, S, Endo, Se. See *Leidapoconcha* (p. 26). *Mineralogy.*— ?Bimineralic. See *Leidapoconcha* (p. 26).

Superfamily ARCOIDEA Lamarck, 1809 Family PARALLELODONTIDAE Dall, 1898

Newell in Cox and others (1969, p. 256) pointed out that the phylogeny of this group is not well known, and several decades later, the problems to distinguish their genera still persist, although many authors have recently discussed this topic (see e.g., Damborenea, 1987a; Amler, 1989; Carter, 1990a; Stiller, 2006). The trouble is mainly focused on Parallelodon Meek & Worthen, 1866, Grammatodon Meek & Hayden, 1860, and Cosmetodon Branson, 1942. The general shell shape, the teeth (especially their arrangement), and ornamentation are features that were commonly used as criteria to distinguish these taxa (Manceñido, González, & Damborenea, 1976). Although it seems that the orientation of hinge teeth is a good criterion, it is not enough, since, as has been shown, in some Parallelodon species, the teeth may change their orientation during ontogeny (Newton in Newton & others, 1987; Hautmann, 2001b). In this regard, Stiller (2006, p. 12) concluded that "the convergence direction of the long posterior pseudolaterals appears to be taxonomically more reliable than the direction of the short anterior cardinals; the anterior ends of the posterior teeth intersect the dorsal shell margin in the Parallelodontinae and the ventral margin of the hinge plate in the Grammatodontinae." The difficulty of applying these criteria is that hinge teeth are not observed in most specimens. A thorough review of this family is needed, but it is beyond the scope of this paper.

Genus MACRODONTELLA Assmann, 1916, p. 616

Type species.—Macrodontella lamellosa Assmann, 1916, p. 616.

Remarks.—Although Assmann (1916) included *Macrodontella* in the family Arcidae, we follow Newell in Cox and others (1969) and later authors (e.g., Sha, Chen, & Qi, 1990) and refer it to the Parallelodontidae.

Stratigraphic range.—Middle Triassic (Anisian). Assmann (1916) described this monotypic genus from lower Muschelkalk (probably Anisian) from the Erzführender Dolomit Formation in Silesia (Poland). Cox and others (1969) assigned it a Middle Triassic range. *Macrodontella* was reported from middle Anisian beds from Poland (Malinowskiej, 1979). The genus was also doubtfully recorded from Chinese Anisian deposits (Sha, Chen, & Qi, 1990).

Paleogeographic distribution.—Tethys (Fig. 8).

Tethys domain: Middle Triassic: Poland (Assmann, 1916; Cox & others, 1969), Anisian of Poland (Malinowskiej, 1979), China (?Qinghai province) (Sha, Chen, & Qi, 1990).

Paleoautoecology.—B, E-Se, S, Epi-Endo, Sed; By. Sha, Chen, and Qi (1990) suggested an epibyssate and suspensivorous mode of life; however, Aberhan and others (2004) assigned it a semiinfaunal mode of life. Because it is a monotypic genus with reduced distribution, it is difficult to obtain good illustrations to help settle this question.

Mineralogy.—Aragonitic (Carter, 1990a, p. 189). No data about shell mineralogy-microstructure of *Macrodontella* are available. Data provided for the family Parallelodontidae (Carter, 1990a).

Genus CATELLA Healey, 1908, p. 13

Type species.—Gramatodon (Catella) laticlava Healey, 1908, p. 13. Stratigraphic range.—Upper Triassic (Carnian)-Lower Paleogene (Danian) (Wen & others, 1976; Heinberg, 1999). Healey (1908) described Catella as a subgenus of Grammatodon from the Rhaetian of Burma. Cox and others (1969) assigned it an Upper Triassic-Jurassic range, and Sepkoski (2002) assigned a Triassic (Norian)-Paleocene (Thanetian) range, and presumably got his data from Heinberg (1978), but in this last paper, the figure with the ranges of genera seems to indicate Danian, not Thanetian. The youngest record is Paleocene (Danian) (Heinberg, 1999), because we were not able to corroborate the range offered by Sepkoski (2002). The oldest record considered for almost all authors for Catella is Norian (Wen & others, 1976; Hallam, 1981; J. Zhang, 1983; Hautmann, 2001b). Nevertheless, Guo (1988) proposed a new subgenus, Catella (Oceanopieris), from the Carnian of Yunnan (China), which was considered a junior synomym of Catella by Z. Fang and others (2009). Catella shows a seemingly discontinuous distribution through time. Although it was widely mentioned from the Upper Triassic (see next section), it is not reported again until the Upper Jurassic (X. Li, 1990; Monari, 1994) and later in the Upper Cretaceous (Heinberg, 1999).

Paleogeographic ditribution.—Eastern Tethys (Fig. 8).

Tethys domain: Late Triassic: Carnian of Yunnan (China) (Guo, 1988); Norian of Yunnan (China) (J. Zhang, 1983), Himalaya (southern Tibet) (Wen & others, 1976; J. Yin & Enay, 2000; Hautmann,

2001b); Norian–Rhaetian of Iran (Hautmann, 2001b); Rhaetian of Burma (Healey, 1908), Indochina (Kutassy, 1931).

Paleoautoecology.—B, Se, S, Endo, Sed; By. Several aspects of the shell shape, such as the reduced anterior part of the shell, absence of ventral flattening, external modioliform appearance, and presence of byssal sinus, suggest that *Catella* was probably an endobyssate semiinfaunal bivalve (Heinberg, 1999; Hautmann, 2001b).

Mineralogy.—Aragonitic (Carter, 1990a, p. 184–185). There are no data about *Catella* shell mineralogy or microstructure. Data provided for superfamily Arcoidea.

Genus PARALLELODON Meek & Worthen, 1866, p. 17 nom. nov. pro Macrodon Buckman [Lycett MS] in Murchison, Buckman, & Strickland, 1844, non Schinz, 1822, p. 482, nec Müller, 1842, p. 308

Type species.—Macrodon rugosus Buckman in Murchison, Buckman, & Strickland, 1844, p. 99.

Stratigraphic range.—Middle Devonian–Upper Cretaceous (Amler & Winkler Prins, 1999). Traditionally, all Paleozoic members of the family Parallelodontidae were referred to *Parallelodon* (Newell in Cox & others, 1969); however, Manceñido, González, and Damborenea (1976) and Yancey (1985) noticed that many of these species should be referred to *Grammatodon* (*Cosmetodon*) instead. The same happens with some Mesozoic species referred to *Parallelodon*, which would better be allocated in *Grammatodon* (*Grammatodon*) (Damborenea, 1987a). Many Paleozoic species were described based on poorly preserved material or with little morphological discussion (Amler, 1989; Anelli, Rocha-Campos, & Simões, 2006). In practice, it is hard to distinguish between *Parallelodon* and *Grammatodon* (Boyd & Newell, 1979). For this reason, the range assigned here is provisional for these two genera, awaiting revision of Paleozoic material.

Paleogeographic distribution.—Cosmopolitan.

Paleoautoecology.-B, E, S, Epi, Sed; By. There are endobenthic and epibenthic species within this genus (S. M. Stanley, 1972). Those with a modioliform appearance are presumably endobyssate. Others are quadrangular and morphologically very similar to epifaunal Recent arcids. Some species, such as P. monobensis Nakazawa, 1955, have a large ventral sinus indicating an epibyssate mode of life. Other species, such as P. groeberi Damborenea, 1987a, and P. riccardii Damborenea, 1987a, were also epifaunal and probably attached to hard substrates with a strong byssus, as living Arca species do (S. M. Stanley, 1970; Damborenea, 1987a). Parallelodon riccardii might even have been a nestler (Damborenea, 1987a), as suggested by its elongated and laterally compressed shell (Thomas, 1978). However, P. tenuistriatus (Meek & Worthen, 1866) and P. hirsonensis (Archiac, 1843) were probably endobyssate (Quiroz-Barroso & Perrilliat, 1998; Fürsich & others, 2001). They are often found associated with corals and sponges (Damborenea, 1987a; Newton in Newton & others, 1987). But they could also live attached on rocks in open substrates (Newton in Newton & others, 1987). J. Yin and McRoberts (2006) suggested that representatives of the genus had an epibyssate and suspensivorous mode of life. We assign Parallelodon the predominant mode of life of species attributed to this genus.

Mineralogy.—Aragonitic (Carter, 1990a, p. 189–190). Outer shell layer: aragonite (prismatic). Middle shell layer: aragonite (cross-lamellar). Inner shell layer: aragonite (complex cross-lamellar).

Genus GRAMMATODON Meek & Hayden, 1860, p. 419

Type species.—Arca (Cucullaea) inornata Meek & Hayden, 1859, p. 51.

Remarks.—We regard *Cosmetodon* Branson, 1942, p. 248, as a subgenus of *Grammatodon*, following most authors (Fürsich, 1982; Kelly, 1984; Yancey, 1985; Damborenea, 1987a; Gardner & Campbell, 1997; Ivimey-Cook & others, 1999; Hautmann, 2001b; Nakazawa, 2002; Delvene, 2003). Some authors (Tashiro, 1986; Stiller, 2006) argued that *Cosmetodon* is a separate genus. Two other subgenera are included in our study range: *Grammatodon* and *Indogrammatodon* Cox, 1937b.

Stratigraphic range.—lower Permian (Artinskian)–Upper Cretaceous (Maastrichtian) (Yancey, 1985; Carter, 1990a). Newell in Cox and others (1969) included all Paleozoic members of the family Parallelodontidae within *Parallelodon*, but as stated above, there are certain specimens that should be attributed to *Grammatodon* instead (Manceñido, González, & Damborenea, 1976; Yancey, 1985). In the absence of a good review of Paleozoic members of this family, the first appearance is from the Permian Pacific margin (Manceñido, González, & Damborenea, 1976; Yancey, 1985) and the last appearance is from Upper Cretaceous age (Carter, 1990a). Sepkoski (2002) assigned it a Jurassic (Hettangian)–Cretaceous (?Cenomanian) range, following Cox and others (1969) and Hallam (1977).

Paleogeographic distribution.—Cosmopolitan (Fig. 8).

Tethys domain: late Permian: Malaysia (Nakazawa, 2002), Greece (Clapham & Bottjer, 2007); Middle Triassic: Anisian of Malaysia (Tamura & others, 1975); Late Triassic: Carnian of Malaysia (Tamura & others, 1975); Norian of Iran (Repin, 2001); Norian–Rhaetian of Iran (Hautmann, 2001b); Rhaetian of Burma (Healey, 1908), northeastern England (Penarth Group) (Ivimey-Cook & others, 1999), Tibet (Hautmann & others, 2005), Austria (Tomašových, 2006a); Early Jurassic: Tibet (Gou, 2003); Hettangian of Tibet (Hautmann & others, 2005); Sinemurian of Vietnam (Hayami, 1964; Sato & Westermann, 1991), Portugal and southern England (Liu, 1995), China (Stiller, 2006).

Circumpacific domain: Early Jurassic: western United States (Oregon) (Fraser, Bottjer, & Fischer, 2004); Hettangian of Japan (Hayami, 1958d); Hettangian–Sinemurian of Japan (Hayami, 1975); Sinemurian of Chile (Aberhan, 1994a) and Canada (Aberhan, 1998a).

Austral domain: Middle Triassic: Ladinian of New Zealand (Marwick, 1953); Early Jurassic: Argentina (Damborenea, 1987a; Damborenea & Lanés, 2007); Hettangian–Sinemurian of Neuquén Basin (Damborenea & Manceñido, 2005b).

Boreal domain: late Permian: Norway (Nakazawa, 1999); Early Jurassic: Hettangian of Greenland (Liu, 1995).

Paleoautoecology.—B, E, S, Epi, Sed; By. The assignation of one specific mode of life to this genus is difficult. Duff (1978) and Damborenea (1987a) found several inconsistencies, depending on which features of the shell were observed. According to authors and

species, Grammatodon was interpreted as epibyssate (Fürsich, 1982; Fürsich & others, 2001; Hautmann, 2001b; Delvene, 2003; J. Yin & Grant-Mackie, 2005; Aberhan, Kiessling & Fürsich, 2006; Stiller, 2006; Tomašových, 2006a), semi-infaunal (Fürsich, 1982; Pugaczewska, 1986; Delvene, 2003) or infaunal (Duff, 1978; Damborenea, 1987a; Gardner & Campbell, 1997; Harries & Little, 1999). According to S. M. Stanley (1972), the members of the subfamily Grammatodontinae would be rather epibyssate as suggested by their elongated shell by comparison with living species. However, in the same genus, we find species (such as G. toyorensis Hayami, 1959) with dorsally inflated shells and no evidence of byssal gape, which could be interpreted as shallow burrowers (Damborenea, 1987a); others [such as G. (Cosmetodon) mediodepressum (Krumbeck, 1913)] with an elongated ventral margin, which were probably epibyssate (Hautmann, 2001b), and finally, others [such as G. (Cosmetodon) keyserlingii (d'Orbigny, 1850) or G. (C.) marshallensis (Winchell, 1862)] showing a modioliform shape with an expanded posterior part, which are interpreted as semi-infaunal (S. M. Stanley, 1972; Fürsich, 1982). Having said that, we assign the prevailing inferred mode of life, i.e., epibyssate, to the genus.

Mineralogy.—Aragonitic (Carter, 1990b, p. 326). Outer shell layer: aragonite (cross-lamellar). Middle shell layer: aragonite (?). Inner shell layer: aragonite (cross-lamellar).

Genus BAPRISTODIA Guo, 1988, p. 115

Type species.—Bapristodia serrata Guo, 1988, p. 116.

Stratigraphic range.—Upper Triassic (Norian) (Guo, 1988). Guo (1988) proposed *Bapristodia*, a monospecific genus, from the Maichuqing formation dated as Norian (H. Yao & others, 2007).

Paleogeographic distribution.—Eastern Tethys (Fig. 8).

Tethys domain: Late Triassic: Norian of southwestern China (Yunan) (Guo, 1988).

Paleoautoecology.—B, E, S, Epi, Sed; By. We assign it the most common mode of life in the family Parallelodontidae. The external morphology of *B. serrata* is similar to some epibyssate species of *Grammatodon*, although no evidence of byssal gape is mentioned in the diagnosis offered by Guo (1988) [translated English version in Z. Fang & others, 2009].

Mineralogy.—Aragonitic (Carter, 1990a, p. 184–185). There are no data about the shell mineralogy or microstructure of *Bapristodia*. Data provided for superfamily Arcoidea (Carter, 1990a).

Family CUCULLAEIDAE Stewart, 1930 Genus CUCULLAEA Lamarck, 1801, p. 116

Type species.—Cucullaea auriculifera Lamarck, 1801, p. 116.

Remarks.—One of the subgenera of *Cucullaea* lived during the study interval: *Idonearca* Conrad, 1862, p. 289 (type species, *Cucullaea tippana* Conrad, 1858, p. 328).

Stratigraphic range.—Lower Jurassic (Hettangian)–Holocene (Hayami, 1958d; Beesley, Ross, & Wells, 1998). *Cucullaea* had its origin during the Early Jurassic and reached its greatest diversity during the Late Cretaceous, followed by a gradual decline until the present. Although Cox and others (1969) assigned it a discontinuous range [Jurassic (Liassic)–Cretaceous, Holocene], *Cucullaea* was present during the Cenozoic (Griffin, 1991; Griffin & Nielsen, 2008). The oldest record is from the Hettangian (Hayami, 1958d, 1975). It is

currently represented by a single species, *C. labiata* (Lightfoot, 1786), with an Indo-Pacific distribution (Beesley, Ross, & Wells, 1998).

Paleogeographic distribution.—Circumpacific (Fig. 9). Although during other times this genus was widely distributed, during the Early Jurassic (Hettangian and Sinemurian), it was only found in northwestern Pacific. During the Pliensbachian and Toarcian, it was distributed in the Arctic region (Zakharov & others, 2006), South America (A. F. Leanza, 1940, 1942; Damborenea, 1987a; Aberhan, 1994a), and Europe (Fürsich & others, 2001; Gahr, 2002).

Circumpacific domain: Early Jurassic: Hettangian of Japan (Hayami, 1958d, 1975).

Paleoautoecology.—B, Is-Se, S, SM; Sb. The species referred to *Cucullaea* have a very inflated quadrangular shell, with a truncated posterior end, indicative of a slow shallow-burrowing mode of life, as in modern species of *Anadara* (S. M. Stanley, 1970). The only extant species, *C. labiata*, lives at depths down to 200 m, buried in sand, with the anterior part downward (Beesley, Ross, & Wells, 1998). Damborenea (1987a) noted that shells of *C. jaworskii* A. F. Leanza and *C. rothi* A. F. Leanza (Lower Jurassic) lack epizoan organisms, whereas other epifaunal invertebrates from the same beds bear abundant epifauna. Thus we regard *Cucullaea* as a shallow infaunal or even semi-infaunal bivalve (see Damborenea, 1987a, p. 75).

Mineralogy.—Aragonitic (Carter, 1990b, p. 326). Outer shell layer: aragonite (prismatic or cross-lamellar). Middle shell layer: aragonite (cross-lamellar). Inner shell layer: aragonite (complex cross-lamellar).

Superfamily LIMOPSOIDEA Dall, 1895 Family PHILOBRYIDAE Bernard, 1897 Genus EOPHILOBRYOIDELLA Stiller & Chen, 2004, p. 414

Type species.—Eophilobryoidella sinoanisica Stiller & Chen, 2004, p. 414.

Stratigraphic range.—Middle Triassic (upper Anisian) (Stiller & Chen, 2004). *Eophilobryoidella* is particularly abundant in the middle part of the Leidapo Member of Qingyan formation. Up to now, the family Philobryidae was believed to range from the Eocene to the present time, but this finding extends its stratigraphic range back to the Middle Triassic. Therefore, the idea that this family evolved from the family Limopsidae, which has a Cretaceous origin, is invalidated (Stiller & Chen, 2004; Oliver & Holmes, 2006).

Paleogeographic distribution.—Eastern Tethys (Fig. 9).

Tethys domain: Middle Triassic: late Anisian of southwestern China (Guizhou province) (Stiller & Chen, 2004).

Paleoautoecology.—B, E, S, Epi, Sed; By. Living members of family Philobryidae are suspensivorous and live at depths that can exceed 1000 m, almost always attached (epibyssate) to other organisms (Beesley, Ross, & Wells, 1998). Stiller and Chen (2004) interpreted that *Eophilobryoidella* had a similar mode of life, although a byssal notch is not observed. These authors suggested that species of this genus were epibyssate, because they found epizoan organisms attached to the shells while the bivalve was alive. According to the environment in which these organisms lived, they concluded the species of *Eophilobryoidella* preferred shallow, low energy and normal salinity waters.

Mineralogy.—Aragonitic (Carter, 1990a, p. 195; Carter, 1990b, p. 328). There is no information about *Eophilobryoidella* shell mineral-

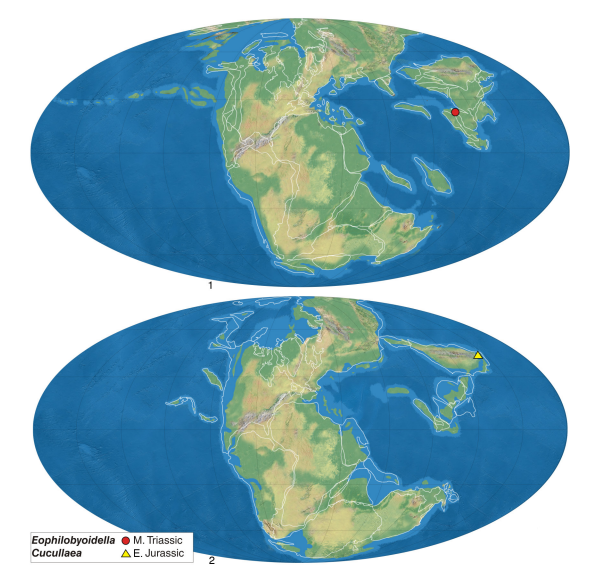


Figure 9. Paleogeographical distribution of Cucullaeidae (Cucullaea) and Philobryidae (Eophilobryoidella). 1, Middle Triassic; 2, Early Jurassic.

ogy or microstructure. Data from Recent Philobryidae specimens are used. Outer shell layer: aragonite (?). Middle shell layer: aragonite (cross-lamellar). Inner shell layer: aragonite (complex cross-lamellar).

Family PICHLERIIDAE Scarlato & Starobogatov, 1979 Genus HOFERIA Bittner, 1894, p. 190

Type species.—Lucina duplicata Münster, 1841, in Goldfuss, 1833–1841, p. 227.

Stratigraphic range.—Upper Triassic (Carnian). There are some *Hoferia* records from the Ladinian (Hallam, 1981; Kobayashi & Tamura, 1983a; Sepkoski, 2002), but none of these figure or indicate the original source of the data, and all of them are referred to the Alps. *Hoferia* was listed from the Cassian Formation in the southern Alps; according to Fürsich and Wendt (1977, fig. 2), this unit is upper Anisian to the lower part of the upper Carnian in age. Nevertheless, Fürsich (in PBDB, 2005) provided data from this paper, and he clearly assigned a Carnian age to *Hoferia* specimens. In addition, Kobayashi and Tamura (1983a) also mentioned *Hoferia*

from the Norian of Yunnan, but they did not provide any bibliographic reference. The only quotation of this genus from Yunnan is Cowper-Reed (1927), but from the Carnian. The last author only had a badly preserved internal mold of a right valve, and he included the specimen in *Hoferia*, because it shows a characteristic anterior lobe. We assign here a Carnian range to *Hoferia*.

Paleogeographic distribution.—Tethys (Fig. 10).

Tethys domain: Late Triassic: Carnian of the Italian Alps (Leonardi, 1943; Fürsich & Wendt, 1977), Yunnan (China) (Cowper-Reed, 1927).

Paleoautoecology.—B, Is, S, Endo, SM; Sb. The family Pichleriidae includes members of both shallow burrowers and those with epibyssate attached. *Hoferia* presents a byssal groove (see diagnosis in Cox & others, 1969, p. 265), and thus it must have been byssate. The globose shell suggests it was an endobyssate shallow burrower that lived near the surface or even semi-infaunally.

Mineralogy.—Aragonitic (Carter, 1990a, p. 196). There are no specific data for *Hoferia*, but we provisionally use those provided by

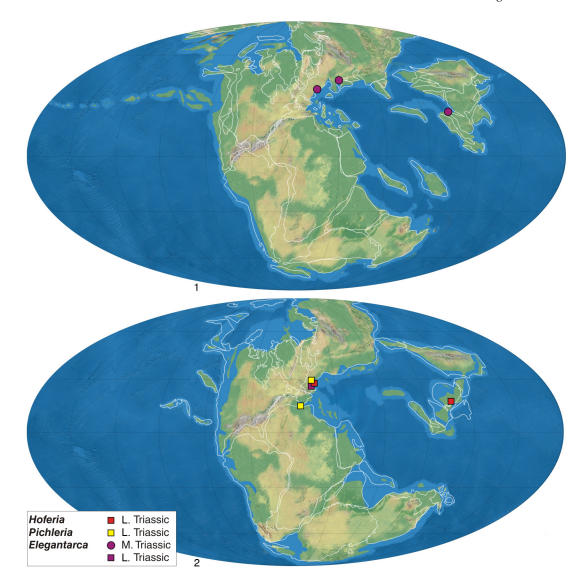


Figure 10. Paleogeographical distribution of Pichleriidae (Hoferia, Pichleria, Elegantarca). 1, Middle Triassic; 2, Late Triassic-Early Jurassic.

Carter (1990a) for the family Pichleriidae, provided by the analysis of *Pichleria*, a genus closely related to *Hoferia* (see above).

Genus PICHLERIA Bittner, 1894, p. 189

Type species.—Cucullaea auingeri Laube, 1865, p. 62.

Stratigraphic range.—Upper Triassic (Carnian) (Cox & others, 1969). Cox and others (1969) assigned *Pichleria* to the Upper Triassic, and Sepkoski (2002) to the Triassic (upper Ladinian–Carnian), following Hallam (1981) (see discussion for *Hoferia*). Wen and others (1976) mentioned *Pichleria* from the Norian of China, but the figured specimens (pl. 7,6–13) are members of family Limidae.

Paleogeographic distribution.—Tethys (Fig. 10).

Tethys domain: Late Triassic: Carnian of southern Alps (Italy) (Bittner, 1894, 1895; Diener, 1923; Leonardi, 1943; Corazzari & Lucchi-Garavello, 1980), southern Tunisia (Desio, Rossi Ronchetti, & Vigano, 1960). Paleoautoecology.—B, Is-Se, S, Sed; ?. The shell morphology indicates that *Pichleria* probably lived semi-infaunally or infaunally near the substrate surface. Byssal notch and sinus appear to be absent.

Mineralogy.—Aragonitic (Carter, 1990a, p. 196). Outer shell layer: aragonite (prismatic). Middle shell layer: aragonite (nacreous). Inner shell layer: aragonite (prismatic).

Genus ELEGANTARCA Tomlin, 1930, p. 23

Type species.—Arcoptera elegantula Bittner, 1895, p. 126.

Remarks.—Cox and others (1969) regarded *Elegantarca* (nom. nov. pro Arcoptera Bittner, 1895, p. 126, non Heilprin, 1887, p. 98) as a synonym of *Hoferia*, but Stiller (personal communication, 2005) argued to maintain them as separate genera, due to differences in orientation, number, and shape of the hinge teeth and other morphological disparities. In his own words: "*Elegantarca* shows some distinct morphological differences to *Hoferia*. Outer shell shape: *Elegantarca* has a large posterodorsal wing separated from the body of shell by a distinct but generally blunt posterior umbonal ridge (this diagonal ridge is lacking in *Hoferia*); *Elegantarca* in many cases is distinctly produced posteroventrally, *Hoferia* generally is shorter and more rounded. However, more important are differences in the hinge structure: *Hoferia* has a hinge with at least 10 short, taxodont teeth, which are radially arranged in two groups (Bittner, 1895; Broili, 1904; Cox & others, 1969); *Elegantarca* has very few, strong, radial teeth below the umbo, and one anterior and one posterior elongated tooth (about parallel to the hinge margin) (Broili 1904). The Chinese *Elegantarca subareata* Chen, Ma, & Zhang, 1974 has a hinge like the bivalves figured by Broili (1904, *Arcoptera*)."

Vokes (1980) regarded *Bittnerella* Dall, 1898, p. 613 (*nom. nov. pro Arcoptera* Bittner, 1895) as a valid name with priority over *Elegantarca*, but *Bittnerella* was included in the synonymy of *Hoferia* by Cox and others (1969), and now the name *Bittnerella* Dagys, 1974, p. 77, is used for a brachiopod genus.

Stratigraphic range.—Middle Triassic (Anisian)–Upper Triassic (Carnian) (Bittner, 1895; Komatsu, Chen, & others, 2004). The oldest known record of *Elegantarca* dates from Anisian times (Komatsu, Chen, & others, 2004; Stiller, personal communication, 2005) and the youngest from the Carnian (Bittner, 1895).

Paleogeographic distribution.—Tethys (Fig. 10).

Tethys domain: Middle Triassic: Anisian of southern China (Anonymous, 1974; Komatsu, Chen, & others, 2004; Stiller, personal communication, 2005), Bosnia (Diener, 1923); Ladinian of southern Alps (Italy) (Diener, 1923); Late Triassic: Carnian of southern Alps (Italy) (Bittner, 1895; Broili, 1904; Waagen, 1907).

Paleoautoecology.—B, Se, S, Endo, Sed; By. See *Hoferia*. Komatsu, Chen, and others (2004) regarded it as endobyssate semi-infaunal.

Mineralogy.—Aragonitic (Carter, 1990a, p. 196). See Hoferia.

Superfamily AMBONYCHIOIDEA Miller, 1877 Family MYALINIDAE Frech, 1891 Genus MYALINA de Koninck, 1842 in 1841–1844, p.125

Type species.—Myalina goldfussiana de Koninck, 1842 in 1841-1844, p. 126.

Stratigraphic range.—Carboniferous (lower Mississippian)–upper Permian, ?Lower Triassic (McRoberts, personal communication, 2005). Both Cox and others (1969) and Sepkoski (2002) assigned a Carboniferous (Lower Mississippian)–upper Permian range to this genus, but previously and subsequently, several authors mentioned *Myalina* from the Lower Triassic (e.g., Kiparisova, 1938; Newell & Kummel, 1942; Ciriacks, 1963; Dagys & Kurushin, 1985; Schubert, 1993; Schubert & Bottjer, 1995; McRoberts & Newell, 2005). All these references are better regarded as belonging to *Promyalina*, *Myalinella*, or even *Promytilus* (McRoberts, personal communication, 2005; McRoberts, 2005). So we leave the Triassic record of this genus as doubtful pending a good review of the problem. According to McRoberts (personal communication, 2005), no myalinid reached the Middle Triassic, since the only genus mentioned for that age (*Aviculomyalina*) should in fact be included in the Pteriidae or Malleidae.

Paleogeographic distribution.—Circumpacific (Fig. 11). Myalina had a cosmopolitan distribution, but from the late Permian, we only find records from the Tethys and Circumpacific domains. The family Myalinidae had significant diversity and abundance during the Carboniferous and Permian, but at the Permian–Triassic extinction, this family was decimated, later to disappear by the end of the Early Triassic (McRoberts, 2005). The doubtful Early Triassic records belong to the Circumpacific and Boreal domains.

Circumpacific domain: late Permian: western United States (Newell, 1942; Walter, 1953; McRoberts & Newell, 2005), Japan (Nakazawa & Newell, 1968; Hayami & Kase, 1977).

Paleoautoecology.—B, E, S, Epi, Sed; By. Species attributed to this genus have different morphologies and, consequently, their mode of life can be semi-infaunal (endobyssate) to epifaunal (epibyssate) (S. M. Stanley, 1972, fig. 12). Upper Permian specimens have reduced anterior lobes and, in some cases, bear a byssal sinus (M. lamellosa McRoberts & Newell, 2005; M. plicata McRoberts & Newell, 2005; M. copei Whitfield, 1902; see diagnosis in McRoberts & Newell, 2005), characters that indicate a byssate mode of life. Nevertheless, their thick and heavy shells were probably not functional for byssus attachment, and they were not active (Newell, 1942). According to McRoberts and Newell (2005), M. lamellosa probably lived lying on its anterior side, with an almost vertical commissure, lightly resting on its left valve. The species had a gregarious mode of life and is found in groups, as are many Recent mussels (Newell, 1942). Substrate type should also be taken into account: in soft substrates, they commonly adopt an endobyssate mode of life to become stable, while on hard substrates, they were frequently epibyssate.

Mineralogy.—Bimineralic (Newell, 1942, p. 33–34; Carter, 1990b, p. 331). Outer shell layer: calcite (prismatic-homogeneous). Inner shell layer: aragonite (nacreous).

Genus MYALINELLA Newell, 1942, 60

Type species .- Myalina meeki Dunbar, 1924, p. 201.

Remarks.—Newell (1942) described *Myalinella* as a subgenus of *Myalina*, pointing out the differences between *Myalinella* and other myalinids, such as *Myalina* (*Myalina*). Later, the same author (in Cox & others, 1969), raised *Myalinella* to genus level.

Stratigraphic range.—Carboniferous (Visean)–Lower Triassic (upper Olenekian) (R. Zhang & Pojeta, 1986; Fraiser & Bottjer, 2007a). Newell in Cox and others (1969) assigned a Carboniferous (Pennsylvanian)–Lower Triassic range to this genus and recorded it from Europe, United States, India, and Greenland. However, R. Zhang and Pojeta (1986) reported the first record of *Myalinella* from the Visean of China. The youngest record is Olenekian (Newell, 1942; Schubert, 1993; Schubert & Bottjer, 1995; Fraiser & Bottjer, 2007a).

Paleogeographic distribution.—Tethys, Boreal, and Circumpacific (Fig. 11). Myalinella had a wide distribution, but, during the late Permian, it seems restricted to few records; for instance, it was extensively mentioned from the western coast of the United States from the Permian until Guadalupian times (e.g., Newell, 1942; Ciriacks, 1963), but, from then on, it is only recorded from the Lower Triassic (e.g., Schubert, 1993), possibly due to lack of upper Permian deposits in this area. Even though during the early Permian it was present in the Tethys (e.g., Zheng, 1993), it seems to have been absent from this domain during the late Permian. However, it was recently reported from the Lower Triassic (Hautmann & others, 2011). Fraiser and Bottjer (2007a) studied several Lower Triassic sections from Japan and Italy, and they did not report Myalinella, but they did find another genus of the same family (*Promyalina*).

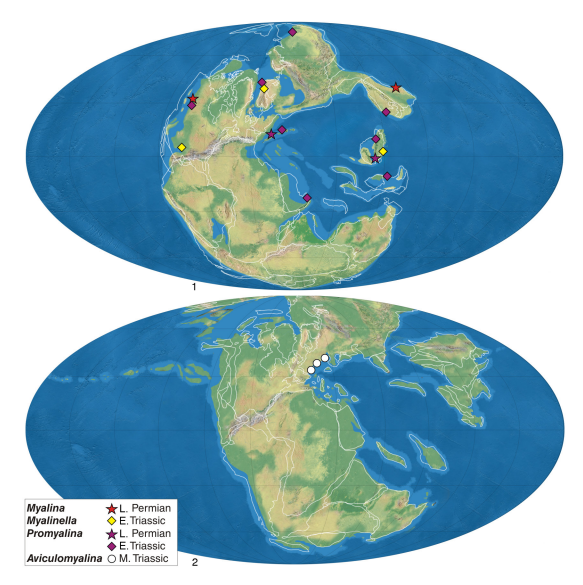


Figure 11. Paleogeographical distribution of Myalinidae (*Myalina, Myalinella, Promyalina, Aviculomyalina*). 1, late Permian–Early Triassic; 2, Middle Triassic.

Tethys domain: Early Triassic: Induan of southern China (Hautmann & others, 2011).

Boreal domain: Early Triassic: Greenland (Newell, 1942).

Circumpacific domain: Early Triassic: United States (Schubert, 1993; Schubert & Bottjer, 1995; Fraiser & Bottjer, 2007a).

Paleoautoecology.—B, Se, S, Endo, Sed; By. Specimens of this genus are usually found with closed valves, suggesting that they lived in low energy areas and/or were buried (Newell, 1942). They tolerated a wide salinity range (Newell, 1942), and they were found in fully marine (Kues, 2004) to estuarine environments (Mack & others, 2003). The shell shows some features indicative of a probable endobyssate and semi-infaunal mode of life: they have an anterior lobe and a byssal sinus, and their shells are small and fragile (Newell, 1942; McRoberts & Newell, 2005).

Mineralogy.—Bimineralic (Newell, 1942, p. 33–34). There are no available data about *Myalinella* shell microstructure. Newell (1942) indicated that, unlike most myalinids, *Myalinella* exhibits the same

structure in both valves. We assign the type present in members of the family Myalinidae.

Genus PROMYALINA Kittl, 1904, p. 690

Type species.—Promyalina hindi Kittl, 1904, p. 690.

Stratigraphic range.—upper Permian (upper Changhsingian)– Lower Triassic (upper Olenekian) (Fraiser & Bottjer, 2007a; He, Feng, & others, 2007). Cox and others (1969) assigned a Lower Triassic range to this genus, and they also doubtfully considered its presence in upper Permian beds. Although some authors (Sepkoski, 2002; McRoberts, 2005; McRoberts & Newell, 2005) only took into account the Lower Triassic records, others (Farabegoli, Perri, & Posenato, 2007, fig. 7; He, Feng, & others, 2007, fig. 5.18) accepted the late Permian records (late Changhsingian) from the Tethys domain. *Promyalina* showed a maximum abundance at the beginning of the Early Triassic, and it went extinct at the end of the same epoch.

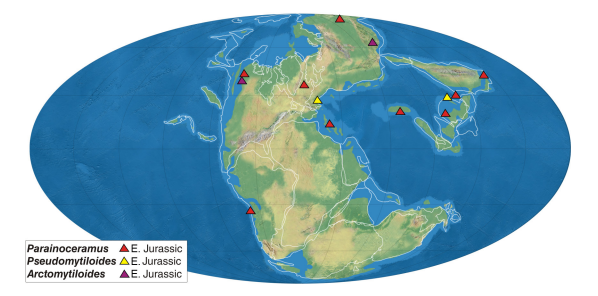


Figure 12. Paleogeographical distribution of Inoceramidae ("Parainoceramus," Pseudomytiloides, Arctomytiloides). Late Triassic-Early Jurassic.

Paleogeographic distribution.—Tethys, Circumpacific, and Boreal (Fig. 11).

Tethys domain: late Permian: Italy (Farabegoli, Perri, & Posenato, 2007), Changhsingian of southern China (He, Feng, & others, 2007); Early Triassic: China (Z. Yang & Yin, 1979; C. Chen, 1982; F. Wu, 1985; Lu & Chen, 1986; Ling, 1988; Komatsu, Huyen, & Chen, 2006, 2007), northern Vietnam (Komatsu, Huyen, & Chen, 2006, 2007); Induan of Oman (Krystyn & others, 2003; Twitchett & others, 2004), south of China (Hautmann & others, 2011); Induan–early Olenekian of Italy (Broglio-Loriga & others, 1990; Fraiser & Bottjer, 2005a, 2007a; Posenato, 2008a).

Circumpacific domain: Early Triassic: early Olenekian of Japan (Nakazawa, 1961; Hayami, 1975; Fraiser & Bottjer, 2007a); Olenekian of western United States (Ciriacks, 1963; Boyd, Nice, & Newell, 1999; Boyer, Bottjer, & Droser, 2004; Fraiser & Bottjer, 2007a).

Boreal domain: Early Triassic: northern Siberia (Dagys & Kurushin, 1985); Induan of Greenland (Wignall, Morante, & Newton, 1998; Wignall & Twitchett, 2002).

Paleoautoecology.—B, Se, S, Endo, Sed; By. Promyalina was one of the more widely distributed genera during the Early Triassic worldwide. Together with Eumorphotis and Unionites, it dominated the bivalve fauna of the seas at the beginning of the Triassic (Fraiser & Bottjer, 2005b, 2007a). It showed a typical opportunistic behavior, being more abundant when conditions were adverse but disappearing as soon as environmental conditions were restored, probably by competition with more specialized taxa. Morphological characteristics indicate a probable semi-infaunal and endobyssate mode of life, similar to Myalina (Schubert, 1993). Posenato (2008a) suggested an epibyssate mode of life.

Mineralogy.—Bimineralic (McRoberts & Newell, 2005). No data are available for shell mineralogy or microstructure of the species of this genus. We use data for the family Myalinidae (see McRoberts & Newell, 2005). Outer shell layer: calcite (prismatic-homogeneous). Inner shell layer: aragonite (nacreous).

Genus AVICULOMYALINA Assmann, 1916, p. 608

Type species.—Aviculomyalina lata Assmann, 1916, p. 608.

Remarks.—Both McRoberts (2005) and Waller (in Waller & Stanley, 2005) pointed out that *Aviculomyalina* could be better located within the Pteriidae or Malleidae. We keep it in Myalinidae until this matter is adequately discussed.

Stratigraphic range.—Middle Triassic (Anisian) (Cox & others, 1969). Assmann (1916) described the genus from the Lower Muschelkalk (probably Anisian), in the "Erzführender Dolomit" Formation of Silesia. Other authors also reported it from Anisian beds (Malinowskiej, 1979; Sepkoski, 2002). The range of this genus could be extended to Carnian and Norian if *Aviculomyalina? williamsi* (McLearn, 1941) were considered to belong to this genus, as proposed by Waller (in Waller & Stanley, 2005), rather than to *Mysidioptera* as proposed by Newton (in Newton & others, 1987).

Paleogeographic distribution.—Tethys (Fig. 11).

Tethys domain: Middle Triassic: Anisian of Poland (Assmann, 1916; Malinowskiej, 1979), Alps (?Switzerland) (Zorn, 1971).

Paleoautoecology.—B, E, S, Epi, Sed; By. An epibyssate mode of life is suggested by some shell features, such as the external shell morphology, the presence of byssal gape, and the flattened anterior margin (Newton in Newton & others, 1987).

Mineralogy.—Bimineralic. There is no information about *Aviculomyalina* shell mineralogy or microstructure, but we regard it equivalent to *Promyalina*.

Family INOCERAMIDAE Giebel, 1852

The family Inoceramidae is especially problematic, due partly to its great morphological variability, and partly to the lack of consensus among specialists about the taxonomically significant characters. It is often difficult to discern between the different genera assigned to this family because, if internal characters are not shown, the external fossil record with a long history but with the serious drawback of a wide disparity of concepts used by successive specialists (Harries & Crampton, 1998).

Genus "PARAINOCERAMUS" Cox, 1954, p. 47

ex Voronetz, 1936, p. 23, 34, nom. nud.

Type species.—Parainoceramus bulkuriensis Voronetz, 1936, p. 24, 34.

Remarks .- The generic name Parainoceramus was proposed by Voronetz (1936, p. 23, 34) on the basis of badly preserved specimens from sediments then dated as Carnian from northern Siberia. The author included four species in this new genus, but he did not designate a type, and thus this name was not available. Years later, Cox (1954) completed the requirements for the validity of the name by designating *P. bulkurensis* Voronetz as the type (ICZN, 1999, Art. 13B, 50). He did not see Voronetz's material, but, nevertheless, he included within *Parainoceramus* two other species that are widely distributed in the European Jurassic: "Crenatula" ventricosa J. de C. Sowerby, 1823, and Inoceramus substriatus Münster, 1835, in Goldfuss, 1833-1841. On the basis of his knowledge of these last species, he emended Voronetz's original diagnosis to include an anterior auricle and anterior teeth on some species. Cox's (1954) concept of the genus Parainoceramus was followed by nearly all later authors dealing with Jurassic material (e.g., Hayami, 1960; Speden, 1970; Duff, 1978; Damborenea, 1987b; J. Chen, 1988; M. A. Conti & Monari, 1991; Monari, 1994), who added more Jurassic species from around the world. Nevertheless, it is all too evident that this was reluctantly done in many instances, in the absence of a better alternative. Another point overlooked in the Treatise (Cox & others, 1969, p. 320) and by later authors is that Emel'yantsev and others (1960; see also Muromtseva, 1979; and Astafieva, 1986) had redated the beds where Voronetz's original material was found to be upper Permian (Wuchiapingian and Changhsingian), and thus the stratigraphic range of Parainoceramus sensu Cox (1954) should be upper Permian (Siberia), Hettangian to Tithonian (cosmopolitan), with no record during the Triassic. A breakthrough was provided by Astafieva (1986, 1993), who revised Voronetz's original material and concluded that the type species should be referred to the Paleozoic genus Kolymia Licharew in Licharew & Einor, 1941. Thus, several widely distributed and common Jurassic species (Parainoceramus sensu Cox non Voronetz) remain without a genus to be referred to. We will provisionally use here the name "Parainoceramus" in this sense, until a proper solution is developed (Ros, Damborenea, & Márquez-Aliaga, 2009), and we record its first appearance in the earliest Jurassic. When the material is not well preserved, it is difficult to distinguish between Parainoceramus in this sense and Pseudomytiloides Koschelkina, 1963 (Aberhan, 1998a; Stiller, 2006).

Stratigraphic range.—Lower Jurassic (Hettangian)–Upper Jurassic (Tithonian) (Escobar, 1980; Kelly, 1984). Both Cox and others (1969) and Sepkoski (2002) regarded the first appearance of *"Parainoceramus"* to be Upper Triassic following Voronetz (1936),

ignoring that Emel'yantsev, Kravtsova, and Puk (1960) had already corrected the dating of the beds from which Voronetz described his specimens from Carnian to upper Permian. We assign the Lower Jurassic (Hettangian) as the oldest record (Escobar, 1980; Damborenea, 1996a), taking into account only the species assigned to this genus *sensu* Cox (1954). The youngest record is from the Tithonian (Kelly, 1984; Fozy, Kázmér, & Szente, 1994; Liu, 1995).

Paleogeographic distribution.—Tethys, Boreal, and Circumpacific (Fig. 12). "*Parainoceramus*" was cosmopolitan during the Early Jurassic (especially during the Pliensbachian), but during the Middle and Late Jurassic, it appears to have had a bipolar distribution (possibly Boreal and Austral domains) (Damborenea, 1996b). For further information, see Damborenea (1987b, p. 142–146).

Tethys domain: Early Jurassic: Tibet (Gou, 1985), China (J. Chen, 1982a, 1988); Hettangian of Vietnam (Hayami, 1964); Hettangian–Sinemurian of China (Stiller, 2006); Sinemurian of southern England (Liu, 1995), Turkey (M. A. Conti & Monari, 1991), China (Y. Wang & Smith, 1986).

Boreal domain: Early Jurassic: Sinemurian of northern Siberia (Hallam, 1977).

Circumpacific domain: Early Jurassic: Japan (Hayami, 1960); Hettangian–Sinemurian of Canada (Aberhan, 1998a); Hettangian of Chile (Escobar, 1980).

Paleoautoecology.—B, E, S, Epi, Sed; By. The shell morphology of species attributed to "Parainoceramus" is variable, and it largely depends on the type of environment. Some species, such as "P." *jinjiensis* Chen, 1988, or "P." subtilis (Lahusen), are mytiliform, and they were probably epibyssate (Duff, 1978; Stiller, 2006), but other species, such as "P." apollo (A. F. Leanza, 1942), are modioliform with a well-developed anterior lobe, and they possibly had an endobenthic mode of life (Damborenea, 1987b). Other genera of the same family, such as Pseudomytiloides, were interpreted as pseudoplanktonic, at least in the early stages (Hayami, 1969a; Etter, 1996). "Parainoceramus" species are found in a wide array of facies types, since they could inhabit different environments, but they are especially abundant in anoxic facies (black shales) (Damborenea 1987b; Harries & others, 1996).

Mineralogy.—Bimineralic (Carter, 1990a, p. 200; Carter, 1990b, p. 330). Outer shell layer: calcite (prismatic). Middle and inner shell layers: aragonite (nacreous).

Genus PSEUDOMYTILOIDES Koschelkina, 1963, p. 129

Type species.—Mytiloides marchaensis Petrova, 1947, p. 130.

Stratigraphic range.—Lower Jurassic (Hettangian)–Middle Jurassic (Aalenian) (Etter, 1996; Stiller, 2006). Cox and others (1969) assigned a Jurassic range to this genus. The oldest record is from Hettangian deposits (Stiller, 2006) and the youngest from Aalenian beds (Etter, 1996). The genus was also mentioned from the Upper Triassic (Norian–Rhaetian) of northeastern Asia (Kurushin, 1990; Polubotko & Repin, 1990), but without indication of the original sources and with no illustrations, so these records remain doubtful.

Paleogeographic distribution.—Tethys (Fig. 12). This genus was especially abundant during the Toarcian. During the earliest Jurassic, it was only reported from the Tethys; but it was also recorded from the Boreal domain by Pliensbachian and Toarcian times (Zakharov & others, 2006). Poulton (1991) reported *Pseudomytiloides* (?) sp. from ?Hettangian beds of Canada, but it was only one specimen questionably referred to this genus.

Tethys domain: Early Jurassic: Hettangian–Sinemurian of China (Stiller, 2006); Sinemurian of southwestern France (Liu, 1995).

Paleoautoecology.-B-Ps, E, S, Epi, Sed-FaM; By. Pseudomytiloides dubius (J. de C. Sowerby, 1823) is particularly linked to the black shale facies, associated with anoxic conditions, since its abundance decreases as soon as the normal environmental conditions are restored after the early Toarcian extinction event (Harries & Little, 1999; Fürsich & others, 2001; Caswell, Coe, & Cohen, 2009). A pseudoplanktonic mode of life was proposed for this species and P. matsumotoi (Hayami, 1960) (Hayami, 1969a; Tanabe 1983; Seilacher, 1990). This was based on many forms of evidence: they are often found in anoxic facies (Hayami, 1969a), attached to pieces of wood (Hayami, 1969a; Tanabe 1983; Seilacher, 1990), and to ammonoid shells and other bivalves (Tanabe, 1983). However, some authors (e.g., Wignall & Simms, 1990; Etter, 1996) suggested that this interpretation is inadequate since the species abundance is too high to be derived only from floating logs. They proposed that *P*. dubius had a benthic epibyssate mode of life instead, but it could occasionally live as facultative pseudoplanktonic, with the capacity to attach to various substrates, such as floating objects, and tolerate low oxygen environments (see also Caswell, Coe, & Cohen, 2009). Some authors proposed that certain species of Pseudomytiloides might contain chemosymbionts that would help them live in these inhospitable settings (Harries & Crampton, 1998). Other species, such as P. yinhangensis Chen, 1988, were found in well-oxygenated, quiet, and near-shore environments (Stiller, 2006). The epibyssate mode of life is clearly suggested by their mytiliform shells and their long and flat anteroventral margin (Tanabe, 1983).

Mineralogy.—Bimineralic (Carter, 1990a, p. 200). There are no specific data for *Pseudomytiloides*. Data used here provided by Carter (1990a) for family Inoceramidae. Outer shell layer: calcite (prismatic). Middle and inner shell layers: aragonite (nacreous).

Genus ARCTOMYTILOIDES Polubotko, 1992, p. 64

Type species.—Pseudomytiloides rassochaensis Polubotko, 1968b, p. 61.

Stratigraphic range.—Lower Jurassic (Sinemurian-?Toarcian) (Aberhan, 1998a). The genus was described by Polubotko (1992). It was reported from the Sinemurian of northeastern Russia, and it includes the following species besides the type: *A. sinuosus* (Polubotko, 1968b), *A. kelimiarensis* Polubotko, 1992, and *A.* (?) turomtchensis Polubotko, 1992. Subsequently, Aberhan (1998a) doubtfully reported it from Toarcian beds.

Paleogeographic distribution.—Boreal and Circumpacific (Fig. 12). Boreal domain: Early Jurassic: Sinemurian of northeastern Russia (Polubotko, 1968b, 1992).

Circumpacific domain: Early Jurassic: Sinemurian-Toarcian of ?Canada (Aberhan, 1998a).

Paleoautoecology.—B, E, S, Epi, Sed; By. Its mytiliform shell indicates an epibyssate mode of life, similar to some living species of the family Mytilidae.

Mineralogy.—Bimineralic (Carter, 1990a, p. 200). No data about *Arctomytiloides* shell mineralogy and microstructure are available. Data provided for the family Inoceramidae (see any genera in this family).

Superfamily PTERIOIDEA Gray, 1847 Family PTERIIDAE Gray, 1847 Genus PTERIA Scopoli, 1777, p. 397

Type species.—Mytilus hirundo Linnaeus, 1758, p. 706.

Remarks.—Pteroperna Morris & Lycett, 1853 in 1851–1855, is a subgenus, and *Rhynchopterus* Gabb, 1864, is a junior synonym of *Pteria s.l.* (see discussion for *Pteroperna* and *Rhynchopterus* in Genera not Included, p. 169 and 170).

Stratigraphic range.—Lower Triassic (Olenekian)–Holocene (Hayami, 1975; Beesley, Ross, & Wells, 1998). Pteria ranges from Triassic to Holocene times (Cox & others, 1969; Sepkoski, 2002). The earliest record found is *P. ussurica* (Kiparisova, 1938) from the Induan (Hayami, 1975). There are some pre-Triassic records (see PBDB, on-line), but most of them were published before Cox and others (1969). M. Wang (1993) described *Pteria? yonganensis* M. Wang, 1993, from upper Permian beds, but the generic assignment was only tentative, since he had few specimens and their differences with *Pteria* were important. Tëmkin (2006) doubted the origin of *Pteria* in the Early Triassic because many so-called winged shells that probably belong to other families (even Bakevelliidae or Isognomonidae) were referred to this genus.

Paleogeographic distribution.—Tethys, Circumpacific, and Boreal (Fig. 13). In the past, *Pteria* was a widespread genus; today it is common in warm seas (Cox & others, 1969; Beesley, Ross, & Wells, 1998), and was apparently not known from the Austral domain.

Tethys domain: Early Triassic: China (Z. Yang & Yin, 1979; S. Yang, Wang, & Hao, 1986; Ling, 1988; L. Li, 1995; Shen, He, & Shi, 1995; Tong & others, 2006; Komatsu, Huyen, & Chen, 2007); Middle Triassic: Anisian of the Alps (Switzerland) (Zorn, 1971), southern China (Komatsu, Chen, & others, 2004; Komatsu, Akasaki, & others, 2004); Anisian–Ladinian of northern Vietnam (Komatsu, Huyen, & Huu, 2010); Late Triassic: Carnian of Italy (Allasinaz, 1966; Fürsich & Wendt, 1977; Corazzari & Lucchi-Garavello, 1980), Germany (Linck, 1972); Norian of China (Lu, 1981); Norian–Rhaetian of Iran (Hautmann, 2001b); Rhaetian of Italy (Allasinaz, 1962; Gelati & Allasinaz, 1964; Gaetani, 1970); Early Jurassic: Hettangian of Italy (Allasinaz, 1962), China (J. Yin & McRoberts, 2006); Hettangian–Sinemurian of Italy (Gaetani, 1970); Sinemurian of eastern Asia (Hallam, 1977), Portugal (Liu, 1995).

Circumpacific domain: Early Triassic: Japan (Nakazawa, 1971; Hayami, 1975; Kashiyama & Oji, 2004); Late Triassic: Norian of Japan (Nakazawa, 1964); Early Jurassic: Hettangian of Japan (Hayami, 1975; Kondo & others, 2006; Fraiser & Bottjer, 2007a), ?Chile (Aberhan, 1994a).

Boreal domain: Late Triassic: Carnian of Primorie (Kiparisova, 1972).

Paleoautoecology.—B, E, S, Epi, Sed; By. Extant *Pteria* species often live attached to corals, usually by a strong byssus (S. M. Stanley, 1970, 1972). In many fossil specimens, the byssal notch is present (e.g., Damborenea, 1987b, *Pteroperna* sp.) and the shell morphology is similar enough to living species shells to assume they had a similar mode of life. They are often found forming groups of several individuals, probably as the result of a gregarious mode of life, as happens in modern species (S. M. Stanley, 1970). *Mineralogy.*—Bimineralic (Carter, 1990b, p. 336, for living species). Outer shell layer: calcite (simple prismatic). Middle and inner shell layers: aragonite (nacreous).

Genus ARCAVICULA Cox, 1964, p. 47

Type species.—Avicula arcuata Münster in Goldfuss, 1835 in 1833–1841, p. 128.

Stratigraphic range.-Lower Triassic (lower Olenekian)-Upper Triassic (?Rhaetian) (Sha & Grant-Mackie, 1996; Newton in Newton & others, 1987). Although many authors assigned it a Middle Triassic (Ladinian)-Upper Triassic (Carnian) range (Cox & others, 1969; Hallam, 1981; Sepkoski, 2002; Tëmkin, 2006), Arcavicula was also mentioned from the Lower Triassic (Sha & Grant-Mackie, 1996) and with some uncertainty from the Norian (Newton in Newton & others, 1987). Newton (in Newton & others, 1987) provisionally referred her specimens to Arcavicula sp. due to the hinge details, but she related them to Rhaetavicula on account of their external similarity. It is also evident that some species were attributed to Pteria regardless of their internal characters. Newton (1988) later confirmed this reference. Laws (1982) mentioned but did not figure Arcavicula sp. from Upper Triassic (upper Norian = Rhaetian, according to Dagys & Dagys, 1994) from Nevada. There are no specific Middle Triassic Arcavicula records, although several authors (e.g., Cox & others, 1969; Hallam, 1981; Sepkoski, 2002; Tëmkin, 2006) mentioned this genus among their materials.

Paleogeographic distribution.—Tethys and Circumpacific (Fig. 13).

Tethys domain: Early Triassic: early Olenekian of China (Sha & Grant-Mackie, 1996); Late Triassic: Carnian of southern Alps and Apennines (Italy) (Broglio-Loriga, Ietto, & Posenato, 1993), Alps and Sicily (Diener, 1923; Kutassy, 1931), early Carnian of Lombardy (Italy) (Allasinaz, 1966), southern Alps (Italy) (Bittner, 1895); Norian of ?China (Kobayashi & Tamura, 1983a).

Circumpacific domain: Late Triassic: Norian of Oregon (United States) (Newton & others, 1987; Newton, 1988); Rhaetian of ?Nevada (United States) (Laws, 1982).

Paleoautoecology.—B, E, S, Epi, Sed; By. The presence of anterior auricle and byssal sinus in some specimens, and their external morphology, indicate that species of this genus most likely lived epibyssate or shallowly buried in the sediment in the adult stage, by comparison with living Pterioida (Newton in Newton & others, 1987).

Mineralogy.—Bimineralic (Carter, 1990b, p. 335). Outer shell layer: calcite (simple prismatic). Middle and inner shell layers: aragonite (nacreous).

Genus RHAETAVICULA Cox, 1962, p. 594

Type species.—Avicula contorta Portlock, 1843, p. 126.

Remarks.—Cox (1962) pointed out the similarities between *Rhaetavicula* and *Oxytoma*, and he proposed that *Rhaetavicula* could even be a member of the family Oxytomidae, but *Rhaetavicula* lacks the deep byssal groove located under the right anterior auricle, which is typical of that family. Based on the information provided by the shell mineralogy of *Rhaetavicula* (calcitic outer layer and alleged aragonitic inner layer), it is probably referable to Pteriidae. Nevertheless, if further studies confirm an inner calcitic layer, it should be referred to the Oxytomidae instead (Cox, 1962). There are no

studies on this subject (Carter, 1990a). Previous to 1962, when Cox described this genus, the type species of *Rhaetavicula* was assigned to different genera: *Avicula, Pseudomonotis, Cassianella,* and *Pteria*.

Stratigraphic range.—Upper Triassic (Rhaetian). *Rhaetavicula contorta* is a Rhaetian guide fossil (see references in paleogeographic distribution). During that stage, it was widely distributed, especially in the Tethys domain.

Paleogeographic distribution.—Tethys and Circumpacific (Fig. 13). Though *Rhaetavicula* was reported from the Austral domain (New Zealand), Cox (1962, p. 594) referred this record to *Oxytoma*. Al-though Hallam (1981, 1990) also listed it from Austral regions, the record will not be taken into account here because this information could not be confirmed.

Tethys domain: Late Triassic: Rhaetian of England (Cox, 1962; Castell & Cox, 1975; Warrington & Ivimey-Cook, 1990; Ivimey-Cook & others, 1999; Wignall & Bond, 2008), Italy (Allasinaz, 1962; Sirna, 1968; Bice & others, 1992; McRoberts, 1994), Hungary (Vörös, 1981), Burma (Vu Khuc & Huyen, 1998), southern Tibet (Hallam & others, 2000; J. Yin & Enay, 2000), Iran (Hautmann, 2001b), western Carpathians (Slovakia) (Tomašových, 2004; Michalík & others, 2007), Alps (Austria) (Tomašových, 2006a, 2006b; McRoberts, 2010), Spain (Goy & Márquez-Aliaga, 1998).

Circumpacific domain: Late Triassic: Rhaetian of Nevada (United States) (Cox & others, 1969; Hallam & Wignall, 2000).

Paleoautoecology.—B, E, S, Epi-Un, Sed; By-R. According to Cox (1962), *Rhaetavicula* lacked a byssal notch, and he assumed that the byssus emerged between the two valves by a narrow gape. Since the shell is strongly inequivalve (convex left valve and flat right valve) and by similarities to living Pterioida, it probably had an epibyssate mode of life. Another possibility is that the byssus was atrophied in adults (and thus the byssal notch is absent), and then it would live reclined on its left valve, similar to members of the family Cassianellidae (Hautmann, 2001b).

Mineralogy.—Bimineralic (Carter, 1990a, p. 206). Outer shell layer: calcite (prismatic). Middle and inner shell layers: aragonite (?).

Genus STEFANINIA Cox in Cox & others, 1969, p. 306 [ex Venzo, 1934, p. 165]

Type species.—Gervilleia? ogilviae Bittner, 1895, p. 88.

Remarks.—Stefaninia was named by Venzo (1934), but his description did not fulfill the nomenclatorial rules (ICZN Code, 1999), since no type species was assigned and no diagnostic features were given (Stenzel, 1971, p. 1215). Cox in Cox and others (1969, p. 306) designated the type species and provided its diagnosis.

Stratigraphic range.—Middle Triassic (upper Ladinian)–Upper Triassic (Carnian). Bittner (1895) described the type species from the Saint Cassian Formation, regarded as Carnian in age (Fürsich & Wendt, 1977). Cox and others (1969) assigned a Ladinian age, probably on the basis of Venzo's paper (1934; see above).

Paleogeographic distribution.—western Tethys (Fig. 13).

Tethys domain: Middle Triassic: late Ladinian of Italy (Cox & others, 1969); Late Triassic: Carnian of Italy (Bittner, 1895).

Paleoautoecology.—B, E, S, Epi, Sed; By. Its external morphology and the presence of byssal notch indicate a probable byssate mode of life. Possibly, like other members of the family, it spent the early

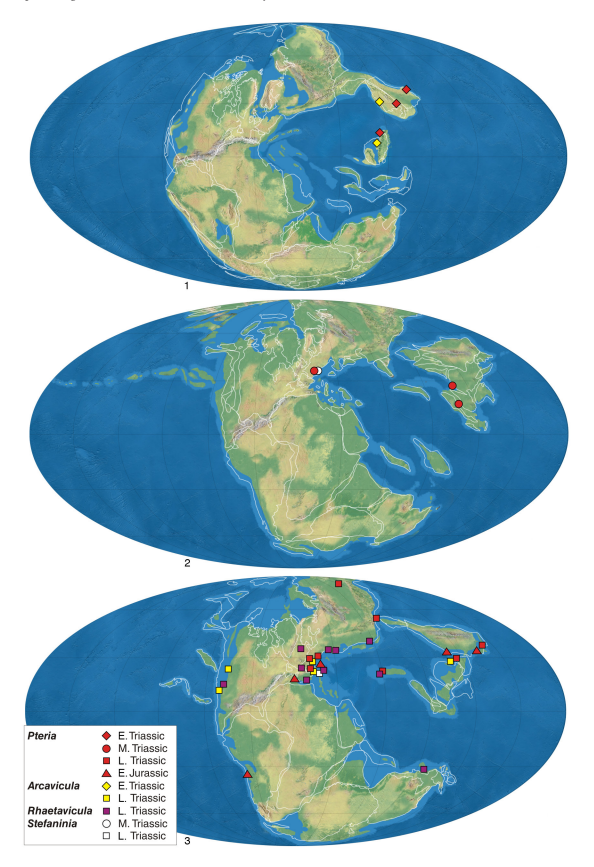


Figure 13. Paleogeographical distribution of Pteriidae (Pteria, Arcavicula, Rhaetavicula, Stefaninia). 1, Early Triassic; 2, Middle Triassic; 3, Late Triassic-Early Jurassic.

stages of development fixed by the byssus, but adults could have lived partially buried in the sediment. There is no information about the type of sediment in which it was found.

Mineralogy.-Bimineralic (Carter, 1990a, p. 206). There is no information about Stefaninia shell mineralogy or microstructure. We use data provided for the family Pteriidae. Outer shell layer: calcite (simple prismatic). Middle and inner shell layer: aragonite (nacreous).

Family BAKEVELLIIDAE King, 1850 Genus BAKEVELLIA King, 1848, p. 10

Neave (1939, p. 385) lists "Miller, 1877, p. 185, pro Bakewellia King, 1848" as author of the genus. Nevertheless, Miller (1877, p. 185) did not indicate he was proposing either an emendation or a replacement name, and already in 1850 King (p. 166-171, 255) spelled it consistently as Bakevellia, although he stated (p. 166, footnote) that the name was dedicated to Mr. Bakewell.

Type species.—Avicula antiqua Münster in Goldfuss, 1835 in 1833–1841, p. 126, non Defrance, 1816.

Remarks.—Several subgenera were proposed within Bakevellia (see Damborenea, 1987b, p. 125-126), but Muster (1995) regarded almost all to be synonyms, considering only two of them to be valid, as also did Cox and others (1969): B. (Bakevellia) King, 1848, and B. (Bakevelloides) Tokuyama, 1959a. The subgenera described for our study interval were Neobakevellia Nakazawa, 1959, Integribakevellia Farsan, 1972, Costibakevellia Farsan, 1972, and Spia Skwarko, 1981 (see list of synonyms for both subgenera in Muster, 1995, p. 29, 42).

Stratigraphic range.-upper Permian-Upper Cretaceous. Cox and others (1969) assigned it a Permian–Upper Cretaceous range. Muster (1995) maintained this range, noting that the first record of the genus is dated as upper Permian. Sepkoski (2002) considered the oldest record to be Carboniferous, but we will not take this into account, since it is based on a personal communication by Yancey to Sepkoski (indicated in Sepkoski, 2002), which has not been published.

Paleogeographic distribution.—Cosmopolitan.

Paleoautoecology.-B, Se, S, Endo, Sed; By. The mode of life of bakevelliids is difficult to identify as that they do not have living representatives to compare with, and the study of the morphology alone does not always provide good results, because morphology traits are sometimes contradictory. It is also helpful to interpret the paleoecology of the environments in which the specimens are found. Most species assigned to *Bakevellia* are almost equivalve, they have a shallow byssal sinus and an anterior lobe, features that indicate an endobyssate way of life, living with the sagittal plane almost vertical (S. M. Stanley, 1972). This interpretation was proposed by Damborenea (1987b) for Bakevellia (Neobakevellia?) pintadae Damborenea, 1987b, and by Aberhan and Muster (1997) for Bakevellia (Bakevellia) waltoni (Lycett, 1863). However, Seilacher (1984) interpreted Bakevellia subcostata (Goldfuss, 1835 in 1833-1841) as reclined and partially buried in the sediment, resting on its left valve, with the commissure plane being almost horizontal.

Mineralogy.—Bimineralic (Márquez-Aliaga & Martínez, 1990a; Carter, 1990b). Outer shell layer: calcite (simple prismatic). Middle and inner shell layers: aragonite (nacreous).

Remarks.—Fürsich and Werner (1988, p. 112) argued that there are no substantial differences between Gervillia Defrance, 1820, and Gervillella to consider them as independent taxa, and they included Gervillella as a subgenus of Gervillia. We follow Freneix (1965) and Muster (1995) in treating them as separated genera.

Stratigraphic range.-Lower Jurassic (Hettangian)-Upper Cretaceous (?) (Aberhan, 1998a; Muster, 1995). Several authors extended the range back to the Triassic (e.g., Gillet, 1924; Hayami, 1957a; Freneix 1965; Cox & others, 1969; Geyer, 1973; Lazo, 2003), but none of them justified this statement, and they did not figure any Triassic specimens. All except Geyer (1973) simply listed the stratigraphic range of several genera. Gever mentioned the presence of Gervillella sp. from the Norian Payandé Formation in Colombia, but he did not figure it. It is possible that some Triassic species assigned to Gervillia should be referred to Gervillella instead, but there is no published reference of their presence in this period. The oldest confirmed record dates from the Hettangian (Aberhan, 1998a), and the youngest from the Upper Cretaceous (Muster, 1995). Sepkoski (2002) assigned the last appearance to the Maastrichtian, but it was not possible to see the original data source. Muster (1995) did not specify the stage, and there is no further information about this topic. However, it is not uncommon to find the genus mentioned from the Lower Cretaceous (Lazo, 2003, 2007a).

Paleogeographic distribution .- western Tethys, Austral, and Circumpacific (Fig. 14). The genus had a particularly wide distribution mainly during the Middle and Late Jurassic (Vörös, 1971; Fürsich & Werner, 1988; Liu, 1995; Muster, 1995; Sha & Grant-Mackie, 1996; Delvene, 2003; Sha, Johnson, & Fürsich, 2004).

Tethys domain: Early Jurassic: Hettangian–Sinemurian of England and Morocco (Liu, 1995).

Austral domain: Early Jurassic: Hettangian–Sinemurian of Southern Andes (Damborenea, 1996a; Damborenea & Lanés, 2007).

Circumpacific domain: Early Jurassic: Hettangian-Sinemurian of Canada (Aberhan & Muster, 1997; Aberhan, 1998a); Sinemurian of Chile (Aberhan, 1994a).

Paleoautoecology .- B, Se, S, Endo, Sed; By. Like most bakevelliids, Gervillella presents features that indicate a semi-infaunal endobyssate mode of life (S. M. Stanley, 1972; Aberhan & Muster, 1997) or so-called mud-sticker (Seilacher, 1984). All species assigned to this genus are almost equivalve, they possess an anterior auricle, and their external morphology is elongate spear-shaped. Thanks to its elongated shape, Gervillella could probably bury deeper than other bakevelliids (S. M. Stanley, 1972; Aberhan & Muster, 1997), similar to members of the family Pinnidae (S. M. Stanley, 1972). Although neither Damborenea (1987b) nor Aberhan and Muster (1997) found evidence in their specimens of a byssal notch, according to Cox (1940), one of the characters that defines the genus is that the anterior auricle extends anteroventrally and is limited in the left valve by a deep groove, which indicates the position of the byssus.

Mineralogy.—Bimineralic (Carter, 1990a, p. 207; Carter, 1990b, p. 336). Outer shell layer: calcite (simple prismatic). Middle and inner shell layers: aragonite (nacreous).

Genus GERVILLELLA Waagen, 1907, p. 98

Type species.—Perna aviculoides J. Sowerby, 1814, p. 147.

Genus GERVILLIA Defrance, 1820, p. 502

Type species.—Gervillia solenoidea Defrance, 1824a, p. 316.

Stratigraphic range.—Middle Triassic (Ladinian)–Upper Cretaceous (Maastrichtian) (Lerman, 1960; Abdel-Gawad, 1986). Both Cox and others (1969) and Muster (1995) indicated its range as beginning at the Upper Triassic, but there are Middle Triassic records of Gervillia, referred to the species G. joleaudi (Schmidt, 1935) from the Anisian of Israel (Lerman, 1960) and Ladinian of Spain (Márquez-Aliaga, 1985). These were not included in Muster's monograph (1995), but Waller and Stanley (2005) indicated that the generic assignation of this species requires revision. However, these authors based their opinion, exposed in the discussion of their new subgenus Gervillaria (Baryvellia), in data from Schmidt (1935), who compared G. joleaudi with Gervillia alberti Credner, 1851. According to Márquez-Aliaga (1985), this last species is a true Bakevellia; therefore, Gervillia joleaudi should be considered as a representative of Gervillia from the Sephardic province of the Tethys domain. With regard to the uppermost stratigraphic occurrence, all agreed that the genus disappeared in the Upper Cretaceous. Within our study interval, we will only consider the subgenus Cultriopsis Cossmann, 1904. Boyd and Newell (1979) doubtfully assigned some of their specimens from the Permian of Tunisia to this subgenus.

Paleogeographic distribution.—Tethys and Circumpacific (Fig. 14). Although Escobar (1980) reported *Gervillia* from Hettangian—Sinemurian beds of Chile, only one of the specimens could be attributed with doubt to this genus (Damborenea, 1987b), so it will not be taken into account in the Austral domain in this temporal range. If the genus is present in this domain, it has occurred since the Pliensbachian.

Tethys domain: Middle Triassic: Anisian of Israel (Lerman, 1960); Ladinian of Spain (Márquez-Aliaga, 1983, 1985; Márquez-Aliaga, Hirsch, & López-Garrido, 1986; Márquez-Aliaga & Martínez, 1990a, 1996; Budurov & others, 1991; Márquez-Aliaga & Montoya, 1991; Martinez & Márquez-Aliaga, 1994; Niemeyer, 2002; Márquez-Aliaga & Ros, 2003); Late Triassic: China (Muster, 1995); Carnian of Italy (Fürsich & Wendt, 1977; Muster, 1995), Spain (Martín-Algarra & others, 1993), Slovenia (Jurkovsek, 1978), China (Wen & others, 1976); Norian–Rhaetian of Iran (Hautmann, 2001b); Rhaetian of Austria (Tanner, Lucas, & Chapman, 2004); Early Jurassic: early Hettangian of Tibet (China) (J. Yin & McRoberts, 2006); Hettangian–Sinemurian of Vietnam (Sato & Westermann, 1991).

Circumpacific domain: Late Triassic: Japan (Muster, 1995); Carnian of Japan (Tamura & others, 1978); Norian of Oregon (Newton, 1986; Newton & others, 1987); Early Jurassic: Hettangian of Japan (Hayami, 1957a, 1964, 1975; Muster, 1995).

Paleoautoecology.—B-Ps, Se-E, S, Endo-Epi, Sed; By. Some species of *Gervillia* are morphologically similar to species of *Gervillela* and *Gervillancea* (see discussion on their mode of life, p. 40, 44). These two genera are interpreted as having a semi-infaunal endobyssate mode of life (Waller & Stanley, 2005). Formerly, Muster (1995) regarded this mode of life to be unlikely for *Gervillia*, since it had a very short ligament area that would not be enough to maintain the shell stability. Seilacher (1984) suggested a pseudoplanktonic mode of life for some species of *Gervillia*, as epibyssate on ammonoids. He called these forms pendent forms. However, other species of *Gervillia* were interpreted as semi-infaunal endobyssate or mud-stickers (Seilacher, Matyja, & Wierzbowski, 1985). These interpretations are

based on the external shell morphology and on the ecological analysis of the depositional environment in which the specimens were found (see Seilacher, 1984; Seilacher, Matyja, & Wierzbowski, 1985). In the Muschelkalk of the Iberian Range (Spain), specimens recorded in marls are common, and they are usually found in semi-infaunal life position. Newton (in Newton & others, 1987) and Damborenea (1987b) interpreted their specimens as epibyssate, but they noted that the shells also had features indicative of a semi-infaunal habit. These are usually found associated with corals.

Mineralogy.—Bimineralic (De Renzi & Márquez-Aliaga, 1980; Carter, 1990a; Márquez-Aliaga & Martínez, 1990a; Martínez & Márquez-Aliaga, 1994). Outer shell layer: calcite (simple prismatic). Middle and inner shell layers: aragonite (nacreous).

Genus HOERNESIA Laube, 1866, p. 52

Type species.—Mytulites socialis Schlotheim, 1823 in 1822–1823, p. 112.

Stratigraphic range.—Lower Triassic (Olenekian)–Upper Triassic (Rhaetian) (Hallam, 1981; Dagys & Kurushin, 1985). Cox and others (1969) indicated a Triassic–Middle Jurassic range, although some authors believed *Hoernesia* disappeared in the Rhaetian (Hallam, 1981, 1990; Hallam & others, 2000). However, Muster (1995, p. 89) extended its range to the Middle Jurassic, because she included *Gervillia radians* Morris & Lycett, 1853 in 1851–1885, in the synonymy list of *Hoernesia socialis* (Schlotheim, 1823 in 1822–1823); besides, she did not consider *Hoernesia* to be present in the Early Triassic. The first record of *Hoernesia* dates from the Early Triassic (Dagys & Kurushin, 1985; Posenato, 2008a).

Paleogeographic distribution.—Tethys and Boreal (Fig. 14).

Tethys domain: Early Triassic: Italy (Neri & Posenato, 1985), Yunnan (China) (Guo, 1985); Middle Triassic: Bulgaria (Stefanov, 1942; Encheva, 1969), Spain (Via, Villalta, & Esteban, 1977; Márquez-Aliaga, 1983, 1985; Márquez-Aliaga & Martínez, 1996; Márquez-Aliaga & others, 2001, 2002, 2004; Márquez-Aliaga & Ros, 2002), Italy (Posenato, 2002; Posenato & others, 2002), Germany (Fuchs & Mader, 1980; Hagdorn, 1982; Hagdorn & Simon, 1983, 1991), Poland (Senkowiczowa, 1985; Kaim, 1997), Hungary (Szente, 1997); Anisian of China (Sha, Chen, & Qi, 1990); Anisian-Ladinian of northern Vietnam (Komatsu, Huyen, & Huu, 2010); Ladinian of the Alps (Austria and Italy) (Arthaber, 1908), Germany (Ürlichs, 1978); Late Triassic: China (Cowper-Reed, 1927), Malaysia (Tamura & others, 1975); Carnian of Italy (Laube, 1865), Slovenia (Jurkovsek, 1978); Norian of China (Lu, 1981); Norian-Rhaetian of Iran (Hautmann, 2001b); Rhaetian of Burma (Healey, 1908), Iran (Repin, 2001), Tibet (J. Yin & Enay, 2000).

Boreal domain: Early Triassic: Olenekian of northern Siberia (Dagys & Kurushin, 1985).

Paleoautoecology.—B, Se, S, Endo, Sed; By. *Hoernesia* is characterized by a strongly inequivalve shell and twisted valves, and thus it was interpreted as a so-called twisted recliner by Seilacher (1984). It also has an umbonal shell thickening, so its life position consisted of this area being introduced into the sediment, with the posterior part of the valves sticking out (Savazzi, 1984; Seilacher, 1990; Muster, 1995). The inferred life position is similar to that of *Gervillaria alaeformis* (J. Sowerby, 1819) (see discussion about the mode of life of this species, below). Seilacher (1990) suggested chemosymbiosis as a functional explanation for this curious life position.

Mineralogy.—Bimineralic (Carter, 1990b, p. 337). Outer shell layer: calcite (simple prismatic). Middle and inner shell layers: aragonite (nacreous).

Genus LANGSONELLA Patte, 1926, p. 139

Type species.—Gervilleia (Cultriopsis) elongata Mansuy, 1919, p. 7. Stratigraphic range.—Middle Triassic (Anisian)–Upper Triassic (Carnian) (Diener, 1923; Komatsu, Huyen, & Huu, 2010). Cox and others (1969) referred this genus to the Triassic without further explanation and indicated it is monospecific. According to Diener (1923), the type species was described by Mansuy from the Carnian of Tonkin, which today covers most of Vietnamese northern regions. Later, Vu Khuc and Huyen (1998) mentioned *L. elongata* (Mansuy, 1919) as being typical from Ladinian beds in the same area, and, recently, Komatsu, Huyen, and Huu (2010) reported it from the Anisian and Ladinian of northern Vietnam.

Paleogeographic distribution.—Eastern Tethys (Fig. 14).

Tethys domain: Middle Triassic: Anisian–Ladinian of northern Vietnam (Komatsu, Huyen, & Huu, 2010); Ladinian of Tonkin (north of Vietnam) (Vu Khuc & Huyen, 1998); Late Triassic: Carnian of Tonkin (Vietnam) (Diener, 1923).

Paleoautoecology.—B, Se, S, Endo, Sed; By. We did not find any figures of the genus; therefore, it is difficult for us to refer it to a particular mode of life, but according to its description in Cox and others (1969), we consider it to be similar to *Hoernesia*.

Mineralogy.—Bimineralic (Carter, 1990a, p. 206–207). There are no data on *Langsonella* shell structure. We used data provided for the family Bakevelliidae. Outer shell layer: calcite (simple prismatic). Middle and inner shell layers: aragonite (nacreous).

Genus CUNEIGERVILLIA Cox, 1954, p. 48

Type species.—Gervillia hagenowii Dunker, 1846, p. 37.

Remarks.—Cox (1954) described Cuneigervillia and included Edentula Waagen, 1907, p. 96 (non Nitzch, 1820, p. 189) as its synonym. Later, in Cox and others (1969), he regarded Edentula [= Waagenorperna Tokuyama, 1959a, p. 151] as a separate genus and included it into the Isognomonidae. In turn, Tokuyama (1959a) proposed the name Waagenoperna to replace Edentula Waagen, 1907. He pointed to significant differences between the *Cuneigervillia* type species designated by Cox (1954) (Gervillia hagenowii Dunker, 1846) and some species attributed to Edentula (E. lateplanata Waagen, 1907, and E. triangularis Kobayashi & Ichikawa, 1952). He designated Edentula lateplanata as the type species of Waagenoperna, maintaining the two genera as distinct taxa, relating G. hagenowii to the Bakevelliidae and E. lateplanata and E. triangularis to the Isognomonidae. Muster (1995) decided to include *Cuneigervillia* with the Isognomonidae, believing that although Cuneigervillia externally seems to be a bakevelliid, it possesses certain characteristics that are typical of the Isognomonidae, such as terminal or subterminal beaks and a toothless adult hinge. It is difficult to decide because both families share many characteristics, but the lack of teeth in the adult stage is not a critical feature because it also occurs in certain bakevelliids; for example, in some species of Bakevellia (Bakevellia) the dentition is obsolete in adults (Cox & others, 1969, p. 306). The Treatise diagnosis states "hinge teeth present at least in lower growth stages" Regarding the beaks, they can either be subterminal (e.g., *Aguilerella*) or terminal (e.g., *Gervillia*). Furthermore, *Cuneigervillia* presents the typical teeth of *Bakevellia* in juvenile stages. Therefore, according to Cox and others (1969), we include *Cuneigervillia* in the Bakevelliidae.

Stratigraphic range.—Lower Jurassic (Hettangian)–Lower Cretaceous (?) (Cox & others, 1969). Cox and others (1969) assigned it a lower Liassic to Lower Cretaceous range, since Tokuyama (1959a) referred several Carnian species to *Waagenoperna* that were initially assigned by Cox (1954) to *Cuneigervillia*.

Paleogeographic distribution.-western Tethys (Fig. 14).

Tethys domain: Early Jurassic: Europe and northern Africa (Cox & others, 1969); Hettangian of France (Freneix & Cubaynes, 1984), south of England (Warrington & Ivimey-Cook, 1990), Spain (Gómez, Goy, & Márquez-Aliaga, 2005; Márquez-Aliaga, Damborenea, & Goy, 2008a, 2008b; Márquez-Aliaga & others, 2010); Hettangian–Pliensbachian of northwestern Europe (Hallam, 1987); Sinemurian of Portugal (Liu, 1995).

Paleoautoecology.—B, Se, S, Endo, Sed; By. *Cuneigervillia* was interpreted as a semi-infaunal endobyssate bivalve (S. M. Stanley, 1972), as were most members of Bakevelliidae.

Mineralogy.—Bimineralic (Carter, 1990a, p. 206–207). There are no data on *Cuneigervillia* mineralogy and shell microstructure. Data provided for the family Bakevelliidae are used here. Outer shell layer: calcite (simple prismatic). Inner shell layer: aragonite (nacreous).

Genus GERVILLARIA Cox, 1954, p. 49

Type species.—Modiola? alaeformis J. Sowerby, 1819, p. 93.

Stratigraphic range.—Middle Triassic (Anisian)–Upper Cretaceous (Turonian) (Komatsu, Chen, & others, 2004; Muster, 1995). Cox and others (1969) assigned this genus a Jurassic–Cretaceous range in Europe, but, since then, new records have extended its stratigraphic range. The oldest record is Anisian (Komatsu, Chen, & others, 2004) and the youngest is Turonian (Muster, 1995).

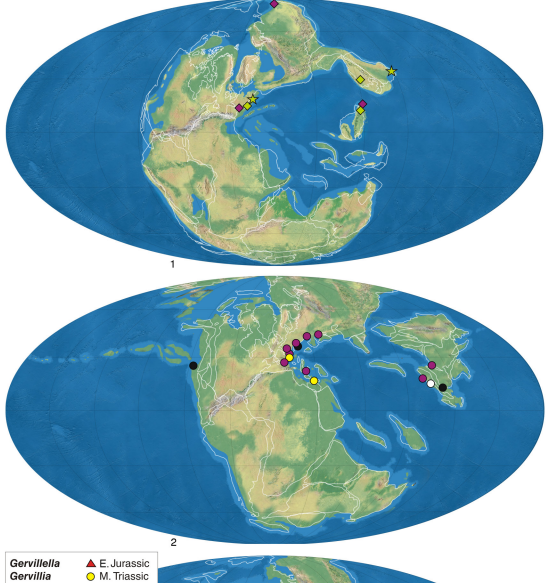
Paleogeographic distribution.—Tethys, Austral, and Circumpacific (Fig. 14).

Tethys domain: Middle Triassic: Muschelkalk of Germany (Muster, 1995); Anisian of Qingyan (southern China) (Komatsu, Chen, & others, 2004); Late Triassic: southwestern China (Komatsu, Chen, & others, 2004); Rhaetian of Lombardy (Italy) (Muster, 1995), Italian Alps and Vietnam (Hautmann, 2001b), western Carpathians (Slovakia) (Tomašových, 2004), Tibet (China) (J. Yin & Grant-Mackie, 2005); Norian–Rhaetian of Iran (Hautmann, 2001b).

Austral domain: Early Jurassic: Sinemurian of the Andean Basin (Aberhan & Fürsich, 1997); Hettangian–Sinemurian of the Andean Basin (Damborenea & Manceñido, 2005b).

Circumpacific domain: Middle Triassic: Ladinian of western Nevada (Waller & Stanley, 2005); Late Triassic: Norian of southeastern Sonora (Mexico) (McRoberts, 1997a); Early Jurassic: Sinemurian of ?western Canada (Aberhan, 1998a), Chile (Aberhan, 1994a).

Paleoautoecology.—B, Se, S, Endo, Sed; By. Some species, such as *Gervillaria alaeformis* (J. Sowerby, 1819) (Muster, 1995, fig. 37) and *Gervillaria pallas* (A. F. Leanza, 1942) (Damborenea, 1987b, fig. 7; Muster, 1995, fig. 43), have a strongly inequivalve and inequilateral shell, with the left valve being more convex



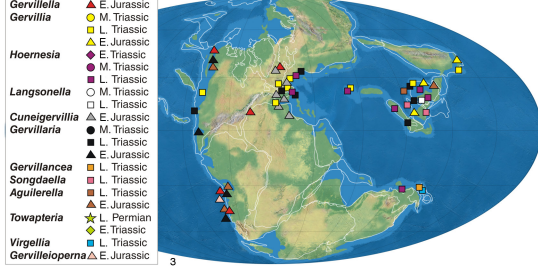


Figure 14. Paleogeographical distribution of Bakevelliidae (Gervillela, Gervillia, Hoernesia, Langsonella, Cuneigervillia, Gervillaria, Gervillancea, Songdaella, Aguilerella, Towapteria, Virgellia, Gervilleioperna). 1, Early Triassic; 2, Middle Triassic; 3, Late Triassic–Early Jurassic.

than the right, twisted valves, and an elongated posterior auricle. These species were interpreted as having a semi-infaunal endobyssate mode of life, probably supplemented by byssal attachment (Damborenea, 1987b; Aberhan & Muster, 1997). Seilacher (1984, fig. 7) proposed an analogous interpretation for a similar species, Hoernesia tortuosa, including it as a twisted recliner. This category was also used by Aberhan and Muster (1997) for their specimens of G. pallas. Gervillaria (Baryvellia) ponderosa Waller in Waller & Stanley, 2005, was also considered semi-infaunal endobyssate, but this species had a peculiar external morphology, which probably means that its life position was also special (Waller & Stanley, 2005) (see discussion on Gervillancea mode of life, below). However, due to the mytiliform appearance of some species referred to Gervillaria, these were interpreted as epibyssate (S. M. Stanley, 1972). Gervillaria ashcroftensis (Crickmay, 1930a) (see Muster, 1995, fig. 39) was also thought to be epibyssate, according to its nearly equivalve shell, umbonal thickening, and flat anteroventral area, among other characteristics (see Aberhan & Muster, 1997).

Mineralogy.—Bimineralic (Carter, 1990a, p. 206-207). There are no data on *Gervillaria* mineralogy or shell microstructure. Data provided for family Bakevelliidae are used here. Outer shell layer: calcite (simple prismatic). Inner shell layer: aragonite (nacreous).

Genus GERVILLANCEA Skwarko, 1967, p. 54

Type species.—Gervillancea coxiella Skwarko, 1967, p. 54.

Stratigraphic range.—Upper Triassic (Carnian–Norian) (Skwarko, 1967). *Gervillancea* is a monospecific genus only known from Upper Triassic of New Guinea (Skwarko, 1967; Muster, 1995; Waller & Stanley, 2005). Although it was described before the publication of the *Treatise* (Cox & others, 1969), it was included neither there nor in Sepkoski (2002).

Paleogeographic distribution.—Southern Tethys (Fig. 14). Gervillancea was endemic in the Australian province (according to Damborenea, 2002b) of the Tethys domain. It was only reported from Papua New Guinea (Skwarko, 1967).

Paleoautoecology.-B, Se, S, Endo, Sed; By. One of the most striking features of this genus is its extremely long anterior auricle, which distinguishes it from almost all other genera of Bakevelliidae. According to Waller and Stanley (2005), there are two species, Gervillaria (Baryvellia) ponderosa Waller in Waller & Stanley, 2005, and Gervillia joleaudi (Schmidt, 1935), that also have this feature. These two species, together with Gervillancea coxiella, may be a good example of evolutionary convergence, but, in fact, Gervillia joleaudi lacks an anterior auricle. The external shape of Gervillancea is very asymmetric and inequivalve. None of the specimens figured by Skwarko (1967) bears a byssal notch, but if the species was byssate, like other bakevelliids, the byssus probably emerged from the shell under the anterior auricle. Taking into account that it probably lived anchored to the substrate with the anterior auricle, a strong byssus was not necessarily needed to maintain stability inside the substrate. Pedal and byssal muscle scars indicate that these muscles were strong and able to aid the shell to penetrate up to a third of its dorsal line into the sediment, since the convexity of the shell increases at this point and thus limits the burial depth (see Waller & Stanley, 2005, p. 27-29). The bivalve most probably introduced

itself into the sediment during the juvenile stages, since the anterior auricle is comparatively thin and thus inadequate to penetrate the sediment in the adult stage.

Mineralogy.—Bimineralic (Carter, 1990a, p. 206–207). No data are available about the shell of *Gervillancea*, but it was probably bimineralic, as in other Bakevillidae (J. D. Taylor, Kennedy, & Hall, 1969). Outer shell layer: calcite (simple prismatic). Inner shell layer: aragonite (nacreous).

Genus SONGDAELLA Vu Khuc, 1977b, p. 50 [180]

Type species.—Songdaella graciosa Vu Khuc, 1977b, p. 51 [182]. Remarks.—Vu Khuc (1977b) assigned Songdaella to the Bakevelliidae, but he indicated that the genus had intermediate characters between this family and the Isognomonidae. Muster (1995) did not include it in her monograph about the family Bakevelliidae and did not comment about its systematic position. In the absence of more information, we include Songdaella in Bakevelliidae.

Stratigraphic range.—Upper Triassic (Norian) (Vu Khuc, 1977b). *Songdaella* was only recorded from Norian beds (Vu Khuc, 1977b; J. Chen, 1982a; Vu Khuc & Huyen, 1998).

Paleogeographic distribution.—Eastern Tethys (Fig. 14). *Songdaella* was endemic to southern East Asia (Vu Khuc & Huyen, 1998).

Tethys domain: Norian of northern Vietnam (Vu Khuc, 1977b) and southern China (J. Chen, 1982a).

Paleoautoecology.—B, E, S, Epi, Sed; By. *Songdaella* is characterized by a mytiliform shell, and some of the specimens figured by Vu Khuc (1977b) are similar to *Mytilus*. The author related his new genus to *Aguilerella* according to its external morphology, which was interpreted as epibyssate by S. M. Stanley (1972), due to its external similarity with *Mytilus* and *Myalina*.

Mineralogy.—Bimineralic (Carter, 1990a, p. 206–207). There are no data about the shell of *Songdaella*. We use the data predominant in the family Bakevelliidae. Outer shell layer: calcite (simple prismatic). Inner shell layer: aragonite (nacreous).

Genus AGUILERELLA Chavan, 1951, p. 211

Type species.—Perna kobyi de Loriol, 1901, p. 99.

Stratigraphic range.—Upper Triassic (Rhaetian)–Lower Cretaceous (Hauterivian) (J. Yin & McRoberts, 2006; Kozai, Ishida, & Kondo, 2006).

Many authors restricted it to a Lower Jurassic–Upper Jurassic range (Cox & others, 1969; Muster, 1995; Sepkoski, 2002). This range was extended due to new records from the Rhaetian of Tibet (J. Yin & McRoberts, 2006) and from the Hauterivian (Kozai, Ishida, & Kondo, 2006).

Paleogeographic distribution.—Eastern Tethys and Circumpacific (Fig. 14). Although during the study interval it was only reported from eastern Tethys and Austral domains, from Toarcian times it extended also to western Tethys and Boreal regions (see Fürsich, 1982; Liu, 1995; Muster, 1995; J. Yin & Grant-Mackie, 2005; Zakharov & others, 2006).

Tethys domain: Late Triassic: Rhaetian of Tibet (China) (J. Yin, H. Yao, & Sha, 2004; J. Yin & McRoberts, 2006); Early Jurassic: Hettangian of China (J. Chen & Liu, 1981; J. Yin & McRoberts, 2006). Circumpacific domain: Early Jurassic: Sinemurian of Chile (Aberhan, 1994a; Aberhan & Fürsich, 1997), Canada (Poulton, 1991); Hettangian–Sinemurian of South America (Damborenea, 1996a).

Paleoautoecology.—B, E, S, Epi, Sed; By. Due to its mytiliform aspect, it was thought to be epibyssate (S. M. Stanley, 1972). Aguilerella is one of the few bakevelliids, together with Songdaella, that are interpreted to be epibyssate due to their triangular form, without anterior lobe and with terminal beaks (Damborenea, 1987b). In some species, a gregarious behavior was observed (Fürsich, 1982; Damborenea, 1987b).

Mineralogy.—Bimineralic (Carter, 1990a, p. 206–207). There are no specific data about the shell of *Aguilerella*. Data provided for the family Bakevelliidae. Outer shell layer: calcite (simple prismatic). Inner shell layer: aragonite (nacreous).

Genus TOWAPTERIA Nakazawa & Newell, 1968, p. 59

Type species.—Towapteria nipponica Nakazawa & Newell, 1968, p. 59.

Stratigraphic range.-lower Permian (Sakmarian)-Lower Triassic (Induan) (Hayami & Kase, 1977; S. Yang, Wang, & Hao, 1986). Nakazawa and Newell (1968) proposed the genus Towapteria with material from the middle Permian of Japan. Cox and others (1969) did not take it into account in the Treatise, probably due to the proximity of publication. Hayami and Kase (1977) assigned it a Sakmarian-upper Permian range with some doubts. Towapteria was later reported from the Tethyan Early Triassic (see paleogeographic distribution below). Nevertheless, Muster (1995) assigned it an upper Permian, ?Upper Triassic, Middle Jurassic discontinuous range, due to the inclusion of some species previously assigned to Gervillia and Costigervillia (see synonymy list in Muster, 1995, p. 92). She did not see the material personally and the addition of most these species to the synonymy list was done doubtfully due to the lack of internal reliable characters for classification. Furthermore, Muster (1995) did not take into account some Tethyan, Early Triassic species, such as T. scythica (Wirth), among others.

Paleogeographic distribution.—Tethys and ?Circumpacific (Fig. 14). Tethys domain: late Permian: Changhsingian of Italy (Farabegoli, Perri, & Posenato, 2007); Early Triassic: Induan of Italy (Broglio-Loriga, Neri, & Posenato, 1980, 1986; Broglio-Loriga, Masetti, & Neri, 1982; Neri, Pasini, & Posenato, 1986; Broglio-Loriga & others, 1988, 1990; Posenato, 1988, 2008a), China (S. Yang, Wang, & Hao, 1986; L. Li, 1995; Tong & Yin, 2002; Waller & Stanley, 2005; Komatsu, Huyen, & Chen, 2007).

Circumpacific domain: late Permian: ?Japan (Nakazawa & Newell, 1968; Hayami & Kase, 1977).

Paleoautoecology.—B, E, S, Epi, Sed; By. It is hard to assign a specific life habit to *Towapteria*, because there are some features indicative of an epibyssate and others of an endobyssate mode of life. Due to its external similarity to *Costigervillia* Cox & Arkell, 1948 in 1848–1850 (a genus not included here because it first appeared in the Middle Jurassic), we can assume that *Towapteria* was endobyssate, but its smooth and lobate anterior auricle and radially ribbed shell indicate otherwise. We suggest it was byssate and probably lived with the anterior part introduced in the sediment.

The ribs probably helped to stabilize the shell, as was postulated for *Costigervilla* by Seilacher (1984).

Mineralogy.—Bimineralic (Carter, 1990a, p. 206–207). No data are available about the shell of *Towapteria*. Data provided for family Bakevelliidae. Outer shell layer: calcite (simple prismatic). Inner shell layer: aragonite (nacreous).

Genus VIRGELLIA Freneix, 1965, p. 61

Type species.—Virgellia coxi Freneix, 1965, p. 64.

Remarks.—Fürsich & Werner (1988) considered *Virgellia* as subgenus of *Gervillia* Defrance, 1820, since, in their opinion, it had intermediate features between *Gervillia* and *Gervillella* (which they also regarded as subgenus of *Gervillia*). Following Freneix (1965) and Muster (1995), we regard *Virgellia* as a separate genus. Freneix (1965) proposed *Virgellia* and originally included in it three species: *V. coxi* Freneix, 1965, *V. fittoni* (Sharpe, 1850), and *V. sobralensis* (Sharpe, 1850). Later, Muster (1995) added *V. simbaiana* (Skwarko, 1967) to *V. coxi* and *V. sobralensis* (see synonym list in Muster, 1995, p. 94–95).

Stratigraphic range.—Upper Triassic (Carnian)–Upper Jurassic (Kimmeridgian) (Muster, 1995). Cox and others (1969) and Sepkoski (2002) did not take *Virgellia* into account. The type species was originally described from Callovian sediments, and the original range assigned to the genus was Bajocian to Kimmeridgian (Freneix, 1965). Later, its range was extended by the inclusion of *V. simbaiana* by Muster (1995). The oldest record of *Virgellia* is from Carnian beds (Skwarko, 1967, 1981) and the youngest from Kimmeridgian beds (Freneix, 1965). It has not been recorded from the Lower Jurassic.

Paleogeographic distribution.—Southern Tethys (Fig. 14). Although during the interval of time under consideration it was only known from southern Tethys, during the Middle and Late Jurassic it was reported also from Tunisia (Freneix, 1965; Holzapel, 1998) and Portugal (Fürsich & Werner, 1988).

Tethys domain: Late Triassic: Carnian–Norian of Papua New Guinea (Skwarko, 1967, 1981; Muster, 1995).

Paleoautoecology.—B, Se, S, Endo, Sed; By. *Virgellia* was externally similar to *Gervillella* and probably had the same mode of life, though slightly less buried into the substrate, as it lacked the *Gervillella* spear shape but had a more developed anterior lobe (Muster, 1995).

Mineralogy.—Bimineralic (Carter, 1990a, p. 206–207). No data about *Virgellia* shell mineralogy or microstructure are available. Data provided for the family Bakevelliidae. Outer shell layer: calcite (simple prismatic). Inner shell layer: aragonite (nacreous).

Genus GERVILLEIOPERNA Krumbeck, 1923b, p. 76

Type species.—Gervilleioperna timoriensis Krumbeck, 1923b, p. 76. *Remarks.*—Although Cox and others (1969) and other authors included *Gervilleioperna* in the Isognomonidae, we assign it to the Bakevelliidae following Damborenea (1987b), noting that it had a pteriform shell and a strong radial carina.

Stratigraphic range.—Lower Jurassic (Sinemurian)—Middle Jurassic (Aalenian) (Aberhan, 1994a; Aberhan & Hillebrandt, 1996). The oldest record of *Gervilleioperna* is from Sinemurian beds (Aberhan, 1994a). During the Toarcian, its distribution became restricted to the Circumpacific domain (in Chile) (Aberhan & Hillebrandt, 1996).

Paleogeographic distribution.—Circumpacific (Fig. 14). Although Gervilleioperna was present beginning in Sinemurian times, it reached a diversity peak in the Pliensbachian and became extinct during the Aalenian. It is especially abundant in the Tethys domain during the Pliensbachian (Accorsi-Benini & Broglio-Loriga, 1975; Buser & Debeljak, 1994; Liu, 1995; Aberhan & Fürsich, 1997; Fraser & Bottjer, 2001a, 2001b; Fraser, Bottjer, & Fischer, 2004), while during the Sinemurian, it was only found in the Circumpacific domain. It was also reported from the northern part of the Austral domain during the Pliensbachian (Damborenea, 1987b). Gervilleioperna was not recorded in high paleolatitudes, and it was therefore restricted to warm waters (Damborenea, 1996a). It had a pan-Tethyan distribution, ranging from the Pacific coast (South America) through southern Europe and northern Africa to the eastern Tethys (Timor).

Circumpacific domain: Early Jurassic: Sinemurian of Chile (Aberhan, 1994a; Aberhan & Muster, 1997).

Paleoautoecology.-B, Se, S, Endo, R, Sed; By. Along with Lithiotis, Lithioperna, Cochlearites, and Mytiloperna, Gervilleioperna was a reef builder, especially during Pliensbachian times, replacing the coral reefs of the Late Triassic (Fraser, Bottjer, & Fischer, 2004). Gervilleioperna was interpreted as being reclined, lying on its left valve (Seilacher, 1984; Damborenea 1987b; Fraser, Bottjer, & Fischer, 2004). Buser and Debeljak (1994) interpreted it as epifaunal epibyssate similarly to Recent Isognomon species, but we believe the semi-infaunal endobyssate option is more reasonable, because its left valve is much heavier than the right and it would have easily sunk into the soft sediment (Damborenea, 1987b). Seilacher (1984) interpreted it as a cup-shaped recliner in soft sediments, similar to Gryphaea. Fraser, Bottjer, and Fischer (2004, fig. 10A) agreed, but they classified it as epifaunal. Aberhan and Hillebrandt (1996) suggested that Gervilleioperna (Gervilleiognoma) aurita Aberhan & Hillebrandt was semiinfaunal endobyssate, lying on the umbonal region and the anterior part of its left valve, and with its commissural plane oblique to the substrate surface. Since a byssal notch is observed (Cox & others, 1969, p. 325), it most likely was endobyssate.

Mineralogy.—Aragonitic (Accorsi-Benini & Broglio-Loriga, 1975; Carter, 1990a; Carter, Barrera, & Tevesz, 1998). Accorsi-Benini and Broglio-Loriga (1975) studied the shell of their specimens of *Gervilleioperna*, but they did not check the mineralogical composition (aragonite or calcite) of the shell layers. Carter (1990a) noted that further analysis is needed to determine whether the outer layer contains prismatic calcite. Outer shell layer: aragonite-calcite (?). Inner shell layer: aragonite (?).

Family CASSIANELLIDAE Ichikawa, 1958 Genus CASSIANELLA Beyrich, 1862, p. 9

Type species.—Avicula gryphaeata Münster in Goldfuss, 1835 in 1833–1841, p. 127.

Stratigraphic range.—?Permian, Middle Triassic (Anisian)–Upper Triassic (Rhaetian). Cox and others (1969) assigned *Cassianella* a Triassic and probably Permian range. Ciriacks (1963) and Waterhouse (1987) mentioned *Cassianella* from the lower and middle Permian (up to Guadalupian), but specimens in both papers were referred to the genus by external morphology only, and in neither reference were internal characters described. Although Ciriacks (1963, p. 31) mentioned *Cassianella* from the Lower Triassic, we did not find any information about this, and he did not figure any specimens. *Cassianella* was common from the Anisian to Rhaetian (see paleo-geographic distribution).

Paleogeographic distribution.—Cosmopolitan (Fig. 15).

Tethys domain: Middle Triassic: Anisian of southern China (Komatsu, Chen, & others, 2004); Anisian-Ladinian of Bulgaria (Stefanov, 1942), Italy (Posenato, 2008a); Ladinian of Spain (Márquez-Aliaga, 1983, 1985; Márquez-Aliaga, García-Forner, & Plasencia, 2002), Slovakia (Kochanová, Mello, & Siblík, 1975), Israel (Lerman, 1960), Italy (Rossi Ronchetti, 1959); Late Triassic: Sumatra (Krumbeck, 1914), Alps (Austria) (Tomašových, 2006a, 2006b), China (Cowper-Reed, 1927; J. Chen, 1982a; Gou, 1993); Carnian of the Italian Alps (Bittner, 1895; Leonardi, 1943; Fürsich & Wendt, 1977), Carpathians (Bittner, 1901a), Turkey (Bittner, 1891), Spain (Márquez-Aliaga & Martínez, 1996), Israel (Lerman, 1960); Norian of western Caucasus (Ruban, 2006a), China (Wen & others, 1976; J. Chen & Yang, 1983), Singapore (Kobayashi & Tamura, 1968a; Norian–Rhaetian of Iran (Hautmann, 2001b); late Rhaetian of Tibet (J. Yin & McRoberts, 2006), Pamira (Polubotko, Payevskaya, & Repin, 2001), Alps (Italy) (Desio, 1929; McRoberts, Newton, & Allasinaz, 1995), India (Healey, 1908).

Circumpacific domain: Late Triassic: Peru (Körner, 1937), Japan (Kobayashi & Ichikawa, 1949a; Tamura, 1990); Norian of Oregon (United States) (Newton, 1986, 1989; Newton & others, 1987), southwestern Alaska (McRoberts & Blodgett, 2000), Canada (Tozer, 1962, 1970); Rhaetian of Nevada (United States) (Silberling, 1961; Laws, 1982); Norian–Rhaetian of Chile (Chong & Hillebrandt, 1985).

Austral domain: Late Triassic: Carnian of New Zealand (Trechmann, 1918; Marwick, 1953); Norian–Rhaetian of Argentina (Riccardi & others, 1997, 2004; Damborenea & Manceñido, 2012).

Boreal domain: Late Triassic: northern Siberia (Kurushin, 1990; Polubotko & Repin, 1990); Carnian of Primorie (Kiparisova, 1972); Norian–Rhaetian of northeastern Russia (Kiparisova, Bychkov, & Polubotko, 1966).

Paleoautoecology.—B, E, S, Un, Sed; R. Cassianellids are generally interpreted as being reclining bivalves, lying on their left valve on the sediment (Fürsich & Wendt, 1977; Laws, 1982; Newton in Newton & others, 1987; Hautmann, 2001b). Some species, such as *C. lingulata* Gabb, 1870, and *C. angusta* Bittner, 1891, were thought by Laws (1982) and Newton (in Newton & others, 1987), respectively, to be byssate, although there is no byssal notch.

Mineralogy.—Aragonitic (Carter, 1990b, p. 338–339; Carter, Barrera, & Tevesz, 1998). Carter, Barrera, and Tevesz (1998) assigned an aragonitic mineralogy for all shell layers to the family Cassianellidae. Previously, Carter (1990b) noted that some species of *Cassianella* [e.g., *C. beyrichi* Bittner, 1895, or *C. inaequiradiata* (Schafhäutl, 1852)] have calcite in the outer shell layer. *Cassianella* is the only genus of this family for which there are studies of the mineralogy and shell microstructure. Outer shell layer: calcite or aragonite (prismatic). Inner shell layer: aragonite (prismatic).

Genus BURCKHARDTIA Frech, 1907, p. 334

Type species.—Cassianella (Burckhardtia) boesei Frech, 1907, p. 334. *Remarks.*—According to Alencaster de Cserna (1961), the type species was first referred to *?Pterinea* by Burckhardt and Scalia (1905).

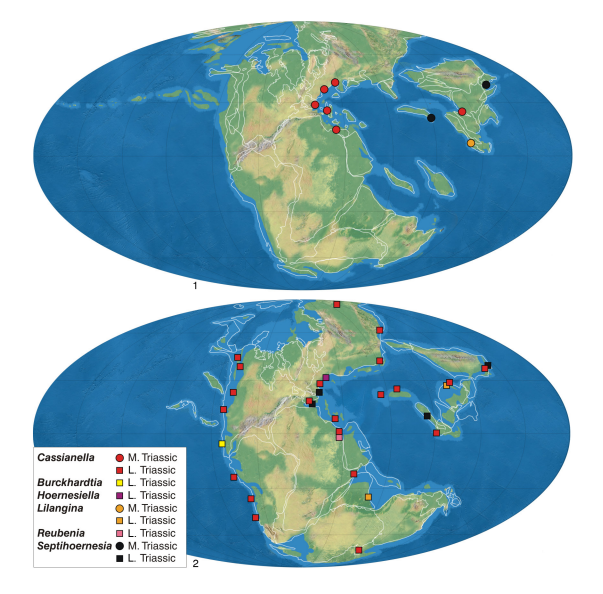


Figure 15. Paleogeographical distribution of Cassianellidae (Cassianella, Burckhardtia, Hoernesiella, Lilangina, Reubenia, Septihoernesia). 1, Middle Triassic; 2, Late Triassic.

Frech (1907) argued that this species was better included in *Cassianella* on the basis of its characteristics, and he named the subgenus *Burckhardtia*. But Alencaster de Cserna (1961) suggested that this species is more related to *Myophoria* Bronn, 1835 in 1834–1838, than to *Cassianella* Beyrich, 1862, and she included it in the first genus. Following Cox and others (1969), we regard it as a separate genus within the family Cassianellidae due to the presence of obtuse wings, a feature not known among myophorids.

Stratigraphic range.—Upper Triassic (Carnian) (Frech, 1907). Frech (1907) described the genus from Carnian beds of Zacatecas (Mexico), and it appears to be endemic in this area and restricted to this age (Burckhardt & Scalia, 1905; Diener, 1923; Alencaster de Cserna, 1961; Cox & others, 1969, Hallam, 1981; Kobayashi & Tamura, 1983a; Barboza-Gudino, Tristán-González, & Torres-Hernández, 1990; Sepkoski, 2002).

Paleogeographic distribution.—Circumpacific (Fig. 15).

Circumpacific domain: Late Triassic: Carnian of Mexico (Burckhardt & Scalia, 1905; Frech, 1907; Alencaster de Cserna, 1961; Barboza-Gudino, Tristán-González, & Torres-Hernández, 1990).

Paleoautoecology.—B, E, S, Un, Sed; R. *Burckhardtia* probably had a mode of life similar to *Cassianella*, but considering that it is almost subequivalve, a semi-infaunal mode of life, similar to *Hoernesia*, would perhaps be more likely.

Mineralogy.—Aragonitic (Carter, 1990a, p. 207; Carter, Barrera, & Tevesz, 1998). Data provided for family Cassianellidae. There are no studies on the shell of *Burckhardtia* (see *Cassianella* mineralogy, p. 46). Outer shell layer: calcite or aragonite (prismatic). Inner shell layer: aragonite (prismatic).

Genus HOERNESIELLA Ichikawa, 1958, p. 195

ex Gugenberger, 1935, p. 250

Type species.—Hoernesiella horrida Gugenberger, 1935, p. 250.

Remarks.—Gugenberger did not designate a type species for his genus *Hoernesiella*. Ichikawa (1958, p. 195) designated *Hoernesiella horrida* as type species, and he claimed the generic authorship under Article 25 c 3 of ICZN (1999). Years later, Cox in Cox and others (1969, p. 312), surely without knowledge of Ichikawa's (1958) paper, noticed the lack of type species, and designated another type species for *Hoernesiella*: *H. carinthiaca* Gugenberger, 1935, p. 250, also claiming the generic name authorship (see Stenzel, 1971, p. 1215). Vokes (1980) attributed the authorship to Ichikawa (1958) by priority of type species designation, and this approach is followed here.

Stratigraphic range.—Upper Triassic (Carnian) (Cox & others, 1969). Cox and others (1969), Stenzel (1971), Hallam (1981), and Sepkoski (2002) assigned it a Carnian range. It was not possible to find more information about this genus.

Paleogeographic distribution.—western Tethys (Fig. 15).

Tethys domain: Late Triassic: Carnian of Carinthia (Austria) (Ichikawa, 1958; Cox & others, 1969).

Paleoautoecology.-B, E, S, Un, Sed; R. Similar to Cassianella.

Mineralogy.—Aragonitic (Carter, 1990a, p. 207; Carter, Barrera, & Tevesz, 1998). Data provided for family Cassianellidae. There are no specific studies on the shell of *Hoernesiella* (see *Cassianella* mineralogy, p. 46). Outer shell layer: calcite or aragonite (prismatic). Inner shell layer: aragonite (prismatic).

Genus LILANGINA Diener, 1908, p. 62

Type species .- Lilangina nobilis Diener, 1908, p. 62.

Stratigraphic range.—Middle Triassic (Anisian)–Upper Triassic (Carnian) (Cox & others, 1969; Komatsu, Huyen, & Huu, 2010). All sources checked assigned *Lilangina* to the Carnian (Diener, 1923; Cox & others, 1969; Hallam, 1981; Kobayashi & Tamura, 1983a). However, recently, Komatsu, Huyen, and Huu (2010) reported it from the Anisian and Ladinian.

Paleogeographic distribution.—Eastern Tethys (Fig. 15).

Tethys domain: Middle Triassic: Anisian–Ladinian of northern Vietnam (Komatsu, Huyen, & Huu, 2010); Late Triassic: Carnian of Kashmir (Diener, 1923; Cox & others, 1969; Kobayashi & Tamura, 1983a), China (Wen & others, 1976; Kobayashi & Tamura, 1983a; Sha, Chen, & Qi, 1990).

Paleoautoecology.-B, E, S, Un, Sed; R. Similar to Cassianella.

Mineralogy.—Aragonitic (Carter, 1990a, p. 207; Carter, Barrera, & Tevesz, 1998). Data provided for family Cassianellidae. There are no specific studies on the shell of *Lilangina* (see *Cassianella* mineralogy, p. 46). Outer shell layer: calcite or aragonite (prismatic). Inner shell layer: aragonite (prismatic).

Genus REUBENIA Cox, 1924, p. 61

Type species.—Reubenia hesbanensis Cox, 1924, p. 63.

Stratigraphic range.—Upper Triassic (Carnian) (Cox, 1924). Cox (1924) described *Reubenia* from the Carnian beds of Jordan, including two species, the type species and *Reubenia attenuata* Cox, 1924. All reviewed literature (Kutassy, 1931; Cox & others, 1969; Hallam, 1981; Sepkoski, 2002) assigned it the same stratigraphic range.

Paleogeographic distribution.—western Tethys (Fig. 15).

Tethys domain: Late Triassic: Carnian of Jordan (Cox, 1924). Paleoautoecology.—B, E, S, Un, Sed; R. Similar to Cassianella. *Mineralogy.*—Aragonitic (Carter, 1990a, p. 207; Carter, Barrera, & Tevesz, 1998). Data provided for family Cassianellidae. There are no specific studies on the shell of *Reubenia* (see *Cassianella* mineralogy, p. 46). Outer shell layer: calcite or aragonite (prismatic). Middle shell layer: aragonite (nacreous). Inner shell layer: aragonite (prismatic).

Genus SEPTIHOERNESIA Cox, 1964, p. 40

Type species.—Gervillia johannisaustriae Klipstein, 1845 in 1843–1845, p. 249.

Stratigraphic range.—Middle Triassic (Ladinian)–Upper Triassic (Carnian) (Allasinaz, 1966; Tamura, 1990). Cox and others (1969) assigned it a Triassic range, without further comments. Sepkoski (2002) assigned it a Triassic (lower Anisian–Carnian) range, allegedly using Hallam (1981) as data source, but Hallam only mentioned it from Ladinian and Carnian times. The published records indicate that the genus was only present in these two Triassic stages.

Paleogeographic distribution.—Tethys and Circumpacific (Fig. 15).

Tethys domain: Middle Triassic: Ladinian of Malaysia (Tamura & others, 1975); Late Triassic: Carnian of Italy (Allasinaz, 1966; Fürsich & Wendt, 1977), Spain (Martín-Algarra, Solé de Porta, & Márquez-Aliaga, 1995), Malaysia (Tamura & others, 1975).

Circumpacific domain: Middle Triassic: Ladinian of Japan (Tamura, 1990); Late Triassic: Carnian of Japan (Tamura, 1990).

Paleoautoecology.-B, E, S, Un, Sed; R. Similar to Cassianella.

Mineralogy.—Aragonitic (Carter, 1990a, p. 207; Carter, Barrera, & Tevesz, 1998). There are no studies on the shell of *Septihoernesia* (see *Cassianella* mineralogy, p. 46). Outer shell layer: calcite or aragonite (prismatic). Inner shell layer: aragonite (prismatic).

Family DATTIDAE Healey, 1908 Genus DATTA Healey, 1908, p. 63

Type species.—Datta oscillaris Healey, 1908, p. 63.

Stratigraphic range.—Upper Triassic (Rhaetian) (Healey, 1908). Healey (1908) described the genus from Rhaetian beds of Burma, and Cox and others (1969) repeated these data. Kobayashi and Tamura (1983a) recorded *Datta* from several Upper Triassic localities but did not mention stages or the original data source. The statement in Damborenea (2002b, p. 56): "... During the Jurassic and Lower Cretaceous, most genera of Anomiidae, Burmesiidae, Ceratomyopsidae, Cuspidariidae [Dattidae] Diceratidae" is an error, and there are no records of *Datta* from the Jurassic.

Paleogeographic distribution.—Southern Tethys (Fig. 16). The original material is from Burma (Healey, 1908). Later, Kobayashi and Tamura (1983a) also reported the genus from the Late Triassic of Kashmir and Yunnan (China), but they did not indicate the original source, and no information related to these records was found.

Paleoautoecology.—B, E, S, ?, ?. It is difficult to assign a mode of life to this genus since the only information available is from a left valve mold with a possible chondrophore. The shell morphological features suggest it was probably epifaunal.

Mineralogy.—Bimineralic (Carter, 1990a, p. 205). Data provided for superfamily Pterioidea. There are no specific studies on *Datta* shell. Outer shell layer: calcite (prismatic). Inner shell layer: aragonite (nacreous).

Family ISOGNOMONIDAE Woodring, 1925

It is often difficult to distinguish between Isognomonidae and Bakevelliidae; good examples are the genera *Isognomon (Mytiloperna)* von Ihering, 1903, and *Gervilleioperna* Krumbeck, 1923b, about which there is no agreement among different authors (see discussions for these taxa, below and p. 45). It is quite evident that these two families are phylogenetically related, and consequently setting limits is complicated. A revision of their diagnostic features is needed to establish a consensus. There are also certain difficulties in distinguishing between Inoceramidae and Isognomonidae (see Crampton, 1988).

Genus ISOGNOMON [Lightfoot, 1786], p. 41, 52

Lightfoot proposed the name in an anonymous catalogue (1786, authorship determined by Dance, 1962, see also Kay, 1965) spelling the name both as *Isognoma* and *Isognomon*, and later Dall, Bartsch, and Rehder (1938, p. 61–62) regarded *Isognomon* as the original spelling and *Isognoma* as a misspelling (first revisers action according to Coan, Valentich Scott, & Bernard, 2000, p. 196).

Type species.—Ostrea isognomon Linnaeus, 1764, p. 533 (=Ostrea isognomum in Linné, 1758), by absolute tautonymy (see discussion in Rehder, 1967, p. 6, and Coan, Valentich Scott, & Bernard, 2000, p. 196). This differs from the interpretation by Cox (in Cox and others, 1969, p. 322), which was followed by most authors.

Remarks.—According to Cox and others (1969), there are two subgenera of Isognomon within our interval of study, I. (Isognomon) (but see below) and I. (Mytiloperna). However, some authors noted that probably the latter is more related to bakevelliids than to isognomonids. Mytiloperna was described by H. von Ihering (1903) as a genus based in Perna americana Forbes; Cox (1940) demoted it to a subgenus of Isognomon Lightfoot, 1786, a position maintained in Cox and others (1969) (see Accorsi-Benini & Broglio-Loriga, 1975, for details). There are several reasons to consider that Mytiloperna does not fit into the Isognomonidae. One is the shell microstructure (Broglio-Loriga & Posenato, 1996). Another is that adults had a hinge with teeth, a feature of Bakevelliidae and not of Isognomonidae, which were toothless in the adult stage (Seilacher, 1984; Aberhan, 1998a). However, while pending a good revision of the family, it is advisable to treat Mytiloperna as an isognomonid (Jaitly, Fürsich, & Heinze, 1995). Additionally, due to a misinterpretation about the correct way of fixation of the type species of Isognomon (see above, and IZCN, 1999, Art. 68.1 and 68.4), most Mesozoic species should be referred to Isognomon (Melina) Retzius, 1788, p. 22, and not to Isognomon (Isognomon), a fact overlooked by most authors, even by those who accepted *I*. sognomon as type of the genus..

Stratigraphic range.—Upper Triassic (Carnian)–Holocene (Cox & others, 1969). Cox and others (1969) assigned an Upper Triassic to Holocene range to *Isognomon (Isognomon)* and an Lower Jurassic to Upper Jurassic range to *Isognomon (Mytiloperna)*. Linck (1972) reported the last subgenus from the Carnian, but he included in *Mytiloperna* (considered at genus level) some modioliform specimens somewhat different from the typical ones. Today there are numerous species living in tropical seas (Beesley, Ross, & Wells, 1998).

Paleogeographic distribution.—Tethys (Fig. 16). During other intervals of geological time, the genus also lived in Circumpacific and Austral regions, especially during the Pliensbachian (see e.g., Damborenea, 1987b; Broglio-Loriga & Posenato, 1996; Aberhan & Fürsich, 1997; Aberhan 1994a, 1998a; Liu, 1999; Fraser, Bottjer, & Fischer, 2004). However, during the temporal interval under consideration, it was only known from the Tethys domain. Holocene species of *Isognomon* are mainly distributed in tropical seas.

Tethys domain: Late Triassic: Carnian of China (Sha, Chen, & Qi, 1990), Germany (Linck, 1972), southern Italian Alps (Fürsich & Wendt, 1977), Italy (Gelati & Allasinaz, 1964); Norian of Austria (Tichy, 1975), Italy (Terranini, 1958); late Norian of southern China (J. Chen, 1982a); Norian–Rhaetian of Iran, Burma, and Vietnam (Hautmann, 2001b; Fürsich & Hautmann, 2005); Rhaetian of Alps (Italy) (Pozzi, Gelati, & Allasinaz, 1962); Early Jurassic: Hettangian of Japan (Kondo & others, 2006), Europe and northeastern Asia (Hallam, 1977); Hettangian–Toarcian of Japan (Hayami, 1957a, 1975); Sinemurian of Morocco (Liu, 1995), northwestern Europe (Hallam, 1987).

Paleoautoecology.—B, Se, S, Endo, Sed; By. A semi-infaunal endobyssate mode of life seems likely for most Mesozoic species, although some Recent species live epifaunally (S. M. Stanley, 1970, 1972). This disparity in mode of life can be recognized by differences in shell morphology. Living species are often very inequivalve, unlike Mesozoic species, which were equivalve or subequivalve (Hayami, 1957a); there are also differences in shell thickness, the umbonal part being thicker than the ventral part in Mesozoic species (see Fürsich, 1980; Seilacher, 1984; Broglio-Loriga & Posenato, 1996; Fraser, Bottjer, & Fischer, 2004).

Fürsich (1980) analyzed some of the fossil species and observed that, if only shell characters were taken into account, his interpretations were wrong and not viable when he could contrast these results with direct observation of individuals found in life position in the field. All fossil species studied in his work of1980 seem to support a semi-infaunal endobyssate mode of life. Furthermore, fossil species are often found forming groups, and thus they were interpreted as gregarious (Fürsich, 1982; Damborenea, 1987b).

With regard to *I. (Mytiloperna)*, several authors studied its morphology in relation to its mode of life (e.g., Seilacher, 1984; Broglio-Loriga & Posenato, 1996; Fraser, Bottjer, & Fischer, 2004). These last two papers distinguished several *Mytiloperna* morphotypes with different life habit interpretations, ranging from epifaunal to semi-infaunal.

Mineralogy.—Bimineralic (Carter, 1990b, p. 339). Accorsi-Benini and Broglio-Loriga (1975) and Broglio-Loriga and Posenato (1996) described a fibrous microstructure in the inner shell layer of *Mytiloperna* sp. specimens. Outer shell layer: calcite (simple prismatic). Inner shell layer: aragonite (nacreous).

Genus LEPROCONCHA Giebel, 1856, p. 67

Type species.—Leproconcha paradoxa Giebel, 1856, p. 67.

Remarks.—When Giebel (1856) described the genus, he did not mention its systematic relations, although he indicated that it was intermediate between Ostreacea and Malleacea. Nevertheless, Cox and others (1969) decided to include it with doubts into the Isog*Stratigraphic range.*—Middle Triassic (Giebel, 1856). Giebel (1856) described *Leproconcha* from Muschelkalk of the Germanic Basin. Diener (1923), Kutassy (1931), Cox and others (1969), and Kobayashi and Tamura (1983a) repeated these data, and no further information was found.

Paleogeographic distribution.-western Tethys (Fig. 16).

Tethys domain: Middle Triassic: Muschelkalk of Germany (Giebel, 1856).

Paleoautoecology.—B, E, S, ?, ?. Probably epifaunal.

Mineralogy.—Unknown. There is no information about mineralogy or microstructure of *Leproconcha* shell. We cannot assign it the mineralogy predominant in the family because we have serious doubts about the family assignation of this genus.

Genus WAAGENOPERNA Tokuyama, 1959a, p. 151

Type species.—Edentula lateplanata Waagen, 1907, p. 97.

Remarks.—Waagenoperna Tokuyama, 1959a was proposed to replace *Edentula* Waagen, 1907 (*non* Nitzsch, 1820), and *Edentula lateplanata* Waagen, 1907, was designated as type species by Tokuyama (1959a). Some years earlier, Cox (1954) had proposed the name *Cuneigervillia* also to replace *Edentula*, choosing *Gervillia hagenowii* Dunker, 1846, as the type species. When Tokuyama (1959a) compared the two type species, he noticed that while *Gervillia hagenowii* is a bakevelliid (in fact, Cox, 1954, included *Cuneigervillia* in the family Bakevelliidae), *Edentula lateplanata* is an isognomonid, like other species attributed to *Edentula* (*E. triangularis* Kobayashi &z Ichikawa, 1952). Lower Jurassic species also referred to *Cuneigervillia* by Cox clearly showed that this genus was not objectively the same as *Edentula* (in fact, they have different type species), and he decided to maintain both names, *Cuneigervillia* and *Waagenoperna* (*=Edentula*), which was followed by Cox and others (1969).

Nakazawa and Newell (1968) proposed a new subgenus within *Waagenoperna*, *W. (Permoperna*), and although some authors (Z. Fang, 1982) treated *Permoperna* at generic level, the original rank is here retained because it does not present substantial differences from *Waagenoperna s.s.*

Tëmkin (2006, p. 270) erroneously indicated that *Waagenoperna* was based on *W. triangularis* (Kobayashi & Ichikawa, 1952).

Stratigraphic range.—lower Permian (Sakmarian)–Upper Triassic (upper Norian) (Hayami & Kase, 1977; J. Chen, 1982a). Cox and others (1969) assigned it a middle Permian–Upper Triassic range. The stratigraphic range should be extended back, since Nakazawa and Newell (1968) reported *Waagenoperna (Permoperna)* from the lower Permian (Sakmarian). Sepkoski's (2002) range starts from the Guadalupian, which is odd considering that he took his data from Hayami and Kase (1977) and Skelton and Benton (1993), who indicated Sakmarian as the first record. Regarding the upper extension of this genus, J. Chen (1982a, p. 303) quoted *Waagenoperna* from

the upper Norian of southern China, indicating that the association to which it belonged is "the uppermost Triassic bivalve zone in this region." There are several reports from Hettangian brackish environments of southern China (Sha & Jiang, 2004; Jiang, Sha, & Pan, 2008), although none was corroborated by illustrations or descriptions of the material.

Paleogeographic distribution.—Tethys and Circumpacific (Fig. 16).

Tethys domain: Late Triassic: Carnian of southern Alps (Broili, 1904; Tokuyama, 1959a), southern China (Gu & others, 1980); late Norian of China (J. Chen, 1982a).

Circumpacific domain: late Permian: Japan (Nakazawa & Newell, 1968; Hayami, 1975); Middle Triassic: Ladinian of Japan (Tokuyama, 1959a; Hayami, 1975); Late Triassic: Carnian–Norian of Japan (Tokuyama, 1959a; Hayami, 1975).

Paleoautoecology.—B, Se, S, Endo, Sed; By. Although *Waagenoperna* had a mytiliform shell and most isognomonids were interpreted as epifaunal bivalves, *Waagenoperna* was thought to be a semi-infaunal endobyssate bivalve, similar to *Pinna* and some pterineids (S. M. Stanley, 1972). This author relied on morphological evidence, such as the strongly prosocline, slightly inflated, subequivalve shell and the presence of an anterior lobe, features not shared with any other member of the family Isognomonidae.

Mineralogy.—Bimineralic (Carter, 1990a, p. 209). There are no data about the *Waagenoperna* shell. Information provided for family Isognomonidae. Outer shell layer: calcite (simple prismatic). Middle and inner shell layers: aragonite (nacreous).

Family POSIDONIIDAE Neumayr, 1891

The Posidoniidae family members are hardly distinguishable from each other because one of the main diagnostic features is the ligament area, and this is only preserved in exceptional cases. Due to the usually very thin shell, all internal shell characters are frequently destroyed during diagenesis. Waller (in Waller & Stanley, 2005) gathered the families Posidoniidae and Halobiidae Kittl, 1912, in the superfamily Posidonioidea Frech, 1909. This arrangement is probably more appropriate than the one followed here, but according to Amler (1999), we consider the first family in the superfamily to be Pterioidea Gray 1847, and the second one to be Halobioidea H. J. Campbell, 1994.

Genus BOSITRA De Gregorio, 1886, p. 11

Type species.—Posidonia ornati Quenstedt, 1851 in 1851–1852, p. 501.

Remarks.—Following Waller (in Waller & Stanley, 2005), we regard *Posidonia* Bronn, 1828, as a Paleozoic genus, referring described species from Lower and Middle Triassic to *Bositra*. We consider *Peribositria* Kurushin & Trushchelev, 1989, to be a synonym of *Bositra* (see discussion for *Peribositria* in Genera not Included, p. 167).

Stratigraphic range.—Lower Triassic (lower Olenekian)–Middle Jurassic (lower Oxfordian) (Waller & Stanley, 2005). Cox and others (1969) assigned it a Jurassic range, but after Waller and Stanley (2005) emended the genus and they transferred species traditionally included in *Posidonia* from Lower and Middle Triassic, the range was extended back to the Triassic. The problem in the

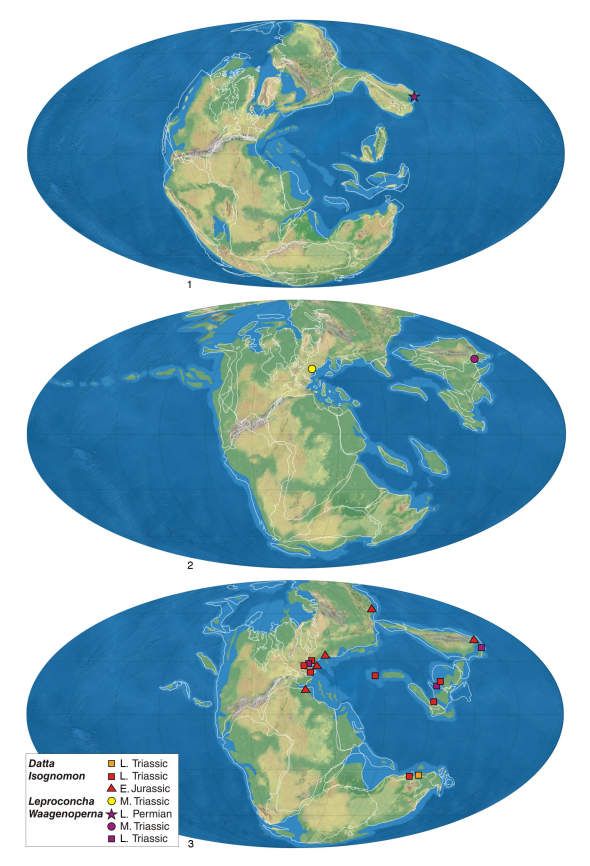


Figure 16. Paleogeographical distribution of Isognomonidae (Isognomon, Leproconcha, Waagenoperna) and Dattidae (Datta). 1, late Permian; 2, Middle Triassic; 3, Late Triassic-Early Jurassic.

Paleontological Contributions, number 8

differentiation of both genera is that the main diagnostic characters are in the ligament area, as well as other internal structures, which are often destroyed during diagenesis (Waller in Waller & Stanley, 2005). Fürsich and Werner (1988) reported *Bositra* from the Upper Jurassic (Kimmeridgian) of Portugal, but their specimens were referred to this genus with some hesitation since the ligament area is not preserved.

Paleogeographic distribution.—Tethys, Circumpacific, and Boreal (Fig. 17). During the Early Triassic, *Bositra* was distributed mainly in the Boreal domain, extending over the Tethys and Circumpacific domains during the Middle Triassic, to become virtually cosmopolitan during the Early Jurassic (Waller & Stanley, 2005), especially during the Toarcian (Damborenea, 1987b; Aberhan, 1994a, 1998a; Monari, 1994; Liu, 1995; Harries & Little, 1999; Gahr, 2002), coinciding with the peak of early Toarcian extinction. We did not take into account the mention in Waterhouse (2000), because Waller and Stanley (2005) affirm the specimens are more clariids than posidoniids.

Tethys domain: Middle Triassic: Slovenia (Jurkovsek, 1984); Anisian of China (Wen & others, 1976; Ling, 1988; Komatsu, Akasaki, & others, 2004a; J. Chen & Stiller, 2007), western Carpathians (Slovakia) (Kochanová, 1985); Anisian–Ladinian of Vietnam (Vu Khuc & Huyen, 1998); Ladinian of Spain (Márquez-Aliaga, 1983, 1985; Márquez-Aliaga & Ros, 2003); Late Triassic: Carnian of China (Wen & others, 1976), Italy (Fürsich & Wendt, 1977); Norian of China (Sha, Chen, & Qi, 1990).

Circumpacific domain: Middle Triassic: Anisian of Japan (Hayami, 1975), of Nevada (United States) (Waller & Stanley, 2005).

Boreal domain: Early Triassic: Siberia (Kiparisova, 1938; Dagys & Kurushin, 1985), Arctic Archipelago (Canada) (Tozer, 1961, 1962, 1970).

Paleoautoecology.—B, E, S, Un, Sed; R. Bositra was interpreted as a pseudoplanktonic bivalve (S. M. Stanley, 1972) or as nektoplanktonic (Jefferies & Minton, 1965; Hayami, 1969a; Duff, 1975), according to its distribution and morphological characteristics. Other authors rejected these interpretations considering the habit of species assigned to Bositra as benthic (M. A. Conti & Monari, 1992; Etter, 1996).

Etter (1996) did a comprehensive study reviewing all possible modes of life ever attributed to *Bositra*, demonstrating that a benthic mode of life is entirely plausible and providing arguments to reject the other two options. Since a byssal notch is not observed, a reclined mode of life would have been the most likely (Waller & Stanley, 2005). The reason for its frequent presence in oxygenpoor sedimentary environments should more probably be related to an opportunistic behavior than to a pseudoplanktonic mode of life (Etter, 1996). An updated discussion on the mode of life for *Bositra* was provided by Caswell, Coe, and Cohen (2009, and see references therein).

Mineralogy.—Bimineralic (Carter, 1990b, p. 340; Waller & Stanley, 2005). Outer shell layer: calcite (homogeneous-prismatic). Inner shell layer: aragonite (nacreous).

Genus AMONOTIS Kittl, 1904, p. 736

Type species.—Amonotis cancellaria Kittl, 1904, p. 736.

Stratigraphic range.—Upper Triassic (Carnian). Amonotis was reported from Carnian beds (Cox & others, 1969; C. Chen & Yu, 1976; Sha, Chen, & Qi, 1990) and apparently also from the Norian, although Norian records lack illustrations or they were dubiously assigned to the genus (Niu, Xu, & Ma, 2003; Tang & others, 2007).

Paleogeographic distribution.—Tethys (Fig. 17). Amonotis was found both in western and eastern Tethys, although some authors (Hallam, 1981; Metwally, 1993) only included European occurrences in their compilations.

Tethys domain: Late Triassic: Carnian of Yugoslavia (Cox & others, 1969), China (C. Chen & Yu, 1976; Sha, Chen, & Qi, 1990).

Paleoautoecology.—B, E, S, Epi, Sed; By. According to S. M. Stanley (1972), almost all posidoniids were epibyssate bivalves, though almost none of the genera shows a sinus or byssal notch, and some of them could be pseudoplanktonic, although there is no evidence that *Amonotis* was one of them. Therefore, we consider *Amonotis* as to be an epibyssate bivalve in agreement with Sha, Chen, and Qi (1990), but the possibility of a reclined habit similar to *Bositra* cannot be ruled out.

Mineralogy.—Bimineralic (Carter, 1990a, p. 211). Data provided for family Posidoniidae. We lack information about the shell of *Amonotis*. Outer shell layer: calcite (prismatic-homogeneous). Inner shell layer: aragonite (nacreous).

Genus VELDIDENELLA Alma, 1925, p. 118

Type species .--- Veldidenella dieneri Alma, 1925, p. 118.

Stratigraphic range.—Middle Triassic (upper Anisian–upper Ladinian) (Kochanová, 1985). Cox and others (1969) assigned it a Upper Triassic range, and both Sepkoski (2002) and other compilation papers (e.g., Metwally, 1993) repeated this information. But according to the information published on this monospecific genus, these data appear to be wrong. According to Kutassy (1931), Alma in 1925 described the type species of this genus from Anisian beds of the northern Alps. Tichy (1970), in his catalog of type specimens housed at the Museum of Natural History in Vienna, also indicated Anisian as the age of the type species. Subsequently, Kochanová (1985) reported *Veldidenella dieneri* from Anisian beds of the western Carpathians and from upper Ladinian beds of the southern Alps.

Paleogeographic distribution.—western Tethys (Fig. 17).

Tethys domain: Middle Triassic: Anisian of northern Alps (Austria) (Kutassy, 1931), Carpathians (Slovakia) (Kochanová, 1985); Ladinian of northern Alps (Austria) (Kochanová, 1985).

Paleoautoecology.-B, E, S, Epi, Sed; By. Similar to Amonotis.

Mineralogy.—Bimineralic (Carter, 1990a, p. 211). There are no data about *Veldidenella* mineralogy or shell microstructure. Data provided for the family Posidoniidae. Outer shell layer: calcite (prismatic-homogeneous). Inner shell layer: aragonite (nacreous).

Genus CAENODIOTIS Monari, 1994, p. 171

nom. nov. pro Diotis Simonelli, 1884, p. 125, non Schmarda, 1859, p. 5

Type species.—Posidonomya janus Meneghini, 1853, p. 27

Remarks.—Damborenea (1987b, p. 191) noticed that the name *Diotis* was used previously for a group of worms (*Diotis* Schmarda, 1859). Years later, Monari (1994) proposed the name *Caenodiotis* to replace *Diotis* Simonelli, 1884, in accordance with Article 52 of ICZN (1999). Following Cox and others (1969) and Monari (1994), we include *Caenodiotis* in Posidoniidae, although given its probable relations to *Posidonotis* (Damborenea, 1987b), it could perhaps be included in Entoliidae.

Stratigraphic range.—Lower Jurassic (Sinemurian–Pliensbachian) (Monari, personal communication, 2007). According to Cox and others (1969), *Diotis* was distributed during the early and middle Early Jurassic, but we could not check exactly which stages. Monari (1994) noted its presence from the Pliensbachian in several Italian localities. He also found it in the Sinemurian of the Umbria-Marche region (Monari, personal communication, 2007). We assign the range Sinemurian–Pliensbachian to this genus, until we can determine the age of the units widely referred to as lower Liassic by Cox and others (1969).

Paleogeographic distribution.—western Tethys (Fig. 17). During the Early Jurassic, especially during the Pliensbachian, the genus was distributed in Italy (Monari, 1994), Hungary (Szente, 1990), and Spain (Jiménez de Cisneros, 1923). During the time interval here analized, we only consider the Sinemurian records.

Tethys domain: Early Jurassic: Sinemurian of Italy (Monari, personal communication, 2007).

Paleoautoecology.-B, E, S, Epi, Sed; By. Similar to Amonotis.

Mineralogy.—Bimineralic (Carter, 1990a, p. 211). There are no data about the *Caenodiotis* shell. Data provided for family Posidoniidae. Outer shell layer: calcite (prismatic-homogeneous). Inner shell layer: aragonite (nacreous).

Genus STEINMANNIA Fischer, 1886 in 1880–1887, p. 960

nom. nov. pro Aulacomya Steinmann, 1881, p. 259, non Mörch, 1853, p. 53

Type species.—Posidonia bronnii Voltz in Zieten, 1833 in 1830–1833, p. 72.

Remarks.—Steinmannia is very similar morphologically to Bositra De Gregorio, 1886, and Posidonia Bronn, 1828, and the observation of the ligament type is key to discriminating between these genera (multivincular in Steinmannia, alivincular in Bositra, and duplivincular in Posidonia), but its preservation is not common (Waller in Waller & Stanley, 2005). Although Guillaume (1928) and later authors (e.g., Cox & others, 1969; Milova, 1988) included Steinmannia in the Inoceramidae due to the ligament type, Waller (in Waller & Stanley, 2005) thought the ligament of Steinmannia was not of the inoceramid type: "the relatively few ligament pits on the ligament area of Steinmannia bronni (three or four according to Guillaume, 1928, p. 221; three to five according to Milova, 1988, p. 63, pl. 2, 1-3) may be a phylogenetically independent multiplication of the simple alivincular ligament of Bositra possibly functionally associated with increase in size and convexity." Following Waller (in Waller & Stanley, 2005), we consider Steinmannia more related to Posidoniidae than Inoceramidae. Some authors included the type species of Steinmannia in Bositra (e.g., Hallam, 1976, 1987; Caswell,

Coe, & Cohen, 2009). *Multisidonia* Polubotko, 1992, p. 60 (type species *M. omolonensis* Polubotko, 1992, p. 60), distinguished by a greater number of ligamental pits, is a possible synonym.

Stratigraphic range.—Lower Jurassic (upper Sinemurian–lower Toarcian) (Guillaume, 1928; Milova, 1988). Cox and others (1969) assigned it a Toarcian range, but Milova (1988) reported a new species, *Steinmannia viligaensis* Milova, from upper Sinemurian beds (also another from the Pliensbachian, *Steinmannia alikiensis* Milova, 1988), and the type of *Multisidonia* is also late Sinemurian in age (Polubotko, 1992). Sepkoski (2002) did not consider this genus.

Paleogeographic distribution.—Boreal (Fig. 17). During the Early Jurassic, especially in the Toarcian, the genus was distributed in the Tethys domain (France, England, Germany, Switzerland) (Guillaume, 1928).

Boreal domain: Early Jurassic: Sinemurian of northeastern Russia (Milova, 1988).

Paleoautoecology.—B, E, S, Un, Sed; R. See mode of life for Bositra (p. 50).

Mineralogy.—Bimineralic (Carter, 1990a, p. 211). There are no data about the shell of *Steinmannia*. Data provided for family Posidoniidae. Outer shell layer: calcite (prismatic-homogeneous). Inner shell layer: aragonite (nacreous).

Genus ELLESMERELLA Waterhouse, 2008, p. 172

Type species .- Posidonia aranea Tozer, 1961, p. 102.

Remarks.—Waterhouse (2008) proposed *Ellesmerella* based on *P. aranea*, included it in the family Aulacomyellidae Ichikawa, 1958, and considered it to be related to *Bositra*. McRoberts (2010) suggested it could be better placed into Posidoniidae or Halobiidae. Provisionally, we include *Ellesmerella* in the Posidoniidae.

Stratigraphic range.—Lower Triassic (upper Olenekian) (Tozer, 1961). The monospecific genus *Ellesmerella* was only reported from the Olenekian stage (Tozer, 1961, 1962, 1970; Vozin & Tikhomirova, 1964; Tozer & Parker, 1968; Waterhouse, 2008; McRoberts, 2010). It has a very short stratigraphical range (uppermost Olenekian) (McRoberts, 2010).

Paleogeographic distribution.—Boreal (Fig. 17).

Boreal domain: Early Triassic: late Olenekian of Arctic Archipelago (Canada) (Tozer, 1961), Svalbard (Norway) (Tozer & Parker, 1968), British Columbia (Tozer, 1962, 1970), northeastern Siberia (Vozin & Tikhomirova, 1964).

Paleoautoecology.-B, E, S, Un, Sed; R. Similar to Bositra.

Mineralogy.—Bimineralic (Carter, 1990a). There are no data about the shell of *Ellesmerella*. Data provided for family Posidoniidae and/ or Halobiidae.

Superfamily PINNOIDEA Leach, 1819 Family PINNIDAE Leach, 1819

In our study interval, there are two genera belonging to Pinnidae: *Pinna* Linnaeus, 1758, and *Atrina* Gray, 1842, p. 83 [1840, p. 151, *nom. nud.*]. These two genera are morphologically very similar in their juvenile stages, but adults are differentiated mainly by the presence of a median shell carina, which is associated with the separation of the internal nacreous layer into two lobes in *Pinna* and is absent in *Atrina* (Cox & others, 1969; Waller & Stanley, 2005). Cox and others

(1969) assigned a Carboniferous–Holocene range to *Pinna (Pinna)* and a Middle Jurassic–Holocene range to *Atrina*. However, many specimens assigned to *Pinna (Pinna)* from beds older than Middle Jurassic do not have this median carina, and thus they should probably be referred to *Atrina* or some Paleozoic genera (*Pteronites* M'Coy, 1844, *Aviculopinna* Meek, 1864, or *Meekopinna* Yancey, 1978) (see Waller & Stanley, 2005, p. 29). In the absence of a review on this subject, we assign provisional stratigraphic ranges for both genera.

Genus PINNA Linnaeus, 1758, p. 707

Type species .- Pinna rudis Linnaeus, 1758, p. 707.

Stratigraphic range.—?Lower Triassic-Holocene (Nakazawa, 1961). Although, as already mentioned, Cox and others (1969) assigned it a range from the Carboniferous, we could not locate any record of specimens attributed to this age except in the Treatise. A specimen of Pinna (Pinna) costata Phillips from the Carboniferous of Belgium was figured in Cox and others (1969, p. 282), but this specimen does not show the typical carina, and therefore it is not supposed to belong to Pinna (Waller & Stanley, 2005). R. Zhang and Yan (1993) reported the genus from the Carboniferous, but they did not illustrate or describe the material. The oldest positive records are from the Lower Triassic. Nakazawa (1961) figured Pinna muikadaniensis Nakazawa, 1961 (p. 267; plate 13,14), and, although in the description he did not mention the median carina, he said: "... parting mediate distinct but weak in the umbonal half and obsolete in the rear part, deviating towards the antero-ventral side . . ." and the figure shows evidence of a true carina. Seguí (1999, p. 21, fig. 1), in the description of his specimen of Pinna bascoi Seguí, 1999, from Ladinian of Spain, said: "There is an edge signal that starting from the apex disappears at about 2/3 of the height. This edge divides the shell into two parts." This edge can be seen in his figures and seems to correspond with the carina. We doubtfully consider Nakazawa's (1961) Lower Triassic record as valid, since the material is not well preserved and we cannot be sure that it actually belongs to Pinna.

Paleogeographic distribution.—Tethys, Circumpacific, and Austral (Fig. 18). Although *Pinna* had a cosmopolitan distribution during our study interval, Recent species are restricted to tropical and sub-tropical seas (Cox & others, 1969). No records were located from the Boreal domain.

Tethys domain: Early Triassic: China (F. Wu, 1985); Middle Triassic: Ladinian of Spain (Seguí, 1999); Late Triassic: southern China (Gou, 1993); Carnian of Slovenia (Jelen, 1988), southern Alps (Italy) (Fürsich & Wendt, 1977), Lombardy (Italy) (Allasinaz, 1964, 1966), Germany (Linck, 1972); Norian of Australia (Grant-Mackie, 1994); Norian–Rhaetian of Iran (Hautmann, 2001b); Rhaetian of Burma (Healey, 1908), Alps (Austria) (Winkler, 1861), Tibet (China) (J. Yin & McRoberts, 2006), Pamir (Afghanistan) (Polubotko, Payevskaya, & Repin, 2001), Hungary (Vörös, 1981), Italy (Allasinaz, 1962; Sirna, 1968); Early Jurassic: Hettangian of Tibet (China) (J. Yin & McRoberts, 2006), England and Morocco (Liu, 1995), Italy (Sirna, 1968), France (Martin, 1860); Sinemurian of Portugal, Spain, England, France, and Morocco (Liu, 1995).

Austral domain: Late Triassic: Carnian of New Zealand (Trechmann, 1918; Marwick, 1953); Early Jurassic: Hettangian–Sinemurian of Argentina (Damborenea, 1996a; Damborenea & Manceñido, 2005b); Sinemurian of Argentina (Damborenea & Lanés, 2007).

Circumpacific domain: Early Triassic: Japan (Nakazawa, 1961; Hayami, 1975); Late Triassic: Norian of Chile (Hayami, Maeda, & Ruiz-Fuller, 1977); Early Jurassic: Sinemurian of Canada (Aberhan, 1998a).

Paleoautoecology.—B, Se, S, Endo, Sed; By. Depending on the species, *Pinna* currently lives more or less with the anterior half of the shell buried in the sediment as endobyssate, with a strong byssus attached to fragments of rocks or other objects, such as sea grass roots (García-March, 2005), although some were found epibyssate on hard substrates (S. M. Stanley, 1970). Regarding fossil species, there are many examples of *Pinna* in upright position, similar to living species (e.g., Fürsich, 1980, 1982; Damborenea, 1987a). It is rare to find complete fossil specimens; only the anterior parts, which are buried in life, are usually found. More detailed information about its mode of life can be found in Yonge (1953) and Seilacher (1984), among many others.

Mineralogy.—Bimineralic (Yonge, 1953; Carter, 1990a; García-March, 2005; García-March, Márquez-Aliaga, & Carter, 2008). Outer shell layer: calcite (prismatic simple). Inner shell layer: aragonite (nacreous).

Genus ATRINA Gray, 1842, p. 83

Gray, 1840, p. 151, nom. nud.

Type species.—Pinna nigra Dillwyn, 1817, p. 325.

Stratigraphic range.—Middle Triassic (Anisian)–Holocene (Stiller, personal communication, 2008). Although Cox and others (1969) assigned it a range from the Middle Jurassic, Waller (in Waller & Stanley, 2005) reported *Atrina* from the Ladinian. Also Stiller, in his doctoral thesis (Stiller, personal communication, 2008), reported it from the Anisian. Although, as noted, it may have appeared before, we will take this well-documented record as its first appearance. In addition, Waller and Stanley (2005, p. 29–30) noted that Carboniferous Pinnidae could possibly be referable to *Atrina* and not to *Pinna*.

Paleogeographic distribution.—Tethys and Circumpacific (Fig. 18). Although in the past *Atrina* was regarded as a cosmopolitan genus, this is not the case for the time interval here considered. Its distribution was surely greater than the one mentioned below, because many species attributed to *Pinna* may belong to *Atrina* instead, but there are no specific data for the study interval.

Circumpacific domain: Middle Triassic: late Ladinian of Nevada (United States) (Waller & Stanley, 2005).

Tethys domain: Middle Triassic: Anisian of China (Stiller, 2001, personal communication, 2008).

Paleoautoecology.—B, Se, S, Endo, Sed; By. *Atrina* living species are semi-infaunal bivalves that live endobyssate, attached by a strong byssus to rock fragments or other materials embedded in the sediment, with the commissure plane being almost vertical, similar to *Pinna*. Unlike *Pinna*, which lives with two-thirds of its shell into the sediment, *Atrina* lives almost completely buried (García-March, Márquez-Aliaga, & Carter, 2008). The same mode of life is assumed for Mesozoic specimens. The external shell spines were interpreted by S. M. Stanley (1970) as protection against breakage of the exposed portion of the shell, rather than a defensive device as in other bivalves.

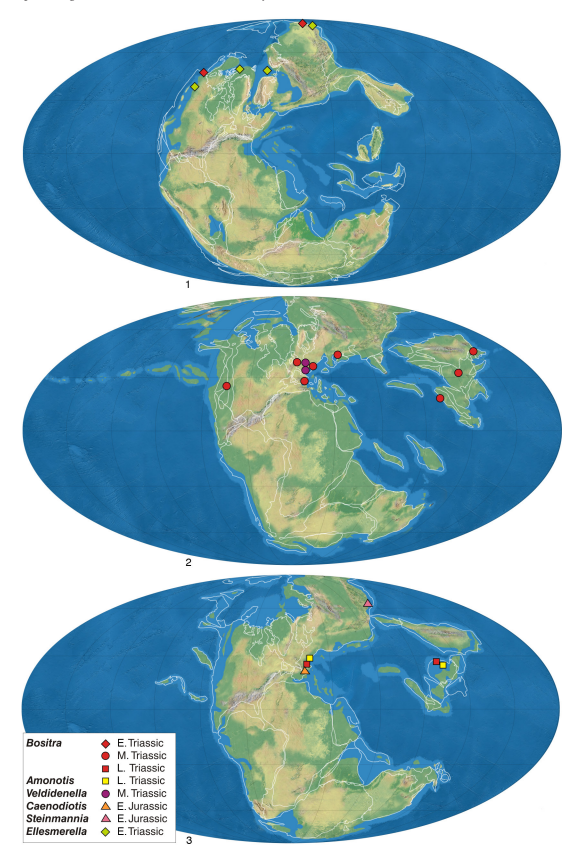


Figure 17. Paleogeographical distribution of Posidoniidae (*Bositna, Amonotis, Veldidenella, Caenodiotis, Steinmannia, Ellesmerella*). 1, Early Triassic; 2, Middle Triassic; 3, Late Triassic–Early Jurassic.

Mineralogy.—Bimineralic (Carter, 1990a; García-March, 2005). Outer shell layer: calcite (simple prismatic). Inner shell layer: aragonite (nacreous).

Superfamily LIMOIDEA Rafinesque, 1815 Family LIMIDAE Rafinesque, 1815 Genus PALAEOLIMA Hind, 1903 in 1896–1905, p. 38

Type species.—Pecten simplex Phillips, 1836, p. 212.

Remarks.—Newell (1999, p. 4) rejected *Palaeolima* because the material of the type species was lost and topotypes were not available. On the other hand, Waller and Stanley (2005, p. 32) stated that "Dickins (1963, p. 91), however, had earlier addressed this problem and designated a neotype of Phillips's species, specifically the specimen figured by Hind (1903 in 1896–1905, p. 19, fig. 26) from Little Island, County Cork, Ireland." According to these authors, *Palaeolima* remains valid.

Stratigraphic range.—Upper Devonian (?Fammenian)–Upper Triassic (Norian) (Lu, 1981; Waller & Stanley, 2005). Cox and others (1969) assigned it a Carboniferous–Late Triassic range. Following Waller and Stanley (2005), the range is extended back to the Upper Devonian. The youngest records are from the Upper Triassic of China (Lu, 1981; J. Chen, 1982a; Lu & Chen, 1986).

Paleogeographic distribution.—Tethys and Circumpacific (Fig. 19). *Palaeolima* had an almost cosmopolitan distribution during the Paleozoic, especially during the Carboniferous–Permian (until the Guadalupian) interval (Yancey, 1985; González, 1992; Hoare, 1993; Nakazawa, 1999, 2002; Sterren, 2000, 2004; Cisterna & Sterren, 2003; Waller & Stanley, 2005). During the Triassic, it is found only from the Tethys and Circumpacific domains.

Tethys domain: late Permian: Kashmir (India) (Brookfield, Twitchett, & Goodings, 2003), China (Y. Zhang, 1981; M. Wang, 1993; L. Li, 1995); Early Triassic: China (Ling, 1988); Middle Triassic: Anisian of southern China (Komatsu, Akasaki, & others, 2004); Late Triassic: China (J. Chen, 1982a; Lu & Chen, 1986; Gou, 1993); Carnian of Italy (Corazzari & Lucchi-Garavello, 1980); Norian of China (Lu, 1981).

Circumpacific domain: Middle Triassic: Ladinian of Nevada (United States) (Waller & Stanley, 2005).

Paleoautoecology.-B, E, S, Epi, Sed; By. Living members of the family Limidae live epibyssate or reclined on the substrate or fixed by a slender byssus that may break, allowing occasional swimming (S. M. Stanley, 1970). From his observations of living species, S. M. Stanley (1970) concluded that good swimmers are often equivalve with symmetrical, equally sized auricles, and they have a large umbonal angle. In terms of external shell morphology, Palaeolima can be compared to the species *Lima lima* (Linnaeus, 1758), which lives epibyssate with a strong byssus. But the species assigned to Palaeolima lack a byssal notch. Other species, such as Lima scabra (Born, 1778) and Lima hians (Gmelin, 1791), are fairly symmetrical, live epibyssate with a weak byssus that can break, and swim occasionally (S. M. Stanley, 1970). Palaeolima has a rather symmetrical shell, and the auricles are of equal size, but the umbonal angle does not normally exceed 80° (from illustrations in the literature). We assume that Palaeolima lived epibyssate, as did P. scabra, although it is very unlikely that it could swim.

Mineralogy.—Bimineralic (Carter, 1990b, p. 345). Outer shell layer: calcite (simple prismatic). Middle and inner shell layers: aragonite (cross-lamellar).

Genus AVICULOLIMA E. Philippi, 1900, p. 622

Type species.—*Aviculolima jaekeli* E. Philippi, 1900, p. 622. *Remarks.*—Although *Aviculolima* is externally similar to *Pteria*, Cox and others (1969) included it in the Limidae with doubts. Having no further information, we follow these authors in their allocation.

Stratigraphic range.—Middle Triassic (Anisian) (Diener, 1923). The only information we have about *Aviculolima* was given by Diener (1923) and Cox and others (1969). In both papers, the authors limit themselves to transcribing data from the original paper in which the genus was proposed. The genus was reported from the Lower Muschelkalk of northern Germany (probably Anisian).

Paleogeographic distribution.—western Tethys (Fig. 19). According to available information, the genus appears to be endemic to northern Germany.

Tethys domain: Middle Triassic: Lower Muschelkalk of northern Germany (Diener, 1923).

Paleoautoecology.—B, E, S, Epi, Sed; By. Given its external resemblance to *Pteria*, probably it was an epifaunal and byssate bivalve, although the generic diagnosis does not mention byssal structures.

Mineralogy.—Bimineralic (Cox & others, 1969; Carter, 1990a, p. 215). There is no information about *Aviculolima* shell. Data provided for family Limidae. Outer shell layer: calcite (prismatic). Inner shell layer: aragonite (non-nacreous).

Genus BADIOTELLA Bittner, 1890, p. 94

Type species.—Badiotella schaurothiana Bittner, 1895, p. 201. (See notes in Bittner, 1895, p. 200, and Cox & others, 1969, p. 386, related to the nomenclatural status of this genus and its type species).

Stratigraphic range.—Middle Triassic (Ladinian)–Upper Triassic (Carnian) (Diener, 1923). Although Cox and others (1969) assigned it a Ladinian range, there is evidence that the genus was also present in the Upper Triassic (see Bittner, 1895; Broili, 1904; Diener, 1923). Sepkoski (2002) assigned it a Ladinian-Carnian range, following the compilation made by Hallam (1981). The youngest records are from Carnian beds (see paleogeographic distribution).

Paleogeographic distribution.—Tethys (Fig. 19).

Tethys domain: Middle Triassic: China (Lu & Chen, 1986); Ladinian of the Alps (Broili, 1904; Cox & others, 1969); Late Triassic: Carnian of the Alps (Bittner, 1895; Broili, 1904; Diener 1923), China (Gou, 1993).

Paleoautoecology.—B, E, S, Epi-Un, Sed; By-R. Like all members of this family, *Badiotella* was an epifaunal bivalve. Byssal structures are not reported in published literature, so it possibly lived slightly reclined or fixed by a weak byssus.

Mineralogy.—Bimineralic (Cox & others, 1969; Carter, 1990a, p. 215). There are no data about *Badiotella* mineralogy or shell microstructure. Information provided for family Limidae. Outer shell layer: calcite (prismatic). Inner shell layer: aragonite (non-nacreous).

Genus LIMATULA Wood, 1839, p. 235

Type species.—Pecten subauriculata Montagu, 1808, p. 63.

Remarks.—Limatula and *Limea* Bronn, 1831, are genera with living representatives and conservative morphology. Even in Recent species, there is some confusion about which taxa should be referred to one genus or the other (Allen, 2004), and this distinction is much more complicated with fossil specimens.

Stratigraphic range.—Middle Triassic (Ladinian)–Recent. Cox and others (1969) assigned it a Triassic–Holocene range. The oldest record is Ladinian (Tamura, 1973); we did not find any records from the Lower Triassic, or from the Lower Jurassic, although from Toarcian times onward, it was fairly common throughout the Jurassic (Hallam, 1976, 1977, 1981; Fürsich, 1982; Pugaczewska, 1986; Komatsu, Saito, & Fürsich, 1993; Liu, 1995; Sha and others, 1998; J. Yin & Grant-Mackie, 2005).

Paleogeographic distribution.—Tethys, Circumpacific, and Austral (Fig. 19).

Tethys domain: Middle Triassic: Ladinian of Malaysia (Tamura, 1973); Late Triassic: Rhaetian of Italy (Chiesa, 1949), Hungary (Vörös, 1981).

Circumpacific domain: Late Triassic: Norian of Japan (Tokuyama, 1959b; doubtful record also in Nakazawa, 1963).

Austral domain: Late Triassic: Carnian of New Zealand (Trechmann, 1918).

Paleoautoecology.—B, E, S, Epi, FaM; By-Sw. Some living species, such as *Limatula strangei* (G. B. Sowerby, 1872), live among rocks and corals, and they are able to swim (Beesley, Ross, & Wells, 1998). They have equivalve and slightly inequilateral shells and subequal auricles. The shell morphology remained practically unchanged from the Triassic to the present (Allen, 2004); we assume that Mesozoic species had similar modes of life. *Limatula* probably lived byssate most of the time and reclined on the substrate, with the anterior part down (Fürsich, 1982).

Mineralogy.—Bimineralic (Carter, 1990b, p. 345). Outer shell layer: calcite (prismatic). Inner shell layer: aragonite (?).

Genus LIMEA Bronn, 1831, p. 115

Type species.—Ostrea strigilata Brocchi, 1814, p. 571.

Remarks.—The only subgenus considered by Cox and others (1969) in our study interval is *Limea (Eolimea)* Cox in Cox and others, 1969, p. 389, with a Middle Triassic range. From the Middle Triassic to the Miocene, when *Limea (Limea)* appears, there is a long time interval without records of this genus. This was noticed by Dhondt (1989), who also noted that *Pseudolimea* Arkell in Douglas & Arkell, 1932, which ranges from Triassic to Cretaceous, was distinguished from *Limea*, such as *Isolimea* or *Eolimea*, are differentiated by their strong ornamentation. Dhondt noted that *Pseudolimea* was very similar to *Limea* and it could fill this time gap, and she included it as a subgenus of *Limea*. This arrangement is followed here.

Stratigraphic range.—Middle Triassic (Anisian)–Holocene (Cox & others, 1969). The stratigraphic range of *Limea* recognized here is the same as in Cox and others (1969). The oldest record is from Anisian times (Kaim, 1997).

Paleogeographic distribution.—Cosmopolitan (Fig. 19).

Tethys domain: Middle Triassic: Poland (Kaim, 1997); Anisian western Caucasus (Russia) (Ruban, 2006a); Ladinian of Spain (Márquez-Aliaga, 1983; Pérez-López, 1991; Pérez-López & others, 1991; López-Gómez & others, 1994; Márquez-Aliaga & Martínez, 1996; Márquez-Aliaga, García-Forner, & Plasencia, 2002), Italy (Rossi Ronchetti, 1959); Late Triassic: Norian of southern China (J. Chen & Yang, 1983); Early Jurassic: Hettangian of northern Alps (Austria) (Golebiowski, 1990); Hettangian–Sinemurian of England (Liu, 1995), Spain (Calzada, 1982); Sinemurian of France, Portugal, and Morocco (Liu, 1995), Turkey (M. A. Conti & Monari, 1991).

Circumpacific domain: Late Triassic: Carnian of Japan (Nakazawa, 1952; Hayami, 1975); Norian of Oregon (United States) (Newton in Newton & others, 1987), Nevada (United States) (Laws, 1982); Norian of Chile (Hayami, Maeda, & Ruiz-Fuller, 1977); Rhaetian of Chile (Chong & Hillebrandt, 1985); Early Jurassic: Hettangian– Sinemurian of western Canada (Aberhan, 1998a; Aberhan, Hrudka, & Poulton, 1998), Chile (Aberhan, 1994a).

Austral domain: Late Triassic: Carnian of New Zealand (Grant-Mackie, 1960); Rhaetian of New Zealand (MacFarlan, 1998) and Argentina (Damborenea & Manceñido, 2012); Early Jurassic: Hettangian–Sinemurian of Neuquén Basin (Argentina) (Damborenea, 1996a; Damborenea & Manceñido, 2005b), New Zealand (MacFarlan, 1998).

Boreal domain: Early Jurassic: Hettangian–Sinemurian of ?Greenland (Liu, 1995).

Paleoautoecology.—B, E, S, Epi, Sed; By. *Limea* is a long-ranging genus that exhibits a conservative morphology throughout its history (Allen, 2004). The Recent species of this genus live mainly in deep water, and it was proposed that the extinct species also lived mostly in this type of environment (Dhondt, 1989). *Limea* species probably lived as epibyssate and reclined on one of the valves (Fürsich, 1982), similar to living species.

Mineralogy.—Bimineralic (Cox & others, 1969; Carter, 1990a, p. 215). There are no data about *Limea* shell. Data provided for family Limidae. Outer shell layer: calcite (prismatic). Inner shell layer: aragonite (non-nacreous).

Genus MYSIDIOPTERA Salomon, 1895, p. 117

Type species.—Mysidioptera ornata Salomon, 1895, p. 117. *Remarks.*—Two subgenera are considered in the study interval,

M. (Mysidioptera) and *M. (Pseudacesta)* Waagen, 1907, p. 113.

Stratigraphic range.—Lower Triassic (Olenekian)–Upper Triassic (Rhaetian). Cox and others (1969) assigned it a Lower Triassic–Upper Triassic range, and this is maintained here. *Mysidioptera* was not abundant in the Triassic, but it had a climax during Ladinian and Carnian; and although it was scarce during the Rhaetian, it lived to the end of the stage, when it became extinct.

Paleogeographic distribution.—Cosmopolitan (Fig. 19).

Tethys domain: Middle Triassic: Anisian of China (Wen & others, 1976; Lu & Chen, 1986; Sha, Chen, & Qi, 1990; Komatsu, Chen, & others, 2004; Komatsu, Akasaki, & others, 2004), Malaysia (Tamura & others, 1975), Swiss Dolomites (Zorn, 1971), Italian Dolomites (Posenato, 2008b), Israel (Lerman, 1960), northern Vietnam (Komatsu, Huyen, & Huu, 2010); Ladinian of Alps (Austria) (Salomon, 1895), north of Vietnam and Thailand (Vu Khuc & Huyen, 1998), Malaysia (Tamura, 1973), Lombardy (Italy) (Rossi Ronchetti, 1959), northern Vietnam (Komatsu, Huyen, & Huu, 2010); Late Triassic: China (Gou, 1993); Carnian of southern Alps (Bittner, 1895, 1900; Salomon, 1895; Broili, 1904; Allasinaz, 1966; Fürsich & Wendt, 1977; Posenato, 2008a, 2008b), Carpathians (Slovakia) (Bujnovsky, Kochanová, & Pevny, 1975; Kochanová, Mello, & Siblík, 1975), Jordan (Cox, 1924); Rhaetian of Tibet (China) (Hallam & others, 2000), East of the Alps (Austria) (Tomašových, 2006a, 2006b).

Circumpacific domain: Early Triassic: Olenekian of Japan (Nakazawa, 1961; Hayami, 1975); Late Triassic: Japan (Hayami, 1975), Norian of Oregon (Newton in Newton & others, 1987), Norian of Sonora (Mexico) (Damborenea in Damborenea & González-León, 1997).

Austral domain: Late Triassic: Carnian of New Zealand (Waterhouse, 1960).

Boreal domain: Late Triassic: Carnian of Arctic area of British Columbia (Canada) (Tozer, 1962, 1970).

Paleoautoecology.—B, E, S, Epi, Sed; By. The external shell morphology of different species attributed to *Mysidioptera* indicates an epibyssate habit, since most show a byssal notch. It could live on both hard and soft substrates (Newton in Newton & others, 1987). They were reported from a variety of facies, as they are thought to have colonized different environments (Damborenea in Damborenea & González-León, 1997).

Mineralogy.—Bimineralic (Cox & others, 1969; Carter, 1990a, p. 215). There are no data about *Mysidioptera* shell mineralogy. Data provided for family Limidae. Outer shell layer: calcite (prismatic). Inner shell layer: aragonite (non-nacreous).

Genus PLAGIOSTOMA J. Sowerby, 1814, p. 175

Type species.—Plagiostoma gigantea J. Sowerby, 1814, p. 176.

Stratigraphic range.—Middle Triassic (Anisian)–Upper Cretaceous (Maastrichtian) (Abdel-Gawad, 1986; Komatsu, Chen, & others, 2004). Cox and others (1969) assigned it a Middle Triassic–Cretaceous range. However, Sepkoski (2002) considered it to be from the Lower Triassic (Induan), indicating that data were taken from Abdel-Gawad (1986), but this author only mentioned *Plagiostoma* from the Cretaceous. This datum is wrong, since *Plagiostoma* was not present before the Middle Triassic (see Paleogeographic distribution, below), and it may have derived from Lower Triassic *Mysidioptera* (Bittner, 1895; Waller & Stanley, 2005). Dagys and Kurushin (1985) reported *Plagiostoma aurita* (Popov) and *Plagiostoma popovi* Kurushin in Dagys & Kurushin, 1985, from the Lower Triassic, but the figured specimens have little in common with the diagnosis of the genus *Plagiostoma*.

Paleogeographic distribution.—Cosmopolitan (Fig. 19).

Tethys domain: Middle Triassic: Hungary (Szente, 1997), Poland (Kaim, 1997); Anisian of Germany (Hautmann, 2006a), southern China (Sha, Chen, & Qi, 1990; Komatsu, Chen, & others, 2004), Switzerland (Zorn, 1971); Ladinian of Sardinia (Italy) (Posenato, 2002; Posenato & others, 2002), Malaysia (Tamura, 1973; Tamura & others, 1975); Late Triassic: China (J. Chen, 1982a; Gou, 1993), Oman (R. Hudson & Jefferies, 1961); Carnian of Malaysia (Tamura & others, 1975), Lombardy (Italy) (Allasinaz, 1966); Norian of Himalaya (Tibet, China) (J. Yin, Enay, & Wan, 1999), southern China (Wen & others, 1976; Sha, Chen, & Qi, 1990), northwestern China (Lu, 1981); Norian–Rhaetian of Iran and the Alps (Hautmann, 2001b); Rhaetian of eastern Alps (Austria) (Tomašových, 2006a, 2006b), Tibet (China) (Hautmann & others, 2005; J. Yin & McRoberts, 2006), Hungary (Vörös, 1981), Lombardy (Italy) (Allasinaz, 1962); Early Jurassic: Hettangian of southern England (Ivimey-Cook & others, 1999), Tibet (China) (Hautmann & others, 2005; J. Yin & McRoberts, 2006), Italy (Gaetani, 1970); Hettangian– Sinemurian of England and France (Liu, 1995), Lombardy (Italy) (Allasinaz, 1962); Sinemurian of Caucasus (southwestern Russia) (Ruban, 2006b), Portugal and Morocco (Liu, 1995).

Circumpacific domain: Middle Triassic: Ladinian of Nevada (United States) (Waller & Stanley, 2005); Late Triassic: Japan (Kobayashi & Ichikawa, 1949a; Tokuyama, 1959a); Carnian of Japan (Hayami, 1975); Norian of Oregon (United States) (Newton in Newton & others, 1987; but see Waller & Stanley, 2005); Early Jurassic: ?Mexico (Damborenea in Damborenea & González-León, 1997); Hettangian–Sinemurian of western Canada (Aberhan, 1998a), Mexico and Texas (Liu, 1995), northern Chile (Aberhan, 1994a); Sinemurian of Japan (Hayami, 1975; Hayami in Sato & Westermann, 1991).

Austral domain: Early Jurassic: Sinemurian of Neuquén Basin (Argentina) (Damborenea, 1996a; Damborenea & Manceñido, 2005b).

Boreal domain: Late Triassic: Carnian of Primorie (Kiparisova, 1972).

Paleoautoecology.—B, E, S, Epi, Sed; By. Like most limids, *Plagiostoma* was probably an epibyssate bivalve. Although no description pointing to a byssal notch was found, we assume that the byssus would have emerged below the anterior auricle. We cannot rule out that it could eventually swim, but this is unlikely due to the shell thickness (Seilacher, 1984); a reclining habit on its broad anterior base is more likely.

Mineralogy.—Bimineralic (Carter, 1990a, p. 215; Carter, 1990b, p. 345). Outer shell layer: calcite (prismatic). Inner shell layer: aragonite (non-nacreous).

Genus SERANIA Krumbeck, 1923a, p. 218

Type species.—Serania seranensis Krumbeck, 1923a, p. 218.

Stratigraphic range.—Upper Triassic (Norian–Rhaetian) (Hautmann, 2001b). According to Kutassy (1931), *Serania* was proposed by Krumbeck (1923a) on the basis of material from Norian beds of Indonesia and Persia. Cox and others (1969) assigned it a Norian age, as did also Sepkoski (2002), based on Hallam's (1981) data. It was subsequently reported also from Rhaetian beds (Hautmann, 2001b).

Paleogeographic distribution.—Eastern Tethys (Fig. 19). Serania was a monospecific genus endemic for the eastern Tethys.

Tethys domain: Late Triassic: Norian of Indonesia and Persia (Kutassy, 1931); Norian–Rhaetian of Iran (Hautmann, 2001b).

Paleoautoecology.—B, E, S, Epi, Sed; By. Like most members of family Limidae, *Serania* was probably an epibyssate bivalve, similar to *Plagiostoma* (Hautmann, 2001b). *Serania seranensis* shows a deep byssal notch (see Cox & others, 1969, p. 392), which implied a strong byssus.

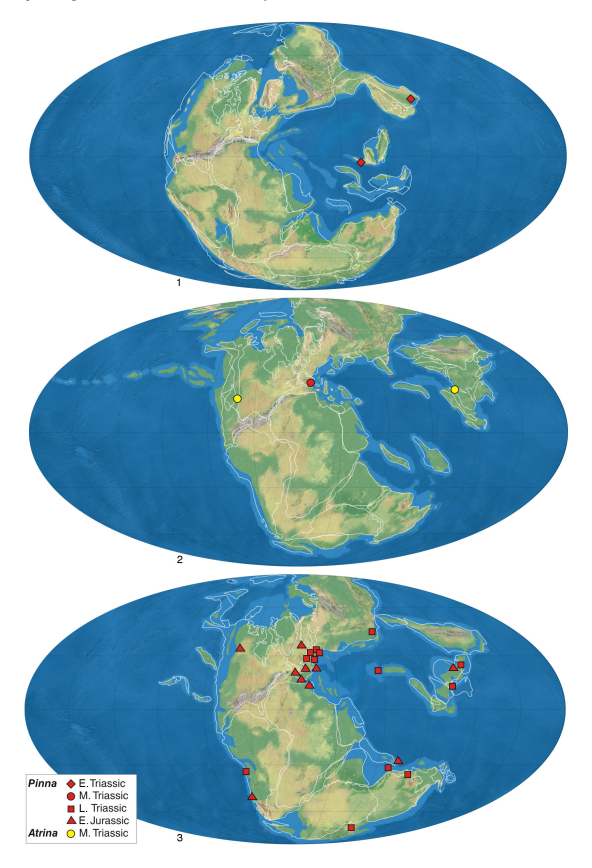


Figure 18. Paleogeographical distribution of Pinnidae (Pinna, Atrina). 1, Early Triassic; 2, Middle Triassic; 3, Late Triassic-Early Jurassic.

Mineralogy.—Bimineralic (Cox & others, 1969; Carter, 1990a, p. 215). There are no data about the shell of *Serania*. Data provided for family Limidae. Outer shell layer: calcite (prismatic). Inner shell layer: aragonite (non-nacreous).

Genus TIROLIDIA Bittner, 1895, p. 202

Type species.—Lima (Tirolidia) haueriana Bittner, 1895, p. 202. Stratigraphic range.—Middle Triassic (Ladinian)–Upper Triassic (Carnian) (Diener, 1923). Cox and others (1969) assigned a Middle Triassic–Upper Triassic range. Bittner (1895) proposed *Tirolidia* from Ladinian and Carnian beds of the southern Alps. Diener (1923) and Kutassy (1931) provided the same data. Little else could be found, except that Hallam (1981) assigned it a Ladinian–Carnian range in western Tethys, and thus this genus seems to be endemic to the southern Alps.

Paleogeographic distribution.—western Tethys (Fig. 19).

Tethys domain: Middle Triassic: Ladinian of Southern Alps (Bittner, 1895); Late Triassic: Carnian of Southern Alps (Bittner, 1895).

Paleoautoecology.—B, E, S, Epi, Sed; By. *Tirolidia* is slightly inequilateral and has unequal auricles, so it is not a good candidate to be a swimmer. We assign it an epibyssate mode of life, similar to other members of this family.

Mineralogy.—Bimineralic (Cox & others, 1969; Carter, 1990a, p. 215). No specific data about *Tirolidia* mineralogy and shell microstructure are known. Data provided for family Limidae. Outer shell layer: calcite (prismatic). Inner shell layer: aragonite (non-nacreous).

Genus ANTIQUILIMA Cox, 1943, p. 179

Type species.—Lima antiquata J. Sowerby, 1818, p. 25.

Stratigraphic range.—Middle Triassic (Ladinian)–Lower Cretaceous (Aptian) (Hayami, 1965; Waller & Stanley, 2005). Cox and others (1969) considered it to be a Jurassic genus (Liassic–Bajocian), but, since then, new records have expanded its range. The oldest record of *Antiquilima* is from Ladinian beds of Nevada (Waller & Stanley, 2005) and the youngest from Lower Cretaceous beds (Aptian) (Hayami, 1965).

Paleogeographic distribution.—Cosmopolitan (Fig. 19). According to Damborenea (2000), *Antiquilima* probably originated in the Eastern Pacific and subsequently it spread to the Tethys, which agrees with the data found about the genus.

Tethys domain: Late Triassic: Norian–Rhaetian of Iran (Hautmann, 2001b); Rhaetian of the Alps (Austria) (Tomašových, 2006a, 2006b), Tibet (China) (J. Yin & McRoberts, 2006), western Carpathians (Slovakia) (Tomašových, 2004); Early Jurassic: Hettangian–Sinemurian of England (Liu, 1995); Sinemurian of France (Vörös, 1971; Liu, 1995), Apennines (Italy) (Monari, 1994).

Circumpacific domain: Middle Triassic: Ladinian of Nevada (United States) (Waller & Stanley, 2005); Late Triassic: Norian of northern Chile (Hayami, Maeda, & Ruiz-Fuller, 1977), Oregon (United States) (Newton, 1986; Newton in Newton & others, 1987); Early Jurassic: Hettangian–Sinemurian of northern Chile (Aberhan, 1994a); Sinemurian of Canada (Aberhan, 1998a), Japan (Hayami, 1975). Austral domain: Early Jurassic: Sinemurian of Argentina (Damborenea, 1996a; Damborenea & Manceñido, 2005b; Damborenea & Lanés, 2007).

Boreal domain: Late Triassic: Carnian of Primorie (Kiparisova, 1972); Norian of northeastern of Asia (Kurushin, 1990; Polubotko & Repin, 1990); Norian–Rhaetian of eastern Siberia (Kiparisova, Bychkov, & Polubotko, 1966), northeastern Russia (Milova, 1976); Early Jurassic: Hettangian of northeastern Asia (Kurushin, 1990; Polubotko & Repin, 1990), northeastern Russia (Milova, 1976).

Paleoautoecology.—B, E, S, Epi, Sed; By. In most species, a byssal notch is present (e.g., specimens described in Hayami, Maeda, & Ruiz-Fuller, 1977; Newton in Newton & others, 1987; Hautmann, 2001b), and most likely it was an epibyssate bivalve as in the rest of the limids. According to Newton (in Newton & others, 1987), *Antiquilima* could sever the byssus and swim for short distances, as do some modern species of the family Limidae. However, due to its external morphology, it was probably not a good swimmer.

Mineralogy.—Bimineralic (Cox & others, 1969; Carter, 1990a, p. 215; Waller & Stanley, 2005). Outer shell layer: calcite (prismatic). Inner shell layer: aragonite (non-nacreous).

Genus CTENOSTREON Eichwald, 1862, p. 374

Type species.—Ostracites pectiniformis von Schlotheim, 1820, p. 231.

Stratigraphic range.—Upper Triassic (upper Rhaetian)–Lower Cretaceous (?Valanginian). Cox and others (1969) assigned it a Jurassic (Liassic)–Lower Cretaceous (Neocomian) range. Sepkoski (2002) considered that it originated in the lower Hettangian following Hallam (1977, 1987). The origin of this genus was regarded as Hettangian for a long time (Hallam, 1977, 1987, 1990), but recently J. Yin, H. Yao, and Sha (2004) and J. Yin and McRoberts (2006) found *Ctenostreon* in layers transitional between the Rhaetian and Hettangian. These Himalayan records were dated as Rhaetian, because they were associated with the ammonoid *Choristoceras* (J. Yin, H. Yao, & Sha, 2004). We ignore to what part of the Neocomian Cox and others (1969) referred for the last record. The youngest record of the genus dates from Valanginian times (Császár & Turnšek, 1996), but specimens were neither figured nor described, so we tentatively consider this as the last appearance.

Paleogeographic distribution.—Tethys, Austral, and Circumpacific (Fig. 19).

Tethys domain: Late Triassic: Rhaetian of southern China (J. Yin, H. Yao, & Sha, 2004; J. Yin & McRoberts, 2006); Early Jurassic: Hettangian of Tibet (China) (J. Yin & McRoberts, 2006; J. Yin & others, 2007).

Austral domain: Early Jurassic: Hettangian–Sinemurian of the Neuquén Basin (Argentina) (Damborenea, 1996a; Damborenea & Manceñido, 2005b).

Circumpacific domain: Early Jurassic: Sinemurian of Japan (Hayami, 1975), Chile (Aberhan, 1994a).

Paleoautoecology.—B, E, S, Epi, Sed; By. *Ctenostreon* is regarded as an epibyssate bivalve, although the byssal notch is not always evident and sometimes it is even absent. Its shell is thick compared with other members of this family; we assume that it would not need a very strong byssus. Seilacher (1984) suggested that it is one of those limids for which a swimming mode of life is excluded, due to its

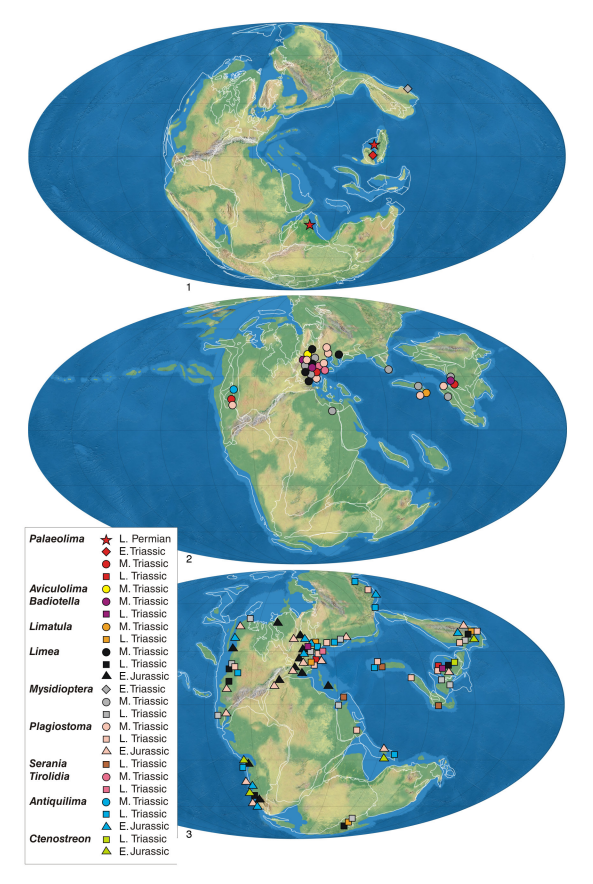


Figure 19. Paleogeographical distribution of Limidae (*Palaeolima, Aviculolima, Badiotella, Limatula, Limea, Mysidioptera, Plagiostoma, Serania, Tirolidia, Antiquilima, Ctenostreon). 1*, late Permian–Early Triassic; 2, Middle Triassic; 3, Late Triassic–Early Jurassic.

thick shell and presence of spines; it was probably a pleurothetic reclined bivalve.

Mineralogy.—Bimineralic (Cox & others, 1969; Carter, 1990a, p. 215). No specific data about *Ctenostreon* mineralogy and shell micro-structure is known. Information provided for family Limidae. Outer shell layer: calcite (prismatic). Inner shell layer: aragonite (non-nacreous).

Superfamily OSTREOIDEA Wilkes, 1810 Family GRYPHAEIDAE Vialov, 1936 Genus GRYPHAEA Lamarck, 1801, p. 398

Type species.—Gryphaea arcuata Lamarck, 1801, p. 398.

Remarks.—Newell and Boyd (1970, 1989, 1995) discussed the external morphological similarity between *Pseudomonotis* and *Gryphaea* (see fig. 47 in Newell & Boyd, 1995). Frequently, these genera can only be distinguished by the shell microstructure and which valve is attached to the substrate (right and left respectively) (Newell & Boyd, 1995). Many of the *Pseudomonotis* records from Triassic and Jurassic could actually belong to *Gryphaea*, since *Pseudomonotis* is now regarded as a strictly Paleozoic genus, and it is not an ostreoid.

Stratigraphic range.—Upper Triassic (Carnian)–Upper Cretaceous (Campanian) (McRoberts, 1992; Newell & Boyd, 1989). Stenzel (1971) considered that *Gryphaea* was present in the Upper Triassic of the Boreal domain and had a worldwide distribution for most of the Jurassic (Hettangian–Oxfordian). New records changed the observed stratigraphic range of this genus: *Gryphaea* was reported from Upper Triassic beds, from the Carnian (McRoberts, 1992) in the Paleopacific eastern margin, in addition to the Boreal regions. It had not been found anywhere in upper Norian and Rhaetian beds, so McRoberts (1992) interpreted it as a Lazarus taxon that reappeared in the Hettangian stage, but Rubilar (1998) reported *Gryphaea* from the Norian–Rhaetian of Chile.

The youngest record is from the Upper Cretaceous (Newell & Boyd, 1989). Although there are some papers that mentioned *Gryphaea* up until the Pleistocene, they are not taken into account as they are bio-stratigraphic studies and they do not describe or figure the listed material.

Paleogeographic distribution.—Cosmopolitan (Fig. 20).

Tethys domain: Early Jurassic: Tibet (China) (Gou, 2003); Hettangian of Italy (Gaetani, 1970); Hettangian–Sinemurian of France (Liu, 1995; Nori & Lathuilière, 2003), England, Spain, Portugal, and Morocco (Liu, 1995).

Circumpacific domain: Late Triassic: Carnian–Norian of Canada, Oregon, and Nevada (United States) (McRoberts, 1992), northern and southern Alaska (McRoberts, 1992; McRoberts & Blodgett, 2000); Norian of Chile (Hayami, Maeda, & Ruiz-Fuller, 1977); Norian–Rhaetian of Chile (Rubilar, 1998); Early Jurassic: Hettangian of Chile (Aberhan, 1994a); Sinemurian of Chile (Steinmann, 1929; Chong & Hillebrandt, 1985; Hillebrandt, 1990; Aberhan, 1994a; Malchus & Aberhan, 1998; Rubilar, 1998), Canada (Poulton, 1991).

Austral domain: Early Jurassic: Sinemurian of Argentina (Damborenea & Manceñido, 2005b; Damborenea & Lanés, 2007).

Boreal domain: Late Triassic: Arctic area of Siberia (Kiparisova, 1954), northeastern Russia (Milova, 1976); Carnian of Arctic Island (Canada) (Tozer, 1970); Carnian–Norian of Primorie (Kiparisova, 1972); Early Jurassic: Hettangian of northeastern Russia (Milova, 1988).

Paleontological Contributions, number 8

Paleoautoecology.—B, E, S, Un, Sed; R. Some species of *Gryphaea* lived cemented to the substrate during the juvenile stages, but they often changed to a reclined life habit in the adult stage (Fürsich & others, 2001). Seilacher (1984) interpreted certain *Gryphaea* species as being cup-shaped recliners, living on soft substrates: they probably rested on their left valve, which is strongly convex and thick, unlike the flat and smooth right valve. The left valve was used to anchor the shell to soft sediments. *Gryphaea* would thus live epifaunally or partially buried. The specimens are usually found in fine-grained sediments (clay, marl) that are characteristic of low-energy marine environments (Lewy, 1976). The shell morphology of *Gryphaea*, as in other ostreids, is strongly influenced by the environment. Nori and Lathuilière (2003) proved that several factors (temperature, oxygen level, and humidity) were responsible for the different morphologies.

Mineralogy.—?Bimineralic (Carter, 1990a, p. 232). Although the shell microstructure of Triassic specimens is not known, Jurassic specimens have a prismatic outer shell layer and a foliated inner shell layer, both being calcitic (Carter, 1990a). However, McRoberts and Carter (1994) found that middle and inner layers of *G. nevadensis* were originally of nacreous microstructure (aragonite).

Family OSTREIDAE Wilkes, 1810 Genus UMBROSTREA Hautmann, 2001a, p. 359

Type species.—Umbrostrea emamii Hautmann, 2001a, p. 361. Remarks.—Hautmann (2001a) proposed Umbrostrea to include some specimens from the Upper Triassic of Iran that attached by the left valve (consensual basis for defining the true oysters), built reefs, and possessed an inner foliated shell microstructure and aragonitic inner shell layer (data considered as preliminary by the author). He proposed two new species, Umbrostrea emamii and Umbrostrea iranica, and tentatively considered Umbrostrea? aff. parasitica (Krumbeck, 1913) within the genus. Subsequently, Márquez-Aliaga and others (2005) examined a sample of hundreds of specimens attributed to Enantiostreon difforme (Goldfuss, 1833 in 1833–1841) [=Ostracites cristadifformis Schlotheim, 1820] and Enantiostreon spondyloides (Schlotheim, 1820) from the Lower and Upper Muschelkalk (Middle Triassic, Anisian-Ladinian) of the Germanic Basin, from levels equivalent to those from where the Enantiostreon species were described, attributed by these authors to real oysters. The authors accepted that the first record of ostreids from those levels was by Seilacher (1954), who classified some specimens attached by their left valve to Plagiostoma shells as Alectryonia (=Lopha), but with Enantiostreon morphology. Seilacher (1954) relied on the kind of so-called twisting of the valves and on the antimarginal pattern of the shell folds. This last criterion was developed by Checa and Jiménez-Jiménez (2003b) for cemented bivalves, and it is characteristic of ostreids. In the same paper, Márquez-Aliaga and others (2005) studied the Hispanic Muschelkalk (Ladinian) specimens attributed to E. difformis by Márquez-Aliaga (1985), on which microstructural studies were performed (De Renzi & Márquez-Aliaga, 1980; Márquez-Aliaga & Martínez, 1990b; Márquez-Aliaga & Márquez, 2000). In these studies, the authors verified the presence of foliated and calcitic outer shell layer and an inner shell layer replaced by sparite of possible aragonitic origin; this type of microstructure is characteristic of ostreids. However, the absence of internal features in all the studied

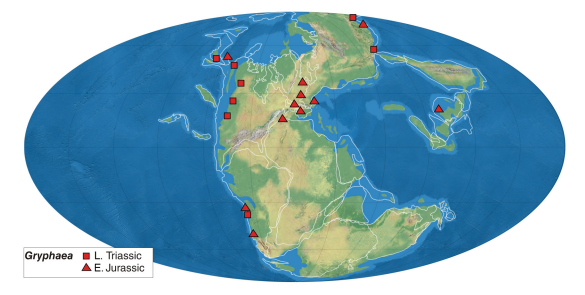


Figure 20. Paleogeographical distribution of Gryphaeidae (Gryphaea). Late Triassic-Early Jurassic.

specimens did not solve the controversial problem of the origin of the oysters. Thus, several replies to this proposal were generated, including other evolutionary aspects (see discussion in Márquez-Aliaga & others, 2005; Hautmann, 2006b; Checa & others, 2006; and Malchus, 2008). Other authors, like Ivimey-Cook and others (1999) and J. Yin and McRoberts (2006), preferred to include the species *Enantiostreon difforme* within *Terquemia* Cox, 1964. Here we provisionally accept the criteria of Márquez-Aliaga and others (2005).

Stratigraphic range.—Middle Triassic (Anisian)–Upper Triassic (Rhaetian) (Hautmann, 2001a; Márquez-Aliaga & others, 2005). Hautmann (2001a, 2001b) assigned it a Norian–Rhaetian range. Accepting the species assigned by Márquez-Aliaga and others (2005) into *Umbrostrea*, the range extends from the Middle Triassic, with the origin of *U. cristadifformis* and *U. spondyloides* being in Anisian times.

Paleogeographic distribution.—Tethys (Fig. 21).

Tethys domain: Middle Triassic: Anisian of Poland (Kaim, 1997), Bulgaria (Budurov & others, 1993), Hungary (Szente, 1997); Ladinian of Sardinia (Márquez-Aliaga & others, 2000; Posenato, 2002), Spain (Márquez-Aliaga, 1985; Márquez-Aliaga & Martínez, 1996); Late Triassic: Norian of Indonesia (Diener, 1923); Norian–Rhaetian of Iran (Hautmann, 2001a, 2001b; Fürsich & Hautmann, 2005), Rhaetian of England (Ivimey-Cook & others, 1999), ?Rhaetian of Tibet (China) (J. Yin & McRoberts, 2006).

Paleoautoecology.—B, E, S, C, Sed; C. Although Hautmann (2001a, 2001b) considered that *Umbrostrea* lived cemented to the substrate by the left valve, the species *U. cristadifformis* and *U. spondyloides* could attach by either valve (Márquez-Aliaga & others, 2005). It lived forming small reefs in fully marine environments and was associated with corals, brachiopods, and other bivalves (Hautmann, 2001a).

Mineralogy.—Bimineralic (De Renzi & Márquez-Aliaga, 1980; Carter, Barrera, & Tevesz, 1998; Hautmann, 2001a). In the diagnosis of *Umbrostrea*, Hautmann (2001a) indicated that the species attributed to this genus are characterized by a regular simple prismatic outer shell layer of calcite, a middle shell layer of foliated calcite, and an aragonitic inner shell layer of unknown microstructure. *Umbrostrea cristadifformis* had a foliated calcitic outer shell layer and an aragonitic inner shell layer (De Renzi & Márquez-Aliaga, 1980). According to Carter, Barrera, and Tevesz (1998), *U. spondyloides* (upper Muschelkalk, Ladinian, southwestern Germany) had an aragonitic inner shell layer and calcitic middle and upper shell layers, the last with regular to homogeneous and irregular prismatic microstructures. Outer shell layer: calcite (simple prismatic–foliated). Middle shell layer: calcite (?foliated). Inner shell layer: aragonite (?nacreous).

Genus ACTINOSTREON Bayle, 1878, expl. pl. 132

Type species.—Ostrea solitaria J. de C. Sowerby, 1824, p. 105.

Remarks.—Palaeolopha Malchus, 1990, is regarded a junior synonym of *Actinostreon* (see discussion under *Palaeolopha*, Genera not Included, p. 166).

Stenzel (1971) considered *Actinostreon* as a subgenus of *Lopha*, and this was followed by most authors. However, Malchus (1990) included *Actinostreon*, together with his new genus *Palaeolopha*, in his new family Palaeolophidae, and he regarded *Actinostreon* as an independent genus different from *Lopha*. Checa and Jiménez-Jiménez (2003b) included the species *Enantiostreon difforme* (Goldfuss, 1833 in 1833–1841) [=*Ostracites cristadifformis* Schlotheim, 1823 in 1822–1823] in *Actinostreon*, since Malchus (1990) included this species in *Palaeolopha*, and they followed the synonymy proposed by Hautmann (2001a, p. 359), i.e. *Actinostreon =Palaeolopha* Malchus, 1990). Subsequently, Márquez-Aliaga and others (2005) included the species *cristadifformis* in *Umbrostrea* Hautmann, 2001a.

Stratigraphic range.—Upper Triassic (Rhaetian)–Upper Cretaceous (Maastrichtian) (Chiplonkar & Badve, 1977; Hautmann, 2001a). Stenzel (1971) assigned it a Jurassic–Cretaceous range; these data were incorporated by Sepkoski (2002), who added Maastrichtian

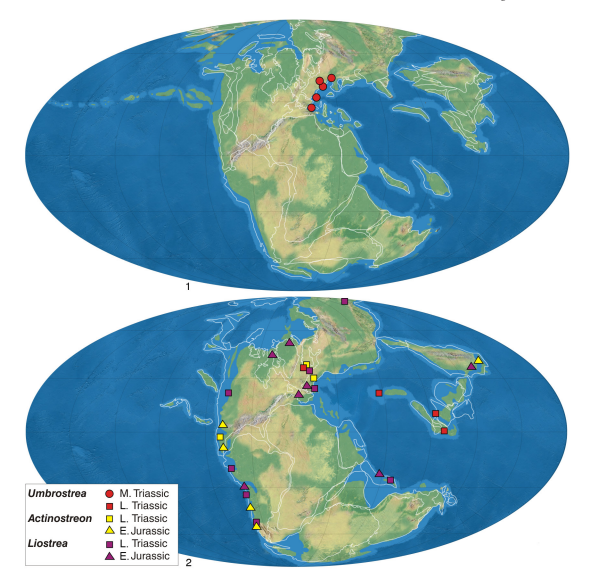


Figure 21. Paleogeographical distribution of Ostreidae (Umbrostrea, Actinostreon, Liostrea). 1, Middle Triassic; 2, Late Triassic-Early Jurassic.

as the last appearance, but he did not indicate the original source. The oldest records are from Rhaetian (Ivimey-Cook & others, 1999; Hautmann, 2001a). *Actinostreon* was very well represented throughout the Jurassic, and there are very few records from the Cretaceous. The youngest record is the species *Lopha (Actinostreon) diluvian* (Linnaeus) from Maastrichtian beds (Chiplonkar & Badve, 1977), quoted also by Ayyasami (2006) from the Turonian, both in southern India, although the latter is a biostratigraphic paper. However, this species is frequently assigned to *Lopha*, although according to Malchus (1990), *Lopha* had a Tertiary origin.

Paleogeographic distribution.—Tethys, Circumpacific, and Austral (Fig. 21).

Tethys domain: Late Triassic: Rhaetian of England (Penarth Group) (Ivimey-Cook & others, 1999), Austria (Hautmann, 2001a).

Circumpacific domain: Late Triassic: Norian of Mexico (Damborenea in Damborenea & González-León, 1997); Early Jurassic: Hettangian–Sinemurian of Mexico and Texas (United States) (Liu, 1995), Andes (Chile and Argentina) (Damborenea, 1996a, 2000); Sinemurian of Chile (Aberhan, 1994a; Sha, Smith, & Fürsich, 2002), Japan (Toyora Group) (Hayami, 1975).

Austral domain: Early Jurassic: Hettangian–Toarcian of the Andes (Chile and Argentina) (Damborenea, 1996a, 2000).

Paleoautoecology.—B, E, S, C, Sed; C. Actinostreon was a cemented bivalve that attached itself to the substrate by the left valve. Usually it formed clusters in high-energy marine environments (Sha, Smith, & Fürsich, 2002). It could attach to inorganic substrates and also to the shells of other organisms. Most often it attached by cementation to individuals of the previous generation, but it was also found on other bivalves (e.g., *Modiolus* in Ivimey-Cook & others, 1999) or solitary (Machalski, 1998). According to Sha (2002), ostreids have planktotrophic larvae that are responsible for their wide dispersion. *Mineralogy.*—Calcitic (Carter, 1990a; Hautmann, 2001a). According to Carter (1990a), *Lopha haidingeriana* (Emmrich, 1853) had predominantly foliated middle and inner shell layers, but a thin prismatic outer shell layer may also be present. Hautmann (2001a) found no trace of an aragonitic inner shell layer in one of his *Actinostreon haidingerianum* (Emmrich, 1853) specimens, and in a tangential section, he observed thin layers of foliated structure. The aragonite is limited to the miostracum and ligostracum (Hautmann, 2001a). The shells show a typical structure with biconvex chambers (Malchus, 1998). Outer shell layer: calcite (?prismatic). Middle and inner shell layers: calcite (regular foliated).

Genus LIOSTREA Douvillé, 1904, p. 273

Type species.—Ostrea sublamellosa Dunker, 1846, p. 41.

Stratigraphic range.—Upper Triassic (Carnian)–Upper Cretaceous (Cenomanian) (Hayami, 1975; Carter, 1990a). Stenzel (1971) reported *Liostrea* as being present in the Norian of Siberia and from the Rhaetian to the Jurassic of Europe. Subsequently, the genus was reported from Carnian beds of Japan (Hayami, 1975). The oldest record is from the Ladinian (Waller in Waller & Stanley, 2005), but, in these specimens, some diagnostic characters are not seen, and thus a definite identification is not possible. This record is regarded as dubious *Liostrea* until more material is found in the area. If confirmed, the origin of *Liostrea* goes back to the Middle Triassic. According to Carter (1990a), the youngest record is *Liostrea* oxiana Romer from the Cretaceous (Cenomanian) (Seeling & Bengtson, 1999).

Paleogeographic distribution.—Cosmopolitan (Fig. 21).

Tethys domain: Late Triassic: Carnian of China (J. Chen, 1982a), ?Italy (Gaetani, 1970); Rhaetian of Tibet (China) (?Hautmann & others, 2005; J. Yin & McRoberts, 2006), England (Ivimey-Cook & others, 1999), Italy (Gaetani, 1970); Early Jurassic: Hettangian of Tibet (China) (?Hautmann & others, 2005; J. Yin & McRoberts, 2006), England (Liu, 1995; Ivimey-Cook & others, 1999), Italy (Gaetani, 1970); Sinemurian of England and Portugal (Liu, 1995).

Circumpacific domain: Late Triassic: Carnian of ?Peru (Cox, 1949); Norian of Oregon (United States) (Newton, 1986; Newton in Newton & others, 1987); Norian–Rhaetian of Chile (Rubilar, 1998); Early Jurassic: Sinemurian of Japan (Hayami, 1975), Chile (Rubilar, 1998).

Austral domain: Late Triassic: Rhaetian of Argentina (Riccardi & others, 2004; Damborenea & Manceñido, 2012).

Boreal domain: Late Triassic: Norian of Siberia (Stenzel, 1971); Early Jurassic: Hettangian–Sinemurian of Greenland (Liu, 1995).

Paleoautoecology.—B-Ps, E, S, C, Sed-FaM; C. Liostrea cemented to the substrate by the left valve, like the other ostreids. Unlike Gryphaea, it has a large cementation area. Liostrea cemented itself to hard substrates, bivalve shells, or other organisms (Newton in Newton & others, 1987). It was usually found forming reefs during the Jurassic (Fürsich, Palmer, & Goodyear, 1994). However, the species Liostrea erina (d'Orbigny) was found cemented to ammonoids (Leioceras) in the Middle Jurassic Opalinum Clay (Switzerland), so it is supposed to have been pseudoplanktonic (Etter, 1996). This author found evidence indicating that the cementation was achieved when the ammonoids were still alive (see fig. 4 in Etter, 1996). Other species, such as L. plastica (Trautschold) from the Upper Jurassic of Greenland, were also found cemented to ammonoids, but it was not possible to determine whether the cementation was pre- or postmortem for the ammonites (Fürsich, 1982).

Mineralogy.—?Calcitic (Carter, Barrera, & Tevesz, 1998). There are no conclusive studies on *Liostrea* mineralogy or shell microstructure. According to Carter, Barrera, and Tevesz (1998), the mineralogy of the different shell layers of members of family Ostreidae is calcitic.

Superfamily DIMYOIDEA Fischer, 1886 in 1880–1887 Family DIMYIDAE Fischer, 1886 in 1880–1887 Genus ATRETA Etallon, 1862, p. 191

Type species.—Ostrea blandina d'Orbigny, 1850, p. 375 (designated by Cox, 1964, p. 45).

Remarks.—Dimyodon Munier-Chalmas in Fischer, 1886 in 1880–1887, p. 937, is considered to be a junior synonym of *Atreta* (see discussion for *Dimyodon*, Genera not Included, p. 160). Although for a long time it was regarded as a plicatulid, both Fürsich and Werner (1988) and Hodges (1991), analyzing specimens of *Atreta unguis* (Loriol *ex* Merian, 1900) and *Atreta intusstriata* (Emmrich, 1853), respectively, demonstrated the presence of dimyarian structures typical of family Dimyidae.

Stratigraphic range.—Upper Triassic (Carnian)–Upper Cretaceous (Maastrichtian) (Bittner, 1895; Abdel-Gawad, 1986). Cox and others (1969) assigned it an Upper Triassic (Carnian)–Upper Cretaceous (Campanian) range. Sepkoski (2002) referred its origin to the Rhaetian, based on data provided by Skelton and Benton (1993). The oldest records of *Atreta* are from Carnian beds, with the species *A. richthofeni* (Bittner, 1895) and *A. subrichthofeni* (Krumbeck, 1924). H. Yin (1985) reported *Dimyodon* from Anisian and Ladinian beds. J. Chen, Stiller, and Komatsu (2006) believed that Anisian specimens of *Dimyodon (D. qingyanensis* Yin in Gan & Yin, 1978) were actually juvenile stages of *Protostrea sinensis* Hsu in Hsu & Chen, 1943 (type of *Protostrea* Chen, 1976). *Atreta nilssoni* (von Hagenow, 1842) is the latest species of the genus, reported from the Maastrichtian (Abdel-Gawad, 1986).

Paleogeographic distribution.—Tethys (Fig. 22). Over the study interval, this genus was only known from the Tethys domain, but in the upper Pliensbachian, it is recorded from Argentina (Damborenea, 2002a). *Atreta* showed dispersal patterns from the western Tethys to Eastern Circumpacific domain across the Hispanic corridor during the Early Jurassic (Damborenea, 2000).

Tethys domain: Late Triassic: Carnian of southern Alps (Bittner, 1895; Fürsich & Wendt, 1977), Timor (Krumberck, 1924); Norian of Oman (Hautmann, 2001a); Norian–Rhaetian of Austria (Tanner, Lucas, & Chapman, 2004), Iran (Hautmann, 2001a, 2001b); Rhaetian of the Alps (Austria) (Tomašových, 2006a, 2006b), western Carpathians (Slovakia) (Tomašových, 2004), Austria and Germany (Hodges, 1991), England (Penarth Group) (Ivimey-Cook & others, 1999), Italy (Allasinaz, 1962); Early Jurassic: Hettangian of northwestern Europe (Hallam, 1987); early Liassic of South Wales (Hodges, 1991), Italy (Allasinaz, 1962).

Paleoautoecology.—B, E, S, C, Sed; C. Atreta was a cemented bivalve (fixed by its right valve) on other live invertebrates, such as sponges (Delvene, 2003; P. D. Taylor & Wilson, 2003), echinoids (Saint-Seine, 1951; Jagt, Neumann, & Schulp, 2007), other bivalves

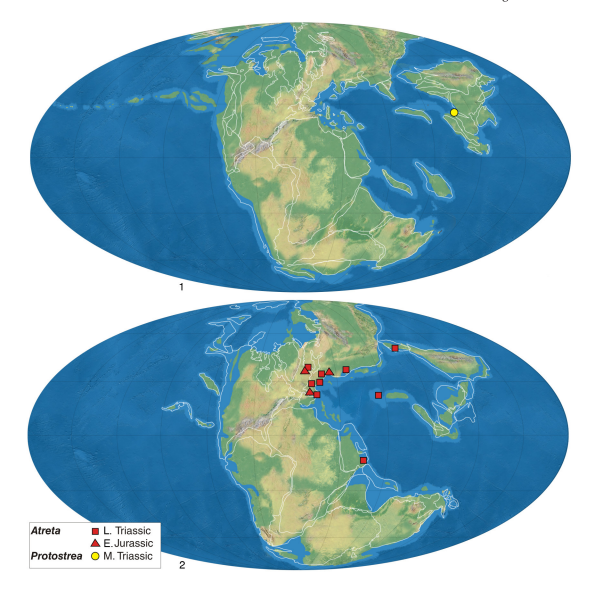


Figure 22. Paleogeographical distribution of Dimyidae (Atreta, Protostrea). 1, Middle Triassic; 2, Late Triassic-Early Jurassic.

such as *Plagiostoma*, *Gryphaea*, *Pinna*, *Antiquilima* (Hodges, 1991), *Indopecten* (Hautmann, 2006a), *Lopha*, *Cardinia*, *Myoconcha*, and corals (Damborenea, 2002a). It is usually associated with other encrusting bivalves such as *Liostrea* (Hodges, 1991) or *Lopha* (Damborenea, 2002a). It was a gregarious bivalve, although it is rare to find it encrusting other individuals of the same species, and specimens are usually oriented with their dorsal part upward on sloping surfaces (Damborenea, 2002a).

Mineralogy.—Bimineralic (Malchus, 2000). Hodges (1991) did not find any shell preserved, and he believed it was likely that it was originally aragonitic. Malchus (2000) studied the microstructure of lower stages of excellently preserved *Atreta* specimens and found a foliated calcitic in outer shell layer and a well-developed cross-lamellar microstructure in the inner shell layer. Hautmann (2001a, 2006a) indicated that his specimens have a foliated calcite microstructure in the outer shell layer, and they did not have the inner one preserved. Outer shell layer: calcite (foliated). Inner shell layer: aragonite (cross-lamellar).

Genus PROTOSTREA Chen in Gu & others, 1976, p. 243

Type species.—Ostrea sinensis Hsu in Hsu & Chen, 1943, p. 136. Remarks.—This genus was also called Proostrea (e.g., Skelton & Benton, 1993; Sepkoski, 2002) or Prostrea (e.g., Kobayashi & Tamura, 1983a) by mistake. Although its type species was originally included in Ostreoidea, following Morris in Skelton and Benton (1993), Komatsu, Akasaki, and others (2004), and J. Chen, Stiller, and Komatsu (2006), we include Protostrea in the Dimyidae (see J. Chen, Stiller, & Komatsu, 2006, for review and emendation of the genus). These authors interpreted Dimyodon qingyanensis Yin in Gan & Yin, 1978, as a juvenile stage of Protostrea sinensis and considered it the oldest member of the family Dimyidae.

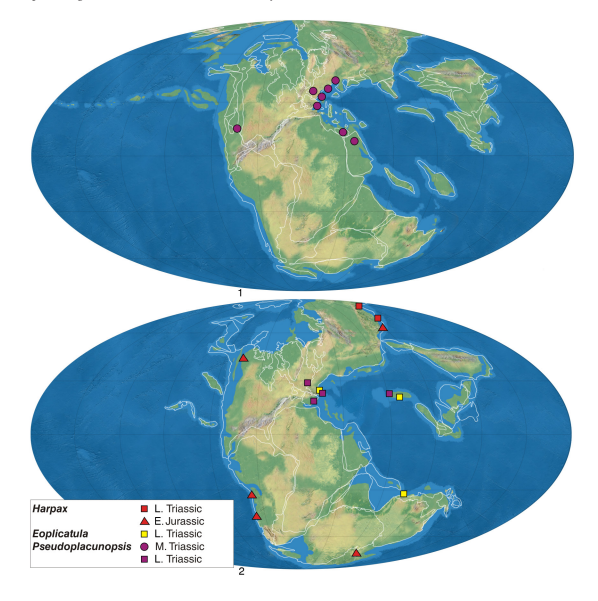


Figure 23. Paleogeographical distribution of Plicatulidae (Harpax, Eoplicatula, Pseudoplacunopsis). 1, Middle Triassic; 2, Late Triassic-Early Jurassic.

Stratigraphic range.—Middle Triassic (Anisian) (J. Chen, Stiller, & Komatsu, 2006). *Protostrea* is a monospecific genus only known from the upper Anisian in the Qingyan formation (Stiller, 2000; Komatsu, Chen, & others, 2004; J. Chen, Stiller, & Komatsu, 2006).

Paleogeographic distribution.—Eastern Tethys (Fig. 22).

Tethys domain: Middle Triassic: Anisian of southern China (Guizhou province) (Stiller, 2000; Komatsu, Akasaki, & others, 2004; J. Chen, Stiller, & Komatsu, 2006).

Paleoautoecology.—B, E, S, C, Sed; C. *Protostrea sinensis* probably lived cemented to the substrate by their right valve by a large cementation area (J. Chen, Stiller, & Komatsu, 2006). Often it is found cemented to other shells and corals (Komatsu, Chen, & others, 2004). *Protostrea* was also a substrate for other organisms, such as crinoids (Stiller, 2000).

Mineralogy.—Bimineralic. Not much is known about the shell mineralogy and microstructure of members of the family Dimyidae.

Waller (1978) indicated that they may have had an inner shell layer of aragonite and cross-lamellar microstructure and that they do not have a simple prismatic calcitic layer. J. Chen, Stiller, and Komatsu (2006, p. 160) studied thin sections of their specimens, which, although recrystallized into calcite, "... the shells originally probably had a mainly crossed-lamellar microstructure (originally aragonitic); in parts (at least of right valves) there are relics of (an) irregular simple-prismatic outer layer (s) (originally calcitic)."

Superfamily PLICATULOIDEA Watson, 1930 Family PLICATULIDAE Watson, 1930 Genus HARPAX Parkinson, 1811, p. 221

Type species.—Harpax parkinsoni Bronn, 1824, p. 52.

Remarks.—Although *Harpax* was considered a junior synonym of *Plicatula* Lamarck, 1801, by Cox and others (1969), some authors still regarded it as a valid subgenus of *Plicatula* (Okuneva, 1985; Dam-

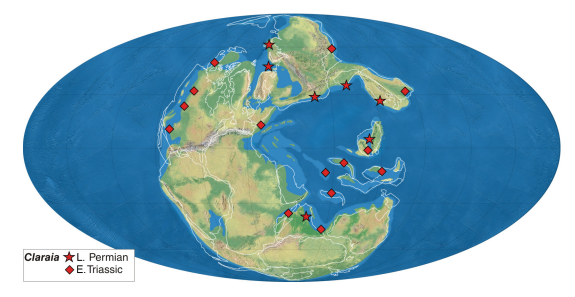


Figure 24. Paleogeographical distribution of Pterinopectinidae (Claraia). late Permian-Early Triassic.

borenea, 1993; Aberhan, 1994a, 1998a; Gahr, 2002, among others). Recently, Damborenea (2002a) validated the genus, distinguishing it from *Plicatula* due to its hinge details, relative convexity of the valves, ornamentation, and ligament (see discussion in Damborenea, 2002a, p. 86–89). The hinge of many species attributed to *Plicatula* is unknown, and so species undoubtedly included in *Harpax* are: *Harpax parkinsoni* Bronn, 1824, *Harpax rapa* (Bayle & Coquand, 1851), *Harpax kolymica* (Polubotko in Kiparisova, Bychkov, & Polubotko, 1966), *Harpax simplex* Milova, 1976, *Harpax spinosa* (J. Sowerby, 1819), and *Harpax auricula* (Eudes-Deslongchamps, 1860), among others.

Stratigraphic range.—Upper Triassic (Norian)–Lower Jurassic (Toarcian) (Damborenea, 1993; Gahr, 2002). It is difficult to assign a specific range to this genus, since diagnostic characters (for example, the hinge) of many species are not known (Damborenea, 2002a). The oldest solid records are from the Norian (Okuneva, 1985; Damborenea, 1993), with the youngest being from the lower Toarcian of Spain and Portugal (Gahr, 2002). Hautmann (2001a, 2001b) considered the genus to be only present in the Lower Jurassic.

Paleogeographic distribution.—Boreal and Austral, ?Tethys (Fig. 23). *Harpax* had a bipolar distribution, at least during the Early Jurassic (Damborenea, 1993, 1996a, 2001). It originated in the Boreal domain during the Late Triassic. Later, during the Pliensbachian–Toarcian, it was also reported from the Tethys domain (Gahr, 2002). With some doubt, it was also reported from the Rhaetian–Hettangian boundary in Tibet (J. Yin & McRoberts, 2006) and from Sinemurian beds of Morocco (Tomašových, 2006c).

Boreal domain: Late Triassic: Norian of Siberia (Okuneva, 1985), northeastern Asia (Polubotko & Repin, 1990); Norian–Rhaetian of Siberia (Kiparisova, Bychkov, & Polubotko, 1966; Polubotko, 1968a; Bychkov & others, 1976); Early Jurassic: northeastern Russia (Milova, 1976); Hettangian of northeastern Asia (Polubotko & Repin, 1990); Hettangian–Sinemurian of Canada (Aberhan, 1998a; Aberhan, Hrudka, & Poulton, 1998). Austral domain: Early Jurassic: Argentina (Damborenea, 1993, 2002a, 2002b); Hettangian–Sinemurian of Chile and Argentina (Damborenea, 1996a), ?New Zealand (Damborenea, 1993); Sinemurian of Chile (Aberhan, 1994a).

Paleoautoecology.—B, E, S, C, Sed; C. The distribution of bipolar (or antitropical) organisms is determined by temperature and substrate availability (Sha, 1996). They are abundant in shallow water areas at high latitudes and in deep water areas at low-latitude seas (Sha & Fürsich, 1994). According to several studies (see Damborenea, 2002a, p. 93), juvenile stages of *Harpax* were often cemented by the right valve to hard substrates (other shells, pebbles, rocks). However, in adult stages, they were often found loose in the sediment, so they had a free mode of life (Damborenea, 2002a). Sha (1996), based on various characters, such as the presence of byssal notch and sinus and pseudoctenolium, believed that in its juvenile stages, it also remained byssate, and he suggested that perhaps it had a pseudoplanktonic mode of life, attaching to floating objects (e.g., wood) or other swimming or nektonic organisms.

Mineralogy.—Bimineralic (Carter, 1990a, p. 226; Carter, Barrera, & Tevesz, 1998, p. 1003). Outer shell layer: calcite (foliated). Middle shell layer: calcite. Inner shell layer: aragonite.

Genus EOPLICATULA Carter, 1990a, p. 221

Type species.—Plicatula imago Bittner, 1895, p. 213.

Stratigraphic range.—Upper Triassic (Carnian–Rhaetian) (Bittner, 1895; Hautmann, 2001a, 2001b). Carter (1990a) proposed *Eoplicatula* as a subgenus of *Plicatula* and only included the type species from the Carnian of Italy. Subsequently, Hautmann (2001a) included the species *Plicatula difficilis* Healey, 1908, from Rhaetian beds of Burma and *Eoplicatula parvadehensis* Hautmann, 2001a, from the Norian of Iran. Hautmann and others (2005) reported *Eoplicatula* from Rhaetian beds of southern Tibet but did not figure or describe the specimens.

Paleogeographic distribution.—Tethys (Fig. 23).

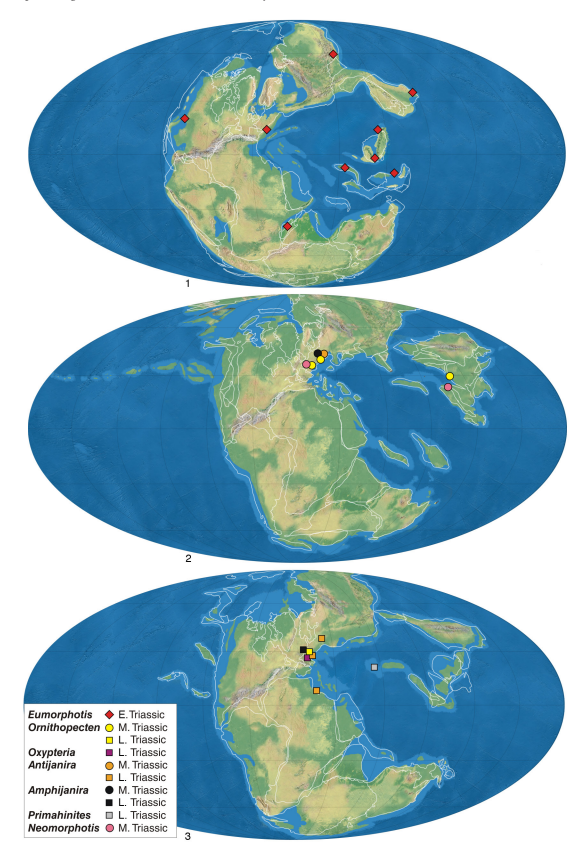


Figure 25. Paleogeographical distribution of Aviculopectinidae (*Eumorphotis, Ornithopecten, Oxypteria, Antijanira, Amphijanira, Primahinnites, Neomorphotis*). 1, Early Triassic; 2, Middle Triassic; 3, Late Triassic–Early Jurassic.

Tethys domain: Late Triassic: Carnian of Italy (Bittner, 1895; Leonardi, 1943; Carter, 1990a); Norian of Iran (Hautmann, 2001a, 2001b; Fürsich & Hautmann, 2005); Rhaetian of Burma (Healey, 1908).

Paleoautoecology.—B, E, S, C, Sed; C. *Eoplicatula* cemented to the substrate by its right valve. According to Hautmann (2001b), it was a reef-builder organism.

Mineralogy.—Bimineralic (Carter, 1990a, p. 223; Hautmann, 2001b). Outer shell layer: calcite (prismatic-foliated). Middle shell layer: aragonite (cross-lamellar). Inner shell layer: aragonite (prismatic–cross-lamellar).

Genus PSEUDOPLACUNOPSIS Bittner, 1895, p. 215

Type species.—Pseudoplacunopsis affixa Bittner, 1895, p. 215. *Remarks.*—After Todd and Palmer (2002), who proposed that

Placunopsis Morris & Lycett, 1853 in 1851-1855, p. 5, is a genus belonging to the Jurassic family Anomiidae, several species that were traditionally attributed to this genus were rejected, as they did not have a byssal foramen, and they were regarded as terquemids (=prospondylids) instead. While Hölder (1990) considered species from Triassic and Cretaceous ages to be within Placunopsis, Todd and Palmer (2002) proposed that their affinities are uncertain, and their knowledge is based on new well-preserved materials. We believe that many of the Triassic species referred to *Placunopsis* and included into the family Terquemiidae Cox, 1964, among them the so-called false oyster, are, in fact, true plicatulids and should be referred to Pseudoplacunopsis. Checa and others (2003) resampled the Middle Triassic (Ladinian) localities studied by Schmidt (1935) and Márquez-Aliaga, Hirsch, and López-Garrido (1986) from the Betic ranges (Jaen), and they obtained several thousand specimens of Placunopsis flabellum Schmidt, 1935, in which only the calcite microstructure of the right valves (the cemented ones) was preserved. In tens of specimens, details of the hinge could be observed, showing an elongated ligament furrow bordered by two crura diverging from the beak and pits corresponding to the other valve crura and inserted below the hinge line. The external ornamentation presented antimarginal thick ribs, and thus the species was referred to Enantiostreon; but hinge characters indicate that P. flabellum was a true plicatulid. Subsequently, one of the authors (Márquez-Aliaga) found identical hinge characters in specimens from Ladinian beds of the Iberian range (Cuenca) attributed to Placunopsis teruelensis Wurm, 1911. This species is ornamented by fine ribs. There are many Middle Triassic nominal species assigned to this genus, which could, in fact, be regarded as synonyms due to the variability of the cemented valve. Among the finely ornamented species, P. plana (Giebel, 1856) from the Germanic Muschelkalk could include as synonyms the following names: alpina Winkler, 1859, schaflautli Winkler, 1859, teruelensis Wurm, 1911, and *filicostata* Hölder, 1990. Within the heavily ornamented species, matercula Quenstedt, 1852 in 1851-1852, could include as a synonym *flabellum* Schmidt (Checa & others, 2003). Recently, Posenato (2008b) developed similar ideas.

Stratigraphic range.—Middle Triassic (Anisian)–Upper Triassic (Rhaetian) (Posenato, 2008b; Márquez-Aliaga, Damborenea, & Goy, 2008a). Cox and others (1969) assigned it an Upper Triassic range, and Hautmann (2001a) also considered that it ranged from the Carnian, but new records, as discussed above, confirmed its

presence in Middle Triassic deposits. Regarding the upper extension of its stratigraphic range, Hautmann (2001a) considered that *Pseudoplacunopsis* lived until Kimmeridgian times, represented by the species *Plicatula ogerieni* Loriol, 1904. Hautmann (2001a) did not make any comments about this species, nor did we find any record of the genus for the Jurassic; and its last appearance seems to be Rhaetian. It is interesting to note that most references to this genus are based on the diagnosis given by Bittner (1895) and Cox (1924), and not on the emended one by Hautmann (2001a), as the hinge and the ligament area are rarely preserved in specimens attributed to this genus.

Paleogeographic distribution.—Tethys and Circumpacific (Fig. 23).

Tethys domain: Middle Triassic: Anisian of Italy (Posenato, 2008b); Ladinian of Spain (Schmidt, 1935; Márquez-Aliaga, 1985; Martínez & Márquez-Aliaga, 1992; López-Gómez & others, 1994; Márquez-Aliaga & Ros, 2002; Márquez-Aliaga, Budurov, & Martínez, 1996; Márquez-Aliaga & others, 2004), Germany (Hagdorn & Simon, 1983; Hölder, 1990), France (Brocard & Philip, 1989), Israel (Lerman, 1960), Poland (Assmann, 1916), Italy (Posenato, 2002), Jordan (Cox, 1924); Late Triassic: Carnian of Spain (Martín-Algarra, Solé de Porta, & Márquez-Aliaga, 1995), Italy (Bittner, 1895; Leonardi, 1943); Norian–Rhaetian of Iran (Hautmann, 2001b); Rhaetian of Austria (Posenato, 2008b), Spain (Márquez-Aliaga, Damborenea, & Goy, 2008b; Márquez-Aliaga & others, 2010).

Circumpacific domain: Middle Triassic: Ladinian of Nevada (United States) (Waller in Waller & Stanley, 2005).

Paleoautoecology.—B, E, S, C, Sed; C. *Pseudoplacunopsis* was a cementing bivalve. It attached to the substrate by its right valve, and according to Hautmann (2001a), it was a reef builder. Márquez-Aliaga and Martínez (1994) studied its behavior as an epizoan organism.

Mineralogy.—Bimineralic (De Renzi & Márquez-Aliaga, 1980; Márquez-Aliaga & Márquez, 2000). Outer shell layer: calcite (foliated). Inner shell layer: aragonite (cross-lamellar).

Superfamily PTERINOPECTINOIDEA Newell, 1938 Family PTERINOPECTINIDAE Newell, 1938 Genus CLARAIA Bittner, 1901a, p. 568

Type species.—Posidonomya clarae Hauer, 1850, p. 112. Remarks.—Several taxa related to Claraia will not be considered in this analysis for various reasons: either they are regarded as synonyms of Claraia or their separation at generic level is not justified. These taxa are: Pseudoclaraia Zhang, 1980, p. 438, 443, Pteroclaraia Guo, 1985, p. 150, 265, Guichiella J. Li & Ding, 1981, p. 328–329, Claraioides Z. Fang, 1993, p. 653, 660, Epiclaraia Gavrilova, 1995, p. 156, and Rugiclaraia Waterhouse, 2000, p. 179 (see discussion for each of them in Genera not Included, p. 156).

Stratigraphic range.—upper Permian (Wuchiapingian)–Lower Triassic (middle Olenekian) (F. Yang, Peng, & Gao, 2001; McRoberts, 2010). For a long time, *Claraia* was regarded as a Lower Triassic index fossil. Cox and others (1969) assigned it a Lower Triassic range with a cosmopolitan distribution. Later, Nakazawa and others (1975) reported *Claraia bioni* Nakazawa in Nakazawa & others, 1975, from upper Permian sediments. Since then, there were many new upper Permian records (H. Yin, 1983; F. Yang, Peng, & Gao, 2001; Z. Fang, 1993, 2003; Gao, Yang, & Peng, 2004; Kotlyar, Zakharov, & Polubotko, 2004; He, Feng, & others, 2007; He, Shi, & others, 2007). Boyd and Newell (1979) reported *Claraia*? *posidoniformis* Termier & Termier, 1977, from Tunisian Guadalupian beds, but they doubted the generic relations of this species, because it shows some features that are not typical of *Claraia*.

Paleogeographic distribution.—Cosmopolitan (Fig. 24). During the late Permian, *Claraia* was widely distributed, mainly in the eastern part of Tethys. During the Early Triassic, it was abundant almost everywhere that beds of this age occur. For this reason, even though it was not reported from certain areas, a cosmopolitan distribution is given.

Tethys domain: late Permian: Kashmir (India) (Nakazawa & others, 1975); Wuchiapingian of China (F. Yang, Peng, & Gao, 2001); Changhsingian of China (Z. Zhang, 1980; H. Yin, 1983, 1990; Z. Fang, 1993, 2003; F. Yang, Peng, & Gao, 2001; Gao, Yang, & Peng, 2004; Z. Chen, Kaiho, & others, 2006; He, Feng, & others, 2007; He, Shi, & others, 2007), northwestern Caucasus (Russia) (Kotlyar, Zakharov, & Polubotko, 2004; Ruban, 2006a); Early Triassic: Pamir (Afghanistan) (Polubotko, Payevskaya, & Repin, 2001), Himalayas (Nepal) (Waterhouse, 2000), Italy (Leonardi, 1935; Broglio-Loriga, Masetti, & Neri, 1982; Neri, Pasini, & Posenato, 1986; Broglio-Loriga & others, 1988, 1990; Posenato, 1988; Posenato, Sciunnach, & Garzanti, 1996; Fraiser & Bottjer, 2007a, 2007b), China (Hsu, 1936–1937; F. Wu, 1985; Z. Li & others, 1986; Lu & Chen, 1986; S. Yang, Wang, & Hao, 1986; Z. Yang & others, 1987; Ling, 1988; M. Wang, 1993; Tong & Yin, 2002), Ussuriland (Russia) (Kiparisova, 1938); Induan of China (C. Chen, 1982; F. Yang, Peng, & Gao, 2001; He, Feng, & others, 2007), Italy (Leonardi, 1960), Malaysia (Ichikawa & Yin, 1966; Tamura & others, 1975), Vietnam (Vu Khuc & Huyen, 1998), Alberta (Canada) (Newell & Boyd, 1995; McRoberts, 2010); Olenekian of Mangyshlak (Kazakhstan) (Gavrilova, 1995), China (H. Yin, 1990; J. Chen & Komatsu, 2002), Pakistan (Nakazawa, 1996), Vietnam (Komatsu & Huyen, 2006).

Circumpacific domain: Early Triassic: Wyoming and Idaho (United States) (Newell & Kummel, 1942), Alberta (Canada) (Newell & Boyd, 1970), Japan (Nakazawa, 1971; Hayami, 1975); Induan of Nevada (United States) (Ciriacks, 1963; Schubert, 1993; Newell & Boyd, 1995; Schubert & Bottjer, 1995; Boyer, Bottjer, & Droser, 2004; Fraiser & Bottjer, 2007a, 2007b).

Boreal domain: late Permian: eastern Greenland (Newell & Boyd, 1995), Nova Zemla (Arctic Ocean) (Muromtseva, 1984); Early Triassic: Queen Elizabeth Islands (Arctic Archipelago, Canada) (Tozer, 1961, 1962, 1970).

Paleoautoecology.—B-Ps, E, S, Epi, Se-FaM. Several modes of life have been attributed to *Claraia*, ranging from benthic epibyssate (Z. Fang, 1993; F. Yang, Peng, & Gao, 2001) to pseudoplanktonic and even occasional swimmer (F. Yang, Peng, & Gao, 2001). *Claraia* shell morphology subtly changed through time. These differences are primarily related to the morphology of the byssal notch, the ornamentation, and the shape of the auricles (F. Yang, Peng, & Gao, 2001; He, Feng, & others, 2007). Permian forms had a more developed and deep byssal notch, shells were small in size, thin, and slightly inequivalve; they were interpreted as living epibyssate with the capacity to swim occasionally (F. Yang, Peng, & Gao, 2001). However, due to the associated fauna, e.g., ammonoids, they were also interpreted as pseudoplanktonic (H. Yin, 1983). Nevertheless, according to F. Yang, Peng, and Gao (2001), features present in Permian forms were unsuited to this mode of life. On the other hand, in the Triassic forms, the byssal notch was shallower, which is associated with increased mobility (Z. Fang, 1993; see also He, Feng, & others, 2007, table 1), and the shells were less ornamented (F. Yang, Peng, & Gao, 2001). These forms were also interpreted by F. Yang, Peng, and Gao (2001) as being pseudoplanktonic bivalves that attached themselves to pieces of wood or algae.

The genus occurred primarily in deep-water Permian deposits, but, by the Triassic, it was in all types of environments, from shallow to deep (F. Yang, Peng, & Gao, 2001). This fact was related to an opportunistic behavior during recovery from the Permian-Triassic (P/T) extinction event (Schubert & Bottjer, 1995; Rodland & Bottjer, 2001). This success during the Triassic appears to be related to the morphological change, since Permian forms with a deep byssal notch did not survive the P/T event; however, the forms with a shallower byssal notch diversified enormously [from 3 species in late Permian to over 30 in the Triassic (He, Feng, & others, 2007)]. According to F. Yang, Peng, and Gao (2001), this was also related to the mode of life of *Claraia* larvae, which probably had a veliger planktonic stage. The deep-water habitats in which mostly Permian forms were found were interpreted by Gao, Yang, and Peng (2004) as potential refuges for those forms that survived and reached the Early Triassic. It is hard to assign a unique mode of life to all species of *Claraia*, since they have a wide range of morphological features that suggests that some species could have been epibenthic, pseudoplanktonic, and even occasional swimmers (He, Feng, & others, 2007).

Mineralogy.—Bimineralic (Boyd & Newell, 1976; Newell & Boyd, 1985, 1995; Carter, 1990a, 1990b). Outer shell layer: calcite (prismatic). Inner shell layer: aragonite (?).

Superfamily AVICULOPECTINOIDEA Meek & Hayden, 1864 Family AVICULOPECTINIDAE Meek & Hayden, 1864 Genus EUMORPHOTIS Bittner, 1901a, p. 566

Type species.—Pseudomonotis telleri Bittner, 1898, p. 710.

Stratigraphic range.-Lower Triassic (Induan-Olenekian) (Broglio-Loriga & Mirabella, 1986). Cox and others (1969) assigned it a Lower Triassic-Upper Triassic range. However, Broglio-Loriga and Mirabella (1986) did a comprehensive study on Eumorphotis, and they noted that Middle and Upper Triassic forms were highly dubious, and therefore they restricted the range of *Eumorphotis* to the Lower Triassic. Newell and Boyd (1995) assigned it the same range. Furthermore, these authors argued that Heteropecten Kegel & Costa, 1951, and Eumorphotis were virtually indistinguishable and that the reason for proposing Eumorphotis was more to separate the Paleozoic from the Triassic forms than to recognize significant morphological differences between the two groups. In fact, Newell and Boyd (1995) considered that some specimens attributed by Bittner (1901b) to the Triassic Eumorphotis from eastern Siberia are similar to Heteropecten. Moreover, Eumorphotis was also reported from the upper Permian, but Broglio-Loriga and Mirabella (1986) doubted all these records, because they were based on poorly preserved material. Posenato, Pelikán, and Hips (2005) proposed a new species, Eumorphotis lorigae Posenato, Pelikán, & Hips, 2005, and they referred it to the upper Permian (upper Changhsingian), but, as indicated by the authors, this age is only provisionally based on bivalves and brachiopods, and thus we will not take this record into account until new data allow a more precise age determination.

Paleogeographic distribution.—Tethys and Circumpacific (Fig. 25). Tethys domain: Early Triassic: Italy (Bittner, 1912; Leonardi, 1935; Broglio-Loriga, Masetti, & Neri, 1982; Neri & Posenato, 1985; Broglio-Loriga & Mirabella, 1986, and references therein; Neri, Pasini, & Posenato, 1986; Broglio-Loriga & others, 1990), Ussuriland (Russia) (Kiparisova, 1938), Vietnam (Vu Khuc & Huyen, 1998), Pakistan (Nakazawa, 1996), Malaysia (Ichikawa & Yin, 1966), China (Hsu, 1936–1937; Z. Yang & Yin, 1979; C. Chen, 1982; F. Wu, 1985; S. Yang, Wang, & Hao, 1986; Ling, 1988; H. Yin, 1990; Tong & others, 2006); Induan of southern China (Hautmann & others, 2011).

Circumpacific domain: Early Triassic: western United States and Japan (Newell & Kummel, 1942; Ciriacks, 1963; Schubert, 1993; Newell & Boyd, 1995; Boyd, Nice, & Newell, 1999; Fraiser & Bottjer, 2007a), Japan (Nakazawa, 1961, 1971; Hayami, 1975; Kashiyama & Oji, 2004).

Paleoautoecology.—B, E, S, Epi, Sed; By. According to key features present in different species of this genus (elongated anterior auricles, byssal notch in adults), and, following S. M. Stanley's (1970, 1972) criteria, *Eumorphotis* could be an epibyssate bivalve. Together with *Claraia, Promyalina,* and *Unionites, Eumorphotis* was one of the dominant bivalves in the Early Triassic seas (Fraiser & Bottjer, 2007a).

Mineralogy.—Bimineralic (Carter, 1990a, p. 241). There are no available data about *Eumorphotis* shell mineralogy or microstructure. Data provided for family Aviculopectinidae. Outer shell layer: calcite (prismatic-homogeneous-foliated). Inner shell layer: aragonite (nacreous–cross-lamellar).

Genus ORNITHOPECTEN Cox, 1962, p. 596

Type species.—Aviculopecten bosniae Bittner, 1903, p. 592.

Remarks.— Cox (1962) proposed *Ornithopecten* to accommodate several Triassic species that were previously attributed to *Aviculopecten* M'Coy, 1851, p. 171 (which is actually regarded a strictly Paleozoic genus).

Stratigraphic range.—Middle Triassic (Anisian)–Upper Triassic (Carnian) (Diener, 1923; Allasinaz, 1972). Cox and others (1969) assigned it a Middle-Upper Triassic range. According to Diener (1923), the species assigned to Ornithopecten by Cox (1962) were reported from Anisian and Carnian beds. Subsequently, the genus was reported from the Lower Triassic of China, but there are several problems with these records. The only reference we could locate in which the material was described and figured is Z. Yang and others (1987). A new species was described there: Ornithopecten? magnauritus Yin, but this was only doubtfully assigned to Ornithopecten, and, as noted by the authors, it might be better located within Eumorphotis Bittner, 1901a, with which we agree. The other papers where the genus was mentioned from the Triassic age (e.g., Z. Chen, Shi, & Kaiho, 2004; Z. Chen & McNamara, 2006; Z. Chen, Shi, & others, 2006) do not have figures or descriptions; furthermore, they do not mention the original source of data, so they are not taken into account.

Paleogeographic distribution.—Tethys (Fig. 25).

Tethys domain: Middle Triassic: Anisian of Yugoslavia (Allasinaz, 1972), China (H. Yin, 1985; J. Chen & Stiller, 2007), Alps (Diener, 1923); Ladinian of China (H. Yin, 1985), southern Alps (Bittner, 1895; Diener, 1923); Late Triassic: Carnian of the Alps (Diener, 1923).

Paleoautoecology.—B, E, S, Epi, Sed; By. Following the guidelines provided by S. M. Stanley (1970, 1972), *Ornithopecten* was most likely an epibyssate bivalve, and the byssus was placed under the anterior auricle.

Mineralogy.—Bimineralic (Carter, 1990a, p. 241). There are no data about *Ornithopecten* shell. Data provided for family Aviculopectinidae. Outer shell layer: calcite (prismatic-homogeneous-foliated). Inner shell layer: aragonite (nacreous–cross-lamellar).

Genus OXYPTERIA Waagen, 1907, p. 93

Type species.—Aviculopecten (Oxypteria) bittneri Waagen, 1907, p. 93.

Stratigraphic range.—Upper Triassic (Carnian) (Cox & others, 1969). Waagen (1907) proposed the genus on the basis of material from Carnian beds of southern Tyrol. The only other references that could be located are Diener (1923) and Cox and others (1969), who repeated the information in Waagen.

Paleogeographic distribution.—western Tethys (Fig. 25).

Tethys domain: Late Triassic: Carnian of southern Tirol (Italian Alps) (Waagen, 1907; Diener, 1923; Cox & others, 1969).

Paleoautoecology.—B, E, S, Epi, Sed; By. Since all that is known of this monospecific genus is a left valve, it is difficult to speculate how it lived. We assign it the dominant mode of life in the family Aviculopectinidae.

Mineralogy.—Bimineralic (Carter, 1990a, p. 241). There are no data about *Oxypteria* shell. Data provided for family Aviculopectinidae. Outer shell layer: calcite (prismatic-homogeneous-foliated). Inner shell layer: aragonite (nacreous–cross-lamellar).

Genus ANTIJANIRA Bittner, 1901c, p. 49

Type species.—Pecten hungaricus Bittner, 1901c, p. 48.

Remarks.—According to Allasinaz (1972), Bittner proposed the name *Antijanira* to accommodate a group of Triassic species with a particular ornamentation type. However, Bittner did not provide a true diagnosis, nor did he indicate the similarities and differences with other taxa (Allasinaz, 1972). Probably for this reason, Newell and Boyd (1995) placed *Antijanira* in synonymy with *Leptochondria* Bittner, 1891, p. 101. Allasinaz (1972) provided an adequate diagnosis and discussed its similarities with other taxa, so the genus will be considered valid here and included in the Aviculopectinidae, according to this author. Cox and others (1969) and Carter (1990a) regarded it a member of Pectinidae, as did other authors (Kobayashi & Tamura, 1983a; Gou, 1993), who also considered it a subgenus of *Chlamys* Röding in Bolten, 1798. However, Johnson and Simms (1989) suggest allocation in Aviculopectinidae is supported by the shell structure and the aviculopectinid-type ligament.

Stratigraphic range.—Middle Triassic (Anisian)–Upper Triassic (Carnian) (Allasinaz, 1972). Cox and others (1969) assigned it an Upper Triassic range in Hungary and Sepkoski (2002) a Triassic (Anisian–Carnian) range, taking data from Hallam (1981), who considered it to be present only in the European Carnian, and Hayami (1975), who reported it from Anisian and Ladinian beds. Although the Anisian records appeared in Bittner (1903), Cox and others (1969) did not take them into account. On the other hand, Allasinaz (1972) did consider them, and we follow him.

Paleogeographic distribution.—western Tethys (Fig. 25). The distribution of this genus was limited to the Tethys domain. Waller and Stanley (1998) found a fragment of scallop that could be attributed to Antijanira amphidoxa (Bittner, 1903) from Middle Triassic beds of Nevada, but Waller and Stanley (2005) later included this specimen in Oxytoma (Oxytoma) grantsvillensis Waller in Waller & Stanley, 2005. Although Antijanira was reported from the Upper Triassic of China (Kobayashi & Tamura, 1983a; Gou, 1993), we cannot be sure of its presence in this area, since the only available described and figured specimens appeared in Gou (1993), and neither the auricles nor the ligament area are seen in them; moreover, neither the description nor the ornamentation match with the diagnosis given by Allasinaz (1972). Z. Fang and others (2009) tentatively suggested Halobia (Enormihalobia) Yin & Gan in Gan & Yin, 1978, p. 352, is a junior synonym of Antijanira. If this synonymy is accepted, the distribution of Antijanira extended to the Eastern Tethys (Carnian of Guizhou province).

Tethys domain: Middle Triassic: Anisian of Yugoslavia (Bittner, 1903; Allasinaz, 1972); Late Triassic: Carnian of Alps (Italy) (Bittner, 1895, 1901a; Allasinaz, 1972; Johnson & Simms, 1989), Tripoli-Garian region (Libya) (Desio, Rossi Ronchetti, & Vigano, 1960), Hungary (Bittner, 1912).

Paleoautoecology.—B, E, S, Epi, Sed; By. According to S. M. Stanley (1970), bivalves that are able to swim possess a symmetrical shell, equal auricles, and an umbonal angle greater than 105°. In the specimens described by Allasinaz, the umbonal angle is usually about 90°, but the angles given by Allasinaz (1972, p. 275) for some specimens from Zardini's collection often exceed 100° and even reach 115°. However, the auricles are not equal (see description in Allasinaz, 1972, p. 271), and in the anterior one, there is a deep byssal groove; also, not all species have a perfectly symmetrical shell, so it is likely that Antijanira species lived epibyssate with the sagittal plane in a horizontal position, interpreted by S. M. Stanley (1970) as being very stable since it increases the surface area in contact with the substrate. In addition, the auricles of these species have different convexity, suggesting they were not well adapted for swimming (S. M. Stanley, 1970).

Mineralogy.—Bimineralic (Allasinaz, 1972; Carter, 1990a, p. 255, 262). Allasinaz (1972) described the shell microstructure of *Antijanira* with an external shell layer of prismatic calcite in the right valves and fibrous in the left. Carter (1990a) described the microstructure of the group *Antijanira* as grade 2. Although there are differences between the valves, the outer layer is always calcitic and the middle and inner layers are aragonitic. Outer shell layer: calcite (prismatic-homogeneous). Middle and inner shell layers: aragonite (cross-lamellar).

Genus AMPHIJANIRA Bittner, 1901c, p. 49

Type species.—Pecten janirula Bittner, 1895, p. 160.

Stratigraphic range.—Middle Triassic (Anisian)–Upper Triassic (Carnian) (Allasinaz, 1972). The range provided by both Cox and others (1969) and Sepkoski (2002) is similar to *Antijanira* (see stratigraphic range for *Antijanira*). Following Allasinaz (1972), we assign a Anisian–Carnian range.

Paleogeographic distribution.—western Tethys (Fig. 25).

Tethys domain: Middle Triassic: Anisian of Yugoslavia (Bittner, 1903; Allasinaz, 1972); Late Triassic: Carnian of the Alps (Italy) (Bittner, 1895; Allasinaz, 1972).

Paleoautoecology.—B, E, S, Epi, Sed; By. *Amphijanira* had an equivalve shell, the auricles were very different, with the posterior one being smaller and separated from the shell and the anterior one with a pronounced byssal notch (Allasinaz, 1972), and thus it is very unlikely that it could swim. It probably lived attached by the byssus.

Mineralogy.—Bimineralic (Carter, 1990a, p. 255, 262). Although there are no specific data for *Amphijanira* shells, we assign the same data as *Antijanira*, following Carter, who provided the same information for the entire *Antijanira* Group. Outer shell layer: calcite (prismatic-homogeneous). Middle shell layer: aragonite (crosslamellar). Inner shell layer: aragonite (cross-lamellar).

Genus PRIMAHINNITES Repin, 1996, p. 367

Type species.—Primahinnites iranica Repin, 1996, p. 367.

Remarks.—Repin (1996) included *Primahinnites* within the family Prospondylidae, but he only had a complete right valve and five fragments of right and left valves. Hautmann (2001b) obtained new and better preserved material, in which he observed certain key features, such as the ctenolium, and he included the genus in the family Aviculopectinidae; indeed, he emended the diagnosis. Repin's (1996) allocation was erroneous because none of his specimens had the cementation area preserved, but they had a byssal notch instead, described as being small (Hautmann, 2001b).

Stratigraphic range.—Upper Triassic (Norian–Rhaetian) (Hautmann, 2001b). Although originally Repin (1996) reported *Primahin-nites* only from the upper Norian, later Hautmann (2001b) reported it from the Rhaetian as well.

Paleogeographic distribution.—Tethys (Fig. 25).

Tethys domain: Late Triassic: Norian of Iran (Repin, 1996; Hautmann, 2001b); Rhaetian of Iran (Hautmann, 2001b).

Paleoautoecology.—B, E, S, Epi, Sed; By. Hautmann (2001b) described a well-developed byssal notch below the anterior auricle of the right valve, so he considered it to be an epibyssate bivalve.

Mineralogy.—Bimineralic (Carter, 1990a, p. 241). There are no data about *Primahinnites* shell mineralogy. Data provided for family Aviculopectinidae. Outer shell layer: calcite (prismatic-homogeneous-foliated). Inner shell layer: aragonite (nacreous–cross-lamellar).

Genus NEOMORPHOTIS H. Yin & Yin, 1983, p. 155

Type species.—Neomorphotis gigantea H. Yin & Yin, 1983, p. 155. This species was regarded as a synonym of *Eumorphotis buhaheensis* Lu by Fang & others, 2009, p. 36.

Remarks.—Neomorphotis was originally included in the family Pectinidae, but H. Yin (1985) and Posenato (2008b) transferred it to Aviculopectinidae. Due to its relationship with *Eumorphotis*, this seems appropriate.

Stratigraphic range.—Middle Triassic (Anisian) (Posenato, 2008b). According to Z. Fang and others (2009), the genus was proposed by H. Yin and Yin in 1983 from Middle Triassic beds of China. All records are from the Anisian (e.g., Lu & Chen, 1986; Ling, 1988; Sha, Chen, & Qi, 1990; Posenato, 2008b), but we are not certain which species were included in the genus by the authors, and thus the range will remain temporarily as Anisian until we have access to more information.

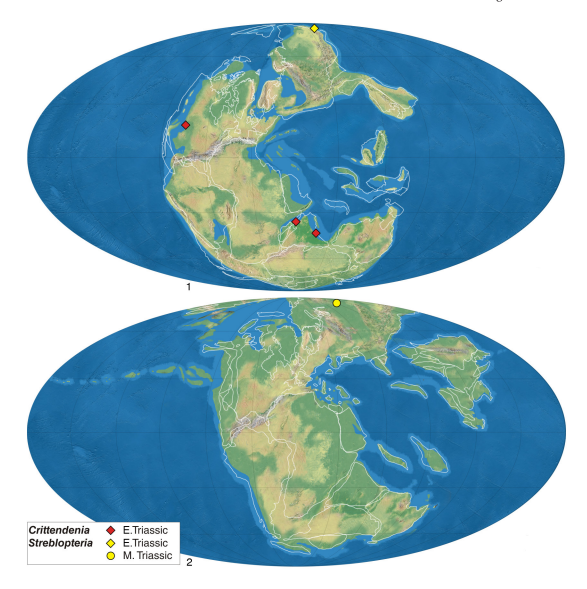


Figure 26. Paleogeographical distribution of Deltopectinidae (Crittendenia, Streblopteria). 1, Early Triassic; 2, Middle Triassic.

Posenato (2008b) raised the possibility that *Pseudomonotis beneckei* Bittner, 1900 (according to this author, included by H. Yin & Yin in *Neomorphotis*) is a junior synonym *Neomorphotis compta* (Goldfuss, 1833 in 1833–1841). If we accept this synonymy, the genus would be also present in the Lower Triassic, as *P. beneckei* was mentioned for this age by several authors. However, we included this species in *Eumorphotis* following Broglio-Loriga and Mirabella (1986). H. Yin (1985) considered that the genus was also present in the Olenekian, data incorporated by Sepkoski (2002).

Paleogeographic distribution.—Tethys (Fig. 25).

Tethys domain: Middle Triassic: Anisian of the Dolomites (Italy) (Posenato, 2008a, 2008b), southern China (Lu & Chen, 1986; Ling, 1988; Sha, Chen, & Qi, 1990).

Paleoautoecology.—B, E, S, Epi, Sed; By. At least N. compta, according to the description offered by Posenato (2008b), prob-

ably was an epibyssate bivalve, since it has an inequivalve shell and a deep byssal notch in the right valve. Moreover, taphonomic evidence also supports this mode of life (see Posenato, 2008b, p. 101).

Mineralogy.—Bimineralic (Posenato, 2008b). Posenato (2008b, p. 101) indicated about *N. compta* (Goldfuss, 1833 in 1833–1841): "shell wall is thin, bimineralic, and consisting of an outer calcitic layer and an inner, thin, calcitized layer."

Family DELTOPECTINIDAE Dickins, 1957 Genus CRITTENDENIA Newell & Boyd, 1995, p. 52

Type species.—Crittendenia kummeli Newell & Boyd, 1995, p. 53. Remarks.—Newell and Boyd (1995) provisionally included their new genus in Deltopectinidae due to external similarities with Streblopteria M'Coy, 1851, in the absence of characters from

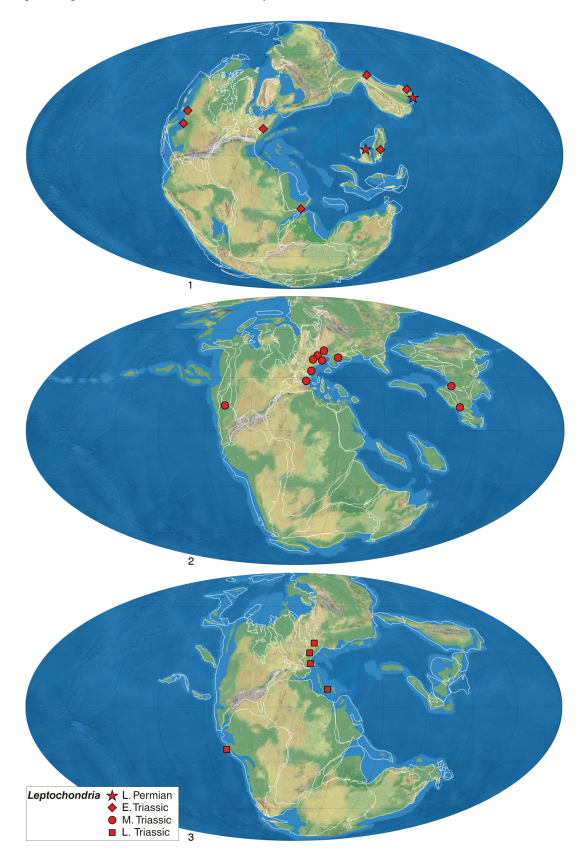


Figure 27. Paleogeographical distribution of Leptochondriidae (Leptochondria). 1, late Permian-Early Triassic; 2, Middle Triassic; 3, Late Triassic-Early Jurassic.

the ligament area and microstructure of the shell. Besides the type species, recorded from the Thaynes Formation in Nevada, they also included *Pseudomonotis decidens* Bittner, 1899, which was referred to *Claraia* and *Streblopteria* by other authors (see Newell & Boyd, 1995, p. 52–53). Gavrilova (1996), ignoring Newell and Boyd's paper, proposed a new subgenus of *Claraia, Bittnericlaraia* Gavrilova, with *Pseudomonotis decidens* Bittner, 1899, as type.

Newell and Boyd (1995) have a few contradictions, however: in the text, they mentioned P. decidens as being collected in Salt Range, Pakistan, and referred it to their figure 39. In this figure explanation, the given name is Crittendenia kummeli from the Lower Triassic of Salt Range, Pakistan. Furthermore, they considered C. kummeli as being present in Nevada and in Pakistan. This was used by Nakazawa (1996) to regard C. kummeli as a synonym of B. decidens. But taking into account that the ligament area is not known in any of these species, this synonymy is not clearly justified (Waterhouse, 2000). Waterhouse (2000) saw a clear relationship between Claraia and Crittendenia, and he referred the latter to the Pterinopectinidae. Furthermore, this author included in Crittendenia several new species, plus those included by Gavrilova (1996) in Claraia (Bittnericlaraia), but Waterhouse had a concept of the genus that is totally different from the original authors. In the absence of more information about key characters of the genus, as discussed above, we provisionally accept the allocation of Newell and Boyd (1995).

Stratigraphic range.—Lower Triassic (Newell & Boyd, 1995). *Crittendenia* was only reported from Lower Triassic (Newell & Boyd, 1995).

Paleogeographic distribution.—Tethys and Circumpacific (Fig. 26).

Tethys domain: Early Triassic: Pakistan (Newell & Boyd, 1995), Himalayas (Bittner, 1899), Nepal (Waterhouse, 2000).

Circumpacific domain: Early Triassic: Nevada (United States) (Newell & Boyd, 1995; Fraiser & Bottjer, 2007a).

Paleoautoecology.—B-Ps, E, S, Epi, Sed-FaM; By. Newell and Boyd (1995) mentioned a deep byssal notch in *Crittendenia* diagnosis, so this is an epibyssate bivalve. In addition, it was often found in association with ammonoids, and the umbilical area of ammonoids is sometimes xenomorphic on the bivalve shell. Therefore, Newell and Boyd (1995) postulated that *Crittendenia* could have had a pseudo-planktonic (attached to objects by the byssus) or even pseudopelagic mode of life (attached by byssus to the shells of live ammonoids).

Mineralogy.—Bimineralic (Waller, 1978). There are no data on the mineralogy and microstructure of *Crittendenia* shell. Due to the uncertainties about its familial assignation, we cannot use here the predominant data from the family. In the diagnosis provided by Waller (1978) for the order Pectinoida, he indicated that the shell is bimineralic.

Genus STREBLOPTERIA M'Coy, 1851, p. 170

Type species.—Meleagrina laevigata M'Coy, 1844, p. 80.

Stratigraphic range.—Carboniferous (Mississippian)–Middle Triassic (Anisian) (Dagys & Kurushin, 1985; Newell & Boyd, 1995). Streblopteria is a distinctive Paleozoic genus (Newell, 1938; Nakazawa & Newell, 1968; Hayami & Kase, 1977; Waterhouse, 1978; Boyd & Newell, 1979; Newell & Boyd, 1987, among others). However, Newell and Boyd (1995, p. 50) argued that it was also reported from the Middle Triassic of Siberia: "Distribution: Cosmopolitan, Miss.-Perm., M. Trias. of Arctic Siberia (*fide* Kurushin, 1982, p. 60)." In fact, ten years earlier, Dagys and Kurushin (1985) had described and listed the species referred by Newell and Boyd (1995): *Streblopteria newelli* Kurushin, 1982, and a new species, *S. jakutica* Kurushin in Dagys & Kurushin, 1985, the first being reported from the Olenekian and Anisian and the second from the Olenekian.

Paleogeographic distribution.—Boreal (Fig. 26). During the Carboniferous and the Permian, it had a cosmopolitan distribution (Newell & Boyd, 1995). Specifically, it was reported from the upper Permian of Nepal (Waterhouse, 1978), China (H. Yin, 1983; Z. Yang & others, 1987; M. Wang, 1993; He, Feng, & others, 2007), and the boreal region of Russia (Astafieva, 1998).

Boreal domain: Early Triassic: Olenekian of northern Siberia (Dagys & Kurushin, 1985); Middle Triassic: Anisian of northern Siberia (Dagys & Kurushin, 1985).

Paleoautoecology.—B, E, S, Epi, Sed; By. Within pectinoids, two groups can be recognized regarding their mode of life (S. M. Stanley, 1972): epibyssate bivalves, which are characterized by different convexity in both valves, the anterior auricle being more developed than the posterior one, and having a byssal sinus throughout its ontogeny; and others, also epibyssate but which developed swimming abilities, which are more symmetrical with both valves being equally convex, with auricles of the same shape and size, and an umbonal angle greater than 90°. *Streblopteria* features indicate it belongs in the first group.

Mineralogy.—Bimineralic (Waller, 1978). According to Newell and Boyd (1985), the outer shell layer of *Streblopteria* was fibrous prismatic in both valves. The inner shell layers are not known, but Waller (1978), in the diagnosis of the order Pectinioida, indicated that the shell was bimineralic.

Superfamily PSEUDOMONOTOIDEA Newell, 1938 Family LEPTOCHONDRIIDAE Newell & Boyd, 1995 Genus LEPTOCHONDRIA Bittner, 1891, p. 101

Type species.—Pecten aeolicus Bittner, 1891, p. 101.

Stratigraphic range.—middle Permian (Guadalupian)–Upper Triassic (Norian) (Cox, 1949; Newell & Boyd, 1995). Although Cox and others (1969) assigned it a Lower–Upper Triassic range, new records expanded the range of this genus. *Leptochondria* was reported from the middle Permian of Texas and Wyoming (United States) (Boyd & Newell, 1995) and from the upper Permian (Nakazawa & Newell, 1968; He, Feng, & others, 2007). Waller and Stanley (2005) reported *Leptochondria* from upper Permian of Pakistan, allegedly taking their data from Newell and Boyd (1995), but there the genus was mentioned only from the Lower Triassic of Pakistan. The youngest record is Norian (Cox, 1949; Newell & Boyd, 1995).

Paleogeographic distribution.—Tethys and Circumpacific (Fig. 27). Tethys domain: late Permian: Changhsingian of southern China (Y. Zhang, 1981; H. Yin, 1983; He, Feng, & others, 2007); Early Triassic: Induan of Pakistan (Nakazawa, 1996); Olenekian of southern China (Sha, 1995, 1998; Sha & Grant-Mackie, 1996; Sha, Johnson, & Fürsich, 2004), Italy (Neri & Posenato, 1985; Posenato, 2008a); Middle Triassic: Poland (Senkowiczowa, 1985); Anisian of Italy, Yugoslavia, and Bulgaria

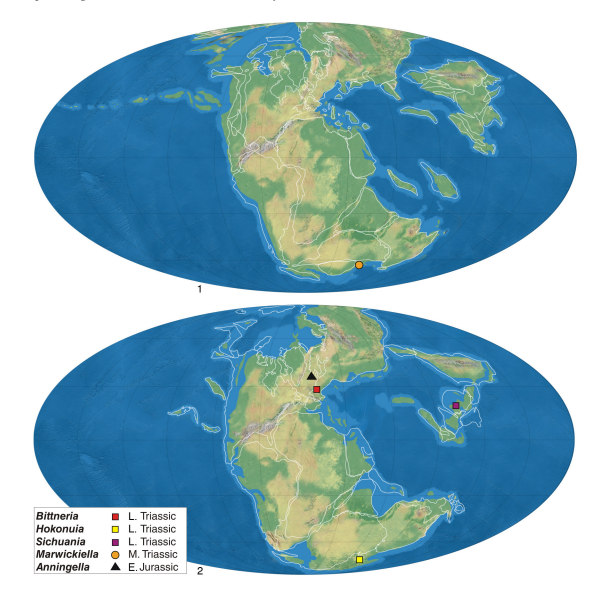


Figure 28. Paleogeographical distribution of Buchiidae (Bittneria, Hokonuia, Sichuania, Marwickiella, Anningella). 1, Middle Triassic; 2, Late Triassic-Early Jurassic.

(Allasinaz, 1972), southern China (Komatsu, Chen, & others, 2004); Anisian–Ladinian of Hungary (Szente, 1997), northern Vietnam (Komatsu, Huyen, & Huu, 2010); Ladinian of Italy (Allasinaz, 1972), Spain (Márquez-Aliaga, 1983, 1985; Márquez-Aliaga, Hirsch, & López-Garrido, 1986; Márquez-Aliaga & Montoya, 1991; Freneix, 1999; Niemeyer, 2002; Márquez-Aliaga & Ros, 2003), Carpathians (Slovakia) (Kochanová, Mello, & Siblík, 1975); Late Triassic: Carnian of Italy and Yugoslavia (Allasinaz, 1972); Norian of Hungary (Allasinaz, 1972), Anatolia (Turkey) (Diener, 1923).

Circumpacific domain: late Permian: Japan (Nakazawa & Newell, 1968; Hayami, 1975; Hayami & Kase, 1977); Early Triassic: Ussuriland (Russia) (Kiparisova, 1938); Olenekian of Nevada (United States) (Newell & Boyd, 1995; Boyer, Bottjer, & Droser, 2004; Fraiser & Bottjer, 2007a), Utah (United States) (Boyer, Bottjer, & Droser, 2004; Fraiser & Bottjer, 2007a), Japan (Nakazawa, 1961, 1971; Fraiser & Bottjer, 2007a); Middle Triassic: Ladinian of Nevada (United States) (Waller & Stanley, 2005); Late Triassic: Norian of Peru (Cox, 1949).

Paleoautoecology.—B, E, S, Epi, Sed; By. According to the diagnosis provided by Newell and Boyd (1995), there is a wide byssal notch in the right valve. It was most likely an epibyssate bivalve. Sha and Grant-Mackie (1996) proposed a possible pseudoplanktonic mode of life for *Leptochondria*.

Mineralogy.—Bimineralic (Waller & Stanley, 2005). Waller and Stanley (2005) assumed bimineralic mineralogy, due to the differential dissolution of the shell layers in their specimens of *Leptochodria shoshonensis* Waller in Waller & Stanley, 2005.

Superfamily MONOTOIDEA Fischer, 1887 in 1880–1887 Family BUCHIIDAE Cox, 1953 Genus BITTNERIA Broili, 1904, p. 168

Type species.—Avicula? efflata Broili, 1904, p. 167.

Remarks.—Broili (1904) proposed the genus *Bittneria* based on *Avicula? efflata*, and he included it in the family Aviculidae, although he noted that it could be considered as being intermediate between *Avicula* and *Pecten*. This allocation is doubtful, because only one left valve was then available. Subsequently, Cox and others (1969) included it in the family Buchiidae, also dubiously. The systematic position of this genus is especially problematic, because the hinge structure is not known (Sha & Fürsich, 1994). These authors related *Bittneria* to *Aucellina* Pompeckj, 1901.

Stratigraphic range.—Upper Triassic (Carnian) (Broili, 1904). The genus was proposed by Broili (1904) as being from Carnian sediments, and little else is known about it. Cox and others (1969) assigned the same range. Subsequently, new material was found at the same stage by Fürsich and Wendt (1977).

Paleogeographic distribution.—western Tethys (Fig. 28). Although J. Chen (1982a) described a new species from the Carnian of southern China (*Bittneria? hunanensis* J. Chen, 1982a), it was only doubtfully assigned to the genus, so we are not considering it.

Tethys domain: Late Triassic: Carnian of southern Alps (Broili, 1904; Waagen, 1907; Fürsich & Wendt, 1977).

Paleoautoecology.—B, E, S, Epi, Sed; By. According to the diagnosis offered by Cox and others (1969), the anterior left auricle is separated from the disk by a deep sinus from where the byssus would probably emerge. *Bittneria* was probably an epibyssate bivalve.

Mineralogy.—?Bimineralic (Carter, 1990a, p. 234). There are no data for the shell of *Bittneria*. We cannot use the characteristic features of the family due to its doubtful systematic allocation. Nevertheless, it is likely that *Bittneria* had a bimineralic shell, as did most members of the Order Pectinioida.

Genus HOKONUIA Trechmann, 1918, p. 202

Type species.—Hokonuia limaeformis Trechmann, 1918, p. 204. *Remarks.*—Although *Hokonuia* was related to the families Pergamidiidae (Trechmann, 1918; Waterhouse, 1960), Myalinidae (Trechmann, 1918; Wilckens, 1927) and Pteriidae (Marwick, 1953), it is now referred to Buchiidae (Cox & others, 1969; H. J. Campbell, 1983; Begg & Campbell, 1985; Sha & Fürsich, 1994).

Stratigraphic range.—Upper Triassic (upper Carnian–Norian) (H. J. Campbell, 1983). The genus was first described from the upper Carnian (Trechmann, 1918). It was later reported from Norian beds (H. J. Campbell, 1983). According to the latest stratigraphic revision (H. J. Campbell & Raine in Cooper, 2004), the type species ranges from Oretian to Warepan (=uppermost Carnian and Norian).

Paleogeographic distribution.—Austral (Fig. 28). J. Chen (1982a) reported *Hokonuia* sp. from the Carnian of southern China, but the figured specimen (pl. II, *14*) is not consistent with the diagnosis of the genus.

Austral domain: Late Triassic: latest Carnian–Norian of New Zealand (Trechmann, 1918; Wilckens, 1927; Marwick, 1953; Waterhouse, 1960; H. J. Campbell, 1983; Grant-Mackie, 1984).

Paleoautoecology.—B, E, S, Epi, Sed; By. Like the other members of the family Buchiidae, *Hokonuia* was an epibyssate bivalve. The byssus

emerged through the deep byssal notch in the right valve (Waterhouse, 1960). Although it was suggested that some buchiids might have a pseudoplanktonic mode of life (Wignall & Simms, 1990), we do not believe *Hokonuia* is among them, since its distribution is very limited.

Mineralogy.—Bimineralic (Waterhouse, 1960; Begg & Campbell, 1985; Carter, 1990a; Carter, Barrera, & Tevesz, 1998, p. 1002). Carter, Barrera, and Tevesz (1998) indicated that the inner shell layer of *Hokonuia* was probably aragonitic, in contrast to other buchiids, which had three calcitic shell layers. Outer shell layer: calcite (prismatic). Middle shell layer: calcite (foliated). Inner shell layer: aragonite (homogeneous).

Genus SICHUANIA Chen in Gu & others, 1976, p. 151

Type species.—Sichuania difformis Chen in Gu & others, 1976, p. 151.

Remarks.—Sichuania Chen, 1976, was also described as a new genus in Wen and others (December 1976). Chen (in Gu & others, 1976) included *Sichuania* in the Buchiidae. This was followed by Sha, Chen, and Qi (1990), but, although *Sichuania* had the typical form of shells of the family Buchiidae, it lacked a right anterior auricle, and, for this reason, Sha and Fürsich (1994) suggested that it belongs neither to Buchiidae nor to Monotoidea. But as these authors did not propose a new assignment, *Sichuania* is here treated in this family, while awaiting more information.

Stratigraphic range.—Upper Triassic (Norian) (Chen in Gu & others, 1976). Sichuania was first described by Chen in Wen and others (1976) from the Norian of China (Sichuan and Yunnan provinces). Later, it was also reported from Rhaetian beds (Hautmann, 2001b), although without a description or the original source data. Waterhouse (1980b) provisionally attributed his supposedly Lower Triassic specimens from New Zealand to Sichuania (?Sichuania marwicki Waterhouse, 1980b). But the sediments in which he found the specimens were not of that age, and the specimens were not well accommodated in this genus (see discussion for Marwickiella, p. 79).

Paleogeographic distribution.—Eastern Tethys (Fig. 28). Hautmann (2001b) mentioned Sichuania from the Norian and Rhaetian beds of Tibet and the Himalayas, but he did not indicate the source of the data. Kobayashi and Tamura (1983a) also quoted it from the Upper Triassic in several Chinese provinces and southern Tibet, but this record does not have a source of original data.

Tethys domain: Late Triassic: Norian of China (Chen in Gu & others, 1976; Sha, Chen, & Qi, 1990).

Paleoautoecology.—B, E, S, Epi, Sed; By. According to the translation of the original generic diagnosis provided by Waterhouse (1980b), the shell is inequivalve, with the left valve being more prominent and convex than the right, and it had a byssal sinus. Due to these characteristics, *Sichuania* should be regarded as an epibyssate bivalve, although Sha, Chen, and Qi (1990) only doubtfully assigned it this mode of life.

Mineralogy.—?Bimineralic (Carter, 1990a, p. 234). There are no data on *Sichuania* shell mineralogy and structure. Since its systematic position is not known, we cannot assign it the dominant mineralogy in the family. Nevertheless, it likely had a bimineralic shell, like most members of the order Pectinioida.

Genus MARWICKIELLA Sha & Fürsich, 1994, p. 21

Type species.—?Sichuania marwicki Waterhouse, 1980b, p. 1.

Remarks.—Waterhouse (1980b) tentatively assigned his specimens to *Sichuania*, thinking that they belonged to the family Buchiidae. We have already seen that *Sichuania* is probably not a buchiid, as it lacked the typical anterior auricle. But Waterhouse's specimens have this auricle and other diagnostic features of the family Buchiidae (Begg & Campbell, 1985; Sha & Fürsich, 1994). Since seemingly this species did not fit into any Buchiidae, Begg and Campbell (1985, p. 739) argued that it represented a new genus, and later Sha and Fürsich (1994) proposed the name *Marwickiella* for it.

Stratigraphic range.—Middle Triassic (Anisian). Although Waterhouse (1980b) originally referred the beds with *?Sichuania marwicki* to the Lower Triassic, Begg (1981) showed that the areas where Waterhouse collected this species were in fact of lower Anisian age.

Paleogeographic distribution.-Austral (Fig. 28).

Austral domain: Middle Triassic: Anisian of New Zealand (Waterhouse, 1980b; Begg, 1981; Begg & Campbell, 1985).

Paleoautoecology.—B, E, S, Epi, Sed; By. According to the description provided by Waterhouse (1980b) for the only species included in the genus, it was probably an epibyssate bivalve.

Mineralogy.—Bimineralic. According to Waterhouse (1980b), his specimens had a thin calcitic shell, although he did not perform microstructural studies of any kind. We assign the dominant mineralogy of family Buchiidae.

Genus ANNINGELLA Cox, 1958, p. 44

nom. nov. pro Anningia Cox, 1936, p. 468, *non* Broom, 1927, p. 227 (Amniota)

Type species.—Anningia carixensis Cox, 1936, p. 468.

Remarks.—Little is known about *Anningella*, because it is only known from its right valve. According to Sha and Fürsich (1994), it is practically indistinguishable from *Chaenocardia* Meek & Worthen, 1869 (a Carboniferous genus not discussed). In their opinion, this genus would be better located in the Asoellidae, but in the absence of any further study, we follow Cox and others (1969) and include it in the Buchiidae.

Stratigraphic range.—Lower Jurassic (Hettangian–Sinemurian) (Hallam, 1987; Warrington & Ivimey-Cook, 1990). Cox (1936) described the genus Anningia (renamed Anningella by Cox [1958]) from the Liassic of Dorset. Subsequently, Cox and others (1969) assigned it a lower Lower Jurassic range. There are some inconsistencies in the literature regarding the stratigraphic range of this genus; most authors reported it from Sinemurian beds of England (Hallam, 1976, 1977, 1987; Liu, 1995; Aberhan, 2001); however, Sepkoski (2002) assigned it a Rhaetian–Sinemurian range, allegedly taking data from Hallam (1977, 1981), but, in this last paper, the genus is not mentioned. Hallam and El Shaarawy (1982) quoted Anningella from the "Penarth group" of Rhaetian age, but later Warrington and Ivimey-Cook (1990) indicated that Anningella had its origin in the *Planorbis* zone (=Hettangian) of the Bristol Channel area, so we believe its presence in the Rhaetian is unlikely.

Paleogeographic distribution.—western Tethys (Fig. 28).

Tethys domain: Early Jurassic: Hettangian of England (Warrington & Ivimey-Cook, 1990); Sinemurian of England (Hallam, 1976, 1977, 1987; Liu, 1995; Aberhan, 2001). *Paleoautoecology.*—B, E, S, Epi, Sed; By. Like the other members of the family Buchiidae, *Anningella* was an epibyssate bivalve.

Mineralogy.—Bimineralic (Carter, 1990a, p. 234). There are no data on the shell of *Anningella*. Nevertheless, it likely had a bimineralic shell, like most members of the Order Pectinioida.

Family MONOTIDAE Fischer, 1887 in 1880–1887 Genus MONOTIS Bronn, 1830a, p. 284

Type species.—Pectinites salinarius von Schlotheim, 1820, p. 230. Remarks.—Although Cox and others (1969) only considered two subgenera of Monotis, M. (Monotis) and M. (Entomonotis) Marwick, 1935, p. 298, in the past three decades many more were named: Pacimonotis Grant-Mackie & Silberling, 1990; Eomonotis Grant-Mackie, 1978a, p. 102; Inflatomonotis Grant-Mackie, 1978a, p. 105; Maorimonotis Grant-Mackie, 1978a, p. 108; 1978d. Although these subgenera group species with different morphotypes and are biostratigraphically useful, they are not based on phylogenetic relationships (McRoberts, Krystyn, & Shea, 2008). However, McRoberts (2010) considered Eomonotis at genus level, following Tozer (1980). Monotis had a wide paleogeographic distribution during the Late Triassic and is a good biochronologic indicator, due to its relatively rapid morphologic change; thus, despite its limited stratigraphic range, many authors studied them from this point of view.

Stratigraphic range.—Upper Triassic (Norian–lower Rhaetian) (McRoberts, Krystyn, & Shea, 2008). Until very recently, *Monotis* was believed to be completely extinguished at the Norian–Rhaetian boundary (Wignall & others, 2007), but McRoberts, Krystyn, and Shea (2008) reported it from the lower Rhaetian.

Paleogeographic distribution.—Cosmopolitan (Fig. 29). McRoberts (1997a) mentioned *Monotis* from the Norian of Mexico, but he neither described nor figured the material.

Tethys domain: Late Triassic: Norian of Slovenia (Jurkovsek, 1982a, 1982b), China (C. Chen & Yu, 1976; J. Chen & Yang, 1983), Afghanistan (Polubotko, Payevskaya, & Repin, 2001), Iran (Westermann & Seyed-Emami, 1981; Hautmann, 2001b), western Caucasus (Russia) (Ruban, 2006a), northern Alps (Austria) (Grant-Mackie & Silberling, 1990; McRoberts, Krystyn, & Shea, 2008); Rhaetian of northern Alps (Austria) (McRoberts, Krystyn, & Shea, 2008; McRoberts, 2010).

Circumpacific domain: Late Triassic: Norian of British Columbia (Ward & others, 2004; Wignall & others, 2007), Alaska (accreted terranes) (Grant-Mackie & Silberling, 1990; Silberling, Grant-Mackie, & Nichols, 1997), Peru (Jaworski, 1922; Steinmann, 1929; Prinz, 1985), Bolivia (Beltan & others, 1987; Suarez-Riglos & Dalenz-Farjat, 1993), Chile (Thiele, 1967; Westermann, 1970; Z. Fang & others, 1998), Nevada (United States) (Grant-Mackie & Silberling, 1990), California (McRoberts, 2010), Japan (Nakazawa, 1964; Hayami, 1975; Ando, 1983, 1984, 1986, 1987; Ando, Noda, & Sato, 1987).

Austral domain: Late Triassic: Norian of New Zealand (Grant-Mackie, 1976, 1978a, 1978b, 1978c, 1978d, 1980a, 1980b; H. J. Campbell, 1983; MacFarlan, 1998).

Boreal domain: Late Triassic: ?Carnian, Norian of Primorie (Kiparisova, 1972); Norian of northeastern Russia (Kiparisova, Bychkov, & Polubotko, 1966; Kurushin, 1990; Klets, 2006), several localities

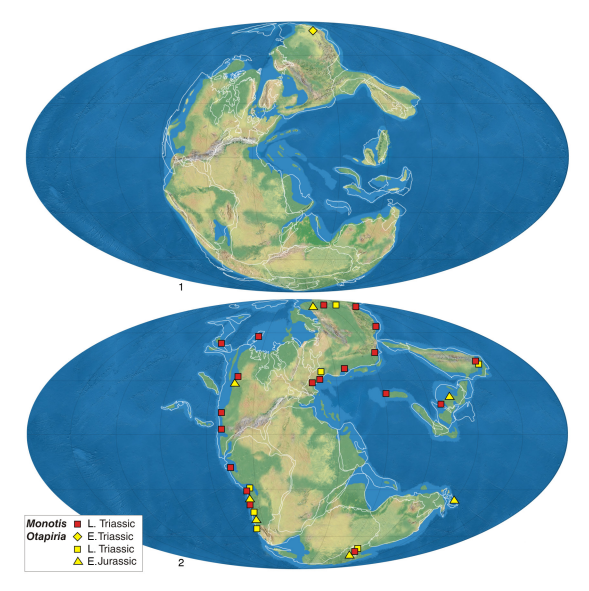


Figure 29. Paleogeographical distribution of Monotiidae (Monotis, Otapiria). 1, Early Triassic; 2, Late Triassic-Early Jurassic.

of Russia (Payevskaya, 1985), Alaska (Arctic terranes) (Silberling, Grant-Mackie, & Nichols, 1997), Arctic Archipelago (Canada) (Tozer, 1970).

Paleoautoecology.—B-Ps, E, S, Epi, Sed-FaM; By. Much has been speculated about the mode of life of *Monotis*. The interpretations range from benthic epibyssate on hard substrate (S. M. Stanley, 1972; Hallam, 1981; McRoberts, Krystyn, & Shea, 2008), byssate on aquatic plants (Ando, 1987; Hautmann, 2001b), pseudoplanktonic (Hayami, 1969a; S. M. Stanley, 1972; Silberling, Grant-Mackie, & Nichols, 1997) and even nektonic (Jefferies & Minton, 1965). These interpretations were based on the *Monotis* shell morphology and on the facies where different species are typically found.

A nektonic mode of life was proposed by Jefferies and Minton (1965) but was rejected by several authors (S. M. Stanley, 1972; Ando, 1987), since it is unlikely that *Monotis* could swim with its inequilateral and inequivalve shell, and some species had a byssal notch (Ando, 1987). According to S. M. Stanley (1972), a benthic

mode of life for some species and a pseudoplanktonic one for others would be more in agreement with the stratigraphic and taphonomic evidence. One of the strongest arguments in favor of a pseudoplanktonic mode of life is the great paleogeographic distribution Monotis had, but this could also be explained by a long-lasting planktotrophic larval stage, which is difficult to corroborate when the protoconch is not preserved. The genus is also usually found in deep-water environments, often deficient in oxygen, but this is not always the case, since Ando (1987) reported that some Japanese species were found in shallow, well-oxygenated environments. A pseudoplanktonic mode of life is also plausible with Monotis morphology, as it had a thin shell and its anterior auricle formed a pseudoctenolium, which indicates that the shell attached to objects by its byssus (Silberling, Grant-Mackie, & Nichols, 1997). Furthermore, the preferential orientation of the shells found in the fossil record, with the concave side upward, suggests they fell floating through the water column toward the bottom, in low-turbulence environments and with little *post mortem* transport (Silberling, Grant-Mackie, & Nichols, 1997). However, McRoberts, Krystyn, and Shea (2008) explain this orientation by suggesting that *Monotis* was epibenthic and lived in this position on the bottom and in cave fissures.

According to criteria by Wignall and Simms (1990) to distinguish between obligate and facultative pseudoplanktonic bivalves, and taking into account everything mentioned above, *Monotis* could be facultative, as it is not usually associated with objects suitable for fixation. As mentioned above, it is often, but not always (see Ando, 1987), found associated with deep facies. Moreover, it is often found in low-oxygen facies, and this could have several explanations, from shells falling into these facies because they had a pseudoplanktonic mode of life, or the presence of some kind of symbiotic organisms that made living in those environments possible, although there is no evidence to support this last possibility (A. G. Fischer & Bottjer, 1995).

According to this evidence, we agree with S. M. Stanley (1972) and consider that some species were epibyssate on the bottom and others might have had a facultative pseudoplanktonic mode of life.

Mineralogy.—?Calcitic (Carter, 1990a, p. 248; Carter, Barrera, & Tevesz, 1998). According to Carter, Barrera, and Tevesz (1998), members of family Monotidae have three calcitic shell layers, although Carter (1990a) noted that the presence of a thin sublayer of aragonite in the inner or middle shell layer of one or both valves was possible. According to the emended diagnosis offered by Carter (1990a) for the family Monotidae, the shell is built of foliated calcite with a thin homogeneous layer within one or both valves.

Genus OTAPIRIA Marwick, 1935, p. 302

Type species.—Pseudomonotis marshalli Trechmann, 1923, p. 270. *Remarks.*—According to Begg and Campbell (1985), Damborenea (1987b, 2002a), Ando (1987), and Carter (1990a), we include *Otapiria* in Monotidae (see Damborenea, 1987b, p. 154, for discussion on this topic); although Cox and others (1969), like other later authors (e.g., J. Yin, H. Yao, & Sha, 2004) included it in Aviculopectinidae. We consider *Lupherella* Imlay, 1967, p. 8, as a subgenus of *Otapiria* and *Pleuromysidia* Ichikawa, 1954, p. 52, as a synonym of *Otapiria* (see discussion in Genera not Included, p. 167).

Stratigraphic range.—Lower Triassic (Olenekian)–Upper Jurassic (Kimmeridgian) (Dagys & Kurushin, 1985; Damborenea, 1987b). Cox and others (1969) assigned it an Upper Triassic (Rhaetian)–Upper Jurassic (Tithonian) range. Damborenea (1987b), in her exhaustive review of the species that were attributed to the genus, recorded Kimmeridgian as the youngest record, as did Sha (1996). It is thought that *Otapiria* originated during the Early Triassic in the Boreal domain (Dagys & Kurushin, 1985), although it was not until Carnian times that this genus began to be abundant (Ando, 1988). Dagys and Kurushin (1985) proposed a new subgenus and new species, *Otapiria (Praeotapiria) bakevelliaeformis* from Lower Triassic beds. According to Ando (1988), this new subgenus is unnecessary, since morphological differences with other species of *Otapiria* are very subtle.

Paleogeographic distribution.—Cosmopolitan (Fig. 29). According to Sha (1996), *Otapiria* originated in the Boreal domain and probably also in the Austral, and its distribution was most likely conditioned by water temperature and substrate. It was especially recorded at high and middle latitudes and adapted to low temperatures (Damborenea,

1993). It is regarded as a bipolar (Damborenea, 1996a) or antitropical taxon (Sha, 1996). According to Sha (1996), during the Early Triassic–Late Jurassic interval, it was widely distributed in Austria, Carpathians, Alaska, northern and northeastern Siberia, Japan, New Caledonia, New Zealand, Peru, Chile, Argentina, Colombia, and Ecuador. It was only known from northeastern Siberia, New Zealand, and Chile from the Late Jurassic. Damborenea (1986) regarded this as a circumpacific genus with sporadic appearances in the Tethys, and this is corroborated by our data.

Tethys domain: Late Triassic: Norian of Austria (Grant-Mackie & Zapfe, 1973; Zapfe, 1973); Rhaetian of Austria (Zapfe, 1973; McRoberts, 2010); Early Jurassic: southeastern China (J. Yin, H. Yao, & Sha, 2004).

Circumpacific domain: Late Triassic: Carnian of Japan (Hayami, 1975; Ando, 1988); Norian of Chile (Chong & Hillebrandt, 1985; Covacevich, Pérez, & Escobar, 1991), Japan (Ando, 1988); Early Jurassic: Hettangian–Sinemurian of Chile (Escobar, 1980); Sinemurian of Canada (Poulton, 1991; Aberhan, 1998a, 1998b), Chile (Covacevich & Escobar, 1979; Aberhan, 1993, 1994a, 1998b).

Austral domain: Late Triassic: ?Andes (Argentina) (Covacevich, Pérez, & Escobar, 1991; Riccardi & others, 1997, 2004); Rhaetian of New Zealand (Marwick, 1953; Grant-Mackie, 1960) and Argentina (Damborenea & Manceñido, 2012); Early Jurassic: Hettangian–Sinemurian Neuquén basin (Argentina) (Damborenea, 1987b, 1993, 1996a, 2002a; Damborenea & Manceñido, 2005b), New Zealand (Marwick, 1953; Grant-Mackie, 1960), New Caledonia (Marwick, 1953).

Boreal domain: Early Triassic: Siberia (Dagys & Kurushin, 1985); Late Triassic: Carnian of Primorie (Kiparisova, 1972); Norian of Siberia (Okuneva, 1985, 1986); Norian–Rhaetian of Siberia (Klets, 2006); Early Jurassic: Hettangian–Sinemurian of Siberia (Polubotko, 1968b).

Paleoautoecology.—B, E, S, Epi, Sed; By. Ando (1988), after studying the environments and taphonomy related to *Otapiria*, concluded that it was probably an epibyssate bivalve. Although wood fragments were found associated with *Otapiria dubia* (Ichikawa, 1954), there was no evidence that the bivalve was attached to them. However, due to the shell orientation, they probably attached to each other by their byssus. Both the sediments in which *Otapiria* is usually found and its orientation indicate that its mode of life was different from *Monotis* (Ando, 1988). An epibyssate mode of life was also proposed for other species of *Otapiria* (Gruber, 1984).

Mineralogy.—?Calcitic (Carter, 1990a). No data are known on *Otapiria* shell microstructure. According to the emended diagnosis offered by Carter (1990a) for the family Monotidae, the shell is composed of foliated calcite with a thin homogeneous outer layer on one or both valves. Carter, Barrera, and Tevesz (1998) assigned a calcitic mineralogy to the three shell layers of family Monotidae members.

Family OXYTOMIDAE Ichikawa, 1958 Genus OXYTOMA Meek, 1864, p. 39

Type species.—Avicula muensteri Bronn, 1830b, p. 164.

Stratigraphic range.—?Lower Triassic, Middle Triassic (Ladinian)–Upper Cretaceous (Maastrichtian) (Abdel-Gawad, 1986; Waller & Stanley, 2005). Cox and others (1969) assigned a Upper Triassic–Upper Cretaceous range. However, over the years, Oxytoma

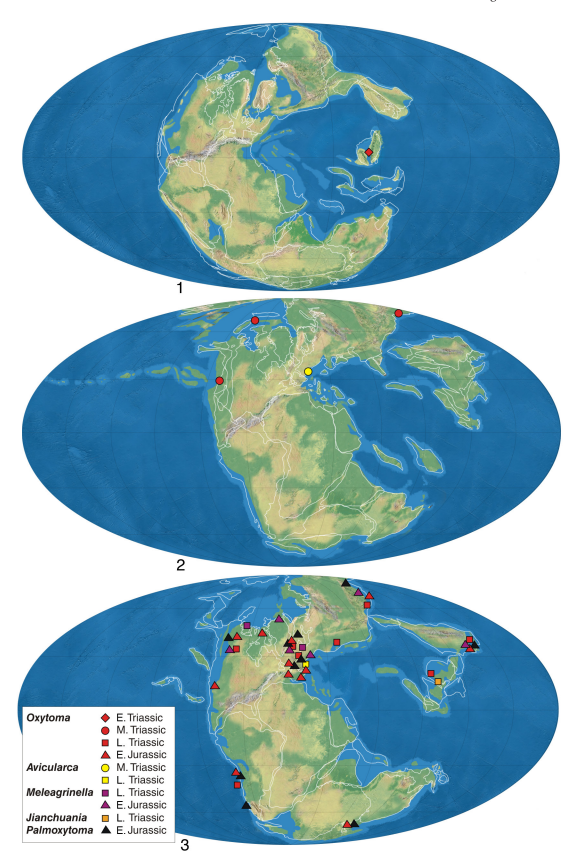


Figure 30. Paleogeographical distribution of Oxytomidae (Oxytoma, Avicularca, Meleagrinella, Jianchuania, Palmoxytoma). 1, Early Triassic; 2, Middle Triassic; 3, Late Triassic-Early Jurassic.

was repeatedly reported from Middle Triassic sediments. Waller (in Waller & Stanley, 2005, p. 38) considered that the oldest records of the genus to be *Oxytoma* sp. aff. *O. inaequivalve* Sowerby var. *intermedia* Emmrich, and *Oxytoma* sp. aff. *Oxytoma mojsisovicsi* Teller, from Triassic beds of Fujian province in southeastern China. They assumed a Lower Triassic age, because these species appeared to be associated with *Eumorphotis*. They also noted that *Oxytoma scythicum* (Wirth, 1936), reported from the Lower Triassic, was later assigned to *Towapteria* (family Bakevelliidae).

Paleogeographic distribution.—Cosmopolitan (Fig. 30).

Tethys domain: Early Triassic: ?southern China (Waller & Stanley, 2005); Late Triassic: China (J. Chen, 1982a); Norian of western Carpathians (Kollarova & Kochanová, 1973); Rhaetian of the Alps (Austria) (Tanner, Lucas, & Chapman, 2004; Tomašových, 2006a), England (Ivimey-Cook & others, 1999); Early Jurassic: Sinemurian of England, France, Spain, and Portugal (Liu, 1995), Italy (Monari, 1994), Hungary (Szente, 1996).

Circumpacific domain: Middle Triassic: Ladinian of Nevada (Waller & Stanley, 2005); Late Triassic: ?Chile (Moscoso & Covacevich, 1982; Damborenea, 1987b); Carnian of Japan (Hayami, 1975; Ando, 1988); Norian of Japan (Nakazawa, 1956, 1963, 1964; Hayami, 1975); Norian or Rhaetian of Chile (Chong & Hillebrandt, 1985); Rhaetian of Canada (Wignall & others, 2007); Early Jurassic: Hettangian of Nevada (United States) (Guex & others, 2003; Lucas & Tanner, 2004); Hettangian–Sinemurian of Chile (Escobar, 1980; Aberhan, 1994a), Canada (Poulton, 1991; Aberhan, 1998a, 1998b; Aberhan, Hrudka, & Poulton, 1998); Sinemurian of Japan (Hayami, 1975).

Austral domain: Early Jurassic: Hettangian–Sinemurian of New Zealand (Marwick, 1953).

Boreal domain: Middle Triassic: Ladinian of Siberia (Klets, 2006), Arctic Archipelago (Canada) (Tozer, 1961); Late Triassic: Primorie (Kiparisova, 1972); Early Jurassic: Hettangian–Sinemurian of Greenland (Liu, 1995), northeastern Russia (Milova, 1988).

Paleoautoecology.—B-Ps, E, S, Epi, Sed-FaM; By. Species belonging to *Oxytoma* had a well-developed byssal notch, suggesting that they were epibyssate bivalves. They had an elongated posterior auricle, and thus *Oxytoma* was compared with *Pteria* regarding its mode of life (Cox & others, 1969), living attached by the byssus to hydrozoa, shells, or other objects (Fürsich, 1980). According to Sha (1991), *Oxytoma* larvae were probably planktotrophic. However, some species, such as *Oxytoma inequivalve* (J. Sowerby, 1819), most likely could also have had a pseudoplanktonic mode of life, because they were found attached to *Echioceras* shells (Sinemurian of Dorset) (Wignall & Simms, 1990, fig. 3). The evidence suggests that *O. inequivalve* specimens were fixed to the ammonoids when they were alive.

Mineralogy.—Calcitic (Carter, 1990a, p. 249). Waller (in Waller & Stanley, 2005) suggested a probably entirely calcitic shell for their specimens of *Oxytoma (Oxytoma) grantsvillensis* Waller in Waller & Stanley, 2005, since there was no evidence of differential recrystallization of the inner shell layer. For the family Oxytomidae, Carter (1990a) indicated that the shells were mostly calcitic, but they could have had a very thin aragonitic middle layer of cross-lamellar microstructure; however, there is no record of this middle shell layer in *Oxytoma*. Outer shell layer: calcite (simple prismatic). Middle and inner shell layers: calcite (foliated).

Genus AVICULARCA Bubnoff, 1821, p. 281

Type species.—Avicula cardiiformis Münster, 1841, p. 78.

Stratigraphic range.—Middle Triassic (Ladinian)–Upper Triassic (Carnian). Cox and others (1969) assigned it an Upper Triassic range in the southern Alps. The type species was reported from sediments of this age (Wissman & Münster, 1841), and later, the genus was mentioned by Laube (1865) as being from the same stage. However, Bubnoff (1821) proposed *Avicularca* as a subgenus of *Avicula*, and he included three new species from the Italian Ladinian. Sepkoski (2002) assigned it a Carnian, ?Rhaetian range, following Crame (1996), but the last author did not adequately substantiate this range.

Paleogeographic distribution.—western Tethys (Fig. 30).

Tethys domain: Middle Triassic: Ladinian of Pedrazzo (Italy) (Kutassy, 1931); Late Triassic: southern Alps (Italy) (Wissman & Münster, 1841; Laube, 1865).

Paleoautoecology.—B, E, S, Epi, Sed; By. According to the generic diagnosis offered by Cox and others (1969, p. 344), *Avicularca* was probably an epibyssate bivalve.

Mineralogy.—Unknown. There are no data about *Avicularca* shell mineralogy. Since the allocation of this genus by Cox and others (1969) was doubtful, we cannot assign the predominant mineralogy of family Oxytomidae.

Genus MELEAGRINELLA Whitfield, 1885, p. 71

Type species.—Avicula curta Hall, 1852, p. 412.

Stratigraphic range.-Upper Triassic (Norian)-Lower Cretaceous (Albian) (Tozer, 1970; Wen, 1999). Although Cox and others (1969) assigned it an Upper Triassic (Rhaetian)–Upper Jurassic range, Meleagrinella was later reported from the Lower Cretaceous: Berriasian (X. Li, 1990), Valanginian (Kaim, 2001), and Albian (Wen, 1999). There are some disagreements regarding the origin of Meleagrinella. Sepkoski (2002) recorded that it ranges from Rhaetian times, based on data provided by Crame (1996), who surely followed Cox and others (1969). Meleagrinella was quoted from the Norian of British Columbia associated with Monotis (Westermann & Verman, 1967; Wignall & others, 2007), but specimens were neither figured nor described. The same occurred in the paper by Klets (2006), who considered Meleagrinella to have originated during the Anisian in the Boreal domain. He probably based this on data in Dagys and Kurushin (1985), who included Avicula polaris Kittl, 1907, and Pseudomonotis tasaryensis Voronetz, 1936, in Meleagrinella, and quoted the Anisian of Siberia from them. Avicula polaris was also reported from Carnian beds of Norway (Diener, 1923). Nevertheless, we leave this record as questionable, since we were unable to confirm this information, as none of these authors mentioned the original records. Tozer (1970) recorded Meleagrinella antiqua Tozer from the Norian of the Arctic Archipelago, and this age will be taken provisionally as the first record of the genus.

Paleogeographic distribution.—Tethys, Circumpacific, and Boreal (Fig. 30). Although *Meleagrinella* was present from the Late Triassic, it started to be more abundant and widely distributed during the Pliensbachian (Marwick, 1953; Duff, 1975; Wen, 1982; Pugaczewska, 1986; Jaitly, 1988; X. Li & Grant-Mackie, 1994; J. Chen, 1999; Harries & Little, 1999; Damborenea, 2002a; Delvene, 2003;

others, 2005; Fürsich & Thomsen, 2005; Zakharov & others, 2006). Tethys domain: Late Triassic: Rhaetian of Austria (Hallam & El Shaarawy, 1982; Early Jurassic: Hettangian–Sinemurian of northwestern Europe (Aberhan, 2001); Sinemurian of Europe (Quenstedt, 1856–1858), England (Liu, 1995).

Circumpacific domain: Early Jurassic: ?Hettangian of Canada (Poulton, 1991); Sinemurian of Japan (Hayami, 1961, 1975), Canada (Poulton, 1991; Aberhan, 1998a, 1998b).

Boreal domain: Late Triassic: Norian of Arctic Archipelago (Canada) (Tozer, 1970); Early Jurassic: Hettangian of northeastern Russia (Sey & others, 1981; Damborenea & others, 1992); Hettangian–Sinemurian of Greenland (Liu, 1995).

Paleoautoecology.—B-Ps, E, S, Epi, Sed-FaM; By. The external morphology of *Meleagrinella* suggests an epibyssate mode of life. Duff (1975) considered that while *Meleagrinella* may not have been strictly benthic, it may have attached to algae or even floating objects such as wood fragments, implying a pseudoplanktonic mode of life. Duff (1975) classified it as pendent. It is often reported from bituminous shales. However, Kaim (2001) found *Meleagrinella* specimens associated with cemented oysters; therefore, in this case, the mode of life was not necessarily pseudoplanktonic.

Mineralogy.—Bimineralic (Carter, 1990a, p. 249). Outer shell layer: calcite (prismatic). Middle shell layer: aragonite (cross-lamellar, in right valve). Inner shell layer: calcite (foliated).

Genus JIANCHUANIA J. Chen & Chen, 1980, p. 57, 59

Type species.—Pteria? problematica J. Chen in Ma & others, 1976, p. 287.

Stratigraphic range.—Upper Triassic (?Rhaetian) (J. Chen & Chen, 1980). J. Chen and Chen (1980) proposed *Jianchuania* and reported it from Upper Triassic beds of Yunnan (China), but they did not specify the stage. However, in the systematic discussion of the genus, they suggested that the specimens described by Healey (1908) as *Conocardium*? sp. and *Conocardium superstes* Healey, 1908, from the Rhaetian of Burma, are very similar and appeared in coeval deposits.

Paleogeographic distribution.—Eastern Tethys (Fig. 30). Jianchuania was endemic to Yunnan province, China.

Tethys domain: Late Triassic: ?Rhaetian of Yunnan (China) (J. Chen & Chen, 1980).

Paleoautoecology.—B, E, S, Epi, Sed; By. *Jianchuania* had a strongly inequilateral and inequivalve shell, with a deep byssal notch and an elongated posterior auricle, similar to *Pteria*. It was probably an epibyssate bivalve.

Mineralogy.—Bimineralic. There are no data about *Jianchuania* shell mineralogy. We therefore used data provided for the family Oxytomidae.

Genus PALMOXYTOMA Cox, 1962, p. 593

Type species.—Pecten cygnipes Young & Bird, 1822, p. 235.

Remarks.—Although Cox (1962) proposed *Palmoxytoma* as subgenus of *Oxytoma* and this was maintained in Cox and others (1969), we consider *Palmoxytoma* to be at the generic level following Damborenea (2002a).

Stratigraphic range.—Lower Jurassic (Hettangian–Pliensbachian) (Cox, 1962). Cox and others (1969) assigned the genus as lower to middle Lower Jurassic range, and this is maintained here. Although there are many papers that regard its origin as Hettangian (see paleogeographic distribution below), Guex and others (2003) and Lucas and Tanner (2004) recorded *Palmoxytoma* from Rhaetian beds of Nevada, but they neither described nor figured the specimens. Some species such as *Oxytoma mojsisovicsi* Teller, 1886, *Oxytoma koniensis* Tuchkov, 1956, and *Oxytoma gizhigensis* Milova, 1976, which are transitional between *Oxytoma* Meek, 1864, and *Palmoxytoma*, were considered to be in *O. (Palmoxytoma)* by some authors (e.g., Milova, 1976; but see Hayami, 1975), and if they were accepted as belonging to *Palmoxytoma*, its range would be extended to Upper Triassic.

Paleogeographic distribution.—Cosmopolitan (Fig. 30). Palmoxytoma had a bipolar distribution during the Hettangian (Damborenea, 1993; Sha, 1996; Aberhan, 1998b, 1999), and it appears to have been restricted to the Boreal domain during the Pliensbachian (Damborenea, 1993). Although it was also present in the Tethys and Circumpacific domains, it was not recorded at low paleolatitudes.

Tethys domain: Early Jurassic: Hettangian of England, Sweden, France, Switzerland (Cox, 1962).

Circumpacific domain: Early Jurassic: Hettangian of Chile (Aberhan, 1994a); Sinemurian of Canada (Poulton, 1991; Aberhan, 1998a), Japan (Hayami, 1975).

Austral domain: Early Jurassic: Hettangian of Argentina (Riccardi & others, 1991; Damborenea, 2002a; Damborenea & Manceñido, 2005b; Damborenea & Lanés, 2007), New Zealand (Trechmann, 1923).

Boreal domain: Early Jurassic: Hettangian of northeastern Siberia (Polubotko, 1968b).

Paleoautoecology.—B, E, S, Epi, Sed; By. We assign a mode of life similar to *Oxytoma*, but there is no evidence for a pseudoplanktonic mode of life.

Mineralogy.—Calcitic (Carter, 1990a, p. 249; Carter, 1990b, p. 371). Carter (1990b, p. 371) indicated that the outer shell layer of the type species of *Palmoxytoma* was built of prismatic calcite. We assume a calcitic mineralogy, as in *Oxytoma*, since there is no evidence of an aragonitic middle shell layer.

Family ASOELLIDAE Begg & Campbell, 1985

Begg and Campbell (1985) proposed the family Asoellidae, naming *Asoella* Tokuyama, 1959c, as type genus and including their new genus *Etalia*.

Genus ASOELLA Tokuyama, 1959c, p. 2

Type species.—Eumorphotis (Asoella) confertoradiata Tokuyama, 1959c, p. 4.

Remarks.—Although Cox and others (1969) included *Asoella* in the Aviculopectinidae following the original reference, Begg and Campbell (1985) proposed the family Asoellidae to accommodate *Asoella, Etalia* Begg & Campbell, 1985, p. 727, and probably also *Aucellina* Pompeckj, 1901, p 365. They related and characterized these three genera in having edentulous hinges and a subumbonal resilifer with anterior and posterior areas.

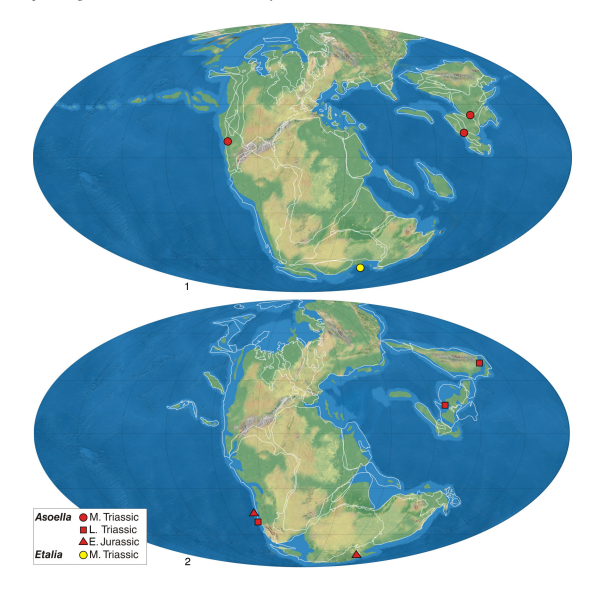


Figure 31. Paleogeographical distribution of Asoellidae (Asoella, Etalia). 1, Middle Triassic; 2, Late Triassic-Early Jurassic.

Stratigraphic range.—Middle Triassic (Anisian)–Lower Jurassic (Pliensbachian) (Sha, Chen, & Qi, 1990; Damborenea, 2002a). Cox and others (1969) assigned this genus a Norian range in Japan, since at that time only that information was available. Sepkoski (2002), based on Hayami (1975) and H. Yin (1985), assigned it an Anisian–Norian range. Regarding the origin of *Asoella*, it seems reasonably acknowledged that it was present in the Anisian of China (Kobayashi & Tamura, 1983b; H. Yin, 1985; Sha, Chen, & Qi, 1990; Tong & Liu, 2000) and Vietnam (Komatsu, Huyen, & Huu, 2010). Tong and Liu (2000) reported *Asoella illyrica* and *Asoella subillyrica* from the Anisian of China, but K. Huang and Opdyke (2000) suggested that the former is currently *Leptochondria illyrica* (Bittner) (see K. Huang & Opdyke, 2000, p. 80; Waller & Stanley, 2005, p. 35). Lu and Chen (1986) doubtfully assigned *Leptochondria subparadoxica* H. Yin & Yin to *Asoella*. As we have observed in these Chinese publica-

tions, there seems to be some confusion between *Leptochondria* and *Asoella*; *Asoella* records from the Middle Triassic of China should be reviewed. In addition, Waller and Stanley (2005) indicated that their Ladinian specimens from the United States, if they really belong to *Asoella*, would be the oldest records of this genus. In principle, since we cannot access all the information related to *Asoella* from China, we provisionally use Anisian as the oldest record. Although *Asoella* was considered to have vanished at the end of the Late Triassic in the past (Hallam, 1981, 1990), in recent years, it was reported from Sinemurian and Pliensbachian beds of South America (Damborenea, 2002a) and Hettangian–Pliensbachian beds of New Zealand (MacFarlan, 1998; N. Hudson, 2003). Its presence in New Zealand is accepted with caution, since the specimens were not described.

Paleogeographic distribution.—Tethys, Circumpacific, and Austral (Fig. 31).

Tethys domain: Middle Triassic: Anisian of southern China (Sha, Chen, & Qi, 1990); Anisian–Ladinian of northern Vietnam (Komatsu, Huyen, & Huu, 2010); Late Triassic: Carnian of China (J. Chen, 1982a; X. Li, Meng, & Wang, 2005).

Circumpacific domain: Middle Triassic: Ladinian of ?Nevada (United States) (Waller & Stanley, 2005); Late Triassic: Carnian– Norian of Japan (Hayami, 1975).

Austral domain: Late Triassic: Argentina (Damborenea & Manceñido, 2012); Early Jurassic: Sinemurian of Argentina (Damborenea, 1996a, 2002a; Damborenea & Manceñido, 2005b; Damborenea & Lanés, 2007); Hettangian–Sinemurian of ?New Zealand (MacFarlan, 1998; N. Hudson, 2003).

Paleoautoecology.—B, E, S, Epi-Un, Sed; By-R. Begg and Campbell (1985, p. 727), in their diagnosis of superfamily Monotoidea, which includes Asoellidae, stated, "Byssate Pectinina with right valve against substrate and with a distinct byssal notch throughout life." Damborenea (2002a) indicated that *Asoella asapha* (A. F. Leanza, 1942) was an epibyssate bivalve, at least in the juvenile stages, but it could have lived reclined on its right valve in the adult stage. Besides, since some species were consistently found in association with plant remains, it is possible that they could attach themselves to plants (Damborenea & Manceñido, 2012).

Mineralogy.—?Calcitic (Begg & Campbell, 1985; Carter, 1990a, p. 248). Begg and Campbell (1985) indicated that the outer shell layer was made of prismatic calcite. Carter (1990a) added that the shell was probably all foliated and calcitic, except the outer shell layer, which probably had a prismatic microstructure.

Genus ETALIA Begg & Campbell, 1985, p. 727

Type species.—Etalia johnstoni Begg & Campbell, 1985, p. 727. *Stratigraphic range.*—Middle Triassic (Anisian) (Begg & Campbell, 1985). *Etalia* was originally reported from the Anisian beds of New Zealand, and no other record of the genus is known. Since it had a restricted stratigraphic range (Etalian, New Zealand regional stage correlated with Anisian), it is a good index fossil for this stage (Begg & Campbell, 1985; H. J. Campbell & Raine in Cooper, 2004).

Paleogeographic distribution.—Austral (Fig. 31).

Austral domain: Middle Triassic: Anisian of Nelson and Southland (New Zealand) (Begg & Campbell, 1985).

Paleoautoecology.—B, E, S, Epi, Sed; By. Begg and Campbell (1985) recorded some specimens in life position and demonstrated that they had a gregarious habit, attaching to each other by the byssus, and reclining the right valves on the substrate, leaving the left valve free to open or close the shell. Authors considered it to be an opportunistic bivalve, a rapid colonizer of shallow-water environments.

Mineralogy.—?Calcitic (Begg & Campbell, 1985; Carter, 1990a, p. 248). Begg and Campbell (1985) indicated that the outer shell layer of the right valves was made of prismatic calcite. Although they did not mention anything about the inner shell layers, they pointed out that the shell was thin and probably calcitic. Carter (1990a), discussing the family Asoellidae, noted that the shell is almost entirely composed of calcite and probably of foliated microstructure, but the outer shell layer of the right valve contatined prismatic calcite.

Family PROSPONDYLIDAE Ptchelincev, 1960

=Terquemiidae Cox, 1964

Due to the wide range of different opinions in the literature about relations of the various genera included in this family, and as it is not an objective of this paper to review all of them, we follow mainly Hautmann (2001a) in his analysis of Prospondylidae, since it seems the most appropriate.

Genus TERQUEMIA Tate in Woodward, 1868, p. 65

nom. nov. pro Carpenteria Eudes-Deslongchamps, 1860, p. 127, non Gray, 1858, p. 269

Type species.—Carpenteria pectiniformis Eudes-Deslongchamps, 1860, p. 130.

Remarks.—Most references to this genus are often based on badly preserved specimens, resulting in a poorly known genus (Damborenea, 2002a; Hautmann & Golej, 2004). The main difficulty is that several genera in this family are externally very similar, and if details of the hinge and other internal characters of the specimens cannot be observed, it is very difficult to know to which genus they belong.

Stratigraphic range.—Lower Jurassic (Sinemurian), ?Upper Jurassic (Hautmann & Golej, 2004). Although Cox and others (1969) assigned it an Upper Triassic–Upper Jurassic range, according to Hautmann (2001a, p. 344): "Most Triassic species assigned to *Terquemia* in lower publications actually belong to *Newaagia* or *Enantiostreon*. Although there are some incompletely preserved specimens which might belong to *Terquemia*, there is no unequivocal record from rocks older than Lower Jurassic."

Ivimey-Cook and others (1999), and J. Yin and McRoberts (2006), reported Terquemia difformis (Schlotheim, 1820) from the Rhaetian of the Penarth Group (England) and from the Rhaetian-Hettangian transition layers in Tibet (China), respectively, but this species was referred to Umbrostrea by Márquez-Aliaga and others (2005), because it had an aragonitic inner shell layer (De Renzi & Márquez-Aliaga, 1980; Carter, 1990a; Carter, Barrera, & Tevesz, 1998) and ligament structure, hinge, and antimarginal ribs, typical of ostreids. J. Yin, Enay, and Wan (1999) reported it from Norian beds of the Himalayas (China), but they did not describe the material, and since we cannot be sure that it is really Terquemia, it will not be taken into account. The same occurs with most of the Triassic records of the genus. We follow Hautmann (2001a) in considering the oldest solid record to be Lower Jurassic. Regarding the youngest record accepted for Terquemia, it is hard to establish due to identification problems. Fürsich and Werner (1988) reported Terquemia from the Kimmeridgian of Portugal, but their specimens were only doubtfully assigned, as the specimens are articulated, and the hinge features, which are key to a proper allocation, cannot be seen. The same occurs with the specimens assigned by Damborenea (2002a) from the Toarcian of South America. No more records of Terquemia from the Upper Jurassic were found, apart from Fürsich and Werner (1988). Tentatively we indicate its range to be until the Upper Jurassic, following Hautmann and Golej (2004).

Paleogeographic distribution.—Tethys (Fig. 32). *Terquemia* was mentioned from the Tethys domain, but the only solid reference is in Hautmann and Golej (2004). Due to pending questions about the relationship of this genus, we cannot provide a complete distribution.

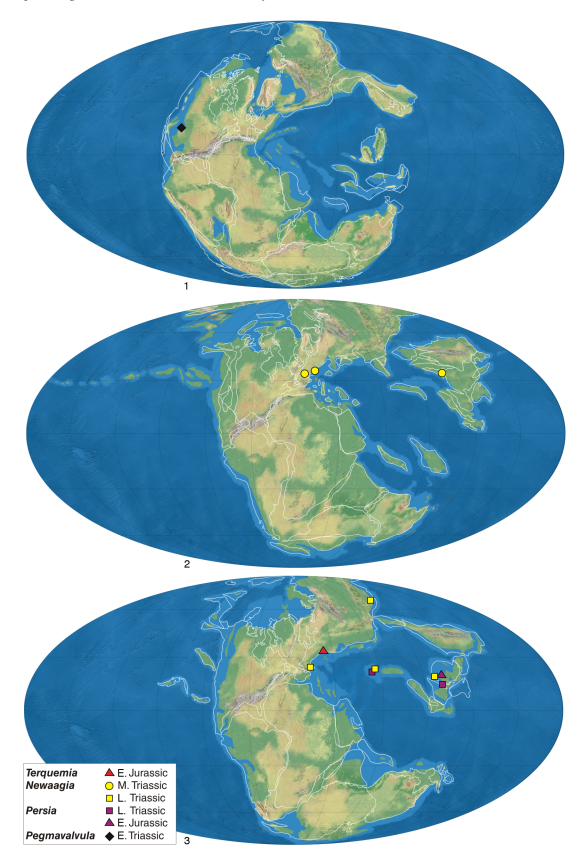


Figure 32. Paleogeographical distribution of Prospondylidae (Terquemia, Newaagia, Persia, Pegmavalvula). 1, Early Triassic; 2, Middle Triassic; 3, Late Triassic-Early Jurassic.

Tethys domain: Early Jurassic: Sinemurian of Western Carpathians (Slovakia) (Hautmann & Golej, 2004).

Paleoautoecology.—B, E, S, C, Sed; C. *Terquemia* was a cemented bivalve that attached to the substrate by its right valve, leaving the left one free. Unlike other cementing bivalves, such as *Persia*, *Terquemia* lacks byssal notch, so it should not have byssate juvenile stages. In many cases, shells of other bivalves were the substrate (Damborenea, 2002a).

Mineralogy.—Bimineralic (Hautmann & Golej, 2004). Hautmann and Golej (2004) described an outer shell layer subdivided into two sublayers, the outer prismatic and the inner foliated (both calcitic), in their specimens of *Terquemia (Dentiterquemia) eudesdeslongchampsi* Hautmann & Golej, 2004. The inner shell layer was recrystallized, but assuming that this layer microstructure was the same as in the rest of the family Prospondylidae, they believed it had a cross-lamellar microstructure (aragonitic).

Genus NEWAAGIA Hertlein, 1952, p. 275

nom. nov. pro Philippiella Waagen, 1907, p. 173, non Pfeffer in von Martens & Pfeffer, 1886, p. 119

Type species.—Spondylus obliquus Münster, 1841, p. 74.

Remarks.—Hertlein (1952) proposed the name *Newaagia* to replace *Philippiella* Waagen, 1907, as the latter name had already been used for another bivalve genus, *Philippiella* Martens & Pfeffer, 1886.

Some Triassic specimens attributed to *Spondylus* could better fit in *Newaagia* (Waller, 2006, p. 334): "So-called *Spondylus* from the Triassic, such as the many species described by Klipstein (1843 in 1843–1845), are spiny, multicostate bivalves cemented by their right valve and having ventrally migrating ligaments that leave a higher ligament area on the right valve than on the left. Although they superficially resemble *Spondylus*, those that I have examined have a pteriid-type resilium and lack hinge teeth. At least some of these are assignable to *Newaagia* Hertlein, 1952..."

Stratigraphic range.—Middle Triassic (Anisian)–Upper Triassic (Rhaetian) (Hautmann, 2001a). Cox and others (1969) assigned it an Upper Triassic (Carnian) range, and, with some doubt, they noted its presence in the Permian. *Newaagia* was later regarded as an exclusively Mesozoic genus (Newell & Boyd, 1970). Sepkoski (2002), following H. Yin (1985), assigned it an Anisian–Carnian range. It was reported from the Anisian (H. Yin & Yin, 1983), and subsequently, from Norian and Rhaetian beds (Hautmann, 2001a, 2001b).

Paleogeographic distribution.—Eastern Tethys and Boreal (Fig. 32). Newaagia was reported from the Norian of China (Sha, Chen, & Qi, 1990), but this was based on only one badly preserved specimen. In addition, it was also reported from the Norian of northeastern Asia (Polubotko & Repin, 1990), but the specimens are not figured or described, and there is no other information about the genus from that area.

Tethys domain: Middle Triassic: Anisian of the Dolomites (Italy) (Posenato, 2008b), Dolomites (Switzerland) (Zorn, 1971), northwestern China (Qinghai province) (H. Yin & Yin, 1983); Late Triassic: Carnian of Italy (Leonardi, 1943; Allasinaz, 1966); Norian–Rhaetian of Iran (Hautmann, 2001a, 2001b; Fürsich & Hautmann, 2005); Rhaetian (transitional layers of Rhaetian–Hettangian) of Tibet (China) (J. Yin & McRoberts, 2006). Boreal domain: Late Triassic: Carnian of Primorie (Kiparisova, 1972).

Paleoautoecology.—B, E, S, C, Sed; C. *Newaagia* was a cemented bivalve, attaching to hard substrates or to other shells by the umbonal area of the right valve (Newell & Boyd, 1970). It was a bioherm builder, according to Hautmann (2001b).

Mineralogy.—Bimineralic (Carter, 1990a, p. 251; Hautmann, 2001a). Outer shell layer: calcite (fibrous prismatic–foliated). Middle shell layer: aragonite (cross-lamellar–complex cross-lamellar). Inner shell layer: aragonite (simple prismatic).

Genus PERSIA Repin, 1996, p. 4 [365 in translation]

Type species.—Persia monstrosa Repin, 1996, p. 4 [365 in translation].

Remarks.—Although *Persia* had some external resemblance to *Newaagia* and *Terquemia*, it is distinguishable from both by differences in the auricles, ligament area, and ornamentation (see Repin, 1996, p. 365). *Persia* was emended by Hautmann (2001a). Although it was originally monospecific, subsequently J. Yin and McRoberts (2006) described a new species: *P. hallami* J. Yin & McRoberts, 2006.

Stratigraphic range.—Upper Triassic (Norian)–Lower Jurassic (lower Hettangian) (Repin, 1996; J. Yin & McRoberts, 2006). The genus was originally reported from the Norian of Iran (Repin, 1996), containing only the type species. Subsequently, it was found in Rhaetian beds of the same area (Hautmann, 2001a, 2001b) and in the Rhaetian–Hettangian transitional layers of Tibet (China) (J. Yin & McRoberts, 2006).

Paleogeographic distribution.—Tethys (Fig. 32).

Tethys domain: Late Triassic: Norian of Central Iran (Repin, 1996; Hautmann, 2001a, 2001b); Rhaetian of central Iran (Hautmann, 2001a, 2001b), Tibet (China) (J. Yin & McRoberts, 2006); Early Jurassic: early Hettangian of Tibet (China) (Tibeticum zone in J. Yin & others, 2007) (J. Yin & McRoberts, 2006).

Paleoautoecology.—B, E, S, C, Sed; C. The presence of a byssal notch between the anterior auricle and the disk may indicate that *Persia* had a byssate state before becoming cemented (Hautmann, 2001a). *Persia* was one of the reef-builder bivalves from the Late Triassic (Fürsich & Hautmann, 2005).

Mineralogy.—Bimineralic. Details of *Persia* shell microstructure are unknown. Probably it had a bimineralic shell, as do other members of family Prospondylidae.

Genus PEGMAVALVULA Newell & Boyd, 1970, p. 263

Type species.—Pegmavalvula gloveri Newell & Boyd, 1970, p. 263. Stratigraphic range.—lower Permian (Artinskian)–Lower Triassic (Olenekian) (Newell & Boyd, 1970, 1995). Although, according to Hautmann (2001a), Pegmavalvula was a Paleozoic genus, there is evidence that at least one species, *P. triassica* Newell & Boyd, 1995, was present in the Lower Triassic (Newell & Boyd, 1995).

Paleogeographic distribution.—Circumpacific (Fig. 32). Pegmavalvula was reported from Guadalupian to Artinskian levels of North America (Newell & Boyd, 1970); it was also reported from Greece in Changhsingian beds (Clapham & Bottjer, 2007), although these authors did not figure or describe the specimens.

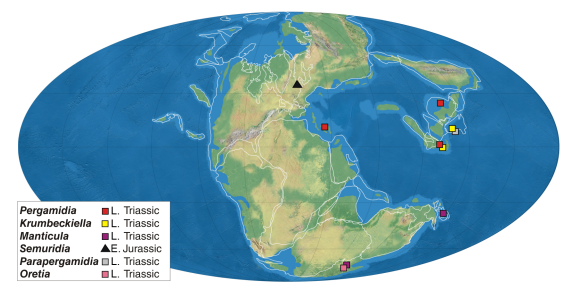


Figure 33. Paleogeographical distribution of Pergamidia, Krumbeckiella, Manticula, Semuridia, Parapergamidia, Oretia). Late Triassic-Early Jurassic.

Circumpacific domain: Early Triassic: Olenekian of Nevada (United States) (Newell & Boyd, 1995).

Paleoautoecology.—B, E, S, C, Sed; C. Species belonging to *Pegmavalvula* were cemented to the substrate with almost the entire surface of the right valve (Newell & Boyd, 1995). Shells had a byssal notch in juvenile stages, but this was closed in the adult stage. This fact was interpreted as evidence that they had an early byssate phase (pectiniform stage) before cementing to the substrate (Newell & Boyd, 1970).

Mineralogy.—Bimineralic. *Pegmavalvula* shell mineralogy is not known. It was probably bimineralic, as were other members of family Prospondylidae.

Family PERGAMIDIIDAE Cox in Cox & others, 1969 Genus PERGAMIDIA Bittner, 1891, p. 103

Type species.—Pergamidia eumenea Bittner, 1891, p. 103. Stratigraphic range.—Upper Triassic (Norian) (L. Lin & others, 2007). Cox and others (1969) assigned it a Norian range, which is confirmed by the reviewed literature. However, Sha and others (2005) indicated that *Pergamidia* lived from Carnian to Norian, but later, in another paper (L. Lin & others, 2007), *Pergamidia* was assigned an exclusively Norian genus.

Paleogeographic distribution.—Tethys (Fig. 33). *Pergamidia* was widely distributed throughout the Paleotethys suture, extending from the Carpathians to the Java Sea (Sha & others, 2005).

Tethys domain: Late Triassic: Norian of Turkey (Bittner, 1891, 1892), Timor (Indonesia) (Krumbeck, 1924), Yunnan province (China) (Cowper-Reed, 1927; Sha & others, 2005), Lungma region (China) (Wen & others, 1976), Qinghai province (China) (Sha, Chen, & Qi, 1990).

Paleoautoecology.—B, E, S, Epi, Sed; By. *Pergamidia* had a mytiliform shell and was monomyarian; it also had a very deep byssal notch, suggesting that it possessed a large byssus to attach itself to the substrate

(Sha & others, 2005). Due to the associated substrate type, its low diversity, and high abundance, it was probably able to put up with extreme conditions, even toxic environments (sulphuric) and those low in oxygen. They were found in large numbers in rift zones and island arcs, which were more or less affected by tectonic movements and volcanic activity. These types of environments imply shallow to deep waters and toxic sulfide seas (Sha & others, 2005). *Pergamidia* was considered an opportunist taxon capable of colonizing highly stressed environments where other organisms could not live, with very unstable populations that went extinct rapidly (Sha & Fürsich, 1994).

Mineralogy.—Bimineralic (Carter, 1990a, p. 196). There are no data about *Pergamidia* shell mineralogy. We assume a bimineralic shell, as in the other members of the order Pterioida.

Genus KRUMBECKIELLA Ichikawa, 1958, p. 196

nom. nov. pro Timoria Krumbeck, 1924, p. 218, *non* Kaye, 1919, p. 93

Type species.—Timoria timorensis Krumbeck, 1924, p. 221.

Remarks.—Krumbeck (1924) proposed the genus *Timoria*, being unaware that this name was already in use for an insect genus, *Timoria* Kaye, 1919. Ichikawa (1958) realized this and renamed the genus as *Krumbeckiella*.

Stratigraphic range.—Upper Triassic (Carnian–upper Rhaetian) (Sha, Chen, & Qi, 1990; J. Yin & McRoberts, 2006). The type species of *Krumbeckiella* was described by Krumbeck (1924) from the Norian of Timor, and this was the stratigraphic range assigned by Cox and others (1969). Subsequently, *Krumbeckiella* was also reported from upper Rhaetian beds (J. Yin & McRoberts, 2006). Sha, Chen, and Qi (1990) reported the genus from the Carnian, and it seems to be fairly common in sediments of that age (X. Wang & others, 2008).

Paleogeographic distribution.—Tethys (Fig. 33). Although the species Krumbeckiella cf. timorensis (Krumbeck, 1924) was men-

tioned from the Circumpacific domain by Newton (in Newton & others, 1987), Waller (in Waller & Stanley, 2005) showed that specimens described by Newton (in Newton & others, 1987) actually belong to *Mysidiella* Cox, 1964, and he renamed the species *Mysidiella newtonae* Waller in Waller & Stanley, 2005. In addition, Waller and Stanley (2005) reported the presence of *Krumbeckiella* at high paleolatitudes in the southern hemisphere, but we did not locate in the literature any reference of the genus from that area.

Tethys domain: Late Triassic: Carnian of Qinghai province (China) (Sha, Chen, & Qi, 1990); Norian of Timor (Indonesia) (Krumbeck, 1924), China (Wen & others, 1976), Tibet (China) (Kobayashi & Tamura, 1983a), Qinghai province (China) (Sha, Chen, & Qi, 1990); Rhaetian of Tibet (China) (J. Yin & McRoberts, 2006; J. Yin & others, 2007).

Paleoautoecology.—B, E, S, Epi, Sed; By. The mode of life of *Krumbeckiella* was probably similar to that of *Pergamidia*.

Mineralogy.—Bimineralic (Carter, 1990a, p. 196). *Krumbeckiella* shell mineralogy is unknown. As for other members of Pterioida, it could have had a bimineralic shell.

Genus MANTICULA Waterhouse, 1960, p. 428

Type species.—Mytilus problematicus Zittel, 1864, p. 28.

Remarks.—Although the type species was related to *Mytilus* Linnaeus, 1758, and *Myalina* de Koninck, 1842 in 1841–1844, Waterhouse (1960) proposed the new genus *Manticula*, characterized by hinge details and shell microstructure. He did not provide a systematic allocation, but Cox and others (1969) referred it to the Pergamidiidae.

Stratigraphic range.—Upper Triassic (?Carnian–Norian), Lower Cretaceous (Berriasian) (Waterhouse, 1960; Crame, 1995). It was originally reported from the Otamitan (then regarded as Carnian) of New Zealand, and this was the stratigraphic range assigned by Cox and others (1969); subsequently, it was also reported from the Norian stage (Freneix & Avias, 1977), and now the Otamitan is correlated with the Norian (H. J. Campbell & Raine in Cooper, 2004). Crame (1995) reported it from the Early Cretaceous (Berriasian) of Antarctica and regarded *Manticula* as a Lazarus taxon, without Jurassic representatives, which survived in Antarctica that acted as a refuge.

Paleogeographic distribution.—Austral (Fig. 38).

Austral domain: Late Triassic: ?Carnian, Norian of New Zealand and New Caledonia (Wilckens, 1927; Waterhouse, 1960; Freneix & Avias, 1977; W. Zhang & Grant-Mackie, 2001).

Paleoautoecology.—B, E, S, Epi, Sed; By. Most probably, *Manticula* was an epibyssate bivalve, as were the other members of the Pergamiidae; this is indicated by its mytiliform shell and the presence of byssal notch.

Mineralogy.—Bimineralic (Waterhouse, 1960; Carter, 1990a, p. 204). Outer shell layer: calcite (prismatic-homogeneous). Inner shell layer: aragonite (cross-lamellar).

Genus SEMURIDIA Melville, 1956, p. 116

Type species.—Semuridia jacksoni Melville, 1956, p. 116.

Remarks.—Semuridia was referred to the Pergamidiidae by Cox and others (1969); Waller and Stanley (2005) believed that it may be included in another clade due to differences in shell microstructure and ligament area, together with other genera of the family such as *Pergamidia*, *Krumbeckiella*, and *Manticula*. Carter (1990a) already indicated that if the inner shell layer of *Semuridia* is really nacreous, as stated by Cox and others (1969), it should be separated from the group at a subfamily or even family level.

Stratigraphic range.—Lower Jurassic (Sinemurian) (Cox & others, 1969). Semuridia was only recorded from Sinemurian beds (Cox & others, 1969; Hallam, 1976, 1977, 1987; Liu, 1995).

Paleogeographic distribution.—western Tethys (Fig. 33).

Tethys domain: Early Jurassic: Sinemurian of England (Cox & others, 1969; Liu, 1995).

Paleoautoecology.—B, E, S, Epi, Sed; By. Like the rest of the Pergamidiidae, *Semuridia* is externally mytiliform, it had a byssal notch, and was monomyarian, so it is interpreted to be an epibyssate bivalve.

Mineralogy.—Bimineralic (Cox & others, 1969, p. 314; Waller & Stanley, 2005). According to Cox and others (1969), *Semuridia* inner shell layer was nacreous (aragonite). Waller and Stanley (2005, p. 9) stated, "its outer shell layer, although not described, appears to be columnar prismatic based on figures of the left valve of *Semuridia dorsetensis* (Cox, 1926) in Cox [and others] (1969, fig. C44.4a)."

Genus PARAPERGAMIDIA L. Lin & others, 2007, p. 110

Type species.—Parapergamidia changtaiensis L. Lin & others, 2007, p. 111.

Stratigraphic range.—Upper Triassic (?upper Carnian–lower Norian) (L. Lin & others, 2007). *Parapergamidia* occurs in the Norian and probably also in the upper Carnian (L. Lin & others, 2007).

Paleogeographic distribution.—Eastern Tethys (Fig. 33).

Tethys domain: Late Triassic: late Carnian–early Norian of western Sichuan (southwestern China) (L. Lin & others, 2007).

Paleoautoecology.—B, E, S, Epi, Sed; By. Externally similar to *Pergamidia*, the main differences are based on the shape of the retractor muscles and the thickness of the shell (L. Lin & others, 2007). The byssal retractor muscle scars of *Parapergamidia* are very prominent and indicate it was fixed by the byssus. It had the same external mytiliform appearance as other members of the family to which it belongs. Its mode of life was probably similar to *Pergamidia*. Interpretation of the facies in which it occurs shows that it lived in deep-water environments (L. Lin & others, 2007). *Mineralogy.*—Bimineralic (Carter, 1990a, p. 196). *Parapergamidia* shell microstructure is unknown. Bimineralic mineralogy is assumed as in the other members of the order Pterioida.

Genus ORETIA Marwick, 1953, p. 62

Type species.—Oretia coxi Marwick, 1953, p. 62.

Remarks.—Marwick (1953) included *Oretia* in the family Pteriidae, but later Cox and others (1969) attributed it to an uncertain family within the superfamily Pectinoidea. Waterhouse (1979a) emended the generic diagnosis, and he tentatively included it in Pergamidiidae, but he also found some similarities with Monotiidae and Mysidiellidae.

Stratigraphic range.—Upper Triassic (lower Norian) (Marwick, 1953; H. J. Campbell & Raine in Cooper, 2004). The genus is restricted to the Oretian (New Zealand local stage), and halobiid and

ammonoid correlations suggest a lowest Norian age (H. J. Campbell & Grant-Mackie, 2000); the stage was correlated with lower Norian (H. J. Campbell & Raine in Cooper, 2004).

Paleogeographic distribution.—Austral (Fig. 33).

Tethys domain: Late Triassic: early Norian of New Zealand (Marwick, 1953; Waterhouse, 1979a).

Paleoautoecology.-B, E, S, Epi, Sed; By. Probably epibyssate.

Mineralogy.—Unknown. *Oretia* shell microstructure is unknown. Since its allocation to the family Pergamidiidae is uncertain, we cannot assign the dominant mineralogy in the family.

Superfamily HALOBIOIDEA Kittl, 1912 Family HALOBIIDAE Kittl, 1912

Over the past three decades, a large number of genera and subgenera related to Daonella and Halobia were proposed (e.g., Polubotko, Payevskaya, & Repin, 2001; Kurushin & Truschelev, 2001). Some authors specializing in this group do not agree (see McRoberts, 1993, 2000, and H. J. Campbell, 1994) which criteria used to characterize these taxa are diagnostic features. The new taxa not taken into account in this paper are: Perihalobia Gruber, 1976; Zittelihalobia Polubotko, 1984; Indigirohalobia Polubotko, 1984; Parahalobia Yin & Hsu, 1938, in C. Chen, 1976; Pacifihalobia Polubotko, 1990; Primahalobia Polubotko, 1988; Comatahalobia Polubotko in Polubotko, Payevskaya, & Repin, 2001; Magnolobia Kurushin & Truschelev, 2001 (see discussion for each of them in Genera not Included, p. 156). McRoberts (1993, p. 201-202) considered Perihalobia, Indigirohalobia, Zittelihalobia, Parahalobia, and Pacifihalobia to be synonyms of Halobia. In his own words: "the characters employed to construct these new taxa are inconsistent with the included taxa, are too narrowly defined to accommodate reasonable amounts of variation, or were erected to fit an *a priori* assumption of inferred phylogenetic relations. Many of the characters used to define the above genera are probably best used in specific rank." When Polubotko (1984) proposed the new genera, he emphasized the features related to the shape and size of the auricles and the type of ornamentation, characters used previously as diagnostic at the species level. In addition, there are a number of Daonella subgenera that were taken into account by Sepkoski (2002), which will not be considered here either: Dipleurites Kittl, 1912; Moussonella Turculet, 1972; Grabella Turculet, 1972; Arzelella Turculet, 1972; Loemmelella Turculet, 1972; Pichlerella Turculet, 1972; and Longidaonella Farsan, 1972; Sepkoski (2002) based these data on H. Yin (1985). Pichlerella and Arzelella are here regarded as subgenera of Daonella (Schatz, 2001b, 2004).

Halobia and *Daonella* were traditionally distinguished by the presence or absence of an anterior auricle, but in the publication of papers by Gruber (1976) and Polubotko (1984, 1988, 1990), there is no consensus about which characters are best suited to distinguish at subgeneric and generic levels (H. J. Campbell, 1994). A thorough review of the group is badly needed to establish the criteria for generic, subgeneric, and specific level discrimination, in order to restore stability, which this group had until the 1970s. In our opinion, the diverse senses in which members of the group were used, depended in part on the taxon concept of different authors. Sometimes, depending on the use to be given to the different taxa proposed, the list tends to swell

as a matter of convenience, for example, in biostratigraphy. We believe that new taxa should be defined as biological concepts, as far as possible.

We include *Daonella* Mojsisovics, 1874; *Aparimella* H. J. Campbell, 1994; *Halobia* Bronn, 1830a; and *Enteropleura* Kittl, 1912, in this family, following H. J. Campbell (1994), McRoberts (2000), and Waller and Stanley (2005). According to McRoberts (2000, p. 600), the first three are distinguished: "*Daonella* lacks an anterior auricle, *Aparimella* possesses an upper anterior auricle, and *Halobia* has a two-fold anterior auricle" (byssal tube of H. J. Campbell, 1994). *Daonella* is similar to *Enteropleura*, but its ornamentation is less marked, and the ligament is alivincular, very similar to *Bositra* De Gregorio, 1886 (Waller & Stanley, 2005).

One of the difficulties with this group is that the shell is very thin and the specimens are often found with the dorsal part broken, so that unless they are very well preserved, the ligament area cannot be observed. Many of the references in the literature were based on shell fragments or on internal and external molds only, and hence the confusion between genera is common.

The great abundance and distribution of halobiids during the Triassic and their high speciation rate made their species very useful as biochronological indicators, which is not common among bivalves.

Genus HALOBIA Bronn, 1830a, p. 282

Type species.—Halobia salinarum Bronn, 1830a, p. 282.

Stratigraphic range.—Upper Triassic (lower Carnian–middle Norian) (McRoberts, 1993, 2000). Although Cox and others (1969) assigned to *Halobia* a Middle–Upper Triassic range, according to McRoberts (2000, p. 602), *Halobia* did not appear until the early Carnian, because, "Earlier reports of Ladinian *Halobia* have now been determined to be either species belonging to other taxa such as *Daonella* and *Aparimella* (e.g., Campbell, 1994), or assigned to younger strata"

Paleogeographic distribution.—Cosmopolitan (Fig. 34). Although *Halobia* was considered to be a taxon with cosmopolitan distribution, according to McRoberts (1997b), it is not known in South America. However, Pérez-Barría (2004, 2006) reported *Halobia* from the Upper Triassic of Chile, but no systematic treatment of the specimens was made.

Tethys domain: Late Triassic: China (Cowper-Reed, 1927; Gou, 1993), Timor and Sumatra (Indonesia) (Krumbeck, 1914, 1924; Gruber in Kristan-Tollman, Barkham, & Gruber, 1987); Carnian of Tibet (China) (Sha, Johnson, & Fürsich, 2004), Qinghai province (China) (Sha, Chen, & Qi, 1990; Sha, 1995, 1998; Sha & Grant-Mackie, 1996), Xizang province (China) (C. Chen, 1982), Yugoslavia (Jurkovsek & Kolar-Jurkovsek, 1986), Italy (Leonardi, 1943; Nicora & others, 2007; McRoberts, 2010); Carnian-Norian of Slovakia (Kochanová, 1987), Sicilia (Italy) (Cafiero & Capoa de Bonardi, 1982; see records for European distribution), Apennines (Italy) (Capoa de Bonardi, 1970), Yugoslavia (Cafiero & Capoa de Bonardi, 1980), China (Wen & others, 1976), ?Singapore (Kobayashi & Tamura, 1968a), Turkey (Allasinaz, Gutnic, & Poisson, 1974); Norian of Tibet (China) (Sha, Johnson, & Fürsich, 2004), Qinghai province (China) (Sha, Chen, & Qi, 1990; Sha, 1995, 1998; Sha & Grant-Mackie, 1996), southern Russia (Okuneva, 1985, 1987), Austria (McRoberts, 2010).

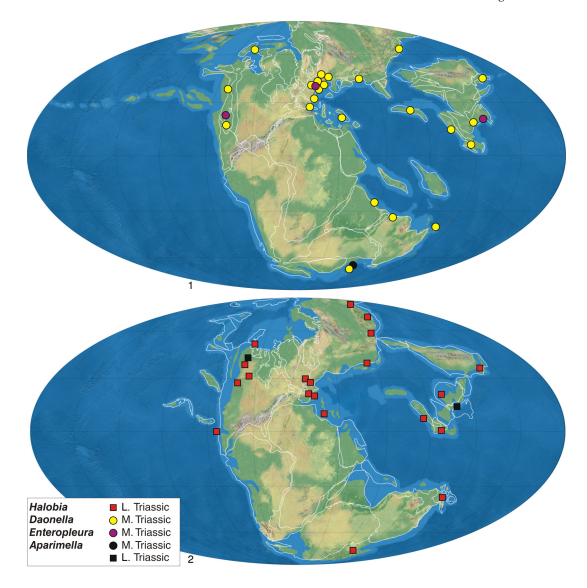


Figure 34. Paleogeographical distribution of Halobiidae (Halobia, Daonella, Enteropleura, Aparimella). 1, Middle Triassic; 2, Late Triassic.

Circumpacific domain: Late Triassic: early Carnian of British Columbia (Canada) (McRoberts, 2000), Japan (Ando, 1988); Carnian–Norian from various localities in North America (J. P. Smith, 1927; McRoberts, 1993, 1997b), Mexico (Lucas & González-León, 1994), Japan (Kobayashi & Ichikawa, 1949a; Nakazawa, 1964; Hayami, 1975; Tamura & others, 1975; Tanaka, 1989); Norian of British Columbia (McRoberts, 2010).

Austral domain: Late Triassic: ?Carnian, Norian of New Zealand and New Caledonia (Trechmann, 1918; Wilckens, 1927; Marwick, 1953; Grant-Mackie, 1960; H. J. Campbell, 1982, 1994).

Boreal domain: Late Triassic: Carnian of Svalbard (H. J. Campbell, 1994), Arctic Canada (McRoberts, 2010), Primorie (Kiparisova, 1972); Carnian–Norian of Arctic zone of Canada (Tozer, 1961, 1962; McRoberts, 1997b), Siberia (McRoberts, 1997b and references therein); Norian of Svalbard (H. J. Campbell, 1994).

Paleoautoecology.—B-Ps, E, S-Ch, Epi-Un, Sed-FaM; By-R. Many authors have speculated about mode of life of halobiids; Jefferies and Minton (1965), Hayami (1969a), S. M. Stanley (1972), Seilacher (1990), H. J. Campbell (1994), Etter (1996), McRoberts (1997b), Waller (in Waller & Stanley, 2005), and Schatz (2005), among many others.

To explain their wide distribution, association with low oxygen facies, morphology, and population structure, different modes of life were postulated: benthic semi-infaunal, benthic epifaunal on the substrate associated with chemosynthetic bacteria, epibyssate on plants or seaweed, pseudoplanktonic fixed to floating objects or other living organisms (such as ammonoids), nektonic (see Schatz, 2005). There are arguments for and against almost all suggested modes of life (good reviews of this topic are in H. J. Campbell, 1994, and Schatz, 2005).

Possibly several modes of life can be assigned to the species of this group (H. J. Campbell, 1994), although Schatz (2005) sup-

ported an epibenthic pleurothetic mode of life on soft substrate and adapted to low oxygen environments for daonellids, as the thin shell could have facilitated the oxygen exchange in these extreme environments. However, other authors, such as McRoberts (1997b) and H. J. Campbell (1994), suggested that a pseudoplanktonic mode of life cannot be ruled out, although Schatz (2005) argued that the morphology exhibited by daonellids and other evidence does not support this hypothesis. The wide distribution of these bivalves may be due to long-term planktotrophic larvae (H. J. Campbell, 1994; McRoberts, 1997b, 2000; Sha, 2003). The external morphology of some species could fit into a swimming mode of life (subcircular form, equivalve, and short hinge line), but the adductor muscle scar is small and falls just below the umbo, and the shells are too thin (Schatz, 2005). The discussion is far from settled, and it is possible that the different morphologies of members of the group indicate slightly different modes of life.

Mineralogy.—Bimineralic (H. J. Campbell, 1994). Outer shell layer: calcite (prismatic). Middle shell layer: aragonite (homogeneous). Inner shell layer: calcite (foliated or lamellar).

Genus DAONELLA Mojsisovics, 1874, p. 7

Type species.—Halobia lommeli Wissmann in Wissmann & Münster, 1841, p. 22.

Stratigraphic range.—Middle Triassic (Anisian–Ladinian). Cox and others (1969) assigned *Daonella* a Triassic range. The range is currently limited to the Middle Triassic. *Daonella pichleri* Mojsisovics, 1874, was mentioned also from the Carnian by Cafiero and Capoa de Bonardi (1980), but Schatz (2001b) reviewed this species and included it within the subgenus *Pichlerella*, and he limited its range to the *archelaus* zone (upper Ladinian). McRoberts (2010) mentioned that several occurrences of *Daonella* from lower Carnian beds are known, but they remain poorly documented.

Paleogeographic distribution.-Cosmopolitan (Fig. 34).

Tethys domain: Middle Triassic: China (Cowper-Reed, 1927; Lu & Chen, 1986); Anisian of southern China (Komatsu, Chen, & others, 2004), Malaysia (Vu Khuc & Huyen, 1998), Slovakia (Kochanová, 1985), Germany (Bartholomä, 1983), Switzerland (Zorn, 1971); Anisian–Ladinian of Italy (Pinna & Teruzzi, 1991; Brack & Rieber, 1993), southern China (J. Chen & others, 1992), Switzerland (Rieber, 1968, 1969); Ladinian of Spain (Schmidt, 1935; Llopis Lladó, 1952; Vía & Villalta, 1975; Vía, Villalta, & Esteban, 1977; Márquez-Aliaga, 1983, 1985; Budurov & others, 1991; Márquez-Aliaga & Martínez, 1996; Márquez-Aliaga & others, 2002, 2004; Márquez-Aliaga & Ros, 2002, 2003), southern Alps (Switzerland) (Schatz, 2001a), Italy (Kittl, 1912; Scandone & Capoa de Bonardi, 1966; Capoa de Bonardi, 1970; Schatz, 2001b), Bosnia, Romania, Turkey, India, Vietnam (Schatz, 2001b, and references therein), Timor (Indonesia) (Krumbeck, 1924), Malaysia (Vu Khuc & Huyen, 1998), Bulgaria (Budurov & others, 1991), southern Russia (Okuneva, 1985), Slovenia (Jurkovsek, 1983, 1984), Yugoslavia (Ramovs & Jurkovsek, 1983a, 1983b; Jurkovsek, 1983), China (Wen & others, 1976), Slovakia (Kochanová, Mello, & Siblík, 1975), Bulgaria (Stefanov, 1963), Timor (Indonesia) (Krumbeck, 1924), Afghanistan (Farsan, 1972, and references therein).

Circumpacific domain: Middle Triassic: Anisian of Japan (Nakazawa, 1961), Nevada (United States) (McRoberts, 2010); Ladinian of Japan (Hayami, 1975; Tamura & others, 1975).

Austral domain: Middle Triassic: Anisian of New Zealand and New Caledonia (H. J. Campbell, 1994).

Boreal domain: Middle Triassic: Anisian of Svalbard (H. J. Campbell, 1994); Anisian–Ladinian of Arctic Archipelago (Canada) (Tozer, 1961, 1962, 1970); Ladinian of northeastern Asia (Kurushin & Truschelev, 2001), Svalbard (H. J. Campbell, 1994; McRoberts, 2010), British Columbia (McRoberts, 2010).

Paleoautoecology.—B-Ps, E, S-Ch, Epi-Un, Sed-FaM; By-R. See Halobia (p. 91).

Mineralogy.—Bimineralic (H. J. Campbell, 1994). Outer shell layer: calcite (prismatic). Middle shell layer: aragonite (homogeneous). Inner shell layer: calcite (foliated or lamellar).

Genus ENTEROPLEURA Kittl, 1912, p. 162

Type species.—Daonella guembeli Mojsisovics, 1874, p. 8.

Remarks.—The systematic position of *Enteropleura* varied throughout its history, from a synonym of *Daonella* (Krumbeck, 1924) to a subgenus of *Daonella* (Capoa de Bonardi, 1970) or valid genus as interpreted by Cox and others (1969) and here. Although the genus was not widely understood, and somewhat neglected, several recent studies (Hopkin & McRoberts, 2005; Waller & Stanley, 2005; J. Chen & Stiller, 2007) clarified its position. Following these papers, the species of the genus are: *E. guembeli* Mojsisovics, 1874; *E. bittneri* Kittl, 1912; *E. lamellosa* (Kittl, 1912); *E. jenksi* Hopkin & McRoberts, 2005; *E. walleri* Chen & Stiller, 2007. The species *boeckhi* Mojsisovics, 1874 (quoted as *Daonella* (*Enteropleura*) *boeckhi* by Capoa de Bonardi, 1970) is regarded as a *Daonella* (J. Chen & Stiller, 2007). *Enteropleura* sp. A Waller in Waller & Stanley, 2005, is the same as *Enteropleura jenksi* Hopkin & McRoberts, 2005.

Stratigraphic range.—Middle Triassic (middle Anisian) (Waller & Stanley, 2005). According to the species included in the genus, its stratigraphic range is middle Anisian and not Anisian–Norian, as assigned by Cox and others (1969). Broglio-Loriga and others (1999) reported *Enteropleura* from the Carnian of the Dolomites (Italy), but this will not be taken into account, because the figured specimen lacks the dorsal part, critical to establish proper relations.

Paleogeographic distribution.—Tethys and Circumpacific (Fig. 34). Tethys domain: Middle Triassic: middle Anisian of Guangxi province (southern China) (J. Chen & Stiller, 2007), Hungary (Kittl, 1912).

Circumpacific domain: Middle Triassic: middle Anisian of Nevada (United States) (Hopkin & McRoberts, 2005; Waller & Stanley, 2005).

Paleoautoecology.—B-Ps, E, S-Ch, Epi-Un, Sed-FaM; By-R. See Halobia (p. 91).

Mineralogy.—Bimineralic (Waller & Stanley, 2005, p. 23–24). According to Waller and Stanley (2005), their specimens of *E. jenksi* provided the following information: Outer shell layer: calcite (prismatic). Middle shell layer: aragonite (?). Inner shell layer: calcite (?).

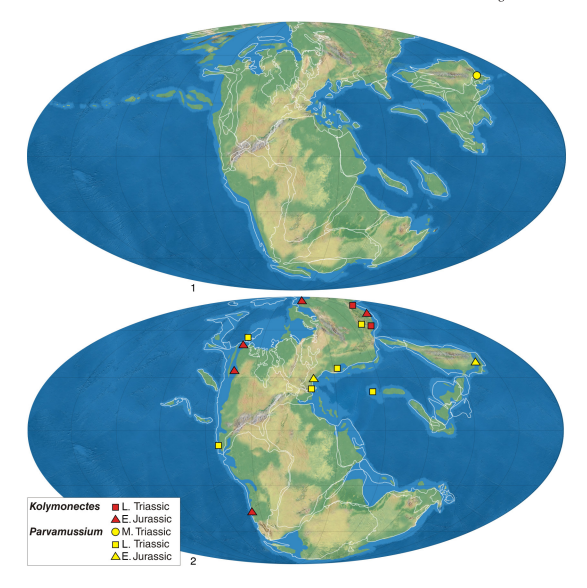


Figure 35. Paleogeographical distribution of Propeamussiidae (Kolymonectes, Parvamussium). 1, Middle Triassic; 2, Late Triassic-Early Jurassic.

Genus APARIMELLA H. J. Campbell, 1994, p. 65

Type species.—Daonella apteryx Marwick, 1953, p. 53.

Stratigraphic range.—Middle Triassic (upper Anisian)–Upper Triassic (lower Carnian) (H. J. Campbell, 1994). The stratigraphic range of *Aparimella* extends from Anisian to Carnian (H. J. Campbell, 1994).

Paleogeographic distribution.—Eastern Tethys, Austral, and Boreal (Fig. 34).

Tethys domain: Late Triassic: early Carnian of Yunnan (southern China) (J. Chen, 1982c).

Austral domain: Middle Triassic: late Anisian of New Zealand (H. J. Campbell, 1994); late Ladinian of New Zealand (Marwick, 1953; H. J. Campbell, 1994).

Boreal domain: Late Triassic: early Carnian of Svalbard (H. J. Campbell, 1994).

Paleoautoecology.—B-Ps, E, S-Ch, Epi-Un, Sed-FaM; By-R. See Halobia (p. 91).

Mineralogy.—Bimineralic (H. J. Campbell, 1994, p. 55). There are no specific data for *Aparimella* shell microstructure. We assign the data provided by H. J. Campbell (1994) for the family Halobiidae. Outer shell layer: calcite (prismatic). Middle shell layer: aragonite (homogeneous). Inner shell layer: calcite (foliated or lamellar).

Superfamily PECTINOIDEA Wilkes, 1810 Family PROPEAMUSSIIDAE Abbot, 1954 Genus KOLYMONECTES Milova & Polubotko in Bychkov & others, 1976, p. 67

Type species.—Aequipecten (?) anjuensis Milova, 1969, p. 182. Remarks.—Milova and Polubotko proposed Kolymonectes as a new genus in two almost simultaneous papers (in Bychkov & others, 1976, July 26; and in Milova, 1976, August 28). Although *Kolymonectes* was regarded a member of the family Entoliidae by some authors (e.g., Polubotko & Milova, 1986), we include it in the family Propeamussiidae, following Damborenea (1998, 2002a), on the basis of the lack of ctenolium and the presence of calcite in the prismatic outer layer of the shell, two diagnostic characters of this family (Damborenea, 1998). This same approach was followed by Aberhan (1998a). However, Waller (2006) included it in the family Entolioididae Waller, 2006. *Kolymonectes* is regarded as a member of a propeamusiid group without internal ribs (Damborenea, 1998).

Stratigraphic range.—Upper Triassic (?Norian)–Lower Jurassic (lower Toarcian) (Damborenea, 2002a). Kolymonectes appeared in the upper Norian of the Boreal domain (Milova & Polubotko in Milova, 1976); Kurushin (1990) and Polubotko and Repin (1990) recorded Kolymonectes from the upper Norian (=?Rhaetian) and Hettangian. Aberhan (1998a) believed that the genus was present until the Middle Jurassic, but he did not indicate the source of this information. The youngest checked records of the genus are upper Pliensbachian (Damborenea, 2002a, 2002b) and lower Toarcian (e.g., Damborenea & others, 1992). Zakharov and others (2006) considered the record to be the lower Toarcian of northern Siberia and the Arctic area, but they did not figure or describe any specimen from than age.

Paleogeographic distribution.—Circumpacific, Austral, and Boreal (Fig. 35). *Kolymonectes* was distributed in the Boreal domain during the Late Triassic and exhibited a bipolar distribution during the Early Jurassic (Damborenea, 1993, 1996a, 1998, 2001), and although it was also reported from the Circumpacific domain, these records are located at high paleolatitudes.

Boreal domain: Late Triassic: northeastern Russia (Damborenea, 1998); Norian of northeastern Asia (Kurushin, 1990; Polubotko & Repin, 1990); late Norian–Rhaetian of Siberia (Milova & Polubotko in Milova, 1976); Early Jurassic: Hettangian–Sinemurian of northeastern Russia (Milova, 1988); ?Hettangian, Sinemurian of Magadan (Russia) (Polubotko & Milova, 1986), Arctic zone of Canada (Aberhan, Hrudka, & Poulton, 1998); Sinemurian of northern Russia (Milova & Polubotko in Milova, 1976).

Circumpacific domain: Early Jurassic: Sinemurian of western Canada (Aberhan, 1998a, 1998b).

Austral domain: Early Jurassic: Hettangian–Sinemurian of Argentina (Damborenea & Manceñido, 2005b); Sinemurian of Argentina (Damborenea, 1998, 2002a).

Paleoautoecology.—B, E, S, Un, FaM; R-Sw. In adult specimens, the byssal gape is not usually observed. *Kolymonectes* probably had an earlier byssate stage, but adults lived reclining on the substrate (Damborenea, 1998, 2002a). According to S. M. Stanley (1972), it was probably a good swimmer, since it had a thin shell, the valves were of the same convexity, the auricles were equal, and the umbonal angle was large enough to allow swimming cycles. Propeamusiids are currently restricted to deep environments, but this was not the case during the Jurassic, as they were present both in deep environments associated with low-oxygen facies (without other benthic organisms), and in coastal environments (Damborenea, 1998, 2002a).

Mineralogy.—Bimineralic (Damborenea, 1998). Damborenea (1998) proposed the presence of a calcitic outer shell layer at least in the right valve, based on indirect evidence, provided by the

shell of specimens of *Kolymonectes weaveri* Damborenea, 1998. She also stated that the observed characters of the shell agree with the distribution of the shell layers in the family Propeamussiidae (see mineralogy for *Parvamussium* below).

Genus PARVAMUSSIUM Sacco, 1897, p. 48

Type species.—Pecten (Pleuronectes) duodecimlamellatum Bronn, 1831, p. 116.

Remarks.—Sacco (1897) distinguished 3 subgenera of Amussium: Propeamussium, species without ornamentation and large size, Eocene-Recent; Parvamussium, species very similar to Propeamussium but smaller, Cretaceous-Recent; and Variamussium, small shells internally ribbed, which include Jurassic forms, some Tertiary, and some modern forms. Subsequently, Cox and others (1969) regarded Variamussium as a junior synonym of Parvamussium, so that the arrangement was as follows: Propeamussium (Propeamussium) and Propeamussium (Parvamussium). Unfortunately, the stratigraphic ranges assigned to the two groups in Cox and others (1969) were mixed, and, from then on, Propeamussium sensu stricto was frequently cited from the Lower Jurassic, but these species should have been included in Parvamussium instead. Both groups are now recognized at the generic level (see Damborenea, 1998, p. 148-149). Parvamussium and Propeamussium can be distinguished as follows (Damborenea, 1998, p. 149): "In Propeamussium, no external sculpture, no byssal notch, equal auricles and a lateral gape, and, in Parvamussium, smaller size, well developed ornamentation on right valve, byssal notch and no lateral gape."

Propeamussium is not included here because its revised stratigraphic range is from the Cretaceous to the present (see discussion in Genera not Included, p. 168).

Filamussium Waller, 2006 (type species: *Pecten schafhäutli* Winkler, 1859) is not included here, as we believe the proposition of this genus was unnecessary in view of the above (see discussion in Genera not Included, p. 161).

Following these arguments, specimens from the Triassic and Lower Jurassic (until Sinemurian) assigned to *Propeamussium (Propeamussium)* by Johnson (1984) and J. Yin and Grant-Mackie (2005), to *Propeamussium (Variamussium)* by Hautmann (2001b), and to *Propeamussium* by McRoberts (1997a), are considered to belong in *Parvamussium*.

Stratigraphic range.—Middle Triassic (Anisian)–Holocene (Nakazawa, 1961; Damborenea, 1998). Cox and others (1969) assigned to *Parvamussium* an Upper Cretaceous–Holocene range, but we explained above why we consider *Parvamussium* to range from the Triassic to the Recent. Although it was more abundant during the Late Triassic, especially during the Norian (see paleogeographic distribution, above), its oldest record is from the Anisian of Japan with *Propeamussium (Variamussium)* n. sp. indet. (Nakazawa, 1961).

Paleogeographic distribution.—Tethys, Circumpacific, and Boreal (Fig. 35). Although during our study interval we only find records from the Tethys and Circumpacific domains, later in the Jurassic the genus was also distributed in South America (Aberhan, 1994a, 1998b; Damborenea, 2002a) and Tibet (J. Yin & Grant- Mackie, 2005).

Tethys domain: Late Triassic: Norian of Iran (Hautmann, 2001b), western Carpathians (Kochanová, 1967; Kollarova & Kochanová, 1973); Rhaetian of Lombardy (Italy) (Allasinaz, 1962), Iran (Haut-

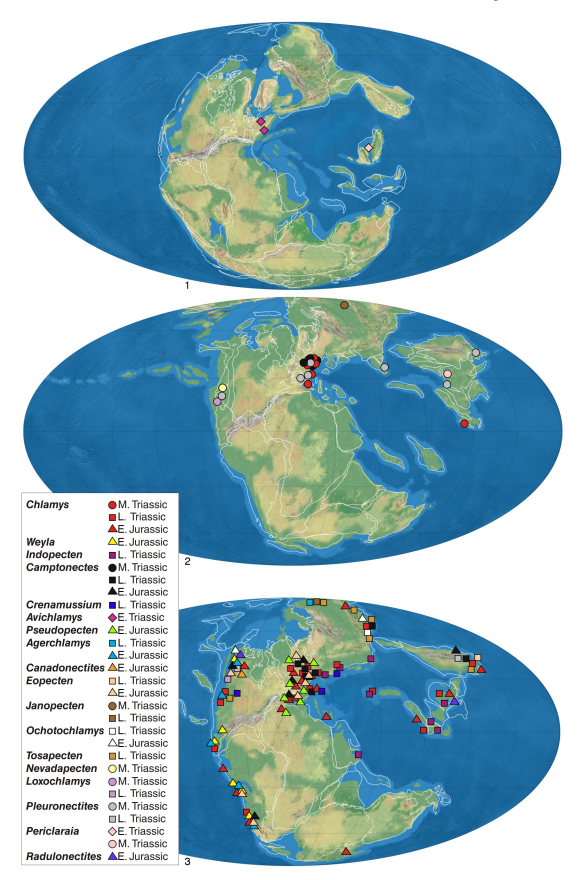


Figure 36. Paleogeographical distribution of Pectinidae (Chlamys, Weyla, Indopecten, Camptonectes, Crenamussium, Avichlamys, Pseudopecten, Agerchlamys, Canadonectites, Eopecten, Janopecten, Ochotochlamys, Tosapecten, Nevadapecten, Loxochlamys, Pleuronectites, Periclaraia, Radulonectites). 1, late Permian–Early Triassic; 2, Middle Triassic; 3, Late Triassic–Early Jurassic.

mann, 2001b); Early Jurassic: Sinemurian of ?Europe (Johnson, 1984).

Circumpacific domain: Middle Triassic: Anisian of Japan (Nakazawa, 1961); Late Triassic: Norian of Sonora (Mexico) (McRoberts, 1997a); Early Jurassic: Japan (Hayami, 1957b).

Boreal domain: Late Triassic: Norian of Arctic zone of Canada (Tozer, 1962).

Paleoautoecology.—B, E, S, Un, FaM; R-Sw. Most Recent species live in deep-water environments, their shells are very fragile, and this is a good place to be safe from predators (Beesley, Ross, & Wells, 1998). Some species can live 600 m deep, although they are also found in shallow water. They live freely reclining (Waller, 2006), and at least juveniles have a byssate stage in (Johnson, 1984; Damborenea, 2002a). Although a pseudoplanktonic mode of life was suggested for some propeamussiids, there is no morphological evidence (Johnson, 1984). Living species, at least, of *Parvamussium* can probably swim for short distances, since it is known that they feed on pelagic organisms (and benthic ones), and they may even actively catch them. A swimming habit is entirely compatible with their morphology (Johnson, 1984).

Mineralogy.—Bimineralic (Waller, 2006; but see Carter, 1990a, p. 256–257). Outer shell layer: calcite (prismatic). Middle shell layer: calcite (foliated). Inner shell layer: aragonite (cross-lamellar).

Family PECTINIDAE Wilkes, 1810 Genus CHLAMYS Röding in Bolten, 1798, p. 161

Type species.—Pecten islandicus Müller, 1776, p. 248.

Remarks.—A synapomorphy of the subfamily Chlamydinae is the presence of aragonite cross-lamellar structure in the shell (Waller & Marincovich, 1992), but this feature only occurs in the specimens younger than the Tertiary. The inclusion of large numbers of Mesozoic species in the genus used in its broadest sense is an unsolved problem, and a review of *Chlamys* and other related genera diagnoses is needed (Damborenea, 2002a).

Although Praechlamys Allasinaz, 1972, was considered as a genus by some authors (e.g., Waller & Marincovich, 1992; Monari, 1994; Szente, 1996; Damborenea, 2002a), it was originally proposed as a subgenus of Chlamys. Other authors (e.g., Posenato, 2008b) still regard it as a subgenus of Chlamys, while Hautmann (2001b) questioned its validity altogether, since, in his opinion, the type of ornamentation is not an important taxonomic character at subgenus level. But Allasinaz (1972) used mainly differences in ornamentation to separate the subgenera C. (Chlamys), C. (Praechlamys), and C. (Granulochlamys). Until this controversy is solved, Praechlamys is here taken in its original sense. It is necessary to establish the diagnostic characters that define each taxon, because the diagnosis given by Allasinaz (1972) was too lax (Damborenea, 2002a). Some of the Triassic species traditionally attributed to Chlamys could fit into Praechlamys if considered at generic level, with an emended diagnosis, but other species are difficult to accommodate in other genera, and therefore a new taxon is required for them (Damborenea, 2002a).

As this discussion is beyond the scope of this paper, *Chlamys* is considered in a broad sense and present in the Triassic, but this is just a temporary solution for a group that has been particularly problematic since its conception.

Stratigraphic range.—Middle Triassic (Anisian)–Holocene (Cox & others, 1969; Waller, 2006). Cox and others (1969) assigned it a Triassic–Holocene range. According to Waller (2006), the oldest member of the subfamily Chlamydiinae is *Praechlamys reticulata* (Schlotheim, 1823 in 1822–1823) from the Anisian of Germany, but Hautmann (2010) stated that this species was not a pectinid. There are older records (e.g., *Praechlamys wuxingensis* Li, in Nanjing Institute of Geology and Mineral Resources, 1982, from the Lower Triassic of China), but those specimens are very doubtful pectinids (see Waller, 2006, p. 331).

Paleogeographic distribution.—Cosmopolitan (Fig. 36).

Tethys domain: Middle Triassic: Anisian of Bosnia (Allasinaz, 1972), Italy (Posenato, 2008b), Switzerland (Zorn, 1971); Ladinian of Italy (Rossi Ronchetti, 1959; Allasinaz, 1972), Germany (Allasinaz, 1972), Spain (Márquez-Aliaga, 1983, 1985; López-Gómez & others, 1994), Slovakia (Kochanová, Mello, & Siblík, 1975), Malaysia (Tamura, 1973); Late Triassic: China (Wen & others, 1976; J. Chen, 1982a; Gou, 1993); Carnian of Italy (Allasinaz, 1966, 1972; Fürsich & Wendt, 1977), Switzerland, Hungary, and Bosnia (Allasinaz, 1972), Carpathians (Hungary) (Turculet, 1988), Spain (Martín-Algarra, Solé de Porta, & Márquez-Aliaga, 1995); Norian-Rhaetian of Iran (Hautmann, 2001b); Rhaetian of Czech Republic (Allasinaz, 1972), Austria (Allasinaz, 1972; Johnson, 1984), England (Johnson, 1984; Ivimey-Cook & others, 1999), Italy (Allasinaz, 1962; Johnson, 1984), Tibet (China) (J. Yin & McRoberts, 2006); Early Jurassic: Hettangian of England and France (Johnson, 1984; Liu, 1995), Italy (Allasinaz, 1962; Gaetani, 1970; Johnson, 1984), Wales (Johnson, 1984), Tibet (China) (J. Yin & McRoberts, 2006); Hettangian-Sinemurian of Spain (Calzada, 1982); Sinemurian of Switzerland (Johnson, 1984), Austria and Hungary (Szente, 1996), England, France, Spain, and Morocco (Liu, 1995), Vietnam (Sato & Westermann, 1991), Turkey (M. A. Conti & Monari, 1991).

Circumpacific domain: Late Triassic: Japan (Nakazawa, 1952; Tokuyama, 1959b); Carnian of Japan (Hayami, 1975); Norian of Oregon (United States) (Newton, 1986; Newton in Newton & others, 1987); Rhaetian of Nevada (United States) (Laws, 1982; Hallam & Wignall, 2000), Sonora (Mexico) (McRoberts, 1997a); Early Jurassic: Hettangian of Peru (Johnson, 1984); Hettangian–Sinemurian of Chile (Aberhan, 1994a); Sinemurian of western Canada (Aberhan, 1998a), Japan (Hayami, 1964, 1975).

Austral domain: Late Triassic: Rhaetian of Argentina (Riccardi & others, 2004); Early Jurassic: Hettangian–Sinemurian of Argentina (Damborenea, 2002a), of New Zealand (Damborenea & Manceñido, 1992); Sinemurian of Argentina (Damborenea & Manceñido, 2005b).

Boreal domain: Late Triassic: Carnian of Primorie (Kiparisova, 1972); Early Jurassic: Hettangian of northeastern Russia (Milova, 1976); Sinemurian of northeastern Russia (Polubotko & Milova, 1986).

Paleoautoecology.—B, E, S, Epi-Un, FaM; By-R-Sw. As inferred by functional morphology (Johnson, 1984), most fossil species probably lived attached by the byssus, at least during juvenile stages, in the same way that living species do. Many of them would be able to swim as well. Depending on the substrate in which they lived, they could spend most of their life as epibyssate on hard substrates. Other species are found in soft substrates, and these have spiny shells and probably lived reclined, using the spines to anchor them. See Johnson (1984) for a broad discussion on the ecology and modes of life of several species.

Mineralogy.—Bimineralic (Carter, 1990a, p. 260). The group is very heterogeneous, and it is difficult to generalize about the microstructure of the shell layers (for details on the various species, see Carter (1990b, p. 381–383).

Genus WEYLA J. Böhm, 1922, p. 138

Type species.—Pecten alatus Buch, 1838, p. 55.

Remarks.—Although Damborenea (1987b) and Damborenea and Manceñido (1988) regarded *Weyla* as a neitheid, this family was not included in the systematic arrangement proposed by Amler (1999), which is followed here. Waller (2006) included this genus in the family Pectinidae by the presence of ctenolium.

Stratigraphic range.—Lower Jurassic (Hettangian-Toarcian) (Damborenea & Manceñido, 1988; Aberhan, 1994a). Cox and others (1969) assigned it a Upper Triassic-Middle Jurassic range, but they included three subgenera: W. (Weyla) from the Lower Jurassic, W. (Pseudovola) Lissajous, 1923, p. 169, from the Middle Jurassic, and W. (Tosapecten) Kobayashi & Ichikawa, 1949b, p. 166, from the Upper Triassic. Currently, Pseudovola and Tosapecten are considered to be separate genera (Hayami, 1975; Damborenea, 1987b), so the remaining range is Lower Jurassic. We include two subgenera of Weyla from the Lower Jurassic, W. (Weyla) and W. (Lywea) Damborenea, 1987b. Damborenea and Manceñido (1988) indicated that the genus was present from Sinemurian to Toarcian, and subsequently it was found in Hettangian deposits (Aberhan, 1994a; Liu, 1995; Damborenea, 1996a). Lucas and Estep (1997, p. 45, fig. 1c and 1d) mentioned and figured Weyla from Carnian beds of Sonora (Mexico), but these specimens were reassigned to *Mysidioptera* by Damborenea in Damborenea and Gonzalez-León (1997); Lucas and Estep (1997) also reported other specimens from the Sinemurian of the same area.

Paleogeographic distribution.—Circumpacific and Austral (Fig. 36). Although the genus is also present in the Tethys domain, it is recorded there only after the beginning of the Pliensbachian (Calzada, 1982; Liu, 1995; Fraser, Bottjer, & Fischer, 2004; Valls, Comas-Rengifo, & Goy, 2004). The genus originated in the Pacific margin and then extended to the western Tethys through the Hispanic Corridor or Proto-Atlantic (Damborenea & Manceñido, 1979, 1988; Aberhan, 2001). See Damborenea and Manceñido (1979) for a complete distribution of the genus.

Circumpacific domain: Early Jurassic: Hettangian–Sinemurian of western Canada (Aberhan, 1998a), Mexico and Texas (Liu, 1995), Chile (Aberhan, 1994a); Sinemurian of Sonora (Mexico) (Damborenea in Damborenea & González-León, 1997; Lucas & Estep, 1997; Scholz, Aberhan, & González-León, 2008), Chile (Escobar, 1980), Peru (Rangel, 1978).

Austral domain: Early Jurassic: Sinemurian of Argentina (Damborenea & Manceñido, 2005b; Damborenea & Lanés, 2007).

Paleoautoecology.—B, Se, S, Un, Sed; R. From the observation of specimens in life position and analysis of the shell morphology (Damborenea & Manceñido, 1979, 1988; Damborenea, 1987b), it was inferred that *Weyla* was sedentary and lived semi-infaunally as a recliner, without byssus attachment in the adult stage. *Mineralogy.*—Bimineralic (Carter, 1990a, p. 260, 263). Little is known about the microstructure of the shell of *Weyla*; the inner shell layer is aragonitic and with cross-lamellar structure. Data provided for the family Pectinidae. Outer shell layer: calcite (prismatic). Inner shell layer: aragonite (cross-lamellar).

Genus INDOPECTEN Douglas, 1929, p. 632

Type species.—Pecten clignetti Krumbeck, 1913, p. 36.

Stratigraphic range.—Upper Triassic (Norian–Rhaetian) (Hautmann, 2001b). Cox and others (1969) assigned it an upper Norian range. New records expanded the range of this genus (see paleogeographic distribution).

Paleogeographic distribution.—Tethys (Fig. 36).

Tethys domain: Late Triassic: Norian of western Carpathians (Ruban, 2006a), China (Wen & others, 1976; J. Chen & Yang, 1983), Oman (R. Hudson & Jefferies, 1961), Armenia (Hautmann, 2001b), Timor (Indonesia) (Krumbeck, 1924), Himalayas (Kutassy, 1931), Thailand (Vu Khuc & Huyen, 1998); Norian–Rhaetian of Iran (Hautmann, 2001b); Rhaetian of Tibet (China) (Hautmann & others, 2005), Pamira (Afghanistan) (Polubotko, Payevskaya, & Repin, 2001).

Paleoautoecology.—B, E, S, Epi-Un, FaM; By-R-Sw. Some of *Indopecten* species, like *I. glaber* Douglas, 1929, could swim, as suggested by their ligament type (so-called alivincular-alate according to Hautmann, 2004), which would be fit enough for the constant opening and closing of the valves required by a swimming activity (Hautmann, 2004). The external morphology is also compatible with a swimming habit, according to S. M. Stanley (1972). It probably lived mostly reclined, but it could swim occasionally. However, other species have a byssal notch, and their shell morphology is not suitable for swimming, so these were probably epibyssate (see Hautmann, 2001b).

Mineralogy.—Aragonitic (Hautmann, 2006a). According to Waller (2006), the *Indopecten* inner shell layers are usually recrystallized, so they were probably aragonitic. Although the microstructure of the outer shell layer is unknown, its mineralogy was calcitic. However, Hautmann (2006a) studied the microstructure of two species of *Indopecten* [*I. serraticostata* (Bittner, 1899) and *I. glaber* Douglas, 1929)] and he concluded that all the shell was composed of a single microstructure (probably cross-lamellar) and was entirely aragonitic.

Genus CAMPTONECTES Agassiz in Meek, 1864, p. 39

Type species.—Pecten lens J. Sowerby, 1818, p. 3.

Remarks.—Cox and others (1969) included three subgenera within *Camptonectes: C. (Camptonectes)*, *C. (Camptochlamys)* Arkell, 1930 in 1929–1937, p. 102, and *C. (Boreionectes)* Zakharov, 1965, p. 72. Subsequently, Allasinaz (1972, p. 316) added a fourth, *C. (Annulinectes)*, and Fürsich (1982, p. 50) another, *C. (Costicamptonectes)*. Kelly (1984) regarded *Boreionectes* as a junior synonym of *Mclearnia* Crickmay, 1930b, p. 45. According to Waller and Marincovich (1992), *Costicamptonectes* is unnecessary, and they raised *Camptochlamys* to generic level. We only consider two subgenera in our study interval: *C. (Camptonectes)*.

Stratigraphic range.—Middle Triassic (Anisian)–Upper Cretaceous (Maastrichtian) (Cox & others, 1969; Allasinaz, 1972). Cox and others

(1969) assigned a Jurassic–Upper Cretaceous (Maastrichtian) range to *C. (Camptonectes*), but some Tertiary and Holocene species are now also referred to this genus by some authors (see Damborenea, 2002a, p. 56). Allasinaz (1972) assigned to *C. (Annulinectes*) an Anisian–Jurassic range, although the species he included in this subgenus were recorded only in Anisian, Ladinian, and Carnian beds. As we could not see the papers that recognize post-Cretaceous *Camptonectes*, we provisionally follow Cox and others (1969) for the end of the range.

Paleogeographic distribution.—Cosmopolitan (Fig. 36).

Tethys domain: Middle Triassic: Anisian of Slovakia (Kochanová, 1985), Romania and Yugoslavia (Allasinaz, 1972); Ladinian of Slovakia (Kochanová, Mello, & Siblík, 1975), Bosnia (Allasinaz, 1972); Late Triassic: Carnian of Slovenia (Jelen, 1988), Hungary and Italy (Allasinaz, 1972); Rhaetian of ?England (Johnson, 1984; Ivimey-Cook & others, 1999); Early Jurassic: Hettangian of England (Johnson, 1984; Liu, 1995), Germany and Switzerland (Johnson, 1984); Sinemurian of England and Portugal (Liu, 1995).

Circumpacific domain: Late Triassic: Carnian of Japan (Nakazawa, 1952; Hayami, 1975); Early Jurassic: Hettangian of Japan (Hayami, 1959; Hayami, 1975; Johnson, 1984); Sinemurian of western Canada (Poulton, 1991; Aberhan, 1998a).

Austral domain: Early Jurassic: Hettangian–Sinemurian of Argentina (Damborenea, 2002a; Damborenea & Manceñido, 2005b).

Boreal domain: Late Triassic: Norian-Rhaetian of northeastern Russia (Milova, 1976).

Paleoautoecology.—B, E, S, Epi, FaM; By-Sw. All species described by Johnson (1984) have a byssal notch and were epibyssate. However, he interpreted that they could swim to escape from predators, on the basis of the analysis of external shell morphology (shell thin, low convexity, and wide umbonal angle). According to Sha (2003), *Camptonectes* had planktotrophic larvae that facilitated its global distribution.

Mineralogy.—Bimineralic (Johnson, 1984; Carter, 1990b, p. 381). Outer shell layer: calcite (foliated-prismatic). Inner shell layer: aragonite (?).

Genus CRENAMUSSIUM Newton in Newton & others, 1987, p. 46

Type species.—Crenamussium concentricum Newton in Newton & others, 1987, p. 50.

Remarks.—Newton (in Newton & others, 1987) included *Crenamussium* in the family Pectinidae. Later, Waller (in Waller & Stanley, 2005) included it in the Entoliidae, and a year later, Waller (2006) suggested that *Crenamussium* is a junior synonym of *Calvaentolium* Romanov, 1985, but he did not justify it. *Calvaentolium* is not included in this paper (see discussion in Genera not Included, p. 158). Newton (in Newton & others, 1987) included the type species in *Crenamussium* and, tentatively, *C. balatonicus* (Bittner, 1901b) (species included in *Pleuronectites* and *Chlamys* by other authors).

Stratigraphic range.—Upper Triassic (Carnian–Norian) (Newton in Newton & others, 1987). According to the species assigned by Newton (in Newton & others, 1987), the stratigraphic range of *Crenamussium* is Carnian–Norian.

Paleogeographic distribution.—Tethys and Circumpacific (Fig. 36). Tethys domain: Late Triassic: Carnian of Italy (Allasinaz, 1972), Hungary (Bittner, 1912), Carpathians (Kiparisova, 1954). Circumpacific domain: Late Triassic: Norian of Oregon (United States) (Newton in Newton & others, 1987).

Paleoautoecology.—B, E, S, Epi, Sed; By. According to the diagnosis given by Newton (in Newton & others, 1987), the shell is subcircular, the auricles are unequal, and a deep byssal notch is observed in the right anterior auricle, thus it probably lived as an epibyssate.

Mineralogy.—Bimineralic (Newton in Newton & others, 1987, p. 50). *Crenamussium* shell microstructure is not well known, but the mineralogy of the shell, as a member of the family Pectinidae, should have at least a calcitic outer layer. In the specimens described by Newton (in Newton & others, 1987), there is evidence of original fibrous calcite in the form of siliceous pseudomorphs.

Genus AVICHLAMYS Allasinaz, 1972, p. 368

Type species.—Pecten csopakensis Frech, 1905, p. 4.

Stratigraphic range.—Lower Triassic (Olenekian) (Posenato, 2008a). Allasinaz (1972) proposed Avichlamys and included two species: Pecten csopakensis Frech, 1905, and Pecten nicolensis Ogilvie Gordon, 1927, both from the Triassic. Subsequently, Neri and Posenato (1985) and Broglio-Loriga and others (1990) also included Chlamys tellinii Tommasi, 1896, which was reported from the Italian Triassic by Leonardi (1935) and Boni (1943).

Paleogeographic distribution.—Eastern Tethys (Fig. 36).

Tethys domain: Early Triassic: Italy (Allasinaz, 1972; Neri & Posenato, 1985; Broglio-Loriga & others, 1990), Hungary (Allasinaz, 1972).

Paleoautoecology.—B, E, S, Epi, Sed; By. According to the generic diagnosis provided by Allasinaz (1972), *Avichlamys* is characterized by a subequivalve shell, with different valve convexity, subequal auricles, and a wide byssal gape in the right auricle. It was probably an epibyssate bivalve.

Mineralogy.—Bimineralic (Carter, 1990a). No data are known about *Avichlamys* shell microstructure. We assume a bimineralic shell, as in the other pectinids, following Carter (1990a).

Genus PSEUDOPECTEN Bayle, 1878, explanation pl. 121

Type species.—Pecten equivalvis J. Sowerby, 1816, p. 83.

Remarks.—Although *Pseudopecten* lacks a ctenolium and should be excluded from the Pectinidae (Damborenea, 2002a), we list it here until it is assigned to another family according to its features.

Stratigraphic range.—Lower Jurassic (Hettangian)–Middle Jurassic (Bajocian) (Johnson, 1984). Cox and others (1969) distinguished two subgenera within *Pseudopecten: P. (Pseudopecten)* and *P. (Echinopecten)* Brasil, 1895. Both were reported from Hettangian beds. However, following Johnson (1984), we regard *P. (Pseudopecten)* as Hettangian and *P. (Echinopecten)* as Toarcian. Therefore, only *P. (Pseudopecten)* is included in our study interval.

Paleogeographic distribution.—western Tethys (Fig. 36). During our study interval, *Pseudopecten* was only distributed in the western Tethys, but from the Pliensbachian onward, it was also reported from South America (Damborenea, 2002a) and Australia (Grant-Mackie, 1994). In addition, Damborenea (2002a) suggested that its presence in the Early Jurassic of Japan and Siberia is also possible.

Tethys domain: Early Jurassic: Hettangian of Spain (Liu, 1995), France, Italy, and Germany (Johnson, 1984); Sinemurian of Spain, Portugal, and Morocco (Liu, 1995), England (Johnson, 1984; Liu, 1995), Italy, France, and Germany (Johnson, 1984).

Paleoautoecology.—B, E, S, Un, FaM; R-Sw. Species attributed to *P. (Pseudopecten)* had a byssal gape at juvenile stages, but it disappeared in adult stages; they lived byssate when young but were later recliners on the substrate, and they could swim actively (see Johnson, 1984, for a complete interpretation of the various species).

Mineralogy.—Bimineralic (Carter, 1990b, p. 388). Outer shell layer: calcite (prismatic + foliated). Inner shell layer: aragonite (cross-lamellar).

Genus AGERCHLAMYS Damborenea, 1993, p. 119

Type species.—Chlamys (Camptochlamys) wunschae Marwick, 1953, p. 98.

Stratigraphic range.—Upper Triassic (Carnian)–Lower Jurassic (Toarcian) (Damborenea, 1993, 2002a). Damborenea (1993) proposed *Agerchlamys*, including several previously described species referred to other genera (see Damborenea, 1993, p. 120, and Damborenea, 2002a, p. 66, for species listed). These species were recorded from the Carnian to the Toarcian, and the author indicated the possibility that the genus may be present up to the Middle Jurassic.

Paleogeographic distribution.—Circumpacific, Austral, and Boreal (Fig. 36). Agerchlamys was distributed through the Austral and Boreal domains and also in the Circumpacific, but always at high latitudes. In the Austral domain (Argentina and New Zealand), it was reported primarily from Pliensbachian beds (Marwick, 1953; Damborenea, 1993, 2002a).

Circumpacific domain: Early Jurassic: Hettangian of Chile (Aberhan, 1994a), Oregon (United States) (D. G. Taylor & Guex, 2002), British Columbia (western Canada) (Wignall & others, 2007); Hettangian–Sinemurian of Canada (Aberhan, 1998a, 1998b), Sonora (Mexico) (Scholz, Aberhan, & González-León, 2008).

Austral domain: Early Jurassic: Hettangian–Sinemurian of Argentina (Damborenea, 2002b; Damborenea & Manceñido, 2005b).

Boreal domain: Late Triassic: Carnian-Norian of Siberia (Kiparisova, Bychkov, & Polubotko, 1966).

Paleoautoecology.—B, E, S, Epi, Sed; By. Due to the presence of a deep byssal gape below the right anterior auricle and a strong ctenolium, it was epibyssate (Damborenea, 1993, 2002a). Although it has a wide umbonal angle, the auricles are of different sizes, so it is not believed that it could swim.

Agerchlamys is usually found in low-energy and well-oxygenated environments, and associated with sponges and other bivalves, especially limids (Damborenea, 2002a).

Mineralogy.—Bimineralic (Carter, 1990a, p. 260). There are no data about *Agerchlamys* shell. We assume bimineralic mineralogy, as in the other members of the family Pectinidae (Carter, 1990a).

Genus CANADONECTITES Aberhan, 1998a, p. 110

Type species.—Canadonectites paucicostatus Aberhan, 1998a, p. 110. Remarks.—Aberhan (1998a) proposed the genus Canadonectites to accommodate specimens with intermediate morphology between Pleuronectites Schlotheim, 1820, and Radulonectites Hayami, 1957c, and differing from both of them by ornamentation features. Stratigraphic range.—Lower Jurassic (Sinemurian–Pliensbachian) (Aberhan, 1998a). It was only reported from Sinemurian and Pliensbachian beds of western Canada (Aberhan, 1998a).

Paleogeographic distribution.—Circumpacific (Fig. 36).

Circumpacific domain: Early Jurassic: Sinemurian of western Canada (Aberhan, 1998a, 2001).

Paleoautoecology.—B, E, S, Epi, Se; By. The mode of life of *Canadonectites* was probably very similar to *Agerchlamys*, since both have a deep byssal notch and ctenolium in the right valve.

Mineralogy.—Bimineralic (Carter, 1990a, p. 260). There are no data on *Canadonectites* shell microstructure. We assume bimineralic mineralogy, as in the other members of the family Pectinidae (Carter, 1990a).

Genus EOPECTEN Douvillé, 1897, p. 203

Type species.—Hinnites tuberculatus Goldfuss (*errore pro Spondylus tuberculosus* Goldfuss), 1835 in 1833–1841, p. 93.

Remarks.—See Johnson (1984, p. 149) and Damborenea (1987b, p. 198) for a discussion about synonymy and the problems related to this genus since its proposal.

Stratigraphic range.—Upper Triassic (Carnian)–Lower Cretaceous (Albian) (Hayami, 1975; Johnson, 1984). Cox and others (1969) assigned it a Jurassic-Lower Cretaceous (Albian) range. Although it seems fairly accepted that it appeared in the Early Jurassic, according to the literature, there are several records from the Carnian of Japan (Kobayashi & Ichikawa, 1949b; Nakazawa, 1952; Hayami, 1975) and from the Norian of Chile (Hayami, Maeda, & Ruiz-Fuller, 1977) that no other author except Hallam (1981) considered. Middle Triassic specimens assigned by Allasinaz (1972) to Radulonectites should be allocated to *Eopecten* instead (Damborenea, 2002a, p. 61). If Radulonectites? flagellum (Stoppani, 1858 in 1858-1860), described by Allasinaz (1972, p. 331), is assigned to Eopecten, the genus was present from Ladinian times. Another species that was referred to Eopecten, originally proposed as Monotis albertii Goldfuss, 1835 in 1833-1841 (Diener, 1923), was reported from the Lower and Middle Triassic of Europe, but it is currently included in Leptochondria (Waller & Stanley, 2005, p. 34).

Paleogeographic distribution.—Tethys, Circumpacific, and Austral (Fig. 36). According to the published records, it seems that *Eopecten* originated in the Late Triassic of Japan, and then it migrated to the western Tethys (Europe) and eastern Paleopacific.

Tethys domain: Early Jurassic: Hettangian of England (Liu, 1995), Belgium and Germany (Johnson, 1984); Sinemurian of Portugal and Spain (Liu, 1995), Germany (Johnson, 1984).

Circumpacific domain: Late Triassic: Carnian of Japan (Kobayashi & Ichikawa, 1949b; Nakazawa, 1952; Hayami, 1975); Early Jurassic: Hettangian of Chile (Aberhan, 1994a); Hettangian–Sinemurian of Canada (Aberhan, 1998a); Sinemurian of Canada (Poulton, 1991).

Austral domain: Early Jurassic: Hettangian–Sinemurian of Argentina (Riccardi & others, 1991; Damborenea, 2002a; Damborenea & Manceñido, 2005b).

Paleoautoecology.—B, E, S, Epi-C, Sed; By-C. Johnson (1984) and Harper and Palmer (1993) analyzed the mode of life of different species of *Eopecten*. The last authors concluded that some species could live cemented to the substrate, while others were epibyssate during most of their life.

Mineralogy.—Bimineralic (Carter, 1990b, p. 388; Harper & Palmer, 1993, p. 67). The shell of *Eopecten* had a foliated outer shell layer, both in the left valve (Carter, 1990a) and in the right one (Harper & Palmer, 1993), and aragonitic middle and inner shell layers with cross-lamellar microstructure (Carter, 1990a).

Genus JANOPECTEN Arkhipov & Trushchelev, 1980, p. 10

Type species.—Janopecten kularensis Arkhipov & Trushchelev, 1980, p. 10.

Stratigraphic range.—Middle Triassic (Anisian)–Upper Triassic (lower Carnian) (Waller in Waller & Stanley, 2005). *Janopecten* was reported from Anisian and Ladinian beds (Dagys & Kurushin, 1985) and from the lower Carnian of the Boreal area (Waller in Waller & Stanley, 2005).

Paleogeographic distribution.—Boreal (Fig. 36).

Boreal domain: Middle Triassic: Anisian of Yakutia (Russia) (Konstantinov, Sobolev, & Yadernkin, 2007); Anisian–Ladinian of Siberia (Arkhipov & Trushchelev, 1980; Dagys & Kurushin, 1985). Late Triassic: early Carnian of Siberia (Arkhipov & Trushchelev, 1980).

Paleoautoecology.—B, E, S, Epi, FaM; By-Sw? At least the type species (Dagys & Kurushin, 1985, pl. 21, *13a*, *13b*) had an equivalve shell, nearly equal auricles, and an umbonal angle large enough to be an occasional swimmer. The shells had a byssal notch throughout their ontogeny (Waller in Waller & Stanley, 2005), and they likely lived epibyssate and occasionally could perform swimming cycles. This is probably true only for the Anisian forms, since, according to Waller (in Waller & Stanley, 2005), Ladinian and Carnian species of *Janopecten* began to develop strongly inequilateral shells.

Mineralogy.—Bimineralic (Carter, 1990a, p. 260). There are no data on *Janopecten* shell microstructure. Bimineralic mineralogy is assumed, as in the other members of the family Pectinidae.

Genus OCHOTOCHLAMYS Milova & Polubotko in Milova, 1976, p. 61

Type species.—Chlamys (Ochotochlamys) gizhigensis Polubotko in Milova, 1976, p. 61.

Remarks.—*Ochotochlamys* was erected as subgenus of *Chlamys* (Milova & Polubotko in Milova, 1976), but it was subsequently raised to genus level (Polubotko & Milova, 1986), which was followed by all later authors.

Stratigraphic range.—Upper Triassic (Norian)–Lower Jurassic (Toarcian) (Milova & Polubotko in Milova, 1976; Aberhan, 1998a). For a long time, *Ochotochlamys* was only reported from Late Triassic of northeastern Asia, but it was later recorded from the Pliensbachian (Polubotko & Milova, 1986; Aberhan 1998a; Damborenea, 2002a) and from the Toarcian (Aberhan, 1998a).

Paleogeographic distribution.—Boreal (Fig. 36). It was originally believed that the genus was restricted to northeastern Asia, but new records from the Pacific margin (Canada and Argentina) (Aberhan, 1998a; Damborenea, 2002b) extended its paleogeographic distribution; but it is only known from high paleolatitudes. The Austral record is Pliensbachian and thus outside our study range (Damborenea, 2002a). It was also mentioned from the Triassic–Jurassic boundary beds of British Columbia (Wignall & others, 2007) and eastern Alberta (Asgar-Deen & others, 2003).

Boreal domain: Late Triassic: Carnian–Norian of northeastern Asia (Polubotko & Milova in Milova, 1976); Early Jurassic: Hettangian of northeastern Russia (Milova, 1988); Hettangian–Sinemurian of western Canada (Aberhan, 1998a, 1998b, 2001); Sinemurian of northeastern Russia (Polubotko & Milova, 1986).

Paleoautoecology.—B, E, S, Epi, Sed; By. According to Aberhan (1998a), a typical feature of the genus is that the right anterior auricle had a byssal notch; a byssal sinus is observed in the left valve, but it is not distinguishable in all specimens. *Ochotochlamys* was probably an epibyssate bivalve.

Mineralogy.—Bimineralic (Carter, 1990a, p. 260). There are no data on *Ochotochlamys* shell microstructure. Bimineralic mineralogy is assumed, as in the other members of the family Pectinidae (Carter, 1990a).

Genus TOSAPECTEN Kobayashi & Ichikawa, 1949b, p. 166

Type species.—Pecten (Velopecten) suzukii Kobayashi, 1931, p. 258. Remarks.—Although Kobayashi and Ichikawa (1949b) proposed Tosapecten within the family Pectinidae, Cox and others (1969) considered it to be a subgenus of Weyla Böhm, 1922. Currently, almost all authors (Hayami, 1975; Milova, 1976; J. Chen, 1982a; Okuneva, 1985; Damborenea, 1987b; Tanaka, 1989; Waller in Waller & Stanley, 2005; Waller, 2006) regard it as a distinct genus, separate from Weyla.

According to Waller (in Waller & Stanley, 2005), *Tosapecten* includes two subgenera, *T. (Tosapecten)* and *T. (Indigiropecten)* Trushchelev, 1984.

Stratigraphic range.—Upper Triassic (Carnian–Rhaetian) (Kobayashi & Ichikawa, 1949b; Milova, 1976). Cox and others (1969) stated that the genus was known from the Upper Triassic of Japan. Subsequently, it was also reported from Siberia (Kobayashi & Tamura, 1983b). It was known throughout the Carnian and Norian in Japan (see paleogeographic distribution, below). We lack information on the Siberian records; we could only check the Norian occurrence in Okuneva (1985). Milova (1976) reported *Tosapecten subhiemalis wittnburgi* n. subsp. from upper Norian–Rhaetian beds. Some biostratigraphic papers mentioned the presence of *Tosapecten* from latest Triassic times (Kurushin, 1990; Polubotko & Repin, 1990; Zakharov & others, 1997), referring to *Tosapecten efimovae* Polubotko, 1966.

Paleogeographic distribution.—Circumpacific and Boreal (Fig. 36). *Tosapecten* was mainly distributed through the northern Circumpacific and Boreal domains.

Circumpacific domain: Late Triassic: Carnian of Japan (Nakazawa, 1952; Ando, 1988); Carnian–Norian of Japan (Kobayashi & Ichikawa, 1949b; Tokuyama, 1959b; Hayami, 1975; Tanaka, 1989; Onoue & Tanaka, 2005); Norian of Japan (Nakazawa, 1963), ?Oregon (United States) (Newton in Newton & others, 1987; Newton, 1988).

Boreal domain: Late Triassic: Carnian of northeastern Russia (Bychkov & others, 1976), Primorie (Kiparisova, 1972); Norian of Siberia (Okuneva, 1985); Norian–Rhaetian of northeastern Russia (Milova, 1976); Rhaetian of northeastern of Siberia (Bychkov & others, 1976; McRoberts, 2010).

Paleoautoecology.—B, E, S, Epi, Sed; By. In all species assigned to *Tosapecten*, a byssal notch is observed, and they have unequal auricles (see description and figures in the published literature, listed above). Like most pectinids, they lived epibyssate.

Mineralogy.—Bimineralic (Carter, 1990a, p. 260). There are no data on *Tosapecten* shell microstructure. Bimineralic mineralogy is assumed, as in the other members of the family Pectinidae.

Genus NEVADAPECTEN Waller in Waller & Stanley, 2005, p. 46

Type species.—Nevadapecten lynnae Waller in Waller & Stanley, 2005, p. 46.

Remarks.—Waller (in Waller & Stanley, 2005) included *Nevadapecten* in the subfamily Tosapectininae and related it with *Tosapecten* and *Janopecten*, considering it to be intermediate between these two genera in several aspects.

Stratigraphic range.—Middle Triassic (upper Ladinian) (Waller in Waller & Stanley, 2005). According to Waller (in Waller & Stanley, 2005), *Nevadapecten* was reported from the upper Ladinian of New Pass Range in Nevada.

Paleogeographic distribution.—Circumpacific (Fig. 36).

Circumpacific domain: Middle Triassic: Ladinian of Nevada (United States) (Waller in Waller & Stanley, 2005).

Paleoautoecology.—B, E, S, Epi, Sed; By. *Nevadapecten* had some features related to a swimming mode of life, such as circular shape, an almost equilateral shell, and a wide umbonal angle, but it had unequal auricles and a byssal gape, which indicates it was an epibyssate bivalve.

Mineralogy.—Bimineralic (Waller in Waller & Stanley, 2005). Outer shell layer: calcite (antimarginal fibrous). Inner shell layer: aragonite.

Genus LOXOCHLAMYS Waller in Waller & Stanley, 2005, p. 40

Type species.—Loxochlamys corallina Waller in Waller & Stanley, 2005, p. 43.

Stratigraphic range.—Middle Triassic (Ladinian)–Upper Triassic (Carnian) (Waller in Waller & Stanley, 2005). Waller (in Waller & Stanley, 2005) included the type species within *Loxochlamys*, from the upper Ladinian, and two other species: *Pecten chiwanae* McLearn, 1941, and *Pecten sasuchan* McLearn, 1941, both from Carnian beds.

Paleogeographic distribution.—Circumpacific (Fig. 36).

Circumpacific domain: Middle Triassic: Ladinian of Nevada (United States) (Waller in Waller & Stanley, 2005); Late Triassic: Carnian of British Columbia (Canada) (McLearn, 1941; Waller in Waller & Stanley, 2005).

Paleoautoecology.—B, E, S, Epi, Sed; By. *Loxochlamys* had a byssal notch and ctenolium throughout its ontogeny; it probably lived epibyssate among corals with which it was usually associated (Waller in Waller & Stanley, 2005).

Mineralogy.—Bimineralic (Waller in Waller & Stanley, 2005). Outer shell layer: calcite (antimarginal fibrous). Inner shell layer: aragonite.

Genus PLEURONECTITES von Schlotheim, 1820, p. 217

Type species.—Pleuronectites laevigatus von Schlotheim, 1820, p. 217.

Stratigraphic range.—Middle Triassic (Anisian)–Upper Triassic (Carnian) (Hayami, 1975; Waller, 2006). Newell and Boyd (1995) extended the range of *Pleuronectites* to the Lower Triassic, with the

species *Pleuronectites meeki* Newell & Boyd, 1995, but Waller (in Waller & Stanley, 2005) regarded this species as an entoliid rather than a pectinid, because it lacks a ctenolium. Waller (2006) indicated that the oldest undoubted *Pleuronectites* is *P. laevigata* Schlotheim, 1820, from the Anisian (see Waller in Waller & Stanley, 2005 and Waller, 2006, for records mentioned from the Lower Triassic that are not taken into account). The youngest record of the genus is from the Carnian with *P. hirabarensis* Amano (Hayami, 1975). Newton (in Newton & others, 1987) mentioned it from the Norian, but this reference is questionable.

Hautmann (2010) considered most species attributed to *Pleuronectites* to be synonyms of *P. laevigatus* (previously referred by Waller in Waller & Stanley, 2005, to *Pecten laterestriatus* Philippi, 1899, and *Pecten schmiederi* Giebel, 1856) or as erroneusly assigned to the genus, so *Pleuronectites* would be a monospecific genus. The author, and previously Waller (in Waller & Stanley, 2005), mentioned *P. balatonicus* (Bittner, 1901c), figured by Allasinaz (1972), and regarded it as entoliid. *Pleuronectites newelli* Waller in Waller & Stanley, 2005, was also regarded as an entoliid by Hautmann (2010). He assigned an Anisian–Ladinian range to *Pleuronectites*, but he did not mention Amano's Carnian species.

Paleogeographic distribution.—Tethys and Circumpacific (Fig. 36). The paleogeographic distribution agrees with Waller (in Waller & Stanley, 2005) and Waller (2006). Although *Pleuronectites* is often quoted as being from China, the specimens are uncertainly or erroneously assigned (Waller in Waller & Stanley, 2005).

Tethys domain: Middle Triassic: Anisian of Germany (Hagdorn, 1982, 1991, 1995); Anisian–Ladinian of Israel (Lerman, 1960), Germany, Hungary, Sardinia (Italy), Afghanistan, and China (Hautmann, 2010); Ladinian of Italy (Allasinaz, 1972), Spain (Márquez-Aliaga, 1983, 1985), Afghanistan (Farsan, 1972).

Circumpacific domain: Middle Triassic: Anisian of Japan (Hayami, 1975); Ladinian of Nevada (United States) (Waller in Waller & Stanley, 2005); Late Triassic: Carnian of Japan (Hayami, 1975).

Paleoautoecology.—B, E, S, Epi, Sed; By. According to the diagnosis offered by Waller (in Waller & Stanley, 2005), *Pleuronectites* probably lived epibyssate, and it had a deep byssal notch below the right anterior auricle, and a ctenolium throughout ontogeny.

Mineralogy.—Bimineralic (Waller in Waller & Stanley, 2005, p. 40). *Pleuronectites newelli* Waller in Waller & Stanley, 2005, shows an originally aragonitic inner shell layer and a calcitic outer shell layer (Waller in Waller & Stanley, 2005), and the same mineralogy was found in *Pleuronectites laevigatus* Schlotheim, 1820 (Hautmann, 2010; Carter & Hautmann, 2011); the microstructure of inner and middle shell layers was probably cross-lamellar, and the outer shell layer was prismatic.

Genus PERICLARAIA J. Li & Ding, 1981, p. 327, 330

Type species.—Periclaraia circularis J. Li & Ding, 1981, p. 327. Remarks.—Periclaraia was proposed by J. Li and Ding (1981) from deposits of Anhui Province (China). These authors included three species: Periclaraia circularis J. Li & Ding, 1981, Periclaraia reticulata J. Li & Ding, 1981, and Periclaraia chaoxianensis J. Li & Ding, 1981. Subsequently, J. Chen and Komatsu (2002) added a new species, Periclaraia jinyaensis Chen & Komatsu, 2002, and they considered that the three species proposed by J. Li and Ding (1981) are variants of the same, as J. Li and Ding (1981) used differences in ornamentation and size of the right anterior auricle to separate them. These variations are regarded as intraspecific by J. Chen and Komatsu (2002). J. Li and Ding (1981) included *Periclaraia* in the family Pectinidae, but other authors (H. Yin, 1985, 1990; Gavrilova, 1995, 1996), based on its external resemblance to the clariids, referred it to the Pteropectinidae. However, J. Chen and Komatsu (2002) argued that since *Periclaraia* had a right valve ctenolium, a diagnostic character of the family Pectinidae, it should be included in this family.

Stratigraphic range.—Lower Triassic (upper Olenekian)–Middle Triassic (lower Anisian) (J. Li & Ding, 1981; J. Chen & Komatsu, 2002). J. Li and Ding (1981) reported *Periclaraia* from upper Olenekian beds of Anhui Province (China). Subsequently, J. Chen and Komatsu (2002) mentioned it from lower Anisian deposits of Guangxi province (China) and doubted the age assignation given by J. Li and Ding (1981). However, *Periclaraia* was quoted in several biostratigraphic papers in Anhui province in beds attributed to the Olenekian (Tong & others, 2004, 2006; Tong, 2005; S. Wu & others, 2005). Curiously, these papers considered *Periclaraia* as endemic to Anhui province, not mentioning J. Chen and Komatsu (2002).

Paleogeographic distribution.—Eastern Tethys (Fig. 36).

Tethys domain: Early Triassic: late Olenekian of Anhui province (China) (J. Li & Ding, 1981); Middle Triassic: early Anisian of Guangxi province (China) (J. Chen & Komatsu, 2002; J. Chen & Stiller, 2007).

Paleoautoecology.—B, E, S, Epi, Sed; By. The features of *Periclaraia* shells, which are inequivalve and inequilateral and had a deep byssal notch, indicate it was probably an epibyssate bivalve.

Mineralogy.—Bimineralic (Carter, Barrera, & Tevesz, 1998). Mineralogical data provided for the family Pectinidae by Carter, Barrera, and Tevesz (1998). Outer shell layer: calcite. Middle and inner shell layers: calcite-aragonite.

Genus RADULONECTITES Hayami, 1957c, p. 89

Type species.—Radulonectites japonicus Hayami, 1957c, p. 90. Remarks.—Hunanonectes Z. Fang, 1978, p. 465, is considered to be a synonym of Radulonectites (see discussion for Hunanonectes in Genera not Included, p. 162).

Stratigraphic range.—Lower Jurassic (Hettangian–Pliensbachian) (Stiller, 2006; Hayami, 1975). Hayami (1957c) proposed the genus from the Japanese Pliensbachian (see also Hayami, 1985) and included the type species and Pecten (Pleuronectites) laterestriatus Philippi, 1899, from the German lower Muschelkalk, and also specimens referred by A. F. Leanza (1942) to Pecten (Camptonectes) lens Sowerby from the Pliensbachian of Argentina. He also provisionally included Chlamys kakanuia Marwick (1956, fig. 1) from the Upper Triassic of New Zealand, a species based on too poorly preserved material to discuss its affinities (Damborenea, 1993, 2002a). Later, Hayami (1975) disregarded these Triassic records and assigned the genus a Pliensbachian range. Cox and others (1969) referred it to the ?Triassic, Lower Jurassic. On the other hand, Sepkoski (2002) assigned it a Triassic (Anisian)-Jurassic (?Pliensbachian) range, mentioning Hayami (1975) and H. Yin (1985) as his sources. H. Yin (1985) mentioned it during Anisian and Ladinian but did not list the original source. Allasinaz (1962, 1972) quoted *Radulonectites* from the European Triassic. The specimens from the Italian Rhaetian referred by Allasinaz (1962) to *Radulonectites* are very poorly preserved; these doubtful records are not taken into account here, since, according to Damborenea (2002a), they belong to *Eopecten* Douvillé, 1897. The same occurs with material from the Triassic of New Zealand, similarly referred by other authors (see Damborenea, 2002a, p. 61). Onoue and Tanaka (2005) mentioned *Radulonectites* sp. from the Japanese Upper Triassic, based on a single deformed specimen, and their incomplete description is not enough to extend the range of this taxon. The oldest solid records are Hettangian (J. Chen, 1982b; Stiller, 2006 [*Hunanonectes*]) and the youngest are Pliensbachian (Hayami, 1957c, 1975, 1985).

Paleogeographic distribution.—Eastern Tethys (Fig. 36). During the Pliensbachian, this genus had a wide distribution (e.g., Siberia, Argentina, Chile) (Hayami, 1975; Polubotko & Milova, 1986; Milova, 1988; Damborenea, 1993, 2002a; Aberhan, 1994a, 1998a; Aberhan & Fürsich, 1997), and during the study interval, it was only reported with certainty from China.

Tethys domain: Early Jurassic: Hettangian–Sinemurian of southern China (J. Chen, 1982b, 1988; Z. Fang, 1978; Stiller, 2006); ?Sinemurian of Canada (Poulton, 1991, according to Aberhan, 1998a).

Paleoautoecology.—B, E, S, Epi, Sed; By. Several morphological traits (shell shape, convexity of valves, and the presence of deep byssal notch) and the encrusted epizoic organisms in several specimens suggest an epibyssate mode of life (Damborenea, 2002a).

Mineralogy.—Bimineralic (Carter, Barrera, & Tevesz, 1998). Mineralogical data provided for the family Pectinidae by Carter, Barrera, and Tevesz (1998). Outer shell layer: calcite. Middle and inner shell layers: calcite-aragonite.

Family ENTOLIIDAE Teppner, 1922

Syncyclonemidae Waller, 1978

In recent years, various genera for different groups of species that were traditionally referred to Entolium were proposed. This is very similar to what happened to the family Halobiidae, but there is no consensus about which characters should be used to discriminate between different taxonomic levels (see Damborenea, 2002a, p. 42-44, for a full discussion on the subject). Features that are distinctive at species level, according to some authors (see Johnson, 1984), are used by others to discriminate at genus level (Damborenea, 2002a): the presence or absence of lateral internal ribs; the presence or absence of byssal notch; and dorsal projection of the auricles. Furthermore, some surface shell structures are used at genus level, even if they are strongly influenced by diagenetic processes (Johnson, 1984; Damborenea, 2002a). In the absence of a good review on the subject, we are not taking into account the genera listed below (see discussion for each of them in Genera not Included, p. 156), since their proposition was based, in most cases, on diagnostic characters that are used by most authors at species level, and they can all be grouped under Staesche's (1926) original concept of Entolium: Costentolium Freneix, 1980, p. 89; Cingentolium Yamani, 1983, p. 6; Neoentolium Romanov, 1985, p. 37; Cornutoentolium Romanov, 1985, p. 52 (Upper Jurassic); Calvaentolium Romanov, 1985, p. 35; and Palaeontolium Romanov, 1985, p. 35. All of these

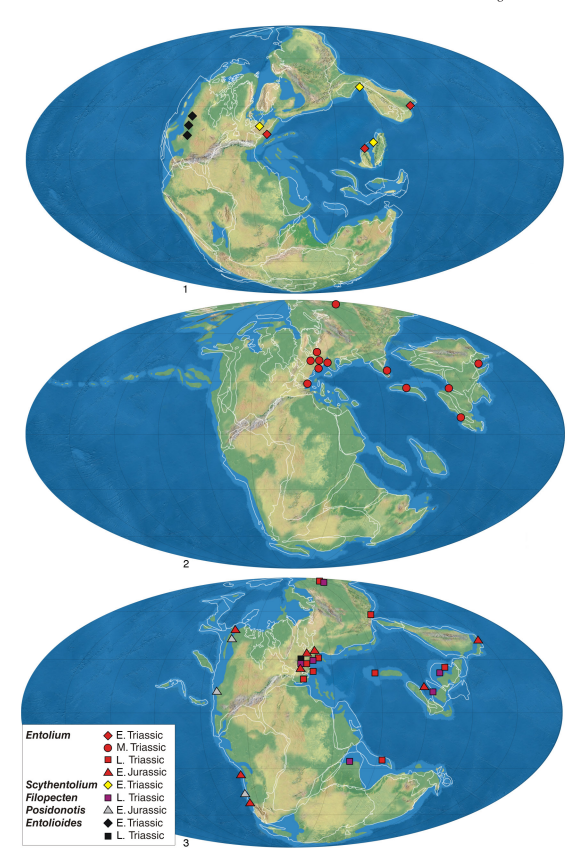


Figure 37. Paleogeographical distribution of Entoliidae (*Entolium, Scythentolium, Filopecten, Posidonotis, Entolioides*). 1, Early Triassic; 2, Middle Triassic; 3, Late Triassic-Early Jurassic.

nominal genera, except *Cornutoentolium*, were recorded as being in our study interval.

Waller (2006) proposed a new family, Entoliolidae Waller, 2006 and included within it the genera Filopecten Allasinaz, 1972, p. 301; Entolioides Allasinaz, 1972, p. 295; Scythentolium Allasinaz, 1972, p. 308; and Calvaentolium (=Crenamussium Newton in Newton & others, 1987, p. 46), thus grouping the old Triassic entoliids with filosus structure and a deep byssal notch, and lacking internal ribs. Waller (2006) regarded this family as the link between Pernopecten and Mesozoic pectinids. On the other hand, other authors (H. Yin, 1983; Nakazawa, 1996) argued that the distinction between Pernopecten and Entolium is just a matter of convenience, using the first name for Paleozoic specimens and the second for Mesozoic ones. In fact, H. Yin (1983) reported Entolium from the upper Permian, like other authors, and Nakazawa (1996) reported Pernopecten from the Lower Triassic. This issue remains unresolved until future research is done. We provisionally follow Newell and Boyd (1995) in their suggestion that *Pernopecten* was a Paleozoic genus.

In view of the significant discrepancies between different authors, this discussion is beyond the purpose of this study, and, while there is no consensus on the diagnostic characters for each taxonomic level, we regard *Entolium* in its original sense (see Damborenea [2002a] for Staesche's concept [1926]).

Genus ENTOLIUM Meek, 1865, p. 478

Type species.—Pecten demissus Meek, 1865, p. 478.

Stratigraphic range.—Lower Triassic–Upper Cretaceous (Maastrichtian) (Allasinaz, 1972; Abdel-Gawad, 1986). Cox and others (1969) assigned it a Middle Triassic–Upper Cretaceous range. The youngest record is from the Maastrichtian (Abdel-Gawad, 1986) and the oldest one from the Lower Triassic (*E. discites* Schlotheim, 1820).

Paleogeographic distribution.—Cosmopolitan (Fig. 37). In the Boreal domain, in addition to the Triassic, it was also recorded in the Early Jurassic (Kurushin, 1990; Polubotko & Repin, 1990), but the specimens were not figured or discussed. Milova (1976) reported it from the Pliensbachian and Milova (1988) from the Toarcian of northeastern Russia.

Tethys domain: Early Triassic: Italy (Allasinaz, 1972; Neri & Posenato, 1985), China (C. Chen, 1982; S. Yang, Wang, & Hao, 1986); Middle Triassic: Hungary (Allasinaz, 1972; Szente, 1997), Germany (Bachmann, 1973; Hagdorn, 1995), Poland (Senkowiczowa, 1985); Anisian of Italy (Allasinaz, 1972; Posenato, 2008b), China (Gu & others, 1976; Sha, Chen, & Qi, 1990; J. Chen, 2003), Bulgaria (Tronkov & Damyanov, 1993), Bosnia and Yugoslavia (Allasinaz, 1972), northern Vietnam (Komatsu, Huyen, & Huu, 2010); Ladinian of Spain (Márquez-Aliaga, 1983, 1985; Márquez-Aliaga & Montoya, 1991; Budurov & others, 1991; López-Gómez & others, 1994), China (Gu & others, 1976), Malaysia (Tamura, 1973), Italy (Rossi Ronchetti, 1959; Allasinaz, 1972), northern Vietnam (Komatsu, Huyen, & Huu, 2010), Afghanistan (Farsan, 1972); Late Triassic: Carnian of the Alps (Allasinaz, 1966, 1972; Fürsich & Wendt, 1977; Hautmann, 2001b), Spain (Martín-Algarra, Solé de Porta, & Márquez-Aliaga, 1995), China (Gu & others, 1976; Sha, Chen, & Qi, 1990); Norian of China (Lu, 1981); Norian-Rhaetian of Iran (Hautmann, 2001b); Rhaetian of ?Tibet ("Lhasa Block") (J. Yin & Grant-Mackie, 2005), Alps (Vöros, 1981; Hautmann, 2001b), Hungary (Vörös, 1981; Hautmann, 2001b), Italy (Sirna, 1968); Early Jurassic: Hettangian of the Alps (Johnson, 1984), Germany and France (Vörös, 1971; Johnson, 1984), Vietnam (Vu Khuc & Huyen in Sato & Westermann, 1991).

Circumpacific domain: Early Triassic: Olenekian of Japan (Nakazawa, 1961; Hayami, 1975; Kashiyama & Oji, 2004); Middle Triassic: Japan (Hayami, 1975; Tamura & others, 1978); Early Jurassic: Hettangian of Chile (Hillebrandt, 1990); Hettangian–Sinemurian of Canada (Aberhan, 1998a; Aberhan, Hrudka, & Poulton, 1998), Chile (Aberhan, 1993, 1994a); Sinemurian of Japan (Hayami, 1975), Canada (Poulton, 1991).

Austral domain: Early Jurassic: Sinemurian of Argentina (Damborenea, 2002a; Damborenea & Manceñido, 2005b).

Boreal domain: Middle Triassic: Anisian of Siberia (Dagys & Kurushin, 1985); Late Triassic: Carnian of Primorie (Kiparisova, 1972); Norian of Siberia (Okuneva, 1985).

Paleoautoecology.—B, E, S, Un, FaM-Sed; R-Sw. Young specimens of *Entolium s.s.* had a byssal notch, but this was lost in adult stages. The life habit also probably changed from epibyssate in early stages to reclined in the upper ones. The low convexity of the shell, its reduced thickness, its circular outline, and the wide umbonal angle are characteristics that indicate it could have been a good swimmer, like some living pectinids (Johnson, 1984). However, in our provisional, broad concept of *Entolium*, there are some species that retained the byssal notch until the adult stages and therefore were epibyssate their entire lives.

Mineralogy.—Bimineralic (Carter, 1990a, p. 257). The family Entoliidae is characterized by bimineralic mineralogy (Carter, 1990a), with different microstructure types in juveniles and adults (homogeneous, simple prismatic, foliated, and fibrous prismatic), and a cross-lamellar inner shell layer (see Allasinaz, 1972; Waller, 1978; Johnson, 1984, among others, for more information).

Genus SCYTHENTOLIUM Allasinaz, 1972, p. 308

Type species.—Pecten tirolicus Wittenburg, 1908, p. 23.

Stratigraphic range.—Lower Triassic (Allasinaz, 1972). Allasinaz (1972) proposed the genus Scythentolium, including several Lower Triassic species: S. eurasiaticum (Wittenburg, 1908); S. kokeni (Wittenburg, 1909); S. longauris (Wittenburg, 1908); S. rombergi (Wittenburg, 1908); S. sojale (Wittenburg, 1908); S. subtile (Wittenburg, 1908); and S. tirolicum (Bittner, 1895).

Paleogeographic distribution.—Tethys (Fig. 37). Although Allasinaz (1972) assigned it a cosmopolitan distribution, the species included within *Scythentolium* were only reported from the southern Alps and Salt Range (Pakistan).

Tethys domain: Early Triassic: Alps (Diener, 1923; Allasinaz, 1972; Neri & Posenato, 1985; Broglio-Loriga & others, 1990; Fraiser & Bottjer, 2007a), China (Lu & Chen, 1986), Pakistan (Diener, 1923; Allasinaz, 1972; Nakazawa, 1996); Induan of southern China (Hautmann & others, 2011).

Paleoautoecology.—B, E, S, Epi, Sed; By. Its external morphology is similar to *Entolium*, but it had a byssal notch in the adult stage and had unequal auricles (Allasinaz, 1972), so we assume an epibyssate mode of life for *Scythentolium*.

Mineralogy.—Bimineralic (Carter, 1990a, p. 257). See mineralogy for *Entolium* (p. 105).

Genus FILOPECTEN Allasinaz, 1972, p. 301

Type species .- Pecten filosus Hauer, 1857, p. 30.

Remarks.—Allasinaz (1972) included in *Filopecten* the following species: *F. filosus* (Hauer); *F. schlosseri* (Woehrmann); *F. incognitus* (Bittner); *F. azzarolae* (Stopanni); *F. helii* (Emmrich); *F. aff. discites* (Cox); *F. fimbriatus* (Mansuy); *F. quotidianus* (Healey); *F. kolymaensis* (Kiparisova); and with doubts, *Pecten rosaliae* (Salomon, 1895). However, Waller (in Waller & Stanley, 2005) suggested that the last species should not be included, since, among other differences, it has a ctenolium, which is absent in all members of family Entoliidae. *Filopecten* is very similar to *Entolium* concerning the shape of the auricles, the general external form, and even the hinge, and they differ in the byssal notch and ornamentation (Allasinaz, 1972). According to Hautmann (2001b), these features should not be used for generic distinction within the family. In fact, Hautmann (2001b) included the species *incognitum* (Bittner, 1901c), which was transferred to *Filopecten* by Allasinaz (1972), into *Entolium (Entolium)*.

Stratigraphic range.—Upper Triassic (Carnian–Rhaetian) (Allasinaz, 1972). According to the species listed by Allasinaz (1972), the genus had an Upper Triassic range.

Paleogeographic distribution.-Tethys and Boreal (Fig. 37).

Tethys domain: Late Triassic: China (Diener, 1923); Carnian of the Alps and Hungary (Allasinaz, 1972); Rhaetian of the Alps and Hungary (Allasinaz, 1972), Burma (Healey, 1908), Indochina (Allasinaz, 1972).

Boreal domain: Late Triassic: Siberia (Allasinaz, 1972).

Paleoautoecology.—B, E, S, Epi, Sed; By. Similar to Scythentolium. Mineralogy.—Bimineralic (Carter, 1990a, p. 257). See mineralogy for Entolium.

Genus POSIDONOTIS Losacco, 1942, p. 11

Type species.—Posidonotis dainelii Losacco, 1942, p. 11.

Remarks.—Following Damborenea (1986, 1987b), we regard *Pectinula* A. F. Leanza, 1943, p. 241, as a junior synonym of *Posidonotis* (see discussion for *Pectinula* in Genera not Included, p. 167), and we include *Posidonotis* in the family Entoliidae, although not all authors agree with this (see Hayami, 1988; Aberhan, 1994a, 1998a; Monari, 1994; Waller, 2006). Cox and others (1969), as well as most mentioned authors, included it within the family Posidoniidae. *Pectinula* was assigned to Pectindae by A. F. Leanza (1943), and this was followed by Cox and others (1969).

Stratigraphic range.—Lower Jurassic (Sinemurian–Toarcian) (Damborenea, 1987b). Cox and others (1969) assigned *Posidonotis* to the Middle Jurassic (Aalenian) and *Pectinula* to the Lower Jurassic. The range assigned by Cox and others (1969) to *Posidonotis* was taken from Losacco (1942), who reported it from Aalenian deposits, but these were later redated as Toarcian (see Damborenea, 1987b, p. 192–193).

Paleogeographic distribution.—Circumpacific (Fig. 37). During our study interval, it was only present on the eastern coast of the Paleopacific, but during the Pliensbachian its distribution was broader (see Damborenea, 1986, 1987b; Hayami, 1988; Monari, 1994).

Circumpacific domain: Early Jurassic: Sinemurian of Chile (Aberhan, 1994a, 1998a), California (United States) and British Columbia (Canada) (Damborenea, 1986, 1987b; Aberhan & Pálfy, 1996).

Paleoautoecology.—B, E, S, Un, Sed; R. Several authors (e.g., Hayami, 1969a, 1988; Hillebrandt, 1981) suggested a pseudoplanktonic mode of life for *Posidonotis*, because it is often found in black shales with no associated benthic fauna. This mode of life is unlikely, because byssal structures are not present in adult specimens, although juveniles had them. In addition, other modes of life have been suggested, as nektoplanktonic, benthic with chemosymbiotic organisms, or teleplanic larvae (see Aberhan & Pálfy, 1996). However, the most plausible mode of life during the adult stages is reclining on soft substrates; in young stages, it was a byssate bivalve (Aberhan & Pálfy, 1996). Some species are interpreted as opportunistic, as they were recorded in great abundance in facies poor in oxygen, where only ammonoids are found (Damborenea, 1987b; Aberhan & Pálfy, 1996).

Mineralogy.—Bimineralic (Carter, Barrera, & Tevesz, 1998). See mineralogy for *Entolium*.

Genus ENTOLIOIDES Allasinaz, 1972, p. 295

Type species.—Pecten zitteli Wöhrmann & Koken, 1892, p. 173. Stratigraphic range.—Lower Triassic–Upper Triassic (Carnian) (Allasinaz, 1972; Newell & Boyd, 1995). Allasinaz (1972) assigned to Entolioides a Middle–Upper Triassic range, but all species he included were only recorded from Carnian deposits [E. deeckei (Parona, 1889); E. lavaredanus (Frech, 1904); E. porschei (Toula, 1913); E. setinus (Gortani, 1902); E. subdemissus (Münster, 1841); and E. zitteli (Wöhrmann & Koken, 1892)], according to the range listed on p. 222 of his monograph. Newell and Boyd (1995) reported the type species (Pecten zitteli) from the Middle Triassic of the Alps, but Allasinaz (1972) reported it from the Carnian of the southern Alps, and we only found it mentioned from this stage. Newell and Boyd (1995) reported the species Entolioides utahensis (Meek, 1877) from the Lower Triassic of the Thaynes Formation.

Paleogeographic distribution.—Tethys and Circumpacific (Fig. 37). Tethys domain: Late Triassic: Carnian of the Alps (Allasinaz, 1972).

Circumpacific domain: Early Triassic: Idaho, Montana, and Nevada (United States) (Newell & Boyd, 1995).

Paleoautoecology.—B, E, S, Epi, FaM; By-Sw. Two groups can be distinguished within pectinoideans regarding their mode of life (S. M. Stanley, 1972): some are epibyssate, and they are characterized by different convexity in both valves, the anterior auricle being more developed, and a byssal sinus throughout their ontogeny; others, with more symmetrical shell with both valves similarly convex, have auricles of the same shape and size, and an umbonal angle greater than 90°, are also epibyssate but with the ability to swim. Both valves in *Entolioides* were nearly equally convex, with subequal auricles, an umbonal angle between 85° and 120°, and a small byssal notch (Allasinaz, 1972). According to these features, *Entolioides* belongs to the second group.

Mineralogy.—Bimineralic (Carter, Barrera, & Tevesz, 1998). See mineralogy for *Entolium* (see p. 105).

Superfamily KALENTEROIDEA Marwick, 1953

According to Z. Fang and Morris (1997) and Damborenea (2004), the classification of the superfamily Kalenteroidea Marwick, 1953, is modified with respect to Amler (1999); we consider the family Permophoridae Poel, 1959, as a synonym of Kalenteridae Marwick, 1953. Several genera assigned to this family are very similar to each other, both externally and internally (hinge details and muscle impressions) (Damborenea, 2004).

Family KALENTERIDAE Marwick, 1953 (=Permophoridae Poel, 1959) Genus PERMOPHORUS Chavan, 1954, p. 200

nom. nov. pro Pleurophorus King, 1844, p. 313, non Mulsant, 1842, p. 312

Type species.—Arca costata Brown, 1844, p. 66.

Stratigraphic range.—Carboniferous (Mississippian)-Lower Triassic (Olenekian) (Hoare, Heaney, & Mapes, 1989; Newell & Boyd, 1999). Cox and others (1969) assigned it a Lower Carboniferous-Permian range. For a long time, it was regarded as an exclusively Paleozoic genus, but recently, it was also reported from the Triassic. We are only taking into account the Triassic record in Newell and Boyd (1999), since other records have some problems we cannot solve now. Newell and Boyd (1999) warned about the misunderstanding of the Permophorus hinge details in Cox and others (1969). Based on this interpretation, Waterhouse (1979b) described Lower Triassic specimens from New Zealand that were subsequently assigned to the Middle Triassic (H. J. Campbell, 1984). Although other species were reported from the Lower Triassic (see Newell & Boyd, 1999), they were based on poorly preserved material. In addition, Permophorus was also reported from the Upper Triassic (Rhaetian) by Ivimey-Cook and others (1999), but these authors pointed out that the assignment was doubtful, since they did not observe the hinge of their specimens. Skwarko (1967) referred his Carnian and Norian specimens from New Guinea to Permophorus? hastatus, but later, Skwarko (1983) designated this as type species of his new genus Somareoides Skwarko, 1983.

Paleogeographic distribution.—Circumpacific (Fig. 38). In our study interval, it was only known from the Circumpacific domain. Fraiser and Bottjer (2007a) also listed it from the Triassic of Italy, but they did not figure or describe the specimens.

Circumpacific domain: late Permian: Japan (Nakazawa & Newell, 1968; Hayami & Kase, 1977); Early Triassic: Olenekian of Utah, Wyoming, Idaho, and Montana (United States) (Newell & Boyd, 1999), ?Idaho (United States) (Ciriacks, 1963), western United States (Boyer, Bottjer, & Droser, 2004; Fraiser & Bottjer, 2007a).

Paleoautoecology.—B, Is, S, Endo-Un, Sed-SM; By-Sb. Permophorus had some characteristics that indicate a shallow burrowing habit. It had an equivalve, inequilateral, and elongated shell, with prosogyrous beaks, lunule, and escutcheon that, according to S. M. Stanley (1975), facilitates burial. No pallial sinus is observed, so if siphonate, siphons would have been very short. According to Quiroz-Barroso and Perrilliat (1998), Permophorus was an endobyssate bivalve, but none of the specimens described in the literature show a byssal notch or gape. However, S. M. Stanley (1972) noted that at least the type species had a reduced anterior part, which suggests the presence of a byssus.

Mineralogy.—Aragonitic (Carter, 1990a, p. 271). Outer shell layer: aragonite (fibrous-prismatic). Middle shell layer: aragonite (cross-lamellar). Inner shell layer: aragonite (homogeneous).

Genus CURIONIA Rossi Ronchetti in Rossi Ronchetti & Allasinaz, 1965, p. 366

Type species.—Myoconcha curionii Hauer, 1857, p. 561.

Stratigraphic range.—Lower Triassic (?)–Upper Triassic (Rhaetian) (Rossi Ronchetti & Allasinaz, 1965). Rossi Ronchetti (in Rossi Ronchetti & Allasinaz, 1965) proposed *Curionia* and assigned it a Triassic (Scythian–Rhaetian) range. When listing the included species, she first mentioned those that were contemporaneous with the type species (Carnian), and then the other Triassic species; so, in her list, the first recorded species is Carnian and the last is Rhaetian. Perhaps because of this arrangement, Cox and others (1969) considered the genus to be present only in the Late Triassic, or maybe they simply disagreed with the species listed by the author of the genus. We assign to *Curionia* the stratigraphic range given by its original authors.

Paleogeographic distribution.—Tethys (Fig. 38). Curionia was only known from the Tethys domain. It was reported from the Early Jurassic of Nevada (Laws, 1982), but Hallam and Wignall (2000) argued that Laws probably confused Curionia with Modiolus, which is very abundant in the area, since Curionia was a European genus that disappeared in the Late Triassic. However, Stiller and Chen (2006) reported it from the Anisian of China.

Tethys domain: Early Triassic: Olenekian of ?Bakony (Hungary) (Frech, 1907; Rossi Ronchetti & Allasinaz, 1965); Middle Triassic: Anisian of China (Stiller & Chen, 2006), Italy (Rossi Ronchetti & Allasinaz, 1965); Muschelkalk of Germany (Rossi Ronchetti & Allasinaz, 1965); Ladinian of Italy (Posenato, 2002); Late Triassic: Carnian of the Alps (Rossi Ronchetti & Allasinaz, 1965); Norian of Italy (Rossi Ronchetti & Allasinaz, 1965); Rhaetian of Italy (Stoppani, 1860–1865; Rossi Ronchetti & Allasinaz, 1965), Iran (Repin, 2001).

Paleoautoecology.—B, Is-Se, S, Endo-Un, Sed-SM; By-Sb. Like all Kalenteridae genera, the external morphology of *Curionia* indicates a shallow infaunal or semi-infaunal mode of life. By analogy with *Modiolus*, it could perhaps have been an endobyssate bivalve.

Mineralogy.—Aragonitic (Schneider & Carter, 2001). Outer shell layer: aragonite (prismatic). Middle shell layer: aragonite (cross-lamellar). Inner shell layer: aragonite (prismatic).

Genus TRIAPHORUS Marwick, 1953, p. 69

Type species.—Pleurophorus zealandicus Trechmann, 1918, p. 212.

Stratigraphic range.—Upper Triassic (Carnian–Norian) (H. J. Campbell, 1984). Marwick (1953) proposed *Triaphorus* from Carnian deposits. Subsequently, Cox and others (1969) assigned it an Upper Triassic range. Grant-Mackie (1960) reported it from Otapirian (=Rhaetian) and Warepan (=Norian) of New Zealand, but he did not figure the specimens and based his record on a personal communication from J. D. Campbell. Moreover, H. J. Campbell

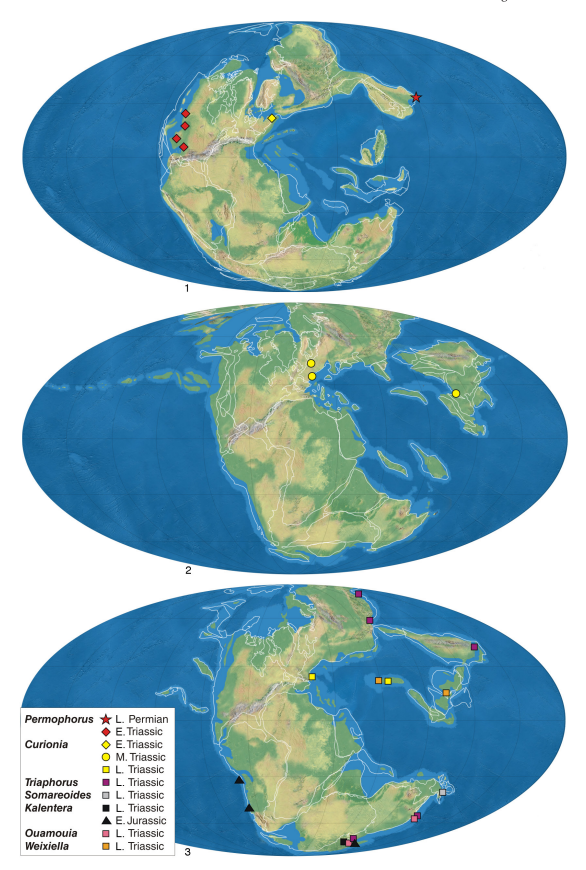


Figure 38. Paleogeographical distribution of Kalenteridae (*Permophorus, Curionia, Triaphorus, Somareoides, Kalentera, Ouanouia, Weixiella*). 1, Early Triassic; Middle Triassic; 3, Late Triassic–Early Jurassic.

(1984) mentioned the genus from the upper ?Carnian–Norian, but neither figured nor described the material, although he mentioned several references. Following H. J. Campbell (1984), we assign it a Carnian–Norian range.

Paleogeographic distribution.—Austral, Boreal, and Circumpacific (Fig. 38). *Triaphorus* was distributed in the Austral and Circumpacific domains and doubtfully in Boreal regions as well. When the specimens are not well preserved and the hinge is not seen, it is difficult to distinguish between *Triaphorus* and *Kalentera* (Damborenea, 2004). This latter genus had a bipolar distribution during the Jurassic, but during the Late Triassic, it was only known from the Austral domain. It is necessary to check if the Boreal records of *Triaphorus* can be confirmed and if the specimens show the diagnostic characters.

Austral domain: Late Triassic: Carnian of New Zealand (Trechmann, 1918; Marwick, 1953); Carnian–Norian of New Zealand and New Caledonia (H. J. Campbell, 1984).

Boreal domain: Late Triassic: northeastern Russia (Kiparisova, Bychkov, & Polubotko, 1966); Carnian of Primorie (Kiparisova, 1972).

Circumpacific domain: Late Triassic: Carnian of Japan (Kobayashi & Ichikawa, 1950; Hayami, 1975).

Paleoautoecology.—B, Is-Se, S, SM; Sb. Triaphorus is externally similar to Permophorus and Kalentera. It had a modioliform shell, and, in the genus diagnosis offered by Marwick (1953), the presence of a pedal muscle scar is pointed out. Since no pallial sinus and no evidence of byssate habit are listed, we assume that it was a shallow burrower belonging to the shallow infauna or semi-infauna (Grant-Mackie, 1960).

Mineralogy.—Aragonitic (Carter, 1990a, p. 270). According to Carter (1990a), although the shell mineralogy of *Triaphorus* and other members of the family Kalenteridae is unknown, its mode of preservation suggests an original aragonitic mineralogy.

Genus SOMAREOIDES Skwarko, 1983, p. 67

Type species.—Permophorus? hastatus Skwarko, 1967, p. 66.

Remarks.—Skwarko (1967) tentatively included the species *hastatus* Skwarko, 1967, in *Permophorus*, as only external structures could be observed in his specimens. The discovery of new material with well-preserved hinge confirmed that this species does not belong to *Permophorus* nor to any previously known genus (Skwarko, 1983).

Stratigraphic range.—Upper Triassic (Carnian) (Skwarko, 1983). Although Skwarko (1967) assigned a Carnian–Norian range to the type species, when he proposed *Somareoides* (Skwarko, 1983), he noted that the most likely age was Carnian.

Paleogeographic distribution.—Southern Tethys (Fig. 38).

Tethys domain: Late Triassic: Carnian of Papua New Guinea (Australian province according to Damborenea, 2002b) (Skwarko, 1967, 1983).

Paleoautoecology.—B, Se, S, Endo-Un, Sed-SM; Sb. According to Skwarko's (1983) description, *Somareoides* had an inequivalve and modioliform shell, with developed anterior part and elongated posterior one. With these characteristics, its mode of life should be similar to other family members. It was probably a semi-infaunal bivalve.

Mineralogy.—Aragonitic (Carter, 1990a, p. 270). See mineralogy for *Triaphorus* (p. 107).

Genus KALENTERA Marwick, 1953, p. 108

Type species.—Kalentera mackayi Marwick, 1953, p. 108.

Stratigraphic range.—Upper Triassic (Norian)-Lower Jurassic (Toarcian) (Grant-Mackie, 1960). Marwick (1953) proposed Kalentera and included two new species, K. mackayi from the Lower Jurassic and K. flemingi from the Temaikan (Bathonian). Later, Grant-Mackie (1960) indicated that the beds that Marwick (1953) referred to as Temaikan were Ururoan instead (= Pliensbachian-Toarcian), and he proposed another new species within the genus, K. marwicki from the Otapirian (=Rhaetian). He also mentioned the genus from Warepan (=Norian) as being associated with Monotis calvata Marwick, 1953. Cox and others (1969) assigned Kalentera a Lower Jurassic range and Sepkoski (2002) assigned it an Upper Triassic (upper Carnian)–Middle Jurassic (Bathonian) range, following Hallam (1977). Hallam mentioned the genus from the Middle Jurassic, but not in the Carnian. H. J. Campbell (1984) assigned it a Norian-Bathonian range. Following Grant-Mackie (1960), we assign it a Norian-Toarcian range.

Paleogeographic distribution.-Austral and Circumpacific (Fig. 38). Although originally thought to be endemic to the Austral domain, new records showed that Kalentera was endemic to the southern domain (Maorian Province) during the Late Triassic and had a bipolar distribution (at high latitudes) during the Early Jurassic (Damborenea, 2001, 2002b, 2004) (see also Triaphorus paleogeographic distribution, p. 107). Boreal records date from the Pliensbachian (Damborenea, 2004; Zakharov & others, 2006), and, therefore, they are not taken into account here. D. G. Taylor and Guex (2002) included their new species K. lawsi Taylor & Guex, 2002, from the Triassic–Jurassic boundary beds of Oregon in Kalentera, but this assignment is tentative, because their specimens did not show hinge details. Damborenea (2004) noticed that those specimens were previously allocated to Curionia sp. by Laws (1982), and, later, D. G. Taylor, Boelling, and Guex (2000) assigned them to Kalentera? sp. However, Hallam and Wignall (2000) considered that Laws's (1982) "Curionia" specimens could probably refer to Modiolus, a genus very abundant in the area. Laws (1982) did not figure the specimens, and those figured by D. G. Taylor and Guex (2002) are unconvincing.

Circumpacific domain: Early Jurassic: Sinemurian of northern Chile (Covacevich, Pérez, & Escobar, 1991).

Austral domain: Late Triassic: Norian–Rhaetian of New Zealand (Grant-Mackie, 1960; MacFarlan, 1998); Early Jurassic: Hettangian– Sinemurian of New Zealand (Marwick, 1953), Argentina (Damborenea, 2004).

Paleoautoecology.—B, Se-Is, S, SM; Sb. *Kalentera* shell morphology indicates that it was probably a shallow burrower that lived wholly or partly buried in the sediment (Grant-Mackie, 1960; H. J. Campbell, 1984; Damborenea, 2004). The inferred environment of the fossilbearing beds indicates that stability and type of substrate limited the distribution of this genus (Damborenea, 2004) in nearshore deposits. According to Grant-Mackie (1960), the absence of excavated galleries in the sediment and of a pallial sinus suggests *Kalentera* lacked siphons (or perhaps they were too short), and therefore it would have been a very shallow burrower.

Mineralogy.—Aragonitic (Carter, 1990a, p. 270; Damborenea, 2004). In *Kalentera* (or other members of the family Kalenteridae), the shell mineralogy is unknown, although, according to Carter (1990a), its mode of preservation suggests an original aragonitic

Genus OUAMOUIA H. J. Campbell, 1984, p. 158

Type species.—Ouamouia grantmackiei H. J. Campbell, 1984, p. 159.

Remarks.—H. J. Campbell (1984) included *Ouamouia* in the family Permophoridae (=Kalenteridae) and related it mainly to *Permophorus* and *Kalentera*. However, Damborenea (2004) argued that *Ouamouia* is quite different from *Kalentera*, and its massive hinge dentition and other characters indicate it was a cardiniid.

Stratigraphic range.—Upper Triassic (Norian–Rhaetian) (H. J. Campbell, 1984). This monospecific genus was described by H. J. Campbell (1984) from Norian–Rhaetian beds.

Paleogeographic distribution.—Austral (Fig. 38).

Austral domain: Late Triassic: Norian–Rhaetian of New Zealand and New Caledonia (H. J. Campbell, 1984).

Paleoautoecology.—B, Is, S, Endo-Un, Sed-SM; By-Sb. Ouamouia grantmackiei had a modioliform shell, undefined pallial sinus, lunule, and ornamented shell. These characteristics match well with a shallow burrower mode of life. Probably it lived buried near the surface and possessed short siphons (see H. J. Campbell, 1984, p. 162). Because it had a thick shell and a massive hinge, it probably lived in high-energy environments (H. J. Campbell, 1984). No structure suggesting it was a byssate bivalve is present, but its modioliform shape may indicate this life habit.

Mineralogy.—Aragonitic. There are no data about *Ouamoia* shell mineralogy or microstructure. The shell was probably entirely aragonitic.

Genus WEIXIELLA Guo & Chen in Guo, 1985, p. 187, 268

Type species.—Weixiella diana Guo & Chen in Guo, 1985, p. 187. Remarks.—Guo and Chen (in Guo, 1985) included Weixiella in the family Pachycardiidae Cox, 1962, due its resemblance to Cardiniodes Kobayashi & Ichikawa, 1952, especially in hinge features. However, Hautmann (2001b) found more similarities with Permophorus Chavan, 1954, and included it in the family Permophoridae (=Kalenteridae), although he also indicated some resemblance to the family Unionidae Fleming, 1828. Z. Fang and others (2009) followed the original paper by Guo (1985) and referred it to the family Pachycardiidae.

Stratigraphic range.—Upper Triassic (Norian–Rhaetian) (Hautmann, 2001b). The genus was described from the Upper Triassic of Yunnan (China). Hautmann (2001b) reported it from the Norian–Rhaetian of Iran and also mentioned it from coeval beds in China, referring to the original description of the genus.

Paleogeographic distribution.—Tethys (Fig. 38).

Tethys domain: Late Triassic: Norian–Rhaetian of China (Guo & Chen in Guo, 1985; Hautmann, 2001b), Iran (Hautmann, 2001b).

Paleoautoecology.—B, Is, S, Endo-Un, Sed-SM; By-Sb. Weixiella was a suspensivorous shallow burrower (Hautmann, 2001b). The external morphology is not very different from the other members of the family Kalenteridae, but the anterior is more lobed. Although structures indicating the presence of byssus were not observed, it might have been endobyssate, as were other members of this family. *Mineralogy.*—Aragonitic. There are no data about *Weixiella* shell structure. We assigned it aragonitic shell mineralogy, as in other members of the family Kalenteridae.

Family MYOCONCHIDAE Newell, 1957 Genus MYOCONCHA J. de C. Sowerby, 1824, p. 103

Type species.—Myoconcha crassa J. de C. Sowerby, 1824, p. 103.

Stratigraphic range.—Upper Triassic (Rhaetian)–Upper Cretaceous (Maastrichtian) (Hodges, 2000). With the inclusion of Triassic species previously assigned to Myoconcha into Curionia Rossi Ronchetti (in Rossi Ronchetti & Allasinaz, 1965, p. 366) and Pseudomyoconcha Rossi Ronchetti (in Rossi Ronchetti & Allasinaz, 1966, p. 1101), Myoconcha was restricted to Jurassic onward. Cox and others (1969) assigned it a Lower Jurassic-Upper Cretaceous range, with a doubtful record from the Permian. Even so, some authors maintained its range from the Upper Triassic. Hautmann (2001b) considered Pseudomyoconcha to be a subgenus of Myoconcha and mentioned it from the Norian and Rhaetian. Hodges (2000) assigned it an Upper Triassic-Upper Cretaceous range and doubtfully extended it to the Permian. Hodges (2000) reported Myoconcha (Myoconcha) psilonoti Quenstedt, 1856 in 1856–1858, from the Rhaetian of England, and it was also mentioned from the Rhaetian of the Apennines (Diener, 1923). Ivimey-Cook and others (1999) mentioned the same species from the Rhaetian of England, but the figured specimen does not show the hinge, a character critical to distinguish it from other Myoconchidae. Other authors (e.g., Zorn, 1971; Busse & Horn, 1978; Malinowskiej, 1979) mentioned Triassic species assigned to Myoconcha that were transferred to other genera by Rossi Ronchetti and Allasinaz (1965, 1966).

The Permian quotations appeared from the subjective synonymy proposed by Newell (1957), who considered Labayophorus Licharew, 1939, upper Permian genus of the Caucasus, as a junior synonym of Myoconcha. This seems to have been accepted with reservations by subsequent authors who indicated the doubtful presence of Myoconcha in the Permian. Rossi Ronchetti and Allasinaz (1966) noticed that the illustration of the right valve of Myoconcha, figured in Newell (1957) and prepared by Cox, is the only existing schematic representation of the right valves of Myoconcha. Labayophorus and Myoconcha are distinguished by the presence of one cardinal tooth in each valve in the former, while the latter has one on the right and two on the left valve. In addition, Rossi Ronchetti and Allasinaz (1966) listed some other differences that could separate the two genera, and they considered Labayophorus to be Paleozoic and Myoconcha to be Mesozoic, predominantly Jurassic. Some Permian records do not seem to be attributable to Myoconcha. Simões and Fittipaldi (1987) reported Myoconcha from the Permian, following Mendes (1944), who recorded a doubtful Myoconcha sp. from sediments originally dated as Triassic but later proved to be Permian (Simões & Fittipaldi, 1987). Later, Mendes (1945) reassigned these specimens to Naiadopsis lamellosus Mendes. Furthermore, Mendes (1944) noticed that Permian specimens attributed to Myoconcha are probably Modiolopsis. Another Permian record is found in Hayasaka (1967) from Japan, but the only available specimen is strongly deformed and incomplete, and its generic assignation is very doubtful. Paleogeographic distribution.—Tethys and Austral (Fig. 39).

Tethys domain: Late Triassic: Rhaetian of England (Ivimey-Cook & others, 1999; Hodges, 2000); Early Jurassic: Hettangian–Sine-

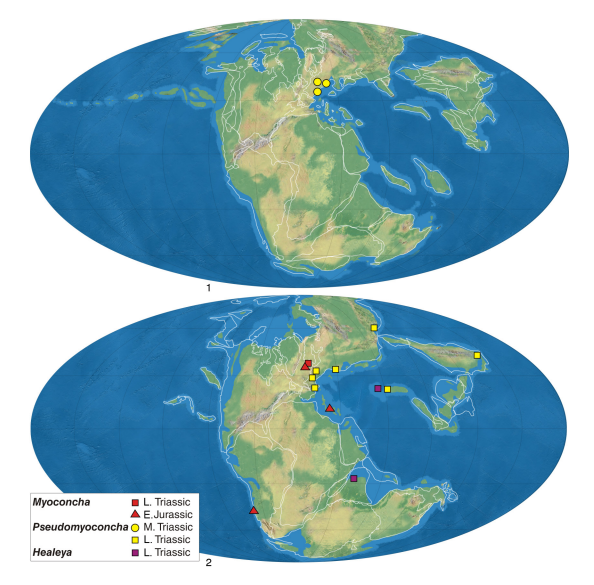


Figure 39. Paleogeographical distribution of Myoconchidae (Myoconcha, Pseudomyoconcha, Healeya). 1, Middle Triassic; 2, Late Triassic-Early Jurassic.

murian of England (Hodges, 2000; Hallam, 1987); Sinemurian of England (Liu, 1995), Turkey (M. A. Conti & Monari, 1991).

Austral domain: Early Jurassic: Hettangian–Sinemurian of Argentina (Damborenea & Manceñido, 2005b).

Paleoautoecology.—B, Se, S, Endo, Sed; By. According to S. M. Stanley (1972), most species belonging to this genus have a modioliform external appearance, have elongated and prosocline shells, with a reduced anterior part and broad byssal gape, indicating an endobyssate semi-infaunal mode of life, similar to *Modiolus*. Other authors agreed with the attribution of this mode of life (Damborenea in Damborenea & González-León, 1997; Hodges, 2000; Delvene, 2001).

Mineralogy.—Aragonitic (Morris, 1978; Z. Fang & Morris, 1997). According to Morris (1978), *Myoconcha decorata* (Münster, 1837, in Goldfuss, 1833–1841) had a homogeneous shell microstructure, but he noticed that the lack of a prismatic outer shell layer may be due to erosion of the shell. Carter (1990a) suggested that the shell should be reviewed to observe if there is cross-lamellar structure. Z. Fang and Morris (1997) found remains of cross-lamellar structure preserved in specimens of *Myoconcha saemanni* Loriol.

Genus PSEUDOMYOCONCHA

Rossi Ronchetti in Rossi Ronchetti & Allasinaz, 1966, p. 1101

Type species.—Myoconcha lombardica Hauer, 1857, p. 559.

Remarks.—Rossi Ronchetti (in Rossi Ronchetti & Allasinaz, 1966) transferred all Triassic species previously assigned to *Myoconcha* to *Pseudomyoconcha*, except for those she included in *Curionia* (see Rossi Ronchetti & Allasinaz, 1965). Some species were introduced in *Pseudomyoconcha* tentatively, because the hinge (a key character) was not observed. She separated these species into two groups: one containing the species that were consistent with the new genus diagnosis, and another with species that did not fit strictly there but were closer to *Pseudomyoconcha* than to *Myoconcha*.

She noticed that the latter group probably could be recognized as a new taxon but could not see details of the hinge and muscle scars. Hautmann (2001b) argued that differences between *Myoconcha* and *Pseudomyoconcha* are very subtle and decided to keep the second as a subgenus of the first.

Stratigraphic range.—Middle Triassic (Ladinian)–Upper Triassic (Rhaetian) (Rossi Ronchetti & Allasinaz, 1966; Hautmann, 2001b). The range assigned by Rossi Ronchetti (in Rossi Ronchetti & Allasinaz, 1966), according to the species included in the new genus, was Ladinian–Norian. Cox and others (1969) assigned it the same stratigraphic range. The genus was extinguished in the Late Triassic (Hallam, 1981, 2002).

Paleogeographic distribution.—Tethys, Boreal, and Circumpacific (Fig. 39). *Pseudomyoconcha* was reported from the Late Triassic of China (Wen & others, 1976; Lu, 1981; Gou, 1993), but the specimens figured in those papers do not seem convincing. In none is the hinge shown, and the external morphology of the members of the families Myoconchidae and Permophoridae are similar. J. Chen (2003, p. 658, fig. 4.4.2) recorded it from the Anisian and throughout the Late Triassic in southern China, but he did not figure it or indicate the original data source. The presence of *Pseudomyoconcha* in southern China is thus still doubtful.

Tethys domain: Middle Triassic: Ladinian of Hungary, Germany, and Italy (Rossi Ronchetti & Allasinaz, 1966); Late Triassic: Carnian of Italy (Rossi Ronchetti & Allasinaz, 1966), Germany (Linck, 1972), Slovenia and Yugoslavia (Jelen, 1988); Carnian–Norian of Hungary (Rossi Ronchetti & Allasinaz, 1966); Norian–Rhaetian of Iran (Hautmann, 2001b).

Circumpacific domain: Late Triassic: Carnian, ?Norian of Japan (Rossi Ronchetti & Allasinaz, 1966); Norian of western Carpathians (Kollarova & Kochanová, 1973).

Boreal domain: Late Triassic: Primorie (Kiparisova, 1972).

Paleoautoecology.—B, Se, S, Endo, Sed; By. According to the generic diagnosis by Rossi Ronchetti (in Rossi Ronchetti & Allasinaz, 1966), the *Pseudomyoconcha* shell is equivalve, strongly inequilateral, modioliform, and had a byssal notch. With these characteristics, it was most likely an endobyssate semi-infaunal bivalve.

Mineralogy.—Aragonitic (Rossi Ronchetti & Allasinaz, 1966; Carter, 1990a, p. 271). Carter (1990a) interpreted the data provided by Rossi Ronchetti and Allasinaz (1966) slightly differently. He identified a fibrous prismatic outer shell layer and a middle shell layer of cross-lamellar structure.

Genus HEALEYA Hautmann, 2001b, p. 108

Type species.—Modiolopsis gonoides Healey, 1908, p. 51.

Remarks.—Hautmann (2001b) proposed the genus *Healeya* within the subfamily Myoconchinae. Subsequently, Hautmann (2008) proposed a new family, Healeyidae, to include *Healeya* and other genera. Given the objections with this new family proposition, we consider *Healeya* in its original allocation (see discussion for the family Mysidiellidae Cox, 1964, p. 22).

Stratigraphic range.—Upper Triassic (Norian-Rhaetian) (Hautmann, 2001b, 2008). *Healeya* is a Upper Triassic monospecific genus. Hautmann (2001b) reported from the Norian-Rhaetian of Iran. The type species was originally described from the Rhaetian of Burma (India) by Healey (1908).

Paleogeographic distribution.—Tethys (Fig. 39). Apart from Iran and Burma, Hautmann (2001b) mentioned the possible occurrence of this genus from the Upper Triassic of Malaysia and Vietnam, due to the doubtful inclusion of two species recorded from that area, in the list of synonyms of the type species.

Tethys domain: Late Triassic: Norian-Rhaetian of Iran (Hautmann, 2001b); Rhaetian of Burma (India) (Healey, 1908).

Paleoautoecology.—B, Se, S, Endo, Sed; By. Hautmann (2001b) found specimens in life position that corroborate what their morphology indicated, i.e., *Healeya* lived semi-infaunally and it was probably an endobyssate bivalve. In the anterior part of the shell, the muscle scar of probably a byssal retractor is observed; the byssus would have been strong and protruded from the anterior part of the shell (Hautmann, 2008).

Mineralogy.—Aragonitic (Hautmann, 2008). Although no studies of *Healeya* shell microstructure were performed, the type of recrystallization indicates an aragonitic mineralogy (Hautmann, 2008).

Family HIPPOPODIIDAE Cox in Cox and others, 1969 Genus HIPPOPODIUM J. Sowerby, 1819, p. 91

Type species.—Hippopodium ponderosum J. Sowerby, 1819, p. 91. Stratigraphic range.—Upper Triassic (?Rhaetian)–Upper Jurassic (Tithonian). Cox and others (1969) assigned it a Lower Jurassic range (Hettangian–Pliensbachian) and also considered it to be from the Upper Jurassic (Tithonian). However, Zapfe (1967) suggested the origin of *Hippopodium* in the Rhaetian of the Alps. Hallam and El Shaarawy (1982) also reported it from the Rhaetian of the Alpine region of northwestern Europe, but they did not figure or describe the specimens, nor did Hallam (1981), who quoted it from the Upper Triassic of western Tethys.

Paleogeographic distribution.—Western Tethys and Boreal (Fig. 40). *Hippopodium* was a characteristic genus of the Boreal domain during the Early Jurassic (Sinemurian and Pliensbachian) (Liu, 1995). This author included England in the Boreal domain during this time. This area was on the boundary between the Boreal and Tethys domains, depending on whether their definition is based on bivalves or ammonoids. Hallam (1977) mentioned it from the European province, and, previously, he stated that although it was present in the Tethys domain, records were always from the north (Hallam, 1972).

Tethys domain: Late Triassic: Rhaetian of the ?Alps (Zapfe, 1967); Early Jurassic: Hettangian–Sinemurian of Germany (Arp, 2007).

Boreal domain: Early Jurassic: Sinemurian of England (Liu, 1995).

Paleoautoecology.—B, Se, S, Endo, Sed; By. *Hippopodium* had a lobate anterior part, thick shell, prosogyrous umbones, and a pallial line without sinus. No structures indicating the presence of byssus were observed. It probably lived semiburied.

Mineralogy.—Aragonitic (Morris, 1978; Carter, 1990a; Z. Fang & Morris, 1997). The species *H. ovale* Moore had a homogeneous microstructure in the whole shell (Morris, 1978). Carter (1990a) noticed the need for more careful studies of the shell of this species

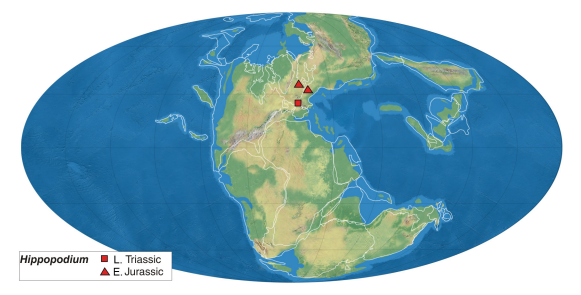


Figure 40. Paleogeographical distribution of Hippopodiidae (Hippopodium). Late Triassic-Early Jurassic.

to exclude the presence of cross-lamellar microstructure. Z. Fang and Morris (1997, p. 57) found, in a *H. ponderosum* Sowerby specimen, "patches of ill preserved but distinct crossed-lamellar structure;" but they did not find any trace of cross-lamellar structure in *H. ovale*, while in other species reported from the same beds, this microstructure was perfectly preserved.

Superfamily TRIGONIOIDEA Lamarck, 1819

For the location of genera in the Trigonioidea families, we mainly follow Cox and others (1969), except for families Minetrigoniidae and Myophoriidae, for which we follow Fleming (1987).

Family TRIGONIIDAE Lamarck, 1819 Genus TRIGONIA Bruguière, 1789 in 1789–1792, p. xiv

Type species.—Venus sulcata Hermann, 1781, p. 127.

Stratigraphic range.—Middle Triassic (Anisian)–Upper Cretaceous (Cenomanian) (H. A. Leanza, 1993). Cox and others (1969) assigned it a Middle Triassic–Upper Cretaceous range. The Anisian species *Trigonia tabacoensis* Barthel, 1958, is the earliest known species in this genus (Fleming, 1964, 1987; Pérez & Reyes, 1991; Francis & Hallam, 2003). It is difficult to establish the top of the range because, in recent years, many Cretaceous genera related to *Trigonia* were proposed, and many of them were based on type species previously referred to *Trigonia*. Of the subgenera considered by Cox and others (1969), only *T. (Trigonia)* is included here, since *Frenguelliella* A. F. Leanza, 1942, is here regarded as a distinct genus, with *Kumatrigonia* Tamura, 1959, p. 213, as its subgenus. We follow Cox and others (1969) for the top of the range (Cenomanian). *T. (Heslingtonia*) Fleming, 1987, p. 22, is also considered as a subgenus within *Trigonia* in our study interval.

Paleogeographic distribution.—Tethys, Circumpacific, and Austral (Fig. 41). Its distribution is limited during the Triassic to Barthel's (1958) record from the Anisian. Pérez and Reyes (1991) also reported *Trigonia* from the Upper Triassic and the Early Jurassic of Peru. In Europe, the family Trigoniidae appears in the Toarcian (Francis & Hallam, 2003). However, Hautmann (2001b) reported the type species of his new subgenus, *Trigonia (Modestella) zlambachensis* Haas, 1909, from the Alpine Rhaetian. Previously, this was reported by Fallahi, Gruber, and Tichy (1983) and Fleming (1987). Moreover, Hautmann (2001b) also quoted this species from the Norian of Vietnam and the Rhaetian of Burma (Malaysia).

Tethys domain: Late Triassic: Norian of Vietnam (Vu Khuc & Huyen, 1998; Hautmann, 2001b; Guo, 1985); Norian–Rhaetian of Iran (Fallahi, Gruber, & Tichy, 1983; Hautmann, 2001b).

Circumpacific domain: Middle Triassic: Anisian of Chile (Barthel, 1958); Late Triassic: Norian of Peru (Pérez & Reyes, 1991); Early Jurassic: Hettangian of Japan (Kobayashi & Kaseno, 1947; Hayami, 1975; Sato & Westermann, 1991); Hettangian–Sinemurian of Peru (Ishikawa & others, 1983; Pérez & Reyes, 1991); Sinemurian of Nevada (United States) (Poulton, 1979).

Austral domain: Middle Triassic: Anisian–Ladinian of New Zealand (Fleming, 1964, 1987).

Paleoautoecology.—B, Is, S, FM; Sb. Many authors have dealt with the mode of life of trigoniids (Tevesz, 1975; S. M. Stanley, 1977, 1978; Kelly, 1995b; Villamil, Kauffman, & Leanza, 1998; Francis & Hallam, 2003). Currently, there is only one genus, Neotrigonia, that lives in Australian waters (Beesley, Ross, & Wells, 1998). Neotrigonia is infaunal, filtering, and a nonsiphonate, fast, shallow-burrowing bivalve, which lives partially buried near the surface of the sediment (Tevesz, 1975). Mesozoic trigoniids probably had this same way of life, by analogy with Neotrigonia. S. M. Stanley (1969, 1970) showed that prosogyrous umbos helped burrowing, but the trigoniids have mostly opisthogyrous or orthogyrous umbos (S. M. Stanley, 1977). However, the external morphology, the varied ornamentation, the strong foot, and the complex hinge teeth are adapted to this mode of life (S. M. Stanley, 1977). Thus, the life position of species of this group is interpreted as being similar to *Neotrigonia*, with the posterior

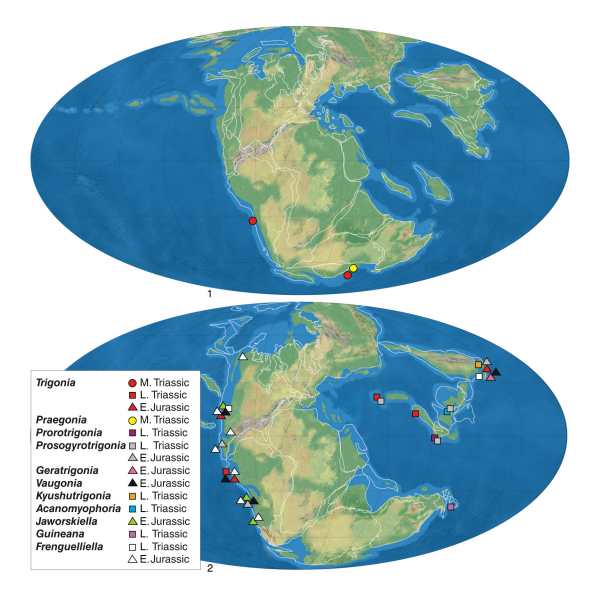


Figure 41. Paleogeographical distribution of Trigoniidae (*Trigonia, Praegonia, Prorotrigonia, Prosogyrotrigonia, Genatrigonia, Vaugonia, Kyushutrigonia, Acanomyophoria, Jaworskiella, Guineana, Frenguelliella). 1,* Middle Triassic; 2, Late Triassic–Early Jurassic.

part near the sediment surface. In many instances, the presence of epibionts on the posterior part was observed, adding to the assumption that some species lived with this part exposed, in a semi-infaunal position (Villamil, Kauffman, & Leanza, 1998). However, no epibionts were observed on the Triassic genera, perhaps due to their small size (as compared with Cretaceous forms). Cretaceous species are larger and tend to have had a more sedentary mode of life (Kelly, 1995b).

We assign an infaunal shallow-burrowing mode of life to all members of the superfamily, although some may live with the posterior part slightly exposed. For more information about the mode of life of this interesting group of burrowers, see Tevesz (1975), S. M. Stanley (1977, 1978), Kelly (1995b), Villamil, Kauffman, and Leanza (1998), or Francis and Hallam (2003), among others. *Mineralogy.*—Aragonitic (J. D. Taylor, Kennedy, & Hall, 1969). The shells of all members of the superfamily Trigonioidea probably were entirely aragonitic, with a prismatic outer shell layer and a nacreous inner shell layer, as in the living species of *Neotrigonia* (J. D. Taylor, Kennedy, & Hall, 1969; Newell & Boyd, 1975).

Genus PRAEGONIA Fleming, 1962, p. 2

Type species.—Praegonia coombsi Fleming, 1962, p. 2.

Stratigraphic range.—Middle Triassic (Ladinian). Fleming (1962) proposed *Praegonia* from the Ladinian of New Zealand. It was only recorded from that time and area (Fleming, 1964, 1987; Cox & others, 1969).

Paleogeographic distribution.—Austral (Fig. 41). Praegonia is monospecific and endemic to the Austral domain.

Austral domain: Middle Triassic: Ladinian of New Zealand (Fleming, 1962, 1964, 1987).

Paleoautoecology.—B, Is, S, FM; Sb. See discussion under *Trigonia* (p. 113).

Mineralogy.—Aragonitic (J. D. Taylor, Kennedy, & Hall, 1969). No details are known about *Praegonia* shell microstructure. See discussion under *Trigonia* (p. 113).

Genus PROROTRIGONIA Cox, 1952, p. 57

Type species.—Trigonia seranensis Krumbeck, 1923a, p. 211.

Stratigraphic range.—Upper Triassic (Norian) (Kutassy, 1931). Cox (1952) erected *Prorotrigonia* and reported it from the Upper Triassic. Cox and others (1969) assigned it an Upper Triassic range. Krumbeck (1923a) described the type species from the Norian of Seram (Indonesia).

Paleogeographic distribution.—Tethys (Fig. 41). Hautmann (2001b) mentioned it from the Himalayas, as well as from southern Indonesia, but he did not refer to the original source. In addition, Tamura and Nishimura (1994) reported *Prorotrigonia* sp. from the Upper Triassic of Japan, but the figure of their specimen is of poor quality and it cannot be assigned with certainty to the genus. In fact, later, Tamura (1996) doubtfully recorded it as *Prorotrigonia* (?) sp. from the Upper Triassic of Japan.

Tethys domain: Late Triassic: Norian of Seram (Indonesia) (Krumbeck, 1923aSepmSepm; Cox, 1952).

Paleoautoecology.—B, Is, S, FM; Sb. See mode of life for *Trigonia*(p. 113).

Mineralogy.—Aragonitic (J. D. Taylor, Kennedy, & Hall, 1969). No details are known about *Prorotrigonia* shell microstructure. See discussion under *Trigonia* (p. 113).

Genus PROSOGYROTRIGONIA Krumbeck, 1924, p. 244

Type species.—Trigonia (Prosogyrotrigonia) timorensis Krumbeck, 1924, p. 245.

Stratigraphic range.—Upper Triassic (Norian)–Lower Jurassic (Sinemurian) (Hayami, 1975; Hautmann, 2001b). Cox and others (1969) assigned it an Upper Triassic range. Subsequent records extended the stratigraphic range of this genus to the Lower Jurassic (Hayami, 1975).

Paleogeographic distribution.—Tethys and Circumpacific (Fig. 41). Prosogyrotrigonia was also mentioned (*Prosogyrotrigonia*? sp.) from the Hettangian of northern Yukon (Canada) (Frebold & Poulton, 1977; Poulton, 1979), and new species were described from the Hettangian–Sinemurian of Chile (Pérez & others, 2008). It was also reported from Tibet (Kobayashi & Tamura, 1983a; Hautmann, 2001b), but these papers did not indicate the original source reference.

Tethys domain: Late Triassic: Norian–Rhaetian of Iran (Fallahi, Gruber, & Tichy, 1983; Hautmann, 2001b), Yunnan (China) (Guo, 1985); Rhaetian of Timor (Indonesia) (Krumbeck, 1924; Kobayashi & Mori, 1954a).

Circumpacific domain: Early Jurassic: Hettangian–Sinemurian of Chile (Pérez & others, 2008); Sinemurian of Japan (Yehara, 1921; Kobayashi & Mori, 1954a; Hayami, 1975; Sato & Westermann, 1991), Sonora (Mexico) (Scholz, Aberhan, & González-León, 2008).

Paleoautoecology.—B, Is, S, FM; Sb. See mode of life for *Trigonia* (p. 113).

Mineralogy.—Aragonitic (J. D. Taylor, Kennedy, & Hall, 1969). No details are known about *Prosogyrotrigonia* shell microstructure. See discussion under *Trigonia* (p. 113).

Genus GERATRIGONIA Kobayashi in Kobayashi & Mori, 1954a, p. 171

Type species .- Trigonia hosourensis Yokoyama, 1904, p. 11.

Stratigraphic range.—Lower Jurassic (Hettangian–Toarcian) (Hayami, 1975). Cox and others (1969) assigned it a Lower Jurassic (lower Lower Jurassic) range, but *Geratrigonia* had been reported from the Toarcian (Kobayashi, 1957). The genus was quite common in the Hettangian of Japan (Hayami, 1959, 1975).

Paleogeographic distribution.—Circumpacific (Fig. 41). Although we consider *Geratrigonia* to be a Japanese endemic genus, it was also mentioned from South America (Pérez & Reyes, 1991), specifically the species *Trigonia (Geratrigonia) kurumensis* Kobayashi, 1954, from the Bata Formation (Colombia), then dated as Lower Jurassic. The specimens were later reassigned to *Vaugonia niranohamensis santamariae* Geyer, 1973, and, moreover, the Bata Formation was redated as Cretaceous (Etayo Serna & others, 2003).

Circumpacific domain: Early Jurassic: Hettangian of Japan (Kobayashi & Mori, 1954a; Hayami, 1959, 1975; Sato & Westermann, 1991; Sugawara & Kondo, 2004; Kondo & others, 2006).

Paleoautoecology.—B, Is, S, FM; Sb. See mode of life for *Trigonia* (p. 113).

Mineralogy.—Aragonitic (J. D. Taylor, Kennedy, & Hall, 1969). No details are known about *Geratrigonia* shell microstructure. See discussion under *Trigonia* (p. 113).

Genus VAUGONIA Crickmay, 1930a, p. 53

Type species.—Vaugonia veronica Crickmay, 1930a, p. 53.

Stratigraphic range.—Lower Jurassic (Hettangian)–Upper Jurassic (Oxfordian) (Hayami, 1975). Crickmay (1932) mentioned Vaugonia from the Middle Jurassic. Subsequently, Kobayashi and Mori (1954b) proposed the new subgenus Hijitrigonia Kobayashi, 1954, from the Jurassic of Japan and indicated that Vaugonia had its origin during the Hettangian in Japan, and later it had a cosmopolitan distribution, probably extending to the Early Cretaceous. Cox and others (1969) considered V. (Hijitrigonia) as a junior synonym of V. (Vaugonia) and assigned a Jurassic range to the genus, including two subgenera: V. (Vaugonia) and V. (Orthotrigonia) Cox, 1952. No evidence of Vaugonia is found after the Jurassic. The youngest record is Oxfordian (Hayami, 1975).

Paleogeographic distribution.—Circumpacific (Fig. 41). Vaugonia originated in Japan during the Hettangian, and later it extended to the rest of the world. However, Francis and Hallam (2003) assumed a South American origin during the Sinemurian. Although during our study interval it was only recorded from the Circumpacific domain, since Pliensbachian times and throughout the Middle Jurassic, it had a cosmopolitan distribution (Fleming, 1964, 1987; Hallam, 1976; Poulton, 1976, 1979, 1991; Ishikawa & others, 1983; Pugaczewska, 1986; H. A. Leanza & Garate Zubillaga, 1987; H. A. Leanza, 1993; H. J. Campbell & Grant-Mackie, 1995; Kelly, 1995a). Circumpacific domain: Early Jurassic: Hettangian–Sinemurian of Japan (Kobayashi & Mori, 1954b; Hayami, 1975; Sato & Westermann, 1991; Sugawara & Kondo, 2004); Sinemurian of Nevada (United States) (Poulton, 1979), Peru (Pérez & Reyes, 1991), Chile (Pérez & others, 2008).

Paleoautoecology.—B, Is, S, FM; Sb. See mode of life for Trigonia. Mineralogy.—Aragonitic (J. D. Taylor, Kennedy, & Hall, 1969).
No details are known about Vaugonia shell microstructure. See discussion under Trigonia (p. 113).

Genus KYUSHUTRIGONIA Tamura & Nishimura, 1994, p. 15

Type species.—Kyushutrigonia hachibarensis Tamura & Nishimura, 1994, p. 18.

Stratigraphic range.—Upper Triassic (Carnian–Norian) (Tamura & Nishimura, 1994; see Onoue & Tanaka, 2005). *Kyushutrigonia* was proposed by Tamura and Nishimura (1994) from Japan (Sambosan Terrane). They indicated it was recorded in the Upper Triassic, but they did not provide the exact age of the association. Onoue and Tanaka (2005) reported an association from the same locality, with bivalves in common, and assigned it a Carnian–Norian age.

Paleogeographic distribution.—Circumpacific (Fig. 41).

Circumpacific domain: Late Triassic: Japan (Tamura & Nishimura, 1994; Tamura, 1996).

Paleoautoecology.—B, Is, S, FM; Sb. See mode of life for Trigonia. Mineralogy.—Aragonitic (J. D. Taylor, Kennedy, & Hall, 1969).
No details are known about Kyushutrigonia shell microstructure. See discussion under Trigonia (p. 113).

Genus ACANOMYPHORIA Guo, 1985, p. 203, 269

Type species.—Acanomyphoria tuberose Guo, 1985, p. 203.

Remarks.—Sichuantrigonia Gou, 1993, was placed in synonymy with *Acanomyophoria* by Z. Fang and others (2009).

Stratigraphic range.—Upper Triassic (Carnian) (Guo, 1985). Guo (1985) proposed Acanomyphoria from Carnian beds of the Weiyuanjiang Formation of Guanfangnabang in Yunnan (China). Gou (1993) proposed Sichuantrigonia for material from the upper member of Hanwang Formation of Maantang area in Jiangyou, Sichuan (China), which was dated as Carnian.

Paleogeographic distribution.—Eastern Tethys (Fig. 41).

Tethys domain: Late Triassic: Carnian of southwestern China (Guo, 1985; Gou, 1993).

Paleoautoecology.—B, Is, S, FM; Sb. See mode of life for *Trigonia* (p. 113).

Mineralogy.—Aragonitic (J. D. Taylor, Kennedy, & Hall, 1969). No details are known about *Acanomyphoria* shell microstructure. See discussion under *Trigonia* (p. 113).

Genus JAWORSKIELLA A. F. Leanza, 1942, p. 144, 166

Type species.—Trigonia burckhardti Jaworski, 1914, p. 299.

Remarks.—Although A. F. Leanza (1942) proposed it as a subgenus of *Trigonia*, Cox and others (1969) and subsequent authors regarded *Jaworskiella* as a separate genus. Later, Reyes and Pérez (1980) proposed a new subgenus, *Quadratojaworskiella* Reyes & Pérez, 1980, and subsequently raised it to generic rank (Pérez & others, 2008). Stratigraphic range.—Lower Jurassic (Hettangian–Pliensbachian) (H. A. Leanza, 1993; Pérez & others, 2008). Cox and others (1969) assigned it a Lower Jurassic (middle Liassic)–Upper Jurassic range, but we only found it recorded from the Lower Jurassic. Poulton (1979) already doubted that the genus was present in the Upper Jurassic, and H. A. Leanza (1993) restricted its range to the Lower Jurassic. It is especially abundant during the Pliensbachian (A. F. Leanza, 1942; H. A. Leanza, 1993; Poulton, 1979; H. A. Leanza & Garate Zubillaga, 1987).

Paleogeographic distribution.—Circumpacific and Austral (Fig. 41). Circumpacific domain: Early Jurassic: Hettangian–Sinemurian of Chile (Pérez & Reyes, 1991; Pérez & others, 2008); Sinemurian

of Nevada (United States) (Poulton, 1979).

Austral domain: Early Jurassic: Sinemurian of Argentina (Pérez & Reyes, 1991; Damborenea & Manceñido, 2005b).

Paleoautoecology.—B, Is, S, FM; Sb. See mode of life for Trigonia. Mineralogy.—Aragonitic (J. D. Taylor, Kennedy, & Hall, 1969).
No details are known about Jaworskiella shell microstructure. See discussion under Trigonia (p. 113).

Genus GUINEANA Skwarko, 1967, p. 59

Type species .- Guineana jimiensis Skwarko, 1967, p. 60.

Stratigraphic range.—Upper Triassic (Carnian–Norian) (Skwarko, 1967). When Skwarko (1967) proposed *Guineana*, he tentatively included in this genus other species distributed throughout the Norian and Rhaetian of western Europe, Asia, and Nevada. We are not taking them into account, as Skwarko simply indicated they were externally similar, but he did not study the hinge details of any of them.

Paleogeographic distribution.—Austral (Fig. 41). According to Damborenea (2002b), *Guineana* was endemic to the Australian province of the South Pacific domain.

Austral domain: Late Triassic: Carnian–Norian of New Guinea (Skwarko, 1967).

Paleoautoecology.—B, Is, S, FM; Sb. See mode of life for *Trigonia* (p. 113).

Mineralogy.—Aragonitic (J. D. Taylor, Kennedy, & Hall, 1969). No details are known about *Guineana* shell microstructure. See discussion under *Trigonia* (p. 113).

Genus FRENGUELLIELLA A. F. Leanza, 1942, p. 164

Type species .- Trigonia inexpectata Jaworski, 1915, p. 377.

Remarks.—A. F. Leanza (1942) proposed *Frenguelliella* as a subgenus of *Trigonia*, and this status was maintained by Cox and others (1969). Poulton (1979) decided to separate it from *Trigonia* and considered it to be a different genus due to the absence of radial ribs in the area. Furthermore, this author regarded *Kumatrigonia* Tamura, 1959, as a subgenus of *Frenguelliella*, as it was originally proposed. Cox and others (1969) considered *Kumatrigonia* as a subgenus of *Trigonia*. We follow Poulton (1979).

Stratigraphic range.—Upper Triassic (Carnian)–Middle Jurassic (Bajocian) (Hayami, 1975; H. A. Leanza, 1996). Cox and others (1969) assigned a Jurassic–Upper Cretaceous and Upper Triassic range to *Trigonia (Frenguelliella)* and *Trigonia (Kumatrigonia)*, respectively. The last refers to *Frenguelliella (Kumatrigonia) tanourensis* Tamura, 1959, from the Carnian of Japan (Hayami, 1975). *Frenguelliella* was

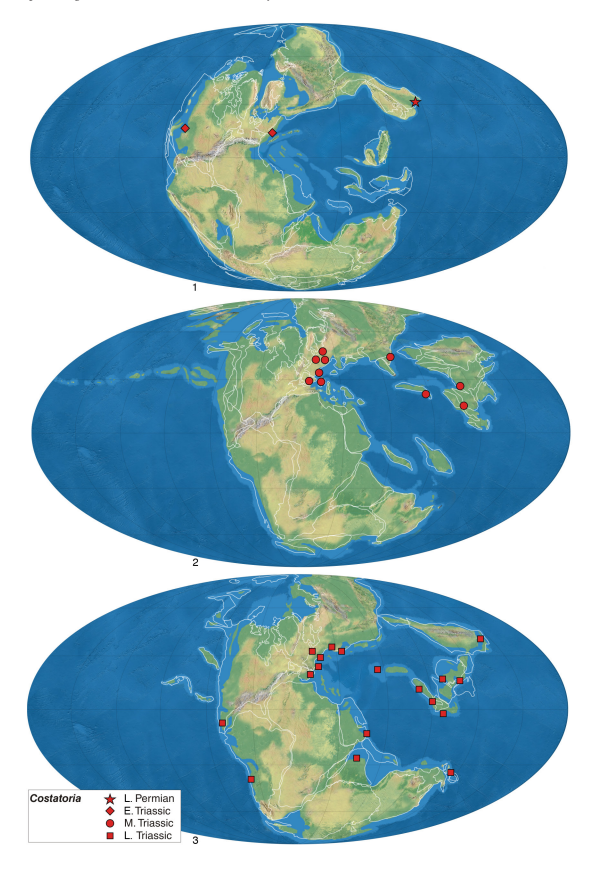


Figure 42. Paleogeographical distribution of Costatoriidae (Costatoria). 1, late Permian-Early Triassic; 2, Middle Triassic; 3, Late Triassic-Early Jurassic.

well distributed during the Jurassic, especially in the Pliensbachian (A. F. Leanza, 1942; Poulton, 1979; Ishikawa & others, 1983; H. A. Leanza & Garate Zubillaga, 1987; Pérez & Reyes, 1991; H. A. Leanza, 1993; Kelly, 1995a; Liu, 1995), but, although it was also recorded from the Upper Cretaceous in several papers (Cox & others, 1969; Poulton 1979; H. A. Leanza, 1993), we did not locate any species from deposits of that age. H. A. Leanza (1996) indicated that *Frenguelliella* was extinct by the Middle Jurassic (Bajocian), and its last species was *F. perezreyesi* H. A. Leanza 1993.

Paleogeographic distribution.—Circumpacific and Austral (Fig. 41). Although Cox and others (1969) considered it to be a cosmopolitan genus, the genus is primarily distributed on the Paleopacific margins. Pérez and Reyes (1991) recorded its presence in Europe, but no record from this area was found.

Circumpacific domain: Late Triassic: Carnian of Japan (Hayami, 1975); Norian of ?Oregon (United States) (Newton in Newton & others, 1987); Early Jurassic: Hettangian–Sinemurian of Texas (United States) (Liu, 1995); Sinemurian of northern Canada and Nevada (United States) (Poulton, 1979), Sonora (Mexico) (Scholz, Aberhan, & González-León, 2008), Peru (Pérez & Reyes, 1991), Chile (Pérez & others, 2008).

Austral domain: Early Jurassic: Sinemurian of Argentina (H. A. Leanza, 1993, 1996; Damborenea, 1996a; Damborenea & Mancenido, 2005b; Damborenea & Lanés, 2007).

Paleoautoecology.—B, Is, S, FM; Sb. See mode of life for *Trigonia* (p. 113).

Mineralogy.—Aragonitic (J. D. Taylor, Kennedy, & Hall, 1969). No details are known about *Frenguelliella* shell microstructure. See discussion under *Trigonia* (p. 113).

Family COSTATORIIDAE Newell & Boyd, 1995 Genus COSTATORIA Waagen, 1907, p. 149

Type species.—Donax costata Zenker, 1833, p. 55.

Stratigraphic range.-upper Permian-Upper Triassic (Rhaetian) (Nakazawa & Newell, 1968; Hautmann, 2001b). Cox and others (1969) reported it from lower Permian of Texas and Wyoming, from the upper Permian of Japan, and assigned it a Triassic cosmopolitan distribution. The lower Permian records are from Ciriacks (1963), with Costatoria sexraditata (Branson, 1930), which is the type species of Procostatoria Newell & Boyd, 1975. The genus is recorded from the upper Permian, with the species C. katsurensis Nakazawa, 1967 (Nakazawa & Newell, 1968; Hayami & Kase, 1977) and C. kobayasii (Kambe, 1957) (Nakazawa, 1960; Hayami, 1975; Hayami & Kase, 1977). Throughout the Triassic, it was recorded from various localities, from the Lower Triassic (Broglio-Loriga & Posenato, 1986) to the Rhaetian (Hautmann, 2001b). Many different species were recorded from the Tethys domain, sometimes based on biostratigraphic criteria; a revision and an evolutionary analysis of the group would be most interesting.

Paleogeographic distribution.—Tethys, Circumpacific, and Austral (Fig. 42). Although Cox and others (1969) regarded *Costatoria* as cosmopolitan during the Triassic, we did not find records from the Boreal domain.

Tethys domain: Early Triassic: Olenekian of Italy (Neri & Posenato, 1985; Broglio-Loriga & Posenato, 1986; Neri, Pasini, & Posenato, 1986; Posenato, 1989; Broglio-Loriga & others, 1990; Fraiser & Bottjer, 2007a); Middle Triassic: Germany (Hagdorn & Simon, 1985; Mahler & Sell, 1991), Poland (Senkowiczowa, 1985), Malaysia (Kobayashi & Tamura, 1968b); Anisian of southern China (Sha, Chen, & Qi, 1990; Tong & Liu, 2000; Komatsu, Chen, & others, 2004), Hungary (Szente, 1997), northern Vietnam (Komatsu, Huyen, & Huu, 2010); Ladinian of Italy (Márquez-Aliaga & Ros, 2002; Posenato, 2002; Márquez-Aliaga & others, 2004), Spain (Márquez-Aliaga, 1983, 1985; Márquez-Aliaga, Hirsch, & López-Garrido, 1986; Budurov & others, 1991; Márquez-Aliaga & Montoya, 1991; López-Gómez & others, 1994; Freneix, 1999; Márquez-Aliaga, García-Forner, & Plasencia, 2002; Márquez-Aliaga & Ros, 2003), northern Vietnam (Komatsu, Huyen, & Huu, 2010), Afghanistan (Farsan, 1972); Late Triassic: Malaysia (Tamura, 1996), China (Sha, Chen, & Qi, 1990; Gou, 1993), Oman (R. Hudson & Jefferies, 1961); Carnian of Italy (Bittner, 1895; Allasinaz, 1966; Márquez-Aliaga & Ros, 2002; Márquez-Aliaga & others, 2004), Spain (Martín-Algarra, Solé de Porta, & Márquez-Aliaga, 1995); Norian of Vietnam (Vu Khuc & Huyen, 1998; Hautmann, 2001b), Germany (Tichy & Schramm, 1983), Austria (Tichy, 1975), Armenia and northern Caucasus (Hautmann, 2001b); Norian-Rhaetian of Iran (Hautmann, 2001b); Rhaetian of Tibet (Hautmann & others, 2005), Myanmar and Sumatra (Hautmann, 2001b).

Circumpacific domain: late Permian: Japan (Nakazawa, 1960; Nakazawa & Newell, 1968; Hayami, 1975; Newell & Boyd, 1975; Hayami & Kase, 1977); Early Triassic: Olenekian of western United States (Fraiser & Bottjer, 2007a); Late Triassic: Carnian–Norian of Japan (Onoue & Tanaka, 2005); Norian of ?Sonora (Mexico) (Damborenea in Damborenea & González-León, 1997); Norian– Rhaetian of Chile (Chong & Hillebrandt, 1985; Hillebrandt, 1990; Pérez & Reyes, 1991).

Austral domain: Late Triassic: Carnian–Norian of New Guinea (Skwarko, 1967).

Paleoautoecology.—B, Is, S, SM; Sb. See mode of life for *Trigonia* (p. 113). Hautmann (2001b) attributed a mode of life similar to the rest of trigonoiods, but he argued that it would be a slow burrower due to the radial ornamentation and an undeveloped foot.

Mineralogy.—Aragonitic (J. D. Taylor, Kennedy, & Hall, 1969). The shells of all members of the superfamily Trigonioidea were probably entirely aragonitic, with a prismatic outer shell layer and a nacreous inner shell layer, as in the living species of *Neotrigonia* (J. D. Taylor, Kennedy, & Hall, 1969; Newell & Boyd, 1975). The last authors noted that *Costatoria harpa* (Münster) may have also had a cross-lamellar structure (see Newell & Boyd, 1975, fig. 8–9).

Family MYOPHORIIDAE Bronn, 1849 in 1848–1849 Genus MYOPHORIA Bronn, 1834 in 1834–1838, p. 54

Type species.—Trigonellites vulgaris Schlotheim, 1822 in 1822–1823, p. 192.

Stratigraphic range.—Middle Triassic (Anisian)–Upper Triassic (Rhaetian) (Newell & Boyd, 1975). Although Cox and others (1969) assigned it a Lower–Upper Triassic range, subsequently Newell and Boyd (1975) reported it only from the Middle and Upper Triassic. No reliable records were found from the Lower Triassic, and we follow Newell and Boyd (1975) in the range assigned. Many Trigonioidea genera were based on species originally described under *Myophoria*,

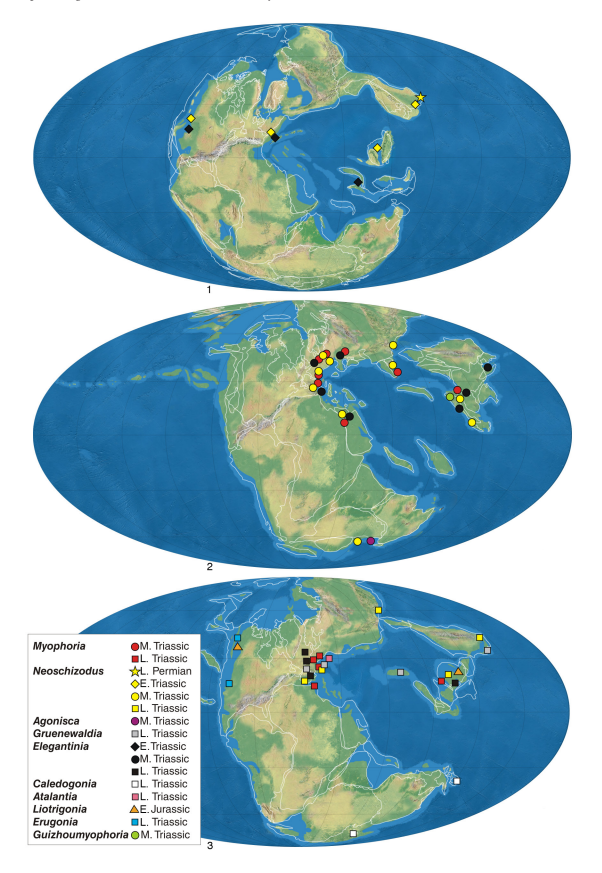


Figure 43. Paleogeographical distribution of Myophoriidae (*Myophoria, Neoschizodus, Agosnisca, Gruenewaldia, Elegantinia, Caledogonia, Atalantia, Liotrigonia, Erugonia, Guizhoumyophoria). 1,* late Permian–Early Triassic; 2, Middle Triassic; 3, Late Triassic–Early Jurassic.

so the stratigraphic and paleogeographic ranges of *Myophoria* may seem larger if a thorough revision of the species is not performed. *Myophoria* does not pass through the Triassic–Jurassic boundary (Hallam, 1981, 1990; Hallam & Wignall, 1997, 2000; Tanner, Lucas, & Chapman, 2004); it is a characteristic Triassic genus.

Paleogeographic distribution.—Tethys (Fig. 43).

Tethys domain: Middle Triassic: Germany (Bachmann, 1973; Baumgarte, 1975; Newell & Boyd, 1975; Fuchs & Mader, 1980; Hagdorn & Simon, 1983; Mahler & Sell, 1991), Poland (Senkowiczowa, 1985), Israel (Lerman, 1960); Anisian of Germany (Klug, Hagdorn, & Monterani, 2005), Spain (Budurov & others, 1991), China (Lu & Chen, 1986), Bulgaria (Encheva, 1969); Ladinian of Germany (Urlichs, 1978; Hagdorn, 1982; Klug, Hagdorn, & Monterani, 2005), Spain (Márquez-Aliaga, 1983, 1985; López-Gómez & others, 1994; Márquez-Aliaga & Martínez, 1996; Márquez-Aliaga & others, 2001, 2002), Italy (Fantini Sestini, 1966; Ürlichs & Tichy, 1998), Afghanistan (Farsan, 1972); Late Triassic: China (Gou, 1993), Yugoslavia (Newell & Boyd, 1975); Carnian of Italy (Fürsich & Wendt, 1977; Urlichs & Tichy, 1998), Slovenia (Boué, 1835, as Cryptina; Jelen, 1988; Jurkovsek, 1978, 1993); Carnian-Rhaetian of Tunisia (Desio, Rossi Ronchetti, & Vigano, 1960), the Alps (Kutassy, 1931); Rhaetian of Europe (Guérin-Franiatte, 1990).

Paleoautoecology.—B, Is, S, FM; Sb. See mode of life for *Trigonia* (p. 113).

Mineralogy.—Aragonitic (J. D. Taylor, Kennedy, & Hall, 1969). No details are known about *Myophoria* shell microstructure. See discussion under *Trigonia* (p. 113).

Genus NEOSCHIZODUS Giebel, 1855, p. 35

Type species.—Lyrodon laevigatum Goldfuss, 1837 in 1833–1841, p. 197.

Remarks.—Only one subgenus is considered in the study interval within *Neoschizodus: Okunominetania* Ichikawa, 1949.

Stratigraphic range.—middle Permian (Guadalupian)–Upper Triassic (Rhaetian) (Newell & Boyd, 1995; Márquez-Aliaga, Plasencia, & Ros, 2005). Cox and others (1969) assigned it a Permian–Upper Triassic range. Subsequently, Newell and Boyd (1975) and Boyd and Newell (1997) reported it from the middle and upper Permian of Japan, the Lower Triassic of Utah (United States), and the German Middle Triassic (Muschelkalk facies). However, *Neoschizodus* probably had a wider range, both stratigraphic and paleogeographic (see paleogeographic distribution). The youngest record is Rhaetian (Márquez-Aliaga, Plasencia, & Ros, 2005).

Paleogeographic distribution.—Cosmpolitan (Fig. 43).

Tethys domain: Early Triassic: China (S. Yang, Wang, & Hao, 1986); Induan of Italy (Fraiser & Bottjer, 2007a), southern China (Hautmann & others, 2011); Olenekian of Italy (Broglio-Loriga, Masetti, & Neri, 1982; Neri & Posenato, 1985; Posenato, 1989; Broglio-Loriga & others, 1990; Fraiser & Bottjer, 2007a); Middle Triassic: China (Ling, 1988), Poland (Senkowiczowa, 1985), Malaysia (Kobayashi & Tamura, 1968b), Israel (Lerman, 1960); lower Muschelkalk (?Anisian) of Germany (Newell & Boyd, 1975); Anisian of Germany (Klug, Hagdorn, & Monterani, 2005), Hungary (Szente, 1997), China (Sha, Chen, & Qi, 1990), northern Vietnam (Komatsu, Huyen, & Huu, 2010); Ladinian of Afghanistan (Farsan, 1972,

1975), Germany (Ürlichs, 1992; Márquez-Aliaga & others, 2002; Klug, Hagdorn, & Monterani, 2005), Spain (Márquez-Aliaga, 1983, 1985; Márquez-Aliaga & Montoya, 1991; Calvet & others, 1994; López-Gómez & others, 1994; Freneix, 1999; Márquez-Aliaga & Ros, 2003), China (Sha, Chen, & Qi, 1990), Italy (Rossi Ronchetti, 1959), northern Vietnam (Komatsu, Huyen, & Huu, 2010); Late Triassic: Carnian of China (Sha, Chen, & Qi, 1990), Italy (Allasinaz, 1962, 1966; Gelati & Allasinaz, 1964); Rhaetian of Italy (Pozzi, Gelati, & Allasinaz, 1962; Sirna, 1968), Spain (Márquez-Aliaga, Plasencia, & Ros, 2005).

Circumpacific domain: late Permian: Japan (Nakazawa, 1960; Nakazawa & Newell, 1968; Hayami & Kase, 1977); Early Triassic: Induan of western United States (Fraiser & Bottjer, 2007a); Olenekian of western United States (Newell & Boyd, 1975; Boyd & Newell, 1997; Schubert, 1993; Fraiser & Bottjer, 2007a), Japan (Nakazawa, 1961; Hayami, 1975; Kashiyama & Oji, 2004; Fraiser & Bottjer, 2007a); Late Triassic: Japan (Nakazawa, 1956; Tamura, 1990); Carnian of Japan (Hayami, 1975).

Austral domain: Middle Triassic: Anisian–Ladinian of New Zealand (Fleming, 1987).

Boreal domain: Middle Triassic: Ladinian of Primorie (Kiparisova, 1972); Late Triassic: of Primorie (Kiparisova, 1972).

Paleoautoecology.—B, Is, S, FM; Sb. See mode of life for *Trigonia* (p. 113).

Mineralogy.—Aragonitic (J. D. Taylor, Kennedy, & Hall, 1969). No details are known about *Neoschizodus* shell microstructure. See discussion under *Trigonia* (p. 113).

Genus AGONISCA Fleming, 1963, p. 843

Type species.—Agonisca corbiensis Fleming, 1963, p. 844.

Stratigraphic range.—Middle Triassic (Ladinian) (Fleming, 1987). Fleming (1963) proposed Agonisca from the Kaihikuan (=Ladinian–Carnian). Cox and others (1969) assigned it the same range. Although originally introduced into the family Trigoniidae, later Fleming (1987) included it in Myophoriidae and indicated its possible occurrence from Etalian (=Anisian–Ladinian) beds (?Agonisca aff. corbiensis Fleming), although the hinge of these specimens was incomplete, and the author pointed out the need for more material to indicate the correct relationships of this taxon.

Paleogeographic distribution.—Austral (Fig. 43).

Austral domain: Middle Triassic: Ladinian of New Zealand (Fleming, 1963, 1964, 1987).

Paleoautoecology.—B, Is, S, FM; Sb. See mode of life for *Trigonia* (p. 113).

Mineralogy.—Aragonitic (J. D. Taylor, Kennedy, & Hall, 1969). No details are known about *Agonisca* shell microstructure. See discussion under *Trigonia* (p. 113).

Genus GRUENEWALDIA Wöhrmann, 1889, p. 215

Type species.—Myophoria woehrmanni Bittner, 1895, p. 106.

Remarks.—Wöhrmann (1889) designated *Cardita decussata* Münster, 1837, in Goldfuss, 1833–1841, p. 185, as type species of *Gruenewaldia*. Bittner (1895) indicated that the identification of Wöhrmann's specimens was incorrect and gave a new name to them: *Myophoria woehrmanni* Bittner, 1895. Cox and others (1969) failed to take this into account, but Hautmann (2001b, 2003) pointed it out and, according to ICZN Article 70 (1999), he designated *Myophoria woehrmanni* Bittner as type species of *Gruenewaldia*, which he included within the family Myophoriidae. Although Cox and others (1969) regarded *Elegantinia* Waagen, 1907, as junior synonym of *Gruenewaldia*, *Elegantinia* is treated here as a valid genus (see discussion under *Elegantinia* below).

Stratigraphic range.—Upper Triassic (Carnian–Rhaetian). Cox and others (1969) assigned it a Middle–Upper Triassic range, but, as mentioned above, the type species designated by Wöhrmann (1889) was not *Cardita decussata* Münster, 1837, in Goldfuss, 1833–1841, from the Ladinian (Diener, 1923; Kutassy, 1931), but *Myophoria woehrmanni* Bittner, 1895, only reported from the Upper Triassic. Newell and Boyd (1975) mentioned the genus from the Middle Triassic of the Cassian Formation, which is currently considered to be Carnian in age (see Fürsich & Wendt, 1977).

Paleogeographic distribution.—Tethys and Circumpacific (Fig. 43). Hautmann (2001b) reported *Gruenewaldia* from Iran and also mentioned it from Yunnan, Tibet, and the Himalayas. Hallam (1981) listed it from the Arctic region, but we did not find any reference from that area.

Tethys domain: Late Triassic: Carnian of the Alps (Laube, 1865; Wöhrmann, 1889; Bittner, 1895; Allasinaz, 1966; Newell & Boyd, 1975; Fürsich & Wendt, 1977), Slovenia (Jelen, 1988); Norian de Iran (Repin, 2001); Norian–Rhaetian of Iran (Hautmann, 2001b, 2003).

Circumpacific domain: Late Triassic: Carnian of Japan (Hayami, 1975; Tamura & others, 1978); Carnian–Norian of Japan (Tamura & Nishimura, 1994; Onoue & Tanaka, 2002, 2005).

 Paleoautoecology.—B, Is, S, FM; Sb. See mode of life for Trigonia. Mineralogy.—Aragonitic (J. D. Taylor, Kennedy, & Hall, 1969).
 No details are known about Gruenewaldia shell microstructure. See discussion under Trigonia (p. 113).

Genus ELEGANTINIA Waagen, 1907, p. 149

Type species.—Lyrodon elegans Dunker, 1851, p. 300.

Remarks.—Cox and others (1969) treated *Elegantinia* as a junior synonym of *Gruenewaldia* Wöhrmann, 1889. However, a year earlier, Kobayashi and Tamura (1968b) had considered both valid and regarded *Lyriomyophoria* Kobayashi, 1954, as a junior synonym of *Elegantinia*, since both genera had the same type species. For years, the vast majority of western authors followed Cox and others (1969), without paying attention to the paper by Kobayashi and Tamura (1968b), until Boyd and Newell (1999) exposed the situation. Subsequently, Hautmann (2003) suggested the separation of *Elegantinia* and *Gruenewaldia* on the basis of hinge and ornamentation differences.

Stratigraphic range.—Lower Triassic (Olenekian)–Upper Triassic (Rhaetian) (Kobayashi & Tamura, 1968b). Kobayashi and Tamura (1968b) indicated a Lower–Upper Triassic (Rhaetian) range for *Elegantinia*, dismissing the Permian records as not belonging to *Elegantinia* (see Kobayashi & Tamura, 1968b, p. 104–105). Nevertheless, they were taken into account by Cox and others (1969) to assign the stratigraphic range to *Lyriomyophoria* [=*Elegantinia*]. Subsequently, several authors (Newell & Boyd, 1975; Boyd & Newell, 1997) regarded *Lyriomyophoria* [=*Elegantinia*] as an exclusively Triassic genus.

Paleogeographic distribution.—Tethys and Circumpacific (Fig. 43). Tethys domain: Early Triassic: Tonkin (Vietnam) (Patte, 1926), Italy (Leonardi, 1935); Middle Triassic: Israel (Lerman, 1960); Anisian of southern China (Komatsu, Chen, & others, 2004); lower Muschelkalk (=?Anisian) of Germany (Brinkmann, 1966; Busse, 1972; Baumgarte, 1975; Newell & Boyd, 1975), Hungary (Szente, 1997); Ladinian of Spain (Márquez-Aliaga, 1983, 1985; Pérez-López, 1991; López-Gómez & others, 1994), Vietnam (Vu Khuc & Huyen, 1998), Sinai (Awad, 1945), Alps (Arthaber, 1908); Late Triassic: Carnian of China (Wen & others, 1976; Sha, Chen, & Qi, 1990; Gou, 1993), Spain (Martín-Algarra & others, 1993; Márquez-Aliaga & Ros, 2002; Márquez-Aliaga & others, 2004); Rhaetian of England (Castell & Cox, 1975; Warrington & Ivimey-Cook, 1990; Ivimey-Cook & others, 1999), Hungary (Vörös, 1981).

Circumpacific domain: Early Triassic: Olenekian of Nevada (United States) (Boyd & Newell, 1997; Fraiser & Bottjer, 2007a); Middle Triassic: Ladinian or Carnian of Japan (Hayami, 1975).

Paleoautoecology.—B, Is, S, FM; Sb. See mode of life for *Trigonia* (p. 113).

Mineralogy.—Aragonitic (J. D. Taylor, Kennedy, & Hall, 1969). No details are known about *Elegantinia* shell microstructure. See discussion under *Trigonia* (p. 113).

Genus CALEDOGONIA Freneix & Avias, 1977, p. 282

Type species.—Caledogonia globosa Freneix & Avias, 1977, p. 286. Stratigraphic range.—Upper Triassic (upper Carnian–lower Norian) (Freneix & Avias, 1977). Caledogonia was reported from the Otamitan (=upper Carnian–lower Norian, according to Freneix & Avias, 1977; now Norian, see H. J. Campbell & Raine in Cooper, 2004) Later, Fleming (1987) treated it as a subgenus of Neoschizodus and reported it from the Norian of New Zealand.

Paleogeographic distribution.—Austral (Fig. 43). Caledogonia was endemic to the Maorian province (Freneix & Avias, 1977; Damborenea, 2002b).

Austral domain: Late Triassic: Carnian-Norian of New Caledonia (Freneix & Avias, 1977); Norian of New Zealand (Fleming, 1987).

Paleoautoecology.—B, Is, S, FM; Sb. See mode of life for *Trigonia* (p. 113). More specific information on the mode of life of *Caledogonia* is provided by Freneix and Avias (1977).

Mineralogy.—Aragonitic (J. D. Taylor, Kennedy, & Hall, 1969). No details are known about *Caledogonia* shell microstructure. See discussion under *Trigonia* (p. 113).

Genus ATALANTIA Termier & Verriez, 1974, p. 158

Type species.—Atalantia trigonioides Termier in Termier & Verriez, 1974, p. 158.

Stratigraphic range.—Upper Triassic (lower Carnian) (Termier & Verriez, 1974). Termier (in Termier & Verriez, 1974) proposed the monospecific genus *Atalantia* from the lower Carnian of southern Atalanti (Greece).

Paleogeographic distribution.—western Tethys (Fig. 43). *Atalantia* is a genus endemic to Greece.

Tethys domain: Late Triassic: early Carnian of Greece (Termier & Verriez, 1974).

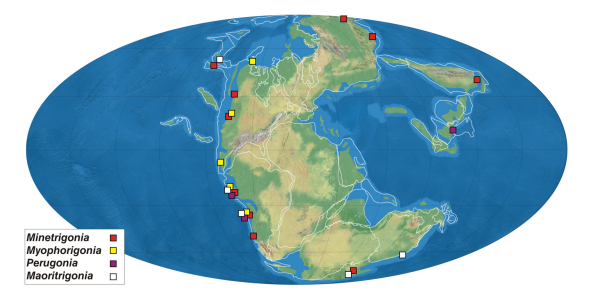


Figure 44. Paleogeographical distribution of Minetrigoniidae (Minetrigonia, Myophorigonia, Perugonia, Maoritrigonia). Late Triassic.

Paleoautoecology.—B, Is, S, FM; Sb. See mode of life for *Trigonia* (p. 113).

Mineralogy.—Aragonitic (J. D. Taylor, Kennedy, & Hall, 1969). No details are known about *Atalantia* shell microstructure. See discussion under *Trigonia* (p. 113).

Genus LIOTRIGONIA Cox, 1952, p. 53

Type species.—Trigonia lingonensis Dumortier, 1869 in 1864–1874, p. 275.

Stratigraphic range.—Lower Jurassic (?Hettangian, Sinemurian–Pliensbachian). Cox (1952) reported it from middle Lias of France and England. Cox and others (1969) assigned it an Lower Jurassic (middle Lower Jurassic) range. Subsequently, new records expanded the range of this genus, since it was quoted from ?Hettangian (Poulton, 1991), Sinemurian (Poulton, 1991; Stiller, 2006), and Pliensbachian (Hallam, 1976, 1977, 1987; Zakharov & others, 2006).

Paleogeographic distribution.—Tethys and Circumpacific (Fig. 43). Although during our study interval it was only known from China and Canada, during the Pliensbachian it was also present in France, England, and Siberia (Cox & others, 1969).

Tethys domain: Early Jurassic: Sinemurian of China (Stiller, 2006). Circumpacific domain: Early Jurassic: ?Hettangian–Sinemurian of Yukon (northwestern Canada) (Poulton, 1991).

Paleoautoecology.—B, Is, S, FM; Sb. See mode of life for *Trigonia* (p. 113).

Mineralogy.—Aragonitic (J. D. Taylor, Kennedy, & Hall, 1969). No details are known about *Liotrigonia* shell microstructure. See discussion under *Trigonia* (p. 113).

Genus ERUGONIA Newton in Newton & others, 1987, p. 63

Type species.—Erugonia canyonensis Newton in Newton & others, 1987, p. 65.

Stratigraphic range.—Upper Triassic (Norian) (Newton in Newton & others, 1987). Erugonia was only quoted from the Norian (Newton in Newton & others, 1987). In PBDB, it is also mentioned from the Rhaetian of Canada, but the review of the original source is pending.

Paleogeographic distribution.—Circumpacific (Fig. 43).

Circumpacific domain: Late Triassic: Norian of Oregon (United States) and British Columbia (western Canada) (Newton in Newton & others, 1987).

Paleoautoecology.—B, Is, S, FM; Sb. See mode of life for *Trigonia* (p. 113).

Mineralogy.—Aragonitic (J. D. Taylor, Kennedy, & Hall, 1969). No details are known about *Erugonia* shell microstructure. See discussion under *Trigonia* (p. 113).

Genus GUIZHOUMYOPHORIA Z. Fang, 2009, p. 998

Type species.—Quadratia quadrata H. Yin, 1974, p. 25.

Remarks.—Guizhoumyophoria was proposed by Z. Fang (2009) as a new name for *Quadratia* H. Yin, 1974, homonym of *Quadratia* Muir-Wood & Cooper, 1960, p. 161 (Brachiopoda: Productellidae). According to Z. Fang and others (2009, p. 56), "This genus is not very well established and all the material is poor."

Stratigraphic range.—Middle Triassic (Anisian) (H. Yin, 1974). Quadratia [Guizhoumyiophoria] was erected by H. Yin (1974) from the Anisian of China, and only the type species was included. The description was made in a restricted publication and in H. Yin in Gan and Yin (1978) was published again. Later, it was reported from the same stage by Komatsu, Chen, and others (2004). Kobayashi and Tamura (1983a) mentioned Quadrata H. Yin, 1974, by mistake and assigned a Middle–Upper Triassic range, but no original source of Upper Triassic records was given, so we are not taking it into account.

Paleogeographic distribution.—Eastern Tethys (Fig. 43).

Tethys domain: Middle Triassic: Anisian of southwestern China (Ghizou province) (H. Yin, 1974; Yin in Gan & Yin, 1978; Komatsu, Chen, & others, 2004; Z. Fang, 2009; Z. Fang & others, 2009).

Paleoautoecology.—B, Is, S, FM; Sb. See mode of life for *Trigonia* (p. 113).

Mineralogy.—Aragonitic (J. D. Taylor, Kennedy, & Hall, 1969). No details are known about *Guizhoumyophoria* shell microstructure. See discussion under *Trigonia* (p. 113).

Family MINETRIGONIIDAE Kobayashi, 1954 Genus MINETRIGONIA Kobayashi & Katayama, 1938, p. 187

Type species.—Trigonia hegiensis Saeki, 1925, p. 35.

Stratigraphic range.—Upper Triassic (Carnian–Rhaetian). Cox and others (1969) assigned it an Upper Triassic range. It is recorded from Carnian (Hayami, 1975) to Rhaetian (Fleming, 1987).

Paleogeographic distribution.—Circumpacific, Austral, and Boreal (Fig. 44). Kobayashi and Tamura (1968b, p. 108–109; 1983b) noticed that many Late Triassic species attributed to *Trigonia* and *Myophoria* actually belonged to *Minetrigonia*; these species were distributed through China, Peru, British Columbia, and New Zealand. Fleming (1987) documented the same situation for some of them known from the boreal area and Nevada (United States). *Minetrigonia* may also have been present in Malaysia (Kobayashi & Tamura, 1968b; Tamura & others, 1975).

Circumpacific domain: Late Triassic: Peru (Rangel, 1978); Carnian of Japan (Nakazawa, 1956; Hayami, 1975), Oregon (United States) (Tamura & McRoberts, 1993); Norian of British Columbia (Canada) (McLearn, 1946), Oregon (United States) (Newton in Newton & others, 1987); Norian or Rhaetian of Chile (Chong & Hillebrandt, 1985).

Austral domain: Late Triassic: Carnian of New Zealand (Trechmann, 1918); ?Norian, Rhaetian of New Zealand (Fleming, 1987); Rhaetian of ?Argentina (Riccardi & others, 1997, 2004; Damborenea & Manceñido, 2012).

Boreal domain: Late Triassic: Norian–Rhaetian of Siberia (Kiparisova, Bychkov, & Polubotko, 1966), northeastern Asia (Kurushin, 1990), Alaska (United States) (McRoberts & Blodgett, 2000).

Paleoautoecology.—B, Is, S, FM; Sb. See mode of life of *Trigonia* (p. 113).

Mineralogy.—Aragonitic (J. D. Taylor, Kennedy, & Hall, 1969). No details are known about *Minetrigonia* shell microstructure. See discussion under *Trigonia* (p. 113).

Genus MYOPHORIGONIA Cox, 1952, p. 52

Type species.—*Myophoria paucicostata* Jaworski, 1922, p. 126. *Stratigraphic range.*—Upper Triassic (Carnian–Rhaetian). Cox (1952) included several species in his new genus and reported it from Upper Triassic of Peru and Bear Islands, Inferior Oolite of England, and doubtfully from the Oxfordian of Syria. Cox and others (1969) assigned it a Upper Triassic–Middle Jurassic (Bajocian) range. Hallam (1981, 1990) indicated *Myophorigonia* did not survive the Triassic–Jurassic boundary, although Newton (1989) considered it to be a survivor of the Triassic–Jurassic extinction event. The oldest record during the Upper Triassic is from the Carnian in the Santa Clara Formation, Sonora (Mexico) (Alencaster de Cserna, 1961). According to Kobayashi and Tamura (1968b), Jurassic records of the genus refer to various species included by Cox (1952), but these need to be better studied to establish their relations. However, Kelly (1995a, p. 80, fig. 17) accepted *Myophorigonia* from Hettangian to Pliensbachian and Bathonian of South America, though these records are now referred to *Groeberella* (see Pérez, Reyes, & Damborenea, 1995). Since we cannot confirm the Lower Jurassic records of the genus, we accept the Rhaetian as the earliest solid record.

Paleogeographic distribution.—Circumpacific (Fig. 44). Although Hallam (1981) mentioned *Myophorigonia* from the Upper Triassic of the Arctic, we did not locate any earlier record from this area. However, Tamura and McRoberts (1993) transferred several Arctic species to *Myophorigonia*. Pérez and Reyes (1994) indicated its presence in the Upper Triassic of Singapore, referring to Kobayashi and Tamura (1968b), but, in this last paper, *Maoritrigonia* is recorded from that area but not *Myophorigonia*. G. D. Stanley and others (1994) and McRoberts (1997a) recorded *Myophorigonia jaworskii* (Steinmann, 1929) from the Norian of Peru, but this is the type species of *Perugonia* Kobayashi & Tamura, 1968a, and they made no comment in this respect.

Circumpacific domain: Late Triassic: Carnian of Sonora (Mexico), Oregon (United States) (Tamura & McRoberts, 1993), Bear Island (Cox, 1952; Kobayashi & Tamura, 1968b); Carnian–Norian of Peru (Körner, 1937); Norian of Chile (Hayami, Maeda, & Ruiz-Fuller, 1977), Sonora (Mexico) (G. D. Stanley & others, 1994; McRoberts, 1997a); Norian–Rhaetian of Peru (Cox, 1949, 1952; Pérez & Reyes, 1991), Chile (Chong & Hillebrandt, 1985; Pérez & Reyes, 1991).

Paleoautoecology.—B, Is, S, FM; Sb. See mode of life for *Trigonia* (p. 113).

Mineralogy.—Aragonitic (Hayami, Maeda, & Ruiz-Fuller, 1977). Hayami, Maeda, and Ruiz-Fuller (1977) described two aragonitic shell layers for *Myophorigonia* sp. aff. *M. paucicostata* (Jaworski, 1922): a prismatic outer layer and an alleged nacreous inner layer.

Genus PERUGONIA Kobayashi & Tamura, 1968a, p. 112

Type species.—Myophoria jaworskii Steinmann, 1929, p. 57.

Remarks.—Myophoria baertli Boit (1966, p. 10), from the same age and region as *M. jaworskii*, is the type species of *Albitrigonia* Prado-Velazco, 1991, p. 444. The relationships within this group of Norian Peruvian species are not well known, and a thorough systematic revision is needed. In the meantime, we regard *Albitrigonia* as a subgenus of *Perugonia*.

Stratigraphic range.—Upper Triassic (?Carnian-?Rhaetian). Perugonia was mainly reported from the Norian, and there are some doubts about whether its range extends to the Carnian (Kobayashi & Tamura, 1968b) or to the Rhaetian (Chong & Hillebrandt, 1985). Pérez and Reyes (1991) recorded *Perugonia* from all the three Upper Triassic stages of Peru and also from the Norian of Chile and Colombia.

Paleogeographic distribution.—Circumpacific (Fig. 44). G. D. Stanley and others (1994) and McRoberts (1997a) reported the type species of *Perugonia* (as *Myophorigonia jaworskii*) from the Norian of Peru. Perez and Reyes (1991) recorded it from the Carnian, Norian, and Rhaetian of Peru, but we have not located original information for the Carnian and Rhaetian.

Circumpacific domain: Late Triassic: Carnian or Norian of Malaysia (Kobayashi & Tamura, 1968b); Norian of Peru (Jaworski, 1922; Steinmann, 1929; Cox, 1949; Rangel, 1978; Maeda & oth-

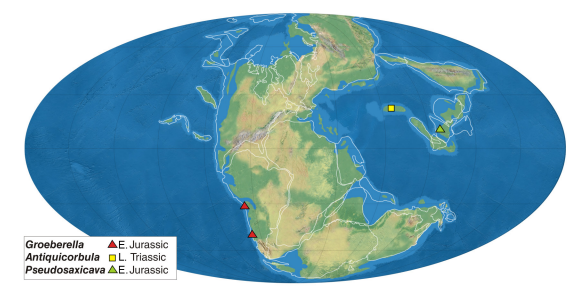


Figure 45. Paleogeographical distribution of Groeberellidae (Groeberella), Corbulidae (Antiquicorbula) and Hiatellidae (Pseudosaxicava). Late Triassic-Early Jurassic.

ers, 1983); Norian or Rhaetian of Chile (Jaworski, 1922; Chong & Hillebrandt, 1985; Hillebrandt, 1990).

Paleoautoecology.—B, Is, S, FM; Sb. See mode of life for *Trigonia* (p. 113).

Mineralogy.—Aragonitic (J. D. Taylor, Kennedy, & Hall, 1969). No details are known about *Perugonia* shell microstructure. See discussion under *Trigonia* (p. 113).

Genus MAORITRIGONIA Fleming, 1962, p. 3

Type species.—Myophoria nuggetensis Trechmann, 1918, p. 210. Stratigraphic range.—Upper Triassic (Carnian–Rhaetian) (Fleming, 1987). Cox and others (1969) assigned it a Carnian–Norian range, as did Fleming (1962), who also noticed possible records of *Maoritrigonia* from the Ladinian. Later, Fleming (1987) reported it from the Rhaetian.

Paleogeographic distribution.—Austral and Circumpacific (Fig. 44). Kobayashi and Tamura (1968b) recorded a possible record of the genus from the Upper Triassic of Malaysia, although they only doubtfully included *Myophoria bittneri* Newton, 1906, into *Maoritrigonia*. Fleming (1962, 1964) and Freneix and Avias (1977) considered *Maoritrigonia* to be endemic to the Maorian province, but later, Fleming (1987) and Pérez and Reyes (1994) reported it from the Upper Triassic of Chile. Damborenea (2002b) indicated that *Maoritrigonia* had a bipolar distribution. The only references we found from northern high latitudes are Polubotko and Repin (1990), who neither described nor illustrated the specimens, and McRoberts and Blodgett (2000).

Austral domain: Late Triassic: Carnian of New Zealand (Trechmann, 1918; Fleming, 1962, 1964, 1987), New Caledonia (Fleming, 1962); Carnian–Norian of New Caledonia (Freneix & Avias, 1977); Norian of New Zealand (Fleming, 1962, 1964, 1987), ?New Caledonia (Fleming, 1987); Rhaetian of New Zealand (Fleming, 1987).

Circumpacific domain: Late Triassic: ?Peru (Ishikawa & others, 1983; Maeda & others, 1983); Norian of southwestern Alaska

(McRoberts & Blodgett, 2000); Norian–Rhaetian of Chile (Fleming, 1987; Pérez & Reyes, 1991, 1994).

Paleoautoecology.—B, Is, S, FM; Sb. See mode of life for *Trigonia* (p. 113).

Mineralogy.—Aragonitic (J. D. Taylor, Kennedy, & Hall, 1969). No details are known about *Maoritrigonia* shell microstructure. See discussion under *Trigonia* (p. 113).

Family GROEBERELLIDAE Pérez, Reyes, & Damborenea, 1995 Genus GROEBERELLA H. A. Leanza, 1993, p. 18

Type species.—Myophoria neuquensis Groeber, 1924, p. 92.

Stratigraphic range.—Lower Jurassic (Sinemurian)–Middle Jurassic (Bajocian) (Pérez, Reyes, & Damborenea, 1995). H. A. Leanza (1993) proposed *Groeberella* and assigned it a Pliensbachian–Bajocian range, tentatively including it in the subfamily Minetrigoniinae. Subsequently, the genus was reported from the Sinemurian and referred to a new family, Groeberellidae, because although it had similarities with both Myophoriidae and Minetrigoniidae, it did not fit into any of them (Pérez, Reyes, & Damborenea, 1995). The genus was especially abundant during the Pliensbachian (H. A. Leanza, 1993, 1996; Pérez, Reyes, & Damborenea, 1995; Scholz, Aberhan, & González-León, 2008).

Paleogeographic distribution.—Austral and Circumpacific (Fig. 45). Austral domain: Early Jurassic: Sinemurian of Argentina (Pérez, Reyes, & Damborenea, 1995; Damborenea & Manceñido, 2005b; Damborenea & Lanés, 2007).

Circumpacific domain: Early Jurassic: Sinemurian of Chile (Pérez, Reyes, & Damborenea, 1995; Pérez & others, 2008).

Paleoautoecology.—B, Is, S, FM; Sb. See mode of life for *Trigonia* (p. 113).

Mineralogy.—Aragonitic (J. D. Taylor, Kennedy, & Hall, 1969). No details are known about *Groeberella* shell microstructure. See discussion under *Trigonia* (p. 113).

Superfamily MYOIDEA Lamarck, 1809 Family CORBULIDAE Lamarck, 1819 Genus ANTIQUICORBULA Hautmann, 2001b, p. 148

Type species.—Antiquicorbula concentrica Hautmann, 2001b, p. 150.

Stratigraphic range.—Upper Triassic (Norian–Rhaetian) (Hautmann, 2001b). *Antiquicorbula* was a monospecific genus only reported from the Upper Triassic of central Iran (Hautmann, 2001b).

Paleogeographic distribution.—Tethys (Fig. 45). Tethys domain: Late Triassic: Norian–Rhaetian of Iran (Haut-

mann, 2001b). *Paleoautoecology.*—B, Is, S, SM; Sb. Hautmann (2001b) compared

the external morphology of *Antiquicorbula* with the living genus *Corbula* Bruguière, 1797, assigning it the same mode of life. S. M. Stanley (1970) studied the behavior of *Corbula caribaea* d'Orbigny, proving that it is a slow, shallow burrower with very short siphons.

Mineralogy.—Aragonitic (J. D. Taylor, Kennedy, & Hall, 1973). There are no data about *Antiquicorbula* mineralogy and shell microstructure. The family Corbulidae is characterized by an entirely aragonitic shell with an outer shell layer of cross-lamellar structure and a complex cross-lamellar inner shell layer (J. D. Taylor, Kennedy, & Hall, 1973; Harper, Palmer, & Hudson, 2002).

Superfamily HIATELLOIDEA Gray, 1824 Family HIATELLIDAE Gray, 1824 Genus PSEUDOSAXICAVA Chavan, 1952, p. 119

Type species.—Pseudosaxicava bernardi Chavan, 1952, p. 119.

Remarks.—Chavan (1952) proposed *Pseudosaxicava* as genus, but later Cox and others (1969) regarded it as a subgenus under *Hiatella* Bosc, 1801, and assigned it an Upper Jurassic range. Many authors (e.g., Kelly, 1980; Vokes, 1980; Pisera, 1987; Fürsich, Palmer & Goodyear, 1994; Schneider & Kaim, 2012) followed Cox and others (1969), but Stiller (2006) regarded it as a genus after studying a rich hiatellid fauna from the Lower Jurassic of China.

Stratigraphic range.—Lower Jurassic (Hettangian)–Upper Jurassic (Tithonian) (Fürsich, Palmer, & Goodyear, 1994; Stiller, 2006). Chavan (1952) described the genus from the lower Kimmeridgian of Normandy (France), and, for a long time, *Pseudosaxicava* was thought to be restricted to the Upper Jurassic (Chavan, 1952; Cox & others, 1969; Kelly, 1980; Pisera, 1987; Wignall, 1990; Fürsich, Palmer, & Goodyear, 1994). However, Sepkoski (2002) assigned it a Middle–Upper Jurassic (upper Tithonian) range, following Fürsich, Palmer, and Goodyear (1994), who only mentioned it from the Tithonian. The Chinese specimens studied by Stiller (2006) show that it was already present in the earliest Jurassic. The top of its range is Tithonian (Portlandian; Fürsich, Palmer, & Goodyear, 1994).

Paleogeographic distribution.—Eastern Tethys (Fig. 45). Although during other times (mainly Late Jurassic), *Pseudosaxicava* was also known from the western Tethys (Chavan, 1952; Kelly, 1980; Pisera, 1987; Fürsich, Palmer, & Goodyear, 1994); during our study interval, it was only reported from the Eastern Tethys.

Tethys domain: Early Jurassic: Hettangian of southern China (Guangdong province) (Gu & others, 1976; J. Chen, 1982b; Stiller, 2006, and references therein); Hettangian–Sinemurian of southern China (Hunan province) (Stiller, 2006). *Paleoautoecology.*—B, E, S, Epi, Sed; By-N. Holocene species of *Hiatella* are chemical borers or byssate nestlers and are characterized by highly variable shell morphology, due to these life habits, with a high percentage of malformed shells. They can actively bore or reoccupy vacant borings built by other organisms or live epibyssate on several substrates, such as rocks, shells, and vegetation (Stiller, 2006). The same habits were suggested for Jurassic species (Kelly, 1980; Fürsich, Palmer, & Goodyear, 1994), although no evidence of boring activity was detected. The Lower Jurassic specimens from China do not show evidence of malformations or boring capacity, and they likely had an epibyssate or nestling mode of life (Stiller, 2006).

Mineralogy.—Aragonitic (J. D. Taylor, Kennedy, & Hall, 1973). There are no data about *Pseudosaxicava* mineralogy and shell microstructure. The Recent genus *Hiatella* is characterized by an entirely aragonitic shell with two shell layers of homogeneus microstructure (J. D. Taylor, Kennedy, & Hall, 1973; Brand & McCarthy, 2005).

Superfamily MEGALODONTOIDEA Morris & Lycett, 1853 in 1851–1855 Family MEGALODONTIDAE Morris & Lycett, 1853 in 1851–1855 Genus CONCHODON Stoppani, 1865 in 1860–1865, p. 246

Lycodus Schafhäutl, 1863, p. 375, non Quenstedt, 1856, p. 240 Type species.—Conchodon infraliassicus Stoppani, 1865 in 1860– 1865, p. 246 (=Lycodus cor Schafhäutl, 1863, p. 375).

Stratigraphic range.—Upper Triassic (?upper Norian, Rhaetian) (Mensink & Tichy, 1977). Cox and others (1969) assigned a Rhaetian range. Mensink and Tichy (1977) reported it from the Norian (=Sevatian) and Rhaetian, but these authors only described material from the Spanish Rhaetian and did not give any specific data from the Norian. Most authors considered *Conchodon* a typical Rhaetian genus, but Végh-Neubrandt (1982) indicated that *Conchodon hungaricus* (Hoernes) had a Norian range, although she expressed some doubts about the reliability of this age.

Paleogeographic distribution.—western Tethys (Fig. 46).

Tethys domain: Late Triassic: ?Norian of Hungary and Bakony (Végh-Neubrandt, 1982); Rhaetian of Poland, Hungary, Bakony, Bavaria, Italy, and Romania (Végh-Neubrandt, 1982), Austria (Végh-Neubrandt, 1982; Hallam & El Shaarawy, 1982), Spain (Mensink & Tichy, 1977; Végh-Neubrandt, 1982), Alps (Italy) (McRoberts, Newton, & Allasinaz, 1995).

Paleoautoecology.—B, E-Se, S, Un, Sed; R. Most megalodontids had very thick, heavy, large, and globose shells with peculiar morphologies. They tended to have the umbonal region thickened; in living position, this area was probably buried into the soft sediment (Posenato & Ietto, 1995). According to Skelton (1978), they may have been byssate in the earlier stages, but adults were reclined on the substrate, probably with part of the shell buried. Pedal muscle scars are present in earlier stages, but the foot was probably atrophied in adulthood. Many megalodontids had a gregarious mode of life. They diversified mainly in tropical environments in shallow, high-energy waters during the Late Triassic (Tichy, 1974; Allasinaz & Zardini, 1977; Freitas, Brunton, & Bernecker, 1993).

The possibility that megalodontids established symbiotic relationships with microorganisms was also postulated, but this is difficult to prove (Seilacher, 1990; Freitas, Brunton, & Bernecker,

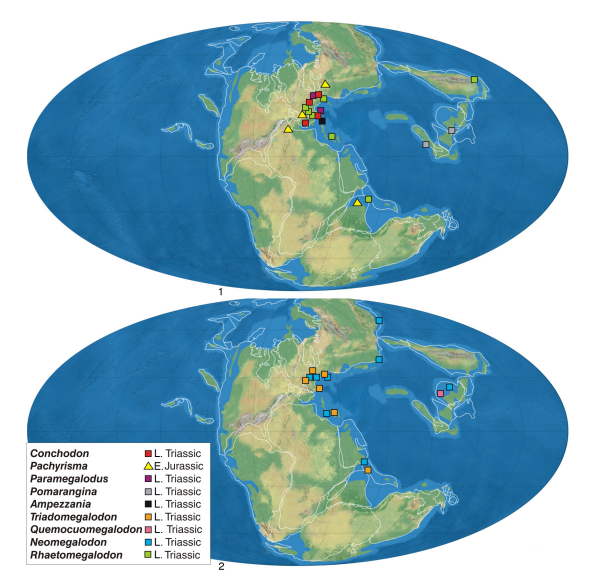


Figure 46. Paleogeographical distribution of Megalodontidae. Late Triassic–Early Jurassic. 1, Conchodon, Pachyrisma, Paramegalodus, Pomarangina, Ampezzania, Rhaetomegalodon; 2, Triadomegalodon, Quemocuomegalodon, Neomegalodon.

1993). The main criteria (see Seilacher, 1990, and Jones & Jacobs, 1992) are the following: they lived in tropical environments; their shells were abnormally large; they showed aberrant morphologies; they had epifaunal habits while coexisting with other infaunal bivalves; and many were gregarious. Although there no isotopic data to support these relationships, photosymbiosis cannot be rejected (see discussion in Freitas, Brunton, & Bernecker, 1993).

Mineralogy.—Aragonitic (Carter, Barrera, & Tevesz, 1998). No details are known about *Conchodon* shell microstructure. Members of the superfamily Megalodontoidea usually appear as molds or with recrystallized shells, which were interpreted as entirely aragonitic (Carter, Barrera, & Tevesz, 1998; Hautmann, 2006a).

Genus PACHYRISMA Morris & Lycett, 1850, p. 399

Type species.—Pachyrisma grande Morris & Lycett, 1850, p. 401.

Remarks.—Pachyrisma Morris & Lycett, 1850, along with *Protodiceras* G. Böhm, 1892 (see discussion in Genera not Included, p. 126), represented the recovery of megalodontids during the Early Jurassic, after extinction of all Triassic genera at the Rhaetian–Hettangian boundary. Some authors (e.g., Végh-Neubrandt, 1982) considered *Pachymegalodon* Gümbel, 1862, at generic level and left *Pachyrisma* as a Middle Jurassic genus. According to Végh-Neubrandt (1982, p. 134), *Protodiceras* arose from the *Dicerocardium* lineage and *Pachymegalodon* from *Triadomegalodon*.

Stratigraphic range.—Lower Jurassic (Sinemurian)–Upper Jurassic (Tithonian) (Cox & others, 1969). Cox and others (1969) considered three subgenera within *Pachyrisma: P. (Pachyrisma)* from the Middle and Upper Jurassic, *P. (Durga)* and *P. (Pachymegalodon)* from the Lower Jurassic (lower Lower Jurassic). After the interval considered in this paper, the genus was particularly abundant in the Pliensbachian (Hallam 1972, 1977; Tichy, 1974;

Buser & Debeljak, 1994; Liu, 1999; Fraser, Bottjer, & Fischer, 2004). However, it was also reported from the Sinemurian: *P.* (*Pachymegalodon*) from Morocco (Liu, 1995); *P.* (*Durga*) nicolisi Böhm, and *P.* (*Pachymegalodon*) chamaeformis (Schlotheim) from Europe (Hallam, 1976); *Pachyrisma* aff. chamaeformis Schlotheim from Pamir (Melnikova, 2006); *Pachymegalodon crassus* Böhm from the lower Lower Jurassic of Italy (Kennedy, Morris, & Taylor, 1970). None of these papers included Sinemurian specimens, but as we lack information about this genus, we provisionally accept them.

Paleogeographic distribution.—Tethys (Fig. 46).

Tethys domain: Early Jurassic: Sinemurian of Morocco (Liu, 1995), Europe (Kennedy, Morris, & Taylor, 1970; Hallam, 1976), Pamir (Melnikova, 2006).

Paleoautoecology.—B, E-Se, S, Un, Sed; R. See mode of life for Conchodon.

Mineralogy.—Aragonitic (Kennedy, Morris, & Taylor, 1970). Kennedy, Morris, and Taylor (1970) studied the shell microstructure of two species attributed to *Pachyrisma: Pachymegalodon crassus* (Böhm, 1884) and *Durga trigonalis* Böhm, 1884. The first shows two aragonitic layers: a homogeneous outer layer, and an inner layer with homogeneous to cross-lamellar structure. The second species has a fully recrystallized shell, but they described two aragonitic homogeneous layers.

Genus PARAMEGALODUS Cox in Cox and others, 1969, p. 746

Type species.—Dicerocardium eupalliatum Frech, 1904, p. 51. Remarks.---Végh-Neubrandt (1982) attributed Paramegalodus to Kutassy, 1953. This will not be followed here, as Kutassy's manuscript about megalodontids was never published (Végh-Neubrandt, 1969), and, although Kutassy (1934) designated a type species, he did not provide a diagnosis. The first diagnosis was done by Cox in Cox and others (1969, p. 746-747), and therefore he is considered to be the author of the genus. Végh-Neubrandt (1982) noticed that the figures in Cox and others (1969, p. 747, fig. E218a-c) belong to Dicerocardium eupalliatum (Vigh, 1914) and not to Dicerocardium eupalliatum Frech, 1904, as indicated in the diagnosis of Paramegalodus. Therefore, the figured specimens belong, in fact, to the type species of Rhaetomegalodon Végh-Neubrandt, 1969 (R. bajotensis Végh-Neubrandt, 1969 [=Megalodus? eupalliatus Vigh, 1914]) rather than to the type species of Paramegalodus [P. eupalliatum (Frech, 1904)].

Allasinaz in Allasinaz and Zardini (1977) found a great number of well-preserved specimens of *Paramegalodus* and offered a new generic diagnosis with clear diagnostic morphological characteristics and clarified the differences between this genus and *Rhaetomegalodon* Végh-Neubrandt, 1969. Indeed, Allasinaz (in Allasinaz & Zardini, 1977) included two new species in *Paramegalodus: P. prolatus* Allasinaz (in Allasinaz & Zardini, 1977), and *P. travenanzesis* Allasinaz (in Allasinaz & Zardini, 1977).

Stratigraphic range.—Upper Triassic (Norian) (Allasinaz & Zardini, 1977). Cox in Cox and others (1969) assigned it a Rhaetian range. Subsequently, Allasinaz in Allasinaz and Zardini (1977) extended its range back to the Norian. However, Végh-Neubrandt (1982) discussed that the Rhaetian specimens referred to *Paramegalodus* actually belong to *Rhaetomegalodon*. This genus was proposed by Végh-Neubrandt (1969) to group the species formerly included in *Paramegalodus* but which are really different from *P. eupalliatum* Frech, 1904 (see discussion in Végh-Neubrandt, 1969, or Végh-Neubrandt, 1982, p. 357). Following this author, records such as *P. belcheri* from the Rhaetian of Japan (Tamura, 1981) should be included in *Rhaetomegalodon*.

Allasinaz (in Allasinaz & Zardini, 1977) assigned a Norian-Rhaetian range to the type species of *Paramegalodus*. This Rhaetian record is based on Végh-Neubrandt (1960, fig. 34), but, later, Végh-Neubrandt (1982) included this specimen in the synonymy of *Rhaetomegalodon bajotiensis bajotiensis* Végh-Neubrandt, 1969 [=*Megalodus* (?) *eupalliatus sensu* Vigh (1914), *non* Frech (1904)], which is the type species of *Rhaetomegalodon*.

Paleogeographic distribution.—Tethys (Fig. 46).

Tethys domain: Late Triassic: Norian of Hungary (Frech, 1904), Italy (Frech, 1904; Allasinaz & Zardini, 1977; Végh-Neubrandt, 1982).

Paleoautoecology.—B, E-Se, S, Un, Sed; R. See mode of life for Conchodon (p. 125).

Mineralogy.—Aragonitic (Carter, Barrera, & Tevesz, 1998). No details are known about *Paramegalodus* shell microstructure. See discussion under *Conchodon* (p. 125).

Genus POMARANGINA Diener, 1908, p. 64

Type species.—Pomarangina haydeni Diener, 1908, p. 66.

Remarks.—Krumbeck (1914) described some specimens he referred to *P. haydeni*, including this species in the family Lucinidae, since he regarded it as being close to *Gonodon* Schafhäultl, 1863. He also figured the hinge of one specimen (plate XVII, fig. 37), which is very similar to the type species of *Schafhaeutlia* Cossman, 1897 (*pro Gonodon* Schafhäultl, 1863). Diener (1915) argued that Krumbeck (1914) misidentified his specimens and referred them to *Krumbeckia tambangensis* Diener, 1915, type species of *Krumbeckia* Diener, 1915, which is considered a synonym of *Schafhaeutlia* by Hautmann (2001b). Cox in Cox and others (1969) included *Pomarangina* in the family Megalodontidae with doubts and indicated that the hinge and muscles of this genus were unknown. Végh-Neubrandt (1982) did not refer to *Pomarangina* in her monograph of Triassic megalodontids, and no other systematic discussion was published.

Stratigraphic range.—Upper Triassic (Carnian). Cox and others (1969) assigned it a Upper Triassic range. The genus was reported from the Carnian (Diener, 1908; Krumbeck, 1914). Kobayashi and Tamura (1983a) mentioned it from the Upper Triassic of Sichuan, but they did not specify the stage.

Paleogeographic distribution.—Eastern Tethys (Fig. 46).

Tethys domain: Late Triassic: Sichuan (China) (Kobayashi & Tamura, 1983a); Carnian of the Himalayas (Diener, 1908).

Paleoautoecology.—B, E-Se, S, Un, Sed; R. The diagnosis offered by Cox and others (1969) indicates that *Pomarangina* was externally similar to other megalodontids. See mode of life for *Conchodon* (p. 125).

Mineralogy.—Aragonitic (Carter, Barrera, & Tevesz, 1998). No details are known about *Pomarangina* shell microstructure. See discussion under *Conchodon* (p. 125).

Genus AMPEZZANIA

Allasinaz in Allasinaz & Zardini, 1977, p. 121

Type species.—Ampezzania zardinii Allasinaz in Allasinaz & Zardini, 1977, p. 122.

Stratigraphic range.—Upper Triassic (Norian) (Allasinaz & Zardini, 1977). The genus was proposed by Allasinaz (in Allasinaz & Zardini, 1977), including two species, *A. zardinii* Allasinaz (in Allasinaz & Zardini, 1977), and *A. subovata* Allasinaz (in Allasinaz & Zardini, 1977), both reported from the Norian.

Paleogeographic distribution.—western Tethys (Fig. 46).

Tethys domain: Late Triassic: Norian of Cortina d'Ampezzo (Italy) (Allasinaz in Allasinaz & Zardini, 1977).

Paleoautoecology.—B, E-Se, S, Un, Sed; R. See mode of life for *Conchodon* (p. 125).

Mineralogy.—Aragonitic (Carter, Barrera, & Tevesz, 1998). No details are known about *Ampezzania* shell microstructure. See discussion under *Conchodon* (p. 125).

Genus TRIADOMEGALODON Végh-Neubrandt, 1974, p. 10

Type species.—Megalodus damesi Hoernes, 1880, p. 121.

Stratigraphic range.—Upper Triassic (Carnian–Rhaetian) (Allasinaz & Zardini, 1977). According to Allasinaz (in Allasinaz & Zardini, 1977), *Triadomegalodon* had a Norian–Rhaetian range. Subsequently, Végh-Neubrandt (1982) reported the genus from Carnian deposits in several localities.

Paleogeographic distribution.—Tethys, ?Circumpacific (Fig. 46). Triadomegalodon was also reported from the Upper Triassic of Japan (Tamura, 1990), but the figured specimens are not conclusive. Yancey and Stanley (1999) reported it from the Norian of the western coast of North America, but this datum is dubious, since, according to these authors, the genus was mentioned by Végh-Neubrandt (1982, p. 204), where the species ?*Triadomegalodon canadensis* (Shimer, 1926), from the Norian of British Columbia (Canada), is referred to an uncertain genus.

Tethys domain: Late Triassic: Carnian of Italy, Slovenia, and Austria (Végh-Neubrandt, 1982); Norian of Italy (Végh-Neubrandt & others, 1976; Allasinaz & Zardini, 1977; Végh-Neubrandt, 1982), Austria (Végh-Neubrandt, 1982), Hungary and Turkey (Végh-Neubrandt & others, 1976), Yugoslavia, Hungary, Bulgaria, and Romania (Végh-Neubrandt, 1982); Rhaetian of Hungary and Turkey (Végh-Neubrandt, 1982), Italy (Végh-Neubrandt, 1982; McRoberts, Newton, & Allasinaz, 1995; Posenato & Ietto, 1995), Austria (Végh-Neubrandt & others, 1976; Végh-Neubrandt, 1982), ?India (Végh-Neubrandt, 1982).

Paleoautoecology.—B, Se, S-Ph, Un, Sed; R. The shell of *Triado-megalodon* is thick and heavy, especially in the dorsal region, and the ventral elongation increased during ontogeny (Posenato & Ietto, 1995). These features are interpreted as adaptations to a semi-infaunal mode of life, the shell resting with its umbonal region buried into the soft substrate and the ventral part exposed (Posenato & Ietto, 1995). In the adult stage, the foot was probably atrophied and the bivalve was sedentary. A symbiotic relationships with zooxanthellae is also very likely (Seilacher, 1990; McRoberts, Newton, & Allasinaz, 1995).

Mineralogy.—Aragonitic (Carter, Barrera, & Tevesz, 1998). No details are known about *Triadomegalodon* shell microstructure. See discussion under *Conchodon* (p. 125).

Genus QUEMOCUOMEGALODON H. Yao, Sha, & Zhang in H. Yao & others, 2003, p. 395, 399

Type species.—*Quemocuomegalodon orientus* H. Yao, Sha, & Zhang in H. Yao & others, 2003, p. 396.

Stratigraphic range.—Upper Triassic (Norian) (H. Yao & others, 2007). H. Yao and others (2003) proposed the genus from Upper Triassic beds of western China, associated with *Palaeocardita*, gastropods, and algae. Later, H. Yao and others (2007) described a richer associated fauna, which included other bivalves and ammonoids and allowed them to date the sediments where *Quemocuomegalodon* was found as Norian. *Quemocuomegalodon* includes three species: *Q. orientus* H. Yao, Sha, & Zhang in H. Yao & others, 2003, *Q. longitatus* H. Yao, Sha, & Zhang in H. Yao & others, 2003, and *Q. circularis* H. Yao & others, 2007.

Paleogeographic distribution.—Eastern Tethys (Fig. 46).

Tethys domain: Late Triassic: Norian of western China (H. Yao & others, 2003, 2007).

Paleoautoecology.—B, E-Se, S, Un, Sed; R. See mode of life for Conchodon (p. 125).

Mineralogy.—Aragonitic (Carter, Barrera, & Tevesz, 1998). No details are known about *Quemocuomegalodon* shell microstructure. See discussion under *Conchodon* (p. 125).

Genus NEOMEGALODON Gümbel, 1862, p. 362

Type species.—Cardium triquetrum Wulfen, 1793, p. 48.

Remarks.—Although Cox and others (1969) considered *Neomegalodon* as a subgenus of *Megalodon*, following Gümbel (1862), Allasinaz (1965) raised it to generic level and included three subgenera: *N. (Neomegalodon)*, *N. (Rossiodus)* Allasinaz, 1965, and *N. (Gemmellarodus)* Di Stefano, 1912. Subsequently, both *Gemmellarodus* and *Rossiodus* were raised to generic level by Allasinaz (in Allasinaz & Zardini, 1977), and followed by Végh-Neubrandt (1982). Neither of these two taxa are included in this study (see discussion in Genera not Included, p. 161, 170). Like most authors, we regard *Neomegalodon* as a distinct genus (Végh-Neubrandt & others, 1976; Allasinaz & Zardini, 1977; Tichy, 1980a, 1980b; Végh-Neubrandt, 1982; Tichy & Schramm, 1983).

Stratigraphic range.—Upper Triassic (Carnian–Rhaetian) (Cox & others, 1969; Allasinaz & Zardini, 1977). *Neomegalodon* was reported exclusively from the Upper Triassic (Cox & others, 1969). There is a record of the genus from Ladinian beds of western North America (Fraser, 1997), but the specimens were not described, and the figures provided are inconclusive.

Paleogeographic distribution.—Tethys and Boreal (Fig. 46). Cox and others (1969) regarded it a cosmopolitan genus, but we only found it recorded in the Tethys domain, the same distribution provided by Allasinaz and Zardini (1977). Damborenea (2002b) indicated that it was never found in South America.

Tethys domain: Late Triassic: Hungary (Frech, 1904); Carnian of the Alps (Austria) (Zapfe, 1972; Tichy, 1980a, 1980b; Végh-Neubrandt, 1982), Italy (Allasinaz, 1965; Allasinaz & Zardini, 1977; Végh-Neubrandt, 1982), Hungary, Bavaria, Slovenia, Slovakia, and India (Végh-Neubrandt, 1982); Carnian–Norian of Slovakia (Bujnovsky & Kochanová, 1973);

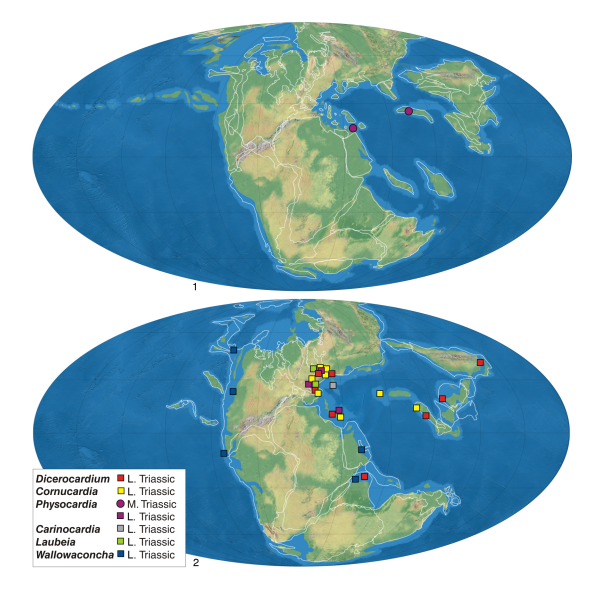


Figure 47. Paleogeographical distribution of Dicerocardiidae (*Dicerocardium, Cornucardia, Physocardia, Carinocardia, Laubeia*) and Wallowaconchidae (*Wallowaconcha*). 1, Middle Triassic; 2, Late Triassic.

Norian of western China (H. Yao & others, 2007), Afghanistan (Polubotko, Payevskaya, & Repin, 2001), Alps (Austria) (Tichy, 1975; Végh-Neubrandt, 1982; Tichy & Schramm, 1983), Italy (Allasinaz & Zardini, 1977; Végh-Neubrandt, 1982), Hungary, Slovenia, Albania, Turkey, Yugoslavia, and India (Végh-Neubrandt, 1982), Turkey (Végh-Neubrandt & others, 1976; Végh-Neubrandt 1982); Rhaetian of Austria (Cox & others, 1969; Végh-Neubrandt, 1982).

Boreal domain: Late Triassic: Carnian of Primorie (Kiparisova, 1972).

Paleoautoecology.—B, E-Se, S, Un, Sed; R. See mode of life for *Conchodon* (p. 125).

Mineralogy.—Aragonitic (Carter, Barrera, & Tevesz, 1998). No details are known about *Neomegalodon* shell microstructure. See discussion under *Conchodon* (p. 125).

Genus RHAETOMEGALODON Végh-Neubrandt, 1969, p. 121

Type species.—Rhaetomegalodon bajotensis Végh-Neubrandt, 1969, p. 123; *=Megalodus ? eupalliatus sensu* Vigh, 1914, p. 603, *non sensu* Frech, 1904.

Remarks.—Végh-Neubrandt (1969) proposed the genus *Rhaeto-megalodon* to group many of the Rhaetian species formerly included in *Paramegalodus*, since she considered them to be different from the type species of *Paramegalodus* and useful biostratigraphically. Allasinaz and Zardini (1977) indicated that Végh-Neubrandt (1969) did not give a precise diagnosis and warned that, given the similarities between *Paramegalodus* and *Rhaetomegalodon*, this could be considered a subgenus of the first (see Allasinaz & Zardini, 1977, p. 46, 109, 117). However, these authors offered a number of differences to distinguish *Rhaetomegalodon* at generic level: "In realtà

questi due generi si differenziano tra loro solo per un diverso modo d'incurvmento dgli umboni: essi sono prima piegati all'esterno e poi verso l'interno in *Rhaetomegalodon*, incurvati prima verso l'avanti e poi con apici tendenti a ruotare verso l'esterno in *Paramegalodus.*" Zapfe (1969) discussed the genus extensively and established the type species but did not give a proper diagnosis of the genus (Allasinaz & Zardini, 1977, p. 117).

Stratigraphic range.—Upper Triassic (Norian–Rhaetian) (Allasinaz & Zardini, 1977). The genus was proposed by Végh-Neubrandt (1969) from the Rhaetian. Subsequently, it was also reported from the Norian (Allasinaz & Zardini, 1977).

Paleogeographic distribution.—western Tethys and Circumpacific (Fig. 46).

Tethys domain: Late Triassic: Norian of Italy (Allasinaz & Zardini, 1977; Végh-Neubrandt, 1982); Rhaetian of Hungary (Végh-Neubrandt, 1969, 1982), Bakony, Poland, and India (Végh-Neubrandt, 1982), Austria (Zapfe, 1969; Végh-Neubrandt, 1982), Turkey (Végh-Neubrandt & others, 1976; Végh-Neubrandt, 1982).

Circumpacific domain: Late Triassic: Rhaetian of Japan (Tamura, 1981.

Paleoautoecology.—B, E-Se, S, Un, Sed; R. See mode of life for *Conchodon* (p. 125).

Mineralogy.—Aragonitic (Carter, Barrera, & Tevesz, 1998). No details are known about *Rhaetomegalodon* shell microstructure. See discussion under *Conchodon* (p. 125).

Family DICEROCARDIIDAE Kutassy, 1934 Genus DICEROCARDIUM Stoppani, 1865 in 1860–1865, p. 248

Type species.—Dicerocardium jani Stoppani, 1865 in 1860–1865, p. 249.

Stratigraphic range.—Upper Triassic (upper Carnian–Rhaetian). Cox and others (1969) assigned it a Norian–Rhaetian range, but later the genus was reported from upper Carnian beds of several localities (Allasinaz & Zardini, 1977; Tamura, 1981; Végh-Neubrandt, 1982).

Paleogeographic distribution.—Tethys and Circumpacific (Fig. 47). Tethys domain: Late Triassic: ?Indonesia (Krumbeck, 1924); Carnian–Norian of Italy (Allasinaz & Zardini, 1977; Végh-Neu-

brandt, 1982); Norian of Hungary, Austria, India, and Slovenia (Végh-Neubrandt, 1982), Turkey (Végh-Neubrandt & others, 1976; Végh-Neubrandt, 1982); Rhaetian of the Alps (Zapfe, 1963, 1981), Himalayas (Stoliczka, 1866, 1871 in 1870–1871; Cox & others, 1969), ?India (Végh-Neubrandt, 1982).

Circumpacific domain: Late Triassic: Carnian–Norian of Japan (Tamura, 1981); Norian of Japan (Tamura, 1990).

Paleoautoecology.—B, E-Se, S-Ph, Un, Sed; R. Dicerocardium is morphologically similar to the living Corculum, which establishes photosymbiotic relationships with algae, and it was proposed that Dicerocardium may also have done this (Seilacher, 1990). The inferred life position of Dicerocardium was very similar to Wallowaconcha Yancey & Stanley, 1999, lying on the substrate with the so-called wings horizontally and in contact with it, maybe somewhat buried.

Mineralogy.—Aragonitic (Carter, Barrera, & Tevesz, 1998). No details are known about *Dicerocardium* shell microstructure. See discussion under *Conchodon* (p. 125).

Genus CORNUCARDIA Koken, 1913, p. 34

Type species.—Craspedodon hornigii Bittner, 1901c, p. 8.

Stratigraphic range.—Upper Triassic (Carnian) (Broglio-Loriga, Ietto, & Posenato, 1993). *Cornucardia* was restricted to the Carnian (Cox & others, 1969).

Paleogeographic distribution.—Tethys (Fig. 47).

Tethys domain: Late Triassic: Iran (Hautmann, 2001b); Carnian of Timor (Indonesia) (Krumberck, 1924; Végh-Neubrandt, 1982), Italy (Leonardi, 1943; Rau & Tongiorgi, 1966; Corazzari & Lucchi-Garavello, 1980; Végh-Neubrandt, 1982; Broglio-Loriga, Ietto, & Posenato, 1993), southern Alps (Bittner, 1901b), northern Alps (Austria) (Zapfe, 1972; Végh-Neubrandt, 1982), Germany (Tichy, 1980a), Hungary (Tichy, 1980a; Végh-Neubrandt, 1982), Slovenia (Jurkovsek, 1978), Turkey (Végh-Neubrandt & others, 1976; Végh-Neubrandt, 1982), Yugoslavia (Végh-Neubrandt, 1982).

Paleoautoecology.—B, E-Se, S, Un, Sed; R. *Cornucardia* had a heavy shell, strongly inflated and coiled umbos, and it most likely lived reclining on the substrate, like most members of the superfamily Megalodontoidea. Since the shell was usually large and heavy, it is possible that it could be partially buried.

Mineralogy.—Aragonitic (Carter, Barrera, & Tevesz, 1998). No details are known about *Cornucardia* shell microstructure. See discussion under *Conchodon* (p. 125).

Genus PHYSOCARDIA Wöhrmann, 1894, p. 671

Type species.—Physocardia ogilviae Wöhrmann, 1894, p. 672.

Stratigraphic range.—Middle Triassic (Anisian)–Upper Triassic (Carnian) (Fantini Sestini, 1984). Cox and others (1969) reported it from the Carnian of the Alps. Subsequently, it was also recorded from the Middle Triassic (middle Anisian) (Fantini Sestini, 1984). This author referred to an Iranian specimen, morphologically very similar to specimens studied in that paper, which could be the oldest of the genus.

Paleogeographic distribution.—western Tethys (Fig. 47).

Tethys domain: Middle Triassic: Anisian of Turkey and Iran (Fantini Sestini, 1984); ?Ladinian of Turkey (Végh-Neubrandt, 1982); Late Triassic: Carnian of the Alps (Frech, 1904; Fürsich & Wendt, 1977; Végh-Neubrandt, 1982), Dolomites (Italy) (Végh-Neubrandt, 1982); Turkey (Végh-Neubrandt & others, 1976; Végh-Neubrandt, 1982).

Paleoautoecology.—B, E-Se S, Un, Sed; R. Similar to Cornucardia. Mineralogy.—Aragonitic (Carter, Barrera, & Tevesz, 1998). No details are known about *Physocardia* shell microstructure. See discussion under Conchodon (p. 125).

Genus CARINOCARDIA Termier & Verriez, 1974, p. 161

Type species.—Carinocardia atalantiensis Termier & Verriez, 1974, p. 161.

Stratigraphic range.—Upper Triassic (lower Carnian) (Termier & Verriez, 1974). Carinocardia was proposed by Termier and Verriez

(1974) from the Upper Triassic (lower Carnian) of Atalanti (Greece). *Paleogeographic distribution.*—western Tethys (Fig. 47).

Tethys domain: Late Triassic: Carnian of Greece (Termier & Verriez, 1974).

Paleoautoecology.-B, E-Se, S, Un, Sed; R. Similar to Cornucardia.

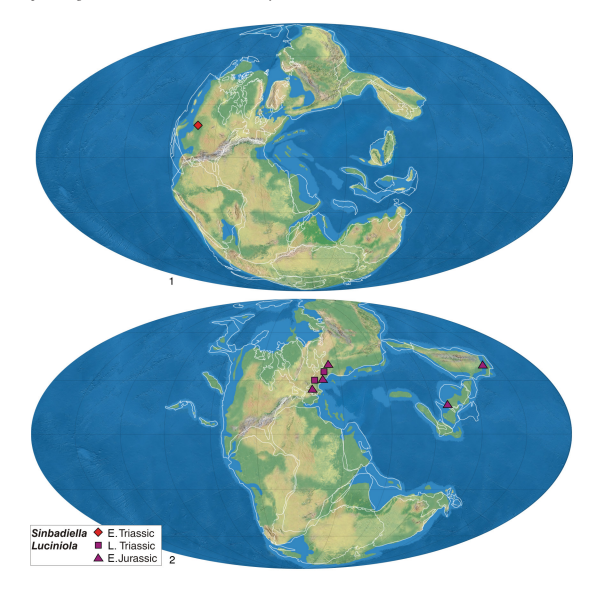


Figure 48. Paleogeographical distribution of Lucinidae (Sinbadiella, Luciniola). 1, Early Triassic; 2, Late Triassic-Early Jurassic.

Mineralogy.—Aragonitic (Carter, Barrera, & Tevesz, 1998). No details are known about *Carinocardia* shell microstructure. See discussion under *Conchodon* (p. 125).

Genus LAUBEIA Bittner, 1895, p. 26

Type species.—Cardita strigilata Klipstein, 1845 in 1843–1845, p. 35.

Remarks.—Allasinaz (1966) included *Laubeia* in the family Megalodontidae. Later, Cox and others (1969) argued that its features did not even fit in the class Bivalvia. Hallam (1981) indicated that *Laubeia* was a bivalve but belonged to an uncertain order. Finally, Végh-Neubrandt (1982) decided to include this monospecific genus in the family Dicerocardiidae.

Stratigraphic range.—Upper Triassic (Carnian) (Cox & others, 1969). Laubeia was a monospecific genus and had a Carnian

range (Bittner, 1895; Allasinaz, 1966; Cox & others, 1969; Végh-Neubrandt, 1982).

Paleogeographic distribution.—western Tethys (Fig. 47).

Tethys domain: Late Triassic: Carnian of southern Tirol (Austria) (Bittner, 1895), southern Alps (Italy) (Allasinaz, 1966; Végh-Neubrandt, 1982).

Paleoautoecology.—B, E-Se, S, Un, Sed; R. Similar to Cornucardia. Mineralogy.—Aragonitic (Carter, Barrera, & Tevesz, 1998). No details are known about Laubeia shell microstructure. See discussion under Conchodon (p. 125).

Family WALLOWACONCHIDAE Yancey & Stanley, 1999 Genus WALLOWACONCHA Yancey & Stanley, 1999, p. 7

Type species.—Wallowaconcha raylenea Yancey & Stanley, 1999, p. 10.

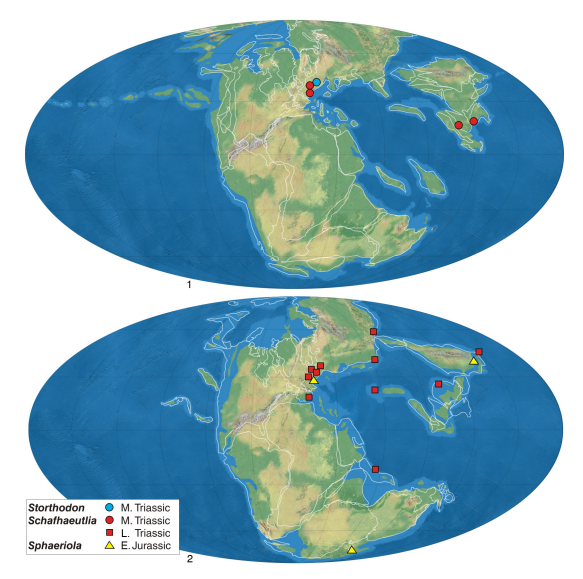


Figure 49. Paleogeographical distribution of Thyasiridae (Storthodon) and Fimbriidae (Schafhaeutlia, Sphaeriola). 1, Middle Triassic; 2, Late Triassic-Early Jurassic.

Stratigraphic range.—Upper Triassic (Norian) (Yancey & others, 2005). Although *Wallowaconcha* was proposed by Yancey and Stanley (1999), it had previously been reported from the Upper Triassic by the same authors (1987, 1996). It had a Norian range (Yancey & others, 2005).

Paleogeographic distribution.—Tethys and Circumpacific (Fig. 47).

Tethys domain: Late Triassic: Norian of United Arab Emirates (Woods & Yancey, 2004; Yancey & others, 2005), Himalaya (northwestern India) (Yancey & others, 2005).

Circumpacific domain: Late Triassic: Norian of Oregon (United States) (Yancey & Stanley, 1987, 1999), Yukon (Canada) (Yancey & Stanley, 1999; Yarnell, Stanley, & Hart, 1999), and Sonora (Mexico) (Yancey & Stanley, 1999).

Paleoautoecology.—B, E-Se, S-Ch-Ph, Un, Sed; R. *Wallowaconcha* lived reclining on the substrate with the commissural plane vertical, with the wings being horizontal on the substrate (see Yancey & Stanley, 1999, fig. 7). It had a gregarious mode of life. Its aberrant morphology indicates that it could establish symbiotic relationships with microorganisms. There are

certain features, such as the heavy shell, large size, and poor capacity to open the valves, that indicate the possibility of chemo- or photosymbiotic relations (see Yancey & Stanley, 1999, p. 19–20). In addition, wallowaconchids lived in tropical environments, another criterion that helps to identify the photosymbiosis in fossil organisms according to Seilacher (1990).

Mineralogy.—Aragonitic (Carter, Barrera, & Tevesz, 1998). The mineralogy and microstructure of *Wallowaconcha* are unknown, but probably part of the shell had a prismatic microstructure (Yancey & Stanley, 1999). Carter, Barrera, and Tevesz (1998) considered that members of the superfamily Megalodontoidea had a completely aragonitic shell.

Superfamily LUCINOIDEA Fleming, 1828 Family LUCINIDAE Fleming, 1828 Genus SINBADIELLA Hautmann & Nützel, 2005, p. 1132

Type species.—Sinbadiella pygmaea Hautmann & Nützel, 2005, p. 1133.

Stratigraphic range.—Lower Triassic (Olenekian) (Hautmann & Nützel, 2005). Hautmann and Nützel (2005) proposed *Sinbadiella* from the Olenekian of the Moenkopi Formation. This is the only record of a bivalve of the order Heterodonta from the Lower Triassic.

Paleogeographic distribution.—Circumpacific (Fig. 48).

Circumpacific domain: Early Triassic: Olenekian of Utah (United States) (Hautmann & Nützel, 2005).

Paleoautoecology.—B, Is, S-?Ch, SM; Sb. Hautmann and Nützel (2005) assumed that *Sinbadiella*, like living lucinids, established chemosymbiotic relations, since the Early Triassic was a time with reduced primary productivity and anoxia-dominated environments after the end of the P/T extinction, although this may be implausible. Most likely, it was a shallow burrower, since the deep burrowers did not appear until later when the recovery of the extinction was complete (Twitchett, 2006).

Mineralogy.—Aragonitic (J. D. Taylor, Kennedy, & Hall, 1973). The shell mineralogy and microstructure of *Sinbadiella* is unknown. J. D. Taylor, Kennedy, and Hall (1973) described three aragonitic shell layers in members of the superfamily Lucinoidea: a prismatic outer layer and middle and inner layers with cross-lamellar microstructure.

Genus LUCINIOLA Skeat & Madsen, 1898, p. 86

Type species.—Venus pumila Münster, 1841, in Goldfuss, 1833–1841, p. 243.

Stratigraphic range.—Upper Triassic (Rhaetian)–Lower Jurassic (Pliensbachian). Cox and others (1969) recorded it from the European Lower Jurassic. Luciniola was long considered to have appeared in the Hettangian just after Triassic–Jurassic extinction event (Hallam, 1987, 1990). However, the species Luciniola stoppaniana (Dittmar, 1864) was also mentioned from the Rhaetian (McRoberts, Newton, & Allasinaz, 1995). This species was originally referred to Lucina Bruguière, 1797, a Cretaceous to Recent (Cox & others, 1969) genus. Luciniola stoppaniana was abundant in the Alpine Rhaetian and in Hungary (Borghi, 1937; Chiesa, 1949; Vörös, 1981; McRoberts, Newton, & Allasinaz, 1995).

Paleogeographic distribution.—Tethys and Circumpacific (Fig. 48). Lucioniola was regarded as a genus endemic in the western Tethys (Cox & others, 1969), but it was subsequently reported from Japan and China.

Tethys domain: Late Triassic: Rhaetian of the Italian Alps (Borghi, 1937; Chiesa, 1949; McRoberts, Newton, & Allasinaz, 1995), Hungary (Vörös, 1981); Early Jurassic: Hettangian of China (Z. Huang, 1986); Hettangian or Sinemurian of China (Stiller, 2006, and references therein); Hettangian–Sinemurian of Europe (Hallam, 1976, 1977, 1987); Sinemurian of China (J. Yin, H. Yao, & Sha, 2004).

Circumpacific domain: Early Jurassic: Sinemurian of Japan (Hayami, 1975).

Paleoautoecology.—B, Is, S-?Ch, SM; Sb. Many living Lucinidae establish chemosymbiotic relationships with sulfate-reducing bacteria (Beesley, Ross, & Wells, 1998). Similar relationships may have been already established by Mesozoic times, and this can explain some of their morphological features (Seilacher, 1990). There are some shell features associated with symbiosis that can be recognized in fossil shells, such as the position of the anterior adductor muscle scars (J. D. Taylor & Glover, 2000, 2006). Lucinidae are infaunal siphonate burrowers. According to Stiller (2006), *Luciniola* was a shallow burrower.

Mineralogy.—Aragonitic (J. D. Taylor, Kennedy, & Hall, 1973). The shell mineralogy and microstructure of *Luciniola* is unknown. See discussion in *Sinbadiella* (p. 132).

Family THYASIRIDAE Dall, 1901 Genus STORTHODON Giebel, 1856, p. 102

Type species.—Storthodon liscaviensis Giebel, 1856, p. 79.

Stratigraphic range.—Middle Triassic (Ladinian) (Hallam, 1981). Storthodon was proposed by Giebel (1856) from the Muschelkalk of Germany. Chavan (in Cox & others, 1969) included it in the family Thyasiridae, but J. D. Taylor and others (2007) and J. D. Taylor, Williams, and Glover (2007) argued that this family had a Early Cretaceous origin and that *Storthodon* did not belong there. They did not give any indication of possible relationships, so we provisionally follow Chavan (in Cox & others, 1969). Giebel (1856) noticed that there was some resemblance between his new genus and *Lucina*, but some aspects, such as the hinge, are not comparable. Hallam (1981) recorded it from Ladinian times.

Paleogeographic distribution.—western Tethys (Fig. 49).

Tethys domain: Middle Triassic: Muschelkalk of Germany (Giebel, 1856).

Paleoautoecology.—Unknown. Living Thyasiridae species normally inhabit deep waters, and, as in Lucinidae, they may have sulphatereducing bacteria in their gills. But it is very unlikely that *Storthodon* belongs to this family.

Mineralogy.—Unknown. The shell mineralogy and microstructure of *Storthodon* are unknown. Because its taxonomic relations are not clear, we cannot use the dominant mineralogy in the family.

Family FIMBRIIDAE Nicol, 1950 Genus SCHAFHAEUTLIA Cossmann, 1897, p. 51

[nom. nov. pro Gonodon Schafhäutl, 1863, p. 382, non Held, 1837, p. 918]

Type species.—Gonodon schafhaeutli Salomon, 1895, p. 169 [=*Gonodon ovatum* Schafhäutl, 1863, p. 382, *non Cyprina ovata* Stoppani, 1860 in 1858–1860, *nec Isocardia ovata* Münster, 1837, in Goldfuss, 1833–1841] (see Hautmann, 2001b, p. 123–124).

Remarks.—Krumbeckia Diener, 1915, was regarded as a synonym of *Schafhaeutlia* by Hautmann (2001a) (see discussion for *Krumbeckia* in Genera not Included, p. 163) and *Isocardioides* Fan, 1963, probably is also a junior synonym of *Schafhaeutlia*, according to Gu and others (1976) and Z. Fang and others (2009).

Stratigraphic range.—Middle Triassic (Anisian)–Upper Triassic (Rhaetian) (Hautmann, 2001b; Komatsu, Chen, & others, 2004). Cox and others (1969) assigned it an Upper Triassic range and a European and South American distribution. Subsequently, the range was expanded by new Middle Triassic records. The range of this genus is most probably artificially extended, as many species were attributed to *Schafhaeutlia* simply on the basis of external form and ornamentation, regardless of internal characters (Monari, 2003), as happened with records of the genus from the eastern coast of the Paleopacific (see paleogeographic distribution).

Paleogeographic distribution.—Tethys, Boreal, and Circumpacific (Fig. 49). For a more complete distribution in the western Tethys, see

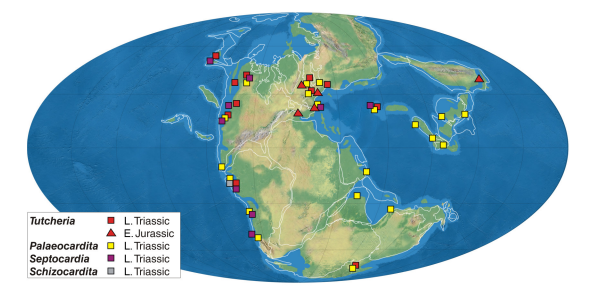


Figure 50. Paleogeographical distribution of Carditidae (Tutcheria, Palaeocardita, Septocardia, Schizocardita). Late Triassic-Early Jurassic.

Monari (2003, p. 882–883). The genus was reported from Rhaetian beds of Nevada (Laws, 1982; Hallam & Wignall, 2000), but none of these papers contain descriptions or figures of the specimens. McRoberts (1997a) recorded *Schafhaeutlia* sp. from the Rhaetian of Sonora (Mexico), but he assigned his two valves to the genus on the basis of external morphology alone, as internal characters were not observed. The same applies to records from the Upper Triassic of Chile (Hayami, Maeda, & Ruiz-Fuller, 1977; Chong & Hillebrandt, 1985) and Peru (Cox, 1949). Therefore, we consider the presence of this genus in the eastern Paleopacific as doubtful.

Tethys domain: Middle Triassic: Anisian of southern China (Sha, Chen, & Qi, 1990; Komatsu, Chen, & others, 2004), southern Alps (Monari, 2003), northern Vietnam (Komatsu, Huyen, & Huu, 2010); Ladinian of Italy (Rossi Ronchetti, 1959), China (Sha, Chen, & Qi, 1990), northern Vietnam (Komatsu, Huyen, & Huu, 2010); Late Triassic: Carnian of the Alps (Italy) (Leonardi, 1943; Allasinaz, 1964, 1966; Gelati & Allasinaz, 1964; Fürsich & Wendt, 1977), Yugoslavia (Jurkovsek & Jelen, 1990), Slovenia (Jelen, 1988), Spain (Pérez-Valera & others, 2007); Norian of western China (H. Yao & others, 2007), Afghanistan (Polubotko, Payevskaya, & Repin, 2001); Norian–Rhaetian of Iran (Hautmann, 2001b); Rhaetian of ?Tibet (J. Yin & McRoberts, 2006), ?Hungary (Vörös, 1981).

Circumpacific domain: Late Triassic: Carnian of Japan (Hayami, 1975).

Boreal domain: Late Triassic: Carnian of Primorie (Kiparisova, 1972).

Paleoautoecology.—B, Is, S, SM; Sb. Living species of *Fimbria*, the only extant genus of Fimbriidae, are slow infaunal burrowers (Beesley, Ross, & Wells, 1998; see discussion in Monari, 2003) and they are found among corals in shallow waters. *Schafhaeutlia globosa* had a globose shell, similar to some species of *Fimbria*, and it was most likely a slow burrower. However, Hautmann (2001b) proposed an epibenthic mode of life for *Schafhaeutlia sphaerioides* (Boettger, 1880), living freely resting on the substrate.

Mineralogy.—Aragonitic (J. D. Taylor, Kennedy, & Hall, 1973). There are no data about *Schafhaeutlia* shell mineralogy or microstructure. J. D. Taylor, Kennedy, and Hall (1973) described three aragonitic shell layers in members of superfamily Lucinoidea: a prismatic outer shell layer and middle and inner shell layers of cross-lamellar microstructure. Monari (2003) studied the shell microstructure of *Cerkesia contiae* Monari, 2003, an Upper Jurassic member of family Fimbriidae from Turkey, and found the same structure.

Genus SPHAERIOLA Stoliczka, 1871 in 1870-1871, p. 247

Type species.—Cardium madridi Archiac, 1843, p. 373.

Stratigraphic range.—Lower Jurassic (Sinemurian)–Middle Jurassic (Callovian) (Monari, 2003). Cox and others (1969) assigned it a Jurassic (Domerian [=Pliensbachian]–Bathonian) range. However, Monari (2003) reviewed the history of *Sphaeriola* and proposed a Sinemurian to Callovian range.

Paleogeographic distribution.—Tethys, Circumpacific, and Austral (Fig. 49). Sphaeriola was doubtfully mentioned from the Hettangian–Sinemurian boundary of the Neuquén Basin (Damborenea & Manceñido, 2005b).

Tethys domain: Early Jurassic: Sinemurian of the southern Alps (Italy) (Parona, 1890; Monari, 2003).

Circumpacific domain: Early Jurassic: Sinemurian of Japan (Hayami, 1975).

Austral domain: Early Jurassic: Hettangian–Sinemurian of New Zealand (Marwick, 1953).

Paleoautoecology.—B, Is, S-Ch, SM; Sb. Living *Fimbria* species are shallow burrowers, but the foot is not as developed as in other Lucinoidea, and they are slow burrowers (Beesley, Ross, & Wells, 1998). Like other Lucinidae, they also have sulphate-reducing bacteria in their gills, at least in *F. fimbriata* (Linnaeus, 1758), studied by J. D. Taylor and Glover (2000). We assume that *Sphaeriola* had a similar mode of life.

Mineralogy.—Aragonitic (J. D. Taylor, Kennedy, & Hall, 1973). There are no data about *Sphaeriola* shell mineralogy or microstructure. See discussion in *Schafhaeutlia* (p. 133).

Superfamily CARDITOIDEA Férussac, 1822 in 1821–1822 Family CARDITIDAE Férussac, 1822 in 1821–1822 Genus TUTCHERIA Cox, 1946, p. 35

Type species.—Cardium submulticostatum d'Orbigny, 1850,p. 235.

Remarks.—Cox (1946) included *Tutcheria* in the family Astartidae. However, Chavan in Cox and others (1969) and Hayami (1969b) transferred it to the Carditidae.

Stratigraphic range.—Upper Triassic (Carnian)–Middle Jurassic (Aalenian) (Cox, 1946). According to Cox (1946), *Tutcheria* had an Upper Triassic (Carnian)–Middle Jurassic (Aalenian) range. The oldest species was *T. parvula* (Trechmann, 1918), and the most recent is *T. aalensis* Cox, 1946, of which only one specimen is known. This range was repeated in Cox and others (1969). Subsequently, it was reported from the Lower Jurassic (Pliensbachian) (Hallam, 1977; Hölder, 1995; Szente, 1997; Harries & Little, 1999).

Paleogeographic distribution.—Tethys, Circumpacific, and Austral (Fig. 50).

Tethys domain: Late Triassic: Norian of the western Carpathians (Kollarova & Kochanová, 1973); Norian–Rhaetian of Iran (Hautmann, 2001b); Rhaetian of the Alps (Hautmann, 2001b), England (Ivimey-Cook & others, 1999); Early Jurassic: Germany (Hölder, 1995); Hettangian of England (Cox, 1946), Italy (Gaetani, 1970); Sinemurian of Portugal (Liu, 1995).

Circumpacific domain: Late Triassic: Norian of southwestern Alaska (United States) (Newton, 1986; McRoberts & Blodgett, 2000), Peru (Körner, 1937; Cox, 1946, 1949), Oregon (United States) (Newton in Newton & others, 1987), Vancouver (Canada) (Tozer, 1962), Nevada (United States) (Laws, 1982); Rhaetian of British Columbia (Canada) (Wignall & others, 2007); Early Jurassic: Sinemurian of Japan (Hayami, 1969b).

Austral domain: Late Triassic: Carnian of New Zealand (Trechmann, 1918; Cox, 1946; Marwick, 1953).

Paleoautoecology.—B, Is, S, SM; Sb. Their rounded outer shapes, the presence of a lunule, and absence of a pallial sinus indicate that *Tutcheria* was a nonsiphonate burrower (Newton in Newton & others, 1987; Hautmann, 2001b). According to the type of facies where it is recorded, it lived in shallow-water sandy substrates (Newton in Newton & others, 1987).

Mineralogy.—Aragonitic (J. D. Taylor, Kennedy, & Hall, 1973). There are no data about *Tutcheria* shell mineralogy or microstructure. Members of the superfamily Carditoidea have a completely aragonitic shell, usually with an outer shell layer of cross-lamellar structure and an inner shell layer of complex cross-lamellar structure (J. D. Taylor, Kennedy, & Hall, 1973).

Genus PALAEOCARDITA Conrad, 1867, p. 11

Type species.—Cardium austriacum Hauer, 1853, p. 736.

Stratigraphic range.—Upper Triassic (Carnian–Rhaetian). Although Cox and others (1969) assigned it a Middle–Upper Triassic range, we only located Upper Triassic records. *Cardita crenata* Münster, 1838, which was reported from Ladinian beds (e.g., Arthaber, 1908), was referred to *Palaeocardita* by Chavan in Cox and others (1969). However, this species is now assigned to *Septocardia* Hall & Whitfield, 1877, for well-justified reasons (see Schneider & Carter, 2001, p. 613).

Paleogeographic distribution.—Tethys, Circumpacific, and Austral (Fig. 50). *Palaeocardita* was also reported from Upper Triassic of Primorie (Kiparisova, 1972), but the figured material is too poor and its record is considered doubtful.

Tethys domain: Late Triassic: China (Cowper-Reed, 1927; Wen & others, 1976; Gou, 1993), Timor (Indonesia) (Krumbeck, 1924); Carnian of Italy (Corazzari & Lucchi Garavello, 1980), the Alps (Italy) (Fürsich & Wendt, 1977); Norian of western China (H. Yao & others, 2007); Norian–Rhaetian of Iran (Hautmann, 2001b), Yunnan province (China) (J. Yin & McRoberts, 2006), Australia (Grant-Mackie, 1994), Oman (Arabia) (R. Hudson & Jefferies, 1961); Rhaetian of the eastern Alps (Austria) (Tomašových, 2006b), Tibet (China) (Hautmann & others, 2005; J. Yin & McRoberts, 2006), Iran (Repin, 2001), England (Ivimey-Cook & others, 1999), ?Vietnam (Vu Khuc & Huyen, 1998), Sumatra (Boettger, 1880), Hungary (Vörös, 1981), the Alps (Italy) (Diener, 1923; Desio, 1929; S. Conti, 1954), Burma (Healey, 1908).

Circumpacific domain: Late Triassic: Norian of Oregon (United States) (Newton in Newton & others, 1987), Peru (Cox, 1949), Sonora (Mexico) (G. D. Stanley & others, 1994; Goodwin, 1997; McRoberts, 1997a; G. D. Stanley, 1997), Chile (Hayami, Maeda, & Ruiz-Fuller, 1977; Chong & Hillebrandt, 1985); Rhaetian of British Columbia (Wignall & others, 2007).

Austral domain: Late Triassic: Carnian–Norian of New Zealand (Trechmann, 1918; Marwick, 1953); Rhaetian of Argentina (Riccardi & others, 1997, 2004; Damborenea & Manceñido, 2012).

Paleoautoecology.—B, Is, S, Endo-Un, Sed-SM; By-Sb. S. M. Stanley (1970) considered two major modes of life in the family Carditidae: shallow burrower and endobyssate infaunal. To differentiate between both types, he proposed to use the degree of elongation (ratio between length and height of the shell). Both interpretations were found in the literature. *Palaeocardita silberlingii* Newton in Newton & others, 1987, and *P. stoecklini* Hautmann, 2001b, interpreted by Newton in Newton and others (1987) and by Hautmann (2001b), respectively, as infaunal bivalves living very near the surface, and endobyssate on rocks or other hard fragments within the sediment, similar to the living *Cardita floridana* Conrad, as discussed by S. M. Stanley (1970). However, *P. iranica* Hautmann, 2001b, was interpreted by its author as a slow, shallow burrower in fine-grained sediments and low-energy environments.

Mineralogy.—Aragonitic (J. D. Taylor, Kennedy, & Hall, 1973). Members of the superfamily Carditoidea have a completely aragonitic shell, usually with an outer shell layer of cross-lamellar structure and an inner shell layer of complex cross-lamellar structure (J. D. Taylor, Kennedy, & Hall, 1973). The same was interpreted by Hayami, Maeda, and Ruiz-Fuller (1977) for their specimens of *P. peruviana* Cox, 1949.

Genus SEPTOCARDIA Hall & Whitfield, 1877, p. 294

Type species.—Septocardia typica Hall & Whitfield, 1877, p. 295. Remarks.—Following Schneider and Carter (2001), who suggested that this genus is closely related to Palaeocardita, we include Septocardia in Carditidae. It includes the species S.

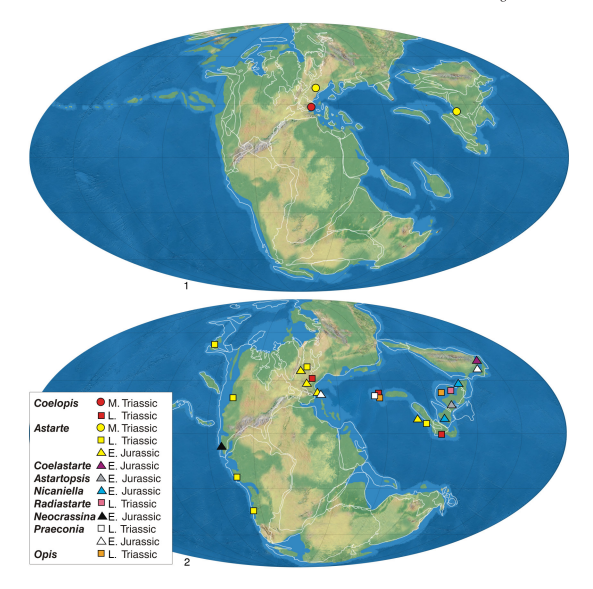


Figure 51. Paleogeographical distribution of Astartidae (Coelopis, Astarte, Coelastarte, Astartopsis, Nicaniella, Radiastarte, Neocrassina, Praeconia, Opis). 1, Middle Triassic; 2, Late Triassic–Early Jurassic.

crenata (Münster, 1838) and *S. pichleri* (Bittner, 1895), which were referred by other authors to *Palaeocardita* (see Schneider & Carter, 2001, p. 613).

Stratigraphic range.—Upper Triassic (Carnian–Rhaetian). Although Arthaber (1908) mentioned the species *Cardita crenata* from the Ladinian of the Alps, Diener (1923) considered this, and all other records of this species, to be Carnian in age. This species was widely distributed during the Late Triassic, but it was not recorded from Ladinian times, as shown by Cox and others (1969, p. 554). Cox and others (1969) reported the genus from the Norian of North and South America (not including *S. crenata* and *S. pichleri*), expanding both the stratigraphic range and the paleogeographic distributions of this genus.

Paleogeographic distribution.—Tethys, Circumpacific, and Austral (Fig. 50). Goodwin (1997) and McRoberts (1997a) reported Sep*tocardia* sp. from the Norian of Sonora (Mexico), but they did not figure the specimens.

Tethys domain: Late Triassic: Carnian of Italy (Bittner, 1895; Schneider & Carter, 2001); Norian of Iran (Repin, 2001).

Circumpacific domain: Late Triassic: Norian of Peru (Cox, 1949), southwestern Alaska (McRoberts & Blodgett, 2000), Oregon (United States) (Newton in Newton & others, 1987), Chile (Hayami, Maeda, & Ruiz-Fuller, 1977; Chong & Hillebrandt, 1985); Rhaetian of British Columbia (Canada) (Wignall & others, 2007), Nevada (United States) (Silberling, 1961; Laws, 1982; Hallam & Wignall, 2000; Guex & others, 2003; Lucas & Tanner, 2004).

Austral domain: Late Triassic: Norian-Rhaetian of Argentina (Riccardi & others, 1997, 2004; Damborenea & Manceñido, 2012). *Paleoautoecology.*—B, Is, S, Endo-Un, Sed-SM; By-Sb. S. M. Stanley (1972) proposed that *S. crenata* was an endobyssate infaunal bivalve, living buried very near the surface. However, as in *Palaeocardita* (already discussed), the interspecific variation within the genus is significant, so other species, such as *S. typica* Hall & Whitfield, 1877, could have been shallow burrowers.

Mineralogy.—Aragonitic (Schneider & Carter, 2001). See Schneider and Carter (2001) for a complete discussion of the microstructure of the species *S. crenata* and *S. pichleri*.

Genus SCHIZOCARDITA Körner, 1937, p. 194

Type species.—Schizocardita cristata Körner, 1937, p. 194.

Stratigraphic range.—Upper Triassic (Norian) (Hallam, 1981). Körner (1937) proposed *Schizocardita* as a subgenus of *Cardita* from the Upper Triassic of Peru. By comparison with European faunas of the Cassian Formation or Raibl Alpine Triassic, Hallam assigned it a Ladinian or Carnian age, since the first formation was then dated as Middle Triassic, but it is now considered to be Carnian in age. Cox (1949) studied the fauna of the same area and assigned it a Norian age. Subsequently, Hallam (1981) assigned the genus a Norian range in South America. Cox and others (1969) considered *Schizocardita* as a genus and assigned it a Triassic range.

Paleogeographic distribution.—Circumpacific (Fig. 50). The genus is endemic to the northern Andes (Kobayashi & Tamura, 1983b; Damborenea, 2002b).

Circumpacific domain: Late Triassic: Norian of Peru (Körner, 1937; Hallam, 1981).

Paleoautoecology.—B, Is-Se, S, SM; Sb. Members of the family Carditidae exhibit a variety of modes of life: shallow infaunal, semi-infaunal, byssate, and even nestler bivalves. Körner (1937) indicated that it is difficult to know how *Schizocardita* lived, because no similar morphology is known. According to its trigoniform external form, it was probably infaunal or semi-infaunal.

Mineralogy.—Aragonitic (J. D. Taylor, Kennedy, & Hall, 1973). There are no data about *Schizocardita* shell mineralogy or microstructure. Members of the superfamily Carditoidea have a completely aragonitic shell, usually with an outer shell layer of cross-lamellar structure and an inner shell layer of complex cross-lamellar structure (J. D. Taylor, Kennedy, & Hall, 1973).

Superfamily CRASSATELLOIDEA Férussac, 1822 in 1821–1822 Family ASTARTIDAE d'Orbigny, 1844 in 1844–1848 Genus COELOPIS Fischer, 1887 in 1880–1887, p. 1019

Type species.—Cardita lunulata J. Sowerby, 1819, p. 55.

Stratigraphic range.—Middle Triassic (Ladinian)–Lower Cretaceous (?) (Cox & others, 1969). Cox and others (1969) included two subgenera within *Coelopis: C. (Coelopis*) from the Middle Triassic to Lower Cretaceous, and *C. (Cryptocoelopis*) Bittner, 1895 from the Middle Triassic (Ladinian). Kutassy (1931) assigned a Carnian age to the records that Cox and others (1969) referred to the Ladinian, and he disregarded the reference of Wurm (1913) from the Ladinian of Menorca in the Balearic Islands. Although Cox and others (1969) recorded *Coelopis* up to the Lower Cretaceous, we could only confirm its presence up to the Upper Jurassic (Hayami, 1975; Delvene, 2000).

Paleogeographic distribution.—Tethys (Fig. 51).

Tethys domain: Middle Triassic: Ladinian of Spain (Wurm, 1913); Late Triassic: Carnian of the Alps (Bittner, 1895; Corazzari & Luchi Garavello, 1980); Norian of Seram (Indonesia) (Hautmann, 2001b); Norian–Rhaetian of Iran (Hautmann, 2001b).

Paleoautoecology.—B, E, S, Epi, Sed; By. Some authors (Delvene, 2000, 2003; Fürsich & others, 2001; Gahr, 2002) interpreted *Coelopis* as a nonsiphonate infaunal bivalve, shallow burrower, like most astartids (S. M. Stanley, 1970). However, Hautmann (2001b) compared the morphology of *Coelopis (Coelopis) aurea* Hautmann, 2001b, and *Coelopis (Coelopis) krumbecki* Hautmann, 2001b, with *Opisoma* Stoliczka, 1871 in 1870–1871, which was interpreted as an epibenthic reclined bivalve that lived with the shell partially buried (see Aberhan & Hillebrandt, 1999). Hautmann (2001b) interpreted these species in the same way, but, given their small size, he supposed they instead lived epibyssate rather than reclined, as does *Opisoma*. The last interpretation seems more accurate, according to the morphology of most species attributed to *Coelopis*.

Mineralogy.—Aragonitic (J. D. Taylor, Kennedy, & Hall, 1973). There are no studies on *Coelopis* shell mineralogy or microstructure. The shell of members of family Astartidae is entirely aragonitic, with an outer cross-lamellar layer and an inner layer formed by myostracaltype prisms and traces of complex cross-lamellar or homogeneous structure (J. D. Taylor, Kennedy, & Hall, 1973).

Genus ASTARTE J. Sowerby, 1816, p. 85

Type species.—Venus scotica Maton & Rackett, 1807, p. 81.

Stratigraphic range.—Middle Triassic (Anisian)–Holocene (Cox & others, 1969; Baumgarte, 1973). Cox and others (1969) assigned it a Jurassic–Holocene range, as did authors who suggested a Hettangian origin for *Astarte* (Hallam, 1987, 1990; Schneider, 1995; Sepkoski, 2002). But other authors considered a Late Triassic origin (Marincovich, Barinov, & Oleinik, 2002). However, *Astarte triasina* Roemer, 1851, from the Middle Triassic (Muschelkalk) (Busse, 1972; Baumgarte, 1973; Busse & Horn, 1978) is the oldest record of this genus. An astartid species was reported from the Upper Triassic of South America (*Astarte inca* Jaworski, 1922), but Riccardi and others (1997, 2004) refer it only doubtfully to *Astarte*.

Paleogeographic distribution.—Tethys, Circumpacific, and ?Austral (Fig. 51).

Tethys domain: Middle Triassic: Muschelkalk of Germany (Busse, 1972; Baumgarte, 1973; Busse & Horn, 1978); Anisian of China (Komatsu, Chen, & others, 2004); Late Triassic: Rhaetian of Tibet (China) (J. Yin & McRoberts, 2006), England (Ivimey-Cook & others, 1999); Early Jurassic: Hettangian of China (Gu & others, 1976; Z. Huang, 1986), Tibet (China) (J. Yin & McRoberts, 2006), England (Liu, 1985), Italy (Gaetani, 1970); Sinemurian of England and France (Liu, 1995).

Circumpacific domain: Late Triassic: Peru (Jaworski, 1922); Norian of ?southwestern Alaska (McRoberts & Blodgett, 2000), Oregon (United States) (Newton & others, 1987).

Austral domain: Late Triassic: Rhaetian of ?Argentina (Riccardi & others, 1997, 2004).

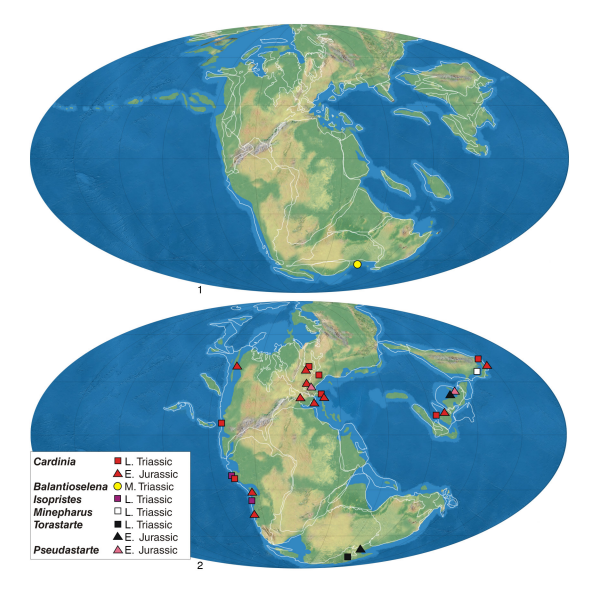


Figure 52. Paleogeographical distribution of Cardiniidae (Cardinia, Balantioselena, Isopristes, Minepharus, Torastarte, Pseudastarte). 1, Middle Triassic; 2, Late Triassic–Early Jurassic.

Paleoautoecology.—B, Is, S, SM; Sb. S. M. Stanley (1970) examined several living *Astarte* species and indicated they are all slow, shallow, nonsiphonate burrowers. We assume a similar mode of life for the Mesozoic species.

Mineralogy.—Aragonitic (J. D. Taylor, Kennedy, & Hall, 1973). J. D. Taylor, Kennedy, and Hall (1973) studied the shell microstructure of several living species, corroborating that all are completely aragonitic, with an outer cross-lamellar layer and an inner layer formed by homogeneous or complex cross-lamellar structure, according to species, and myostracal-type prisms.

Genus COELASTARTE Böhm, 1893, p. 174

Type species.—Astarte excavata J. Sowerby, 1819, p. 57.

Remarks.—Prior to the paper by Hayami (1958d), *Coelastarte* was regarded as a subgenus of *Astarte* J. Sowerby, 1816. Hayami raised it to generic level, because it was easily distinguishable from *Astarte* and other related genera. Cox and others (1969) relegated *Coelastarte*

to a subgenus of *Neocrassina* Fischer, 1886 in 1880–1887. However, Gardner and Campbell (2002) decided to maintain the two taxa as separate and raised *Coelastarte* to generic level. This decision is supported by differences in the hinge of both genera, among other aspects, and this opinion is followed here.

Stratigraphic range.—Lower Jurassic (Hettangian)–Upper Cretaceous (Turonian) (Hayami, 1958d; Cox & others, 1969). Cox and others (1969) assigned it a Middle Jurassic (Aalenian)–Upper Cretaceous (Turonian) range. However, *Coelastarte* originated in the Hettangian of Japan (Hayami, 1958d, 1975; Sato & Westermann, 1991). Although the youngest record we found is from the Upper Jurassic (Tithonian) (Hayami, 1975; Fürsich, 1981), we follow Cox and others (1969), in considering Turonian to be the top of the range, as we could not access the original source of this quotation.

Paleogeographic distribution.—Circumpacific (Fig. 51). Although during our interval it was only recorded from Japan, it had a broader

distribution during the rest of the Jurassic (Cox & others, 1969; Gardner & Campbell, 2002).

Circumpacific domain: Early Jurassic: Hettangian of Japan (Hayami, 1958d, 1975; Sato & Westermann, 1991; Sugawara & Kondo, 2004; Kondo & others, 2006).

Paleoautoecology.—B, Is, S, SM; Sb. Similar to Praeconia.

Mineralogy.—Aragonitic (J. D. Taylor, Kennedy, & Hall, 1973). There are no studies on *Coelastarte* shell mineralogy or microstructure. See discussion in *Astarte* (p. 137).

Genus ASTARTOPSIS de Loriol, 1891, p. 218

Type species.—Astartopsis elongata de Loriol, 1891, p. 219.

Stratigraphic range.—Lower Jurassic (Sinemurian), Upper Jurassic (Oxfordian). According to Cox and others (1969), Astartopsis is known from Jurassic deposits (L. Lusitanian). Septfontaine (1995) noticed that the syntypes of A. elongata are from the Rauracian (=Oxfordian) from Switzerland. In addition to these Upper Jurassic records, the genus was only reported from the Lower Jurassic (Sinemurian) of southern China by J. Chen (1988), who described three species: A. menkei (Dunker) in Xintianmen, Tianmen'ao, and Upper Jinji formations; A. subovalis Chen in Xintianmen Formation, and A. sp. in Upper Jinji Formation.

Paleogeographic distribution.—Eastern Tethys (Fig. 51).

Tethys domain: Early Jurassic: Sinemurian of southern China (J. Chen, 1988).

Paleoautoecology.—B, Is, S, SM; Sb. According to its external shell morphology, *Astartopsis* was probably a shallow burrower.

Mineralogy.—Aragonitic (J. D. Taylor, Kennedy, & Hall, 1973). There are no studies on *Astartopsis* shell mineralogy or microstructure. See discussion in *Astarte* (p. 137).

Genus NICANIELLA Chavan, 1945, p. 43

Type species.—Astarte communis Zittel & Goubert, 1861, p. 201. Stratigraphic range.—Lower Jurassic (Hettangian)–Paleocene (Hayami, 1964; Cox & others, 1969). Cox and others (1969) assigned to Nicaniella a Middle Jurassic–Paleocene range and doubtfully also recorded it from the Pliocene, but the genus had already been reported from the Hettangian of Vietnam (Hayami, 1964). Damborenea and Manceñido (2012) doubtfully referred to Nicaniella specimens identified as N.? cf. inca (Jaworski, 1922) from Rhaetian beds in Argentina.

Paleogeographic distribution.—Eastern Tethys (Fig. 51). *Nicaniella* was originally distributed in the Eastern Tethys, but, starting with Pliensbachian times, its distribution significantly broadened (Hallam, 1976, 1977; Fürsich & others, 1991, 2001, 2005; Liu, 1995; Holzapel, 1998; Harries & Little, 1999; Gahr, 2002; Delvene, 2003; Fürsich & Thomsen, 2005).

Tethys domain: Early Jurassic: Hettangian–Sinemurian of China (Stiller, 2006, and references therein), Vietnam (Hayami, 1964; Sato & Westermann, 1991).

Paleoautoecology.—B, Is, S, SM; Sb. *Nicaniella* is regarded as a slow, shallow burrower, nonsiphonate bivalve according to its external morphology: "strongly inflated shell, marked comarginal lines and thick shell" (Delvene, 2000, p. 132).

Mineralogy.—Aragonitic (J. D. Taylor, Kennedy, & Hall, 1973). There are no studies on *Nicaniella* shell mineralogy or microstructure. See discussion in *Astarte* (p. 137).

Genus RADIASTARTE Chen & Liu in Gu & others, 1976, p. 99

Type species.—Radiastarte yizhangensis Chen & Liu in Gu &

others, 1976, p. 99.

Stratigraphic range.—Upper Triassic (Carnian) (Gu & others, 1976). Chen and Liu (in Gu & others, 1976) proposed *Radiastarte* from the Carnian of Hunnan (China).

Paleogeographic distribution.—Eastern Tethys (Fig. 51). The genus was endemic to China (Kobayashi & Tamura, 1983a).

Tethys domain: Late Triassic: Carnian of Hunnan province (China) (Gu & others, 1976; Kobayashi & Tamura, 1983a).

Paleoautoecology.-B, Is, S, SM; Sb. Similar to Nicaniella.

Mineralogy.—Aragonitic (J. D. Taylor, Kennedy, & Hall, 1973). There are no studies on *Radiastarte* shell mineralogy or microstructure. See discussion in *Astarte* (p. 137).

Genus NEOCRASSINA Fischer, 1887 in 1880–1887, p. 1016

[nom. nov. pro Crassinella Bayle, 1878, expl. pl. 105, non Guppy, 1874, p. 442]

Type species.—Astarte obliqua Deshayes, 1830 in 1830–1832, expl. pl. 22.

Stratigraphic range.—Lower Jurassic (Hettangian)–Lower Cretaceous (Berriasian) (Hayami, 1975; Scholz, Aberhan, & González-León, 2008). Cox and others (1969) included two subgenera within Neocrassina: N. (Neocrassina) and N. (Coelastarte) Böhm, 1893. We only consider the first, since Coelastarte is here regarded as a separate genus (see discussion for Coelastarte, p. 138). The range assigned to N. (Neocrassina) by Cox and others (1969) is Lower Jurassic (upper Liassic)–Lower Cretaceous. Subsequently, Damborenea in Damborenea and González-León (1997) reported some specimens from Lowest Jurassic (?Hettangian, ?Sinemurian), which were doubtfully assigned to this genus, as internal characters were not observed. Later, Scholz, Aberhan, and González-León (2008) confirmed the presence of the genus from the same age and area (N. gueuxi d'Orbigny, 1850).

Paleogeographic distribution.—Circumpacific (Fig. 51). Although during our study interval *Neocrassina* was only reported from the Circumpacific domain, subsequently it extended to the Tethys (Hallam, 1972, 1976, 1977; Pugaczewska, 1986; Liu, 1995; Holzapfel, 1998; Gahr, 2002), the Austral domain (Gardner & Campbell, 1997, 2002) and Boreal domain (Fürsich, 1982).

Circumpacific domain: Early Jurassic: Hettangian–Sinemurian of Sonora (Mexico) (Damborenea in Damborenea & González-León, 1997; Scholz, Aberhan, & González-León, 2008).

Paleoautoecology.—B, Is, S, SM; Sb. By comparison with modern species of the family Astartidae, *Neocrassina* was probably a shallow burrower bivalve. According to the thickness of its shell and its external form, it was probably a slow burrower.

Mineralogy.—Aragonitic (J. D. Taylor, Kennedy, & Hall, 1973). There are no studies on *Neocrassina* shell mineralogy or microstructure. See discussion in *Astarte* (p. 137).

Paleontological Contributions, number 8

Genus PRAECONIA Stoliczka, 1871 in 1870-1871, p. 278

Type species.—Astarte terminalis Roemer, 1842, p. 15.

Stratigraphic range.—Upper Triassic (Norian)–Upper Jurassic (Tithonian) (Liu, 1995; Hautmann, 2001b). Cox and others (1969) assigned it a Jurassic (Lias.–Lusitan.) range. Subsequent findings extended the observed stratigraphic range from Upper Triassic (Hautmann, 2001b) to Upper Jurassic (Hallam, 1976, 1977; Liu, 1995).

Paleogeographic distribution.—Tethys and Circumpacific (Fig. 51). It was recorded from the Triassic–Jurassic transition of southern Tibet by Hautmann and others (2005), but they did not discuss systematically or nor figure the specimens.

Tethys domain: Late Triassic: Norian-Rhaetian of Iran (Hautmann, 2001b); Early Jurassic: Hettangian-Sinemurian of Italy (Gaetani, 1970).

Circumpacific domain: Early Jurassic: Sinemurian of Japan (Hayami, 1975).

Paleoautoecology.—B, Is, S, SM; Sb. The living species of this family are shallow burrowers, not very active, that live in cold water (Beesley, Ross, & Wells, 1998). Probably, *Praeconia* was a filtering shallow burrower, living very near the surface of the sediment, as interpreted by Hautmann (2001b) and Gahr (2002).

Mineralogy.—Aragonitic (J. D. Taylor, Kennedy, & Hall, 1973). There are no studies on *Praeconia* shell mineralogy or microstructure. See discussion in *Astarte* (p. 137).

Genus OPIS Defrance, 1825a, p. 219

Type species.—Trigonia cardissoides Lamarck, 1819, p. 65.

Stratigraphic range.—Upper Triassic (Norian)–Upper Cretaceous (Maastrichtian) (Hautmann, 2001b; Hayami, 1975). Cox and others (1969) considered three subgenera within *Opis*: *O. (Opis)*, *O. (Pachyopis)* Bigot, 1895, and *O. (Trigonopis)* Fischer, 1887 in 1880–1887, assigning a Jurassic to Upper Cretaceous range. Of these three subgenera, the only one reported from our study interval is *O. (Trigonopis)*.

Gardner and Campbell (2002) raised *Trigonopis* to generic level, distinguishing it from *Opis* by its hinge dentition features. This is difficult to apply when the hinge is not observed, for example, in specimens assigned to *O. (Trigonopis)* from the Upper Triassic by Hautmann (2001b).

Paleogeographic distribution.—Eastern Tethys (Fig. 51). Although during the Late Triassic, *Opis* was only reported from the eastern Tethys, in the Jurassic, its distribution was broader (Hayami, 1975; Liu, 1995; Gardner & Campbell, 2002).

Tethys domain: Late Triassic: Norian of Iran and China (Hautmann, 2001b); Rhaetian of Iran (Hautmann, 2001b).

Paleoautoecology.—B, E, S, Un, Sed; R. Considering shell morphology, Hautmann (2001b) assigned *Opis* a mode of life similar to *Coelopis* Fischer, 1887 in 1880–1887, since it is unlikely that these bivalves were burrowers. But the specimens assigned to *O. (Trigonopis)* by Hautmann show no evidence of byssus, therefore he proposed they lived in low-energy environments.

Mineralogy.—Aragonitic (J. D. Taylor, Kennedy, & Hall, 1973). There are no studies on *Opis* shell mineralogy or microstructure. See discussion in *Astarte* (p. 137).

Family CARDINIIDAE Zittel, 1881 Genus CARDINIA Agassiz, 1841 in 1840–1845, p. 220

[nom. conserv. ICZN, 1954, Opinion 292, see discussion in C. Palmer, 1975]

Type species.—Unio listeri J. Sowerby, 1817, p. 123 (ICZN, 1954, Opinion 292).

Stratigraphic range.—Upper Triassic (Carnian)–Lower Jurassic (Toarcian) (Cox & others, 1969). Cox and others (1969) assigned a Carnian–Toarcian range, noting that the Bajocian records of *Cardinia* were not taken into account, because they were based on poorly preserved material.

Paleogeographic distribution.—Cosmopolitan (Fig. 52). Although we did not find any record from the Boreal domain, Cox and others (1969) assigned it a cosmopolitan distribution, and Kobayashi and Tamura (1983b) suggested *Cardinia* originated in Japan and eastern Siberia during the Late Triassic.

Tethys domain: Late Triassic: Rhaetian of Switzerland (Diener, 1923; Kutassy, 1931), Italy (Mariani, 1919), Vietnam (Vu Khuc & Huyen, 1998), England (Ivimey-Cook & others 1999); Early Jurassic: Hettangian of Italy (S. Conti, 1954; Gaetani, 1970); Hettangian–Sinemurian of Vietnam (Hayami, 1964; Sato & Westermann, 1991), England (Palmer, 1975) and Spain (Liu, 1995); Sinemurian of Portugal and France (Liu, 1995), England (Palmer 1975).

Circumpacific domain: Late Triassic: Peru (Rangel, 1978); Carnian of California (United States) (J. P. Smith, 1927), Japan (Nakazawa, 1956; Hayami, 1975); Early Jurassic: Hettangian of Yukon (Canada) (Poulton, 1991); Sinemurian of Japan (Hayami, 1975), ?Chile (Covacevich, Pérez, & Escobar, 1991).

Austral domain: Early Jurassic: Sinemurian of Argentina (Riccardi & others, 1991; Damborenea & Manceñido, 2005b).

Paleoautoecology.—B, Is, S, SM; Sb. The external morphology of *Cardinia* suggests it was a superficial and nonsiphonate burrower, since the pallial line lacks a sinus. The shell thickness and its somewhat globose morphology indicate it was a slow burrower, as were other members of this family.

Mineralogy.—Aragonitic (J. D. Taylor, Kennedy, & Hall, 1973). The shell microstructure of *Cardinia* is not known. Members of the superfamily Crassatelloidea have an entirely aragonitic shell (J. D. Taylor, Kennedy, & Hall, 1973).

Genus BALANTIOSELENA Speden in Gair, Gregg, & Speden, 1962, p. 96

Type species.—Balantioselena gairi Speden in Gair, Gregg, & Speden, 1962, p. 99.

Remarks.—Speden (in Gair, Gregg, & Speden, 1962) proposed *Balantioselena* and included it in the family Astartidae. Subsequently, however, it was referred to the family Cardiniidae, with which is more closely related (Cox & others, 1969; Morris, 1978).

Stratigraphic range.—Middle Triassic (Ladinian) (Speden in Gair, Gregg, & Speden, 1962). Speden (in Gair, Gregg, & Speden, 1962) reported *Balantioselena* from the Ladinian. Subsequently, it was recorded from the same age in several papers without being discussed systematically (J. D. Campbell & Force, 1973; Retallack & Ryburn, 1982; H. J. Campbell, 1987).

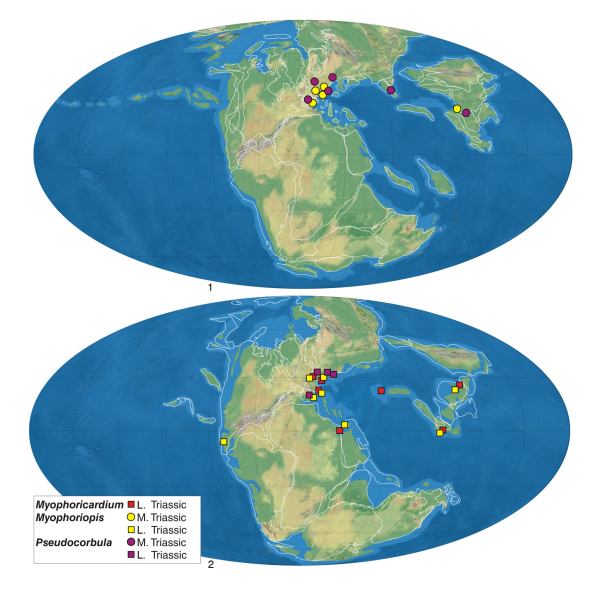


Figure 53. Paleogeographical distribution of Myophoricardiidae (Myophoricardium, Myophoriopis, Pseudocorbula). 1, Middle Triassic; 2, Late Triassic-Early Jurassic.

Paleogeographic distribution.—Austral (Fig. 52).

Austral domain: Middle Triassic: Ladinian of New Zealand (Speden in Gair, Gregg, & Speden, 1962).

Paleoautoecology.—B, Is, S, SM; Sb. Within this family there are genera with compact shapes and others with elongated forms (S. M. Stanley, 1972). The first are often shallow burrowers, while the others would be rather endobyssate. The shell morphology of *Balantioselena* (see description and figures in Speden in Gair, Gregg, & Speden, 1962), its subcircular outline, thick and inflated shell, prosogyrous beaks, and deep lunule indicate it was a nonsiphonate, slow, shallow-burrower bivalve.

Mineralogy.—Aragonitic (J. D. Taylor, Kennedy, & Hall, 1973). The shell microstructure of *Balantioselena* is not known. Members of the superfamily Crassatelloidea have an entirely aragonitic shell (J. D. Taylor, Kennedy, & Hall, 1973).

Genus ISOPRISTES Nicol & Allen, 1953, p. 344

Type species.—Isopristes crassus Nicol & Allen, 1953, p. 345.

Stratigraphic range.—Upper Triassic (Norian). The genus was originally described from the Norian of Peru (Cox & others, 1969) and was subsequently reported from the Norian of Chile (Hayami, Maeda, & Ruiz-Fuller, 1977).

Paleogeographic distribution.—Circumpacific (Fig. 52). *Isopristes* was endemic to the northern Andes (Damborenea, 2002b).

Circumpacific domain: Late Triassic: Norian of Peru (Cox & others, 1969), Chile (Hayami, Maeda, & Ruiz-Fuller, 1977).

Paleoautoecology.—B, Is, S, SM; Sb. Similar to Cardinia.

Mineralogy.—Aragonitic (Hayami, Maeda, & Ruiz-Fuller, 1977). Hayami, Maeda, and Ruiz-Fuller (1977) observed an entirely aragonitic shell with a cross-lamellar outer shell layer and a complex cross-lamellar or homogeneous inner shell layer in *Isopristes* sp. Type species.—Palaeopharus (Minepharus) triadicus Tokuyama, 1958, p. 297.

Remarks.—Minepharus was originally proposed as a subgenus of *Palaeopharus* (family Actinodontophoridae) (Hayami, 1975), but Newell (in Cox & others, 1969) included it, with certain doubts, in the family Cardiniidae. Later, Hayami (1975, p. 170) justified this change of family and discussed Tokuyama's reasons to include it as a subgenus of *Palaeopharus*.

Stratigraphic range.—Upper Triassic (Carnian) (Hayami, 1975). Minepharus was reported from Carnian beds of Japan (Hayami, 1975). Waller (in Waller & Stanley, 2005) found a single valve of what could be a new species of Minepharus, but neither included it in the genus nor described the specimen while waiting for better material.

Paleogeographic distribution.—Circumpacific (Fig. 52).

Circumpacific domain: Late Triassic: Carnian of Japan (Hayami, 1975).

Paleoautoecology.—B, Se, S, Endo, Sed; By. *Minepharus* had an elongated shell, modioliform, like other members of the superfamily Crassatelloidea (S. M. Stanley, 1972). It was likely an endobyssate bivalve, living as did the Paleozoic *Cypricardinia* Hall, 1859 in 1859–1861 (S. M. Stanley, 1972).

Mineralogy.—Aragonitic (J. D. Taylor, Kennedy, & Hall, 1973). The shell microstructure of *Minepharus* is not known. Members of the superfamily Crassatelloidea have an entirely aragonitic shell (J. D. Taylor, Kennedy, & Hall, 1973).

Genus TORASTARTE Marwick, 1953, p. 70

Type species.—Torastarte bensoni Marwick, 1953, p. 70.

Remarks.—Torastarte was originally included in the family Astartidae (Marwick, 1953). Subsequently, specimens with the hinge preserved showed that it was closely related to *Cardinia* Agassiz, 1841 in 1840–1845, and it was assigned to the Cardiniidae (Fleming, 1957), where it was maintained by Cox and others (1969). Hallam (1981) recorded "*Toroastarte*" (typographic error) from the Upper Triassic of New Zealand.

Stratigraphic range.—Upper Triassic (Rhaetian)–Lower Jurassic (Toarcian) (Marwick, 1953; Kear, 1961). Marwick (1953) proposed *Torastarte* from sediments dated as Otapirian (=Rhaetian) from New Zealand. Subsequently, the genus was also reported from the Ururoan (=Hettangian–Toarcian according to N. Hudson [2003]; Pliensbachian–Toarcian according to H. J. Campbell, Raine, & Wilson in Cooper [2004]).

Paleogeographic distribution.—Tethys and Austral (Fig. 52). At first, it was considered endemic to New Zealand, but new records from the eastern Tethys expanded its distribution.

Tethys domain: Early Jurassic: Sinemurian of China (J. Chen, 1988; Stiller, 2006).

Austral domain: Late Triassic: Rhaetian of New Zealand (Marwick, 1953; Fleming, 1957); Early Jurassic: Hettangian–Sinemurian of New Zealand (Kear, 1961).

Paleoautoecology.-B, Is, S, MS; Sb. Similar to Cardinia.

Mineralogy.—Aragonitic (J. D. Taylor, Kennedy, & Hall, 1973). The shell microstructure of *Torastarte* is not known. Members of the superfamily Crassatelloidea have an entirely aragonitic shell (J. D. Taylor, Kennedy, & Hall, 1973).

Genus PSEUDASTARTE Cossmann, 1921, p. 17

Type species.—Astarte (Pseudastarte) emarginata Cossmann, 1921, p. 17.

Stratigraphic range.—Lower Jurassic (Hettangian–Sinemurian) (Stiller, 2006). Cox and others (1969) assigned it a Hettangian range in France, based on the only species then referred to *Pseudastarte*. Subsequently, J. Chen (1988) referred the species *Cardinia exigua* Terquem, 1855, from the Sinemurian of China to *Pseudastarte*. Stiller (2006) recorded it as *Pseudastarte* aff. *exigua* (Terquem, 1855), since he believed that the Chinese specimens were different from the European ones, but new material is needed to see the hinge in the Chinese specimens.

Paleogeographic distribution.—Tethys (Fig. 52).

Tethys domain: Early Jurassic: Hettangian of France (Cossmann, 1921); Hettangian or Sinemurian of China (Stiller, 2006); Sinemurian of China (J. Chen, 1988).

Paleoautoecology.-B, Is, S, SM; Sb. Similar to Cardinia.

Mineralogy.—Aragonitic (J. D. Taylor, Kennedy, & Hall, 1973). *Pseudastarte* shell microstructure is not known. Members of the superfamily Crassatelloidea have an entirely aragonitic shell (J. D. Taylor, Kennedy, & Hall, 1973).

Family MYOPHORICARDIIDAE Chavan in Vokes, 1967 Genus MYOPHORICARDIUM Wöhrmann, 1889, p. 226

Type species.—Myophoricardium lineatum Wöhrmann, 1889, p. 227.

Stratigraphic range.—Upper Triassic (Carnian–Rhaetian) (Hautmann, 2001b). Cox and others (1969) assigned it a Carnian range. Subsequently, new records expanded the observed stratigraphic range up to the Rhaetian (Hautmann, 2001b). It was also reported from the Anisian of China (Komatsu, Chen, & others, 2004), but without description or figures.

Paleogeographic distribution.—Tethys (Fig. 53). The genus was also recorded from the Upper Triassic of Nevada (United States) (Laws, 1982; Hallam & Wignall, 2000), but as the specimens were neither figured nor described, we consider this to be a doubtful record.

Tethys domain: Late Triassic: Carnian of the Alps (Austria) (Wöhrmann, 1889), Italy (Allasinaz, 1966), Slovenia (Jurkovsek, 1978; Jelen, 1988), Yunnan province (China) (Cowper-Reed, 1927), China (Wen & others, 1976), Jordan (Cox, 1924); Norian of Sumatra (Indonesia) (Krumbeck, 1914), China (Gu & others, 1976); Norian–Rhaetian of Iran (Hautmann, 2001b).

Paleoautoecology.—B, Is, S, FM; Sb. Hautmann (2001b) interpreted *M. lineatum* Wöhrmann, 1889, and *M. subquadratum* Hautmann, 2001b, as fast, shallow burrowers with short siphons.

Mineralogy.—Aragonitic (J. D. Taylor, Kennedy, & Hall, 1973). *Myophoricardium* shell microstructure is not known. Members of the superfamily Crassatelloidea have an entirely aragonitic shell (J. D. Taylor, Kennedy, & Hall, 1973).

Genus MYOPHORIOPIS Wöhrmann, 1889, p. 221

Type species.—Myophoria lineata Münster, 1841, p. 88.

Stratigraphic range.—Middle Triassic (Anisian)–Upper Triassic (Rhaetian). Cox and others (1969) assigned a Ladinian–Norian range. However, several reports established its range from the Anisian to Rhaetian (see paleogeographic distribution).

Paleogeographic distribution.—Tethys and Circumpacific (Fig. 53). Tethys domain: Middle Triassic: Anisian of China (Sha, Chen, & Qi, 1990; J. Chen, 2003), the Alps (Switzerland) (Zorn, 1971), Italy (Posenato, 2008a); Ladinian of Austria (Bittner, 1895), Spain (Wurm, 1913; Busnardo, 1970; Márquez-Aliaga, 1983; Calvet & others, 1994); Late Triassic: Carnian of Italy (Bittner, 1895; Broili, 1904; Allasinaz, 1964, 1966; Fürsich & Wendt, 1977), southern Alps (Austria) (Wöhrmann, 1889), China (Gu & others, 1976; Sha, Chen, & Qi, 1990; Sha & Grant-Mackie, 1996), Jordan (Cox, 1924), Slovenia (Boué, 1835, as Corbula); Norian of Sumatra (Indonesia) (Krumbeck, 1914); Rhaetian of Spain (Goy & Márquez-Aliaga, 1998), Hungary (Vörös, 1981), Italy (Chiesa, 1949; Allasinaz, 1962; Gelati & Allasinaz, 1964; Sirna, 1968), eastern Alps (Austria) (Tomašových, 2006a).

Circumpacific domain: Late Triassic: Norian of Sonora (Mexico) (Damborenea in Damborenea & González-León, 1997).

Paleoautoecology.—B, Is, S, FM; Sb. The mode of life of Myophoriopis was probably similar to Myophoricardium.

Mineralogy.—Aragonitic (Schneider & Carter, 2001). Schneider and Carter (2001) studied the shell microstructure of *Myophoriopis carinata* Bittner, 1895, from the Italian Carnian and observed that it was composed of a prismatic outer layer, a cross-lamellar middle layer, and a complex cross-lamellar inner layer.

Genus PSEUDOCORBULA E. Philippi, 1898, p. 168

Type species.—Nucula gregaria Münster, 1837, in Goldfuss, 1833–1841, p. 152.

Stratigraphic range.—Middle Triassic (Anisian)–Upper Triassic (Rhaetian). Cox and others (1969) assigned it a Lower–Upper Triassic range. According to Klets (2006), *Pseudocorbula* originated in the Induan of the Boreal domain. Dagys and Kurushin (1985) mentioned the species *Pseudocorbula gregaroides* Phillipi, 1898, from the Olenekian and Anisian of Siberia, but they did not describe the specimens. According to Diener (1923), this species was reported from the Muschelkalk (currently equivalent to the Middle Triassic of Germany).

We did not locate any Lower Triassic published record. Hautmann and Nützel (2005) indicated that the Triassic record in Cox and others (1969) was questionable since there was no material showing internal characters, and they recognized an Anisian origin for the genus. *Pseudocorbula* is widely recorded from the Middle Triassic, but many species were erroneously attributed to this genus (Márquez-Aliaga, 1985), due to its great interspecific variability. *Pseudocorbula* is only occasionally reported from the Upper Triassic and not in younger sediments, so this is a typically Triassic genus.

Paleogeographic distribution.—Tethys (Fig. 53).

Tethys domain: Middle Triassic: Hungary (Szente, 1997), China (Ling, 1988; Sha, Chen, & Qi, 1990), Poland (Assmann, 1916; Senkowiczowa, 1985), Germany (Philippi, 1898; Walther, 1927), Spain (Wurm, 1911); Anisian of Poland (Kaim, 1997), Bulgaria (Encheva, 1969); Ladinian of Spain (Márquez-Aliaga, 1983, 1985; Márquez-Aliaga, Hirsch, & López-Garrido, 1986; Llompart & others, 1987; López-Gómez & others, 1987, 1994; Márquez-Aliaga & others, 1987, 2004; Márquez-Aliaga & López-Gómez, 1989; Márquez-Aliaga & García-Gil, 1991; Márquez-Aliaga & Montoya, 1991; Márquez-Aliaga & others 2002; Márquez-Aliaga & Ros, 2002, 2003), Germany (Ürlichs, 1992), Afghanistan (Farsan, 1975); Late Triassic: Carnian of Spain (Martín-Algarra & others, 1993; Martín-Algarra, Solé de Porta, & Márquez-Aliaga, 1995; Freneix, 1999), Italy (Allasinaz, 1964); Norian of Germany (Warth, 1994; Arp & others, 2005); Rhaetian of Austria (Tomašových, 2006a), Spain (Goy & Márquez-Aliaga, 1998; Márquez-Aliaga, Plasencia, & Ros, 2005), Hungary (Vörös, 1981), Italy (S. Conti, 1954).

Paleoautoecology.—B, Is, S, FM; Sb. The mode of life of Pseudocorbula was probably similar to that assigned to Myophoricardium.

Mineralogy.—Aragonitic (J. D. Taylor, Kennedy, & Hall, 1973). *Pseudocorbula* shell microstructure is not known. Members of the superfamily Crassatelloidea have an entirely aragonitic shell (J. D. Taylor, Kennedy, & Hall, 1973).

Superfamily CARDIOIDEA Lamarck, 1809 Family CARDIIDAE Lamarck, 1809 Genus PROTOCARDIA von Beyrich, 1845, p. 17

Type species.—Cardium hillanum J. Sowerby, 1813, p. 41.

Stratigraphic range.—Upper Triassic (Norian)–Upper Cretaceous (Maastrichtian) (Cox & others, 1969; Hautmann, 2001b). Cox and others (1969) assigned it an Upper Triassic–Upper Cretaceous range. This is also recognized here, since the post-Maastrichtian records are dubious (e.g., Marincovich, 1993). Schneider (1995) considered that it ranged from the Rhaetian, but Hautmann (2001b) reported it from Norian beds.

Paleogeographic distribution.—Tethys and Circumpacific (Fig. 54).

Tethys domain: Late Triassic: Norian–Rhaetian of China (Gu & others, 1976; Guo, 1988), Iran (Hautmann, 2001b); Rhaetian of eastern Alps (Austria) (Tomašových, 2006b), Burma (Healey, 1908), Spain (Márquez-Aliaga & Ros, 2002; Márquez-Aliaga & others, 2004; Márquez-Aliaga, Plasencia, & Ros, 2005), Tibet (China) (J. Yin & Grant-Mackie, 2005), England (Castell & Cox, 1975; Ivimey-Cook & others, 1999), Italy (Allasinaz, 1962; Gelati & Allasinaz, 1964; Sirna, 1968), Oman (Arabia) (R. Hudson & Jefferies, 1961); Early Jurassic: Hettangian of England (Liu, 1995; Ivimey-Cook & others, 1999), China (Z. Huang, 1986); Hettangian–Sinemurian of China (Stiller, 2006, and references therein); Sinemurian of China (J. Chen, 1988), France and Portugal (Liu, 1995).

Circumpacific domain: Early Jurassic: Sinemurian of Japan (Hayami, 1975).

Paleoautoecology.—B, Is, S, SM; Sb. Living Cardiidae are shallow burrowers; they have short siphons and a large foot with which they can burrow rapidly (Beesley, Ross, & Wells, 1998). *Protocardia* most likely had the same mode of life, but its globose shell indicates a slow burrower (Fürsich, 1982).

Mineralogy.—Aragonitic (Schneider & Carter, 2001). Schneider and Carter (2001) studied the shell microstructure of two species of *Protocardia* from the Upper Jurassic and Cretaceous. Although the two species differ in some details, they interpreted the shell as being completely aragonitic with a prismatic outer layer, a cross-lamellar middle layer, and a complex cross-lamellar inner layer.

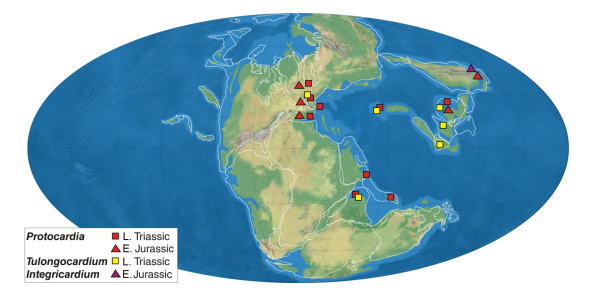


Figure 54. Paleogeographical distribution of Cardiidae (Protocardia, Tulongocardium, Integricardium). Late Triassic-Early Jurassic.

Genus TULONGOCARDIUM Chen, Chen, & Zhang in Wen & others, 1976, p. 31

Type species.—Cardium (Tulongocardium) pluriradiatum Chen, Chen, & Zhang in Wen & others, 1976, p. 32.

Remarks.—Although *Tulongocardium* was originally proposed as a subgenus of *Cardium* (Wen & others, 1976) and some authors (e.g., Z. Fang & others 2009) still maintain this taxonomic position, following Schneider (1995), we consider it as a different genus, with *Vietnamicardium* Vu Khuc, 1977a, p. 678, as the junior synonym (see discussion for *Vietnamicardium* in Genera not Included, p. 172). However, Hautmann (2001b) considered *Vietnamicardium* as a valid genus. See Schneider (1995, p. 322) for a thorough discussion.

Stratigraphic range.—Upper Triassic (Norian–Rhaetian) (Wen & others, 1976). The genus *Tulongocardium* was originally reported from the Norian of China, and its range was later extended to the Rhaetian when several species were reassigned to it (see Wen & others, 1976, p. 31–32).

Paleogeographic distribution.—Tethys (Fig. 54). Kobayashi and Tamura (1983a) also reported it from the Himalayas, southern Tibet, and Malaysia, but systematic treatment was not provided.

Tethys domain: Late Triassic: Sumatra (Krumbeck, 1914), China (Gou, 1993); Norian of China (Wen & others, 1976), Vietnam (Vu Khuc, 1977a; Vu Khuc & Huyen, 1998), Iran (Hautmann, 2001b), China (Sha, Chen, & Qi, 1990); Rhaetian of Germany (Kutassy, 1931), the Alps (Diener, 1923), Burma (Healey, 1908), Vietnam (Vu Khuc & Huyen, 1998).

Paleoautoecology.—B, Is, S, SM; Sb. Similar to Protocardia.

Mineralogy.—Aragonitic (J. D. Taylor, Kennedy, & Hall, 1973). There are no data about *Tulongocardium* shell microstructure. The shells of the living species of the superfamily Cardioidea are totally aragonitic with a cross-lamellar outer layer and complex cross-lamellar inner layer (J. D. Taylor, Kennedy, & Hall, 1973).

Genus INTEGRICARDIUM Rollier, 1912 in 1911–1918, p. 127

Type species.—Cardium dupinianum d'Orbigny, 1844 in 1844–1848, p. 26.

Remarks.—The genus *Yokoyamaina* Hayami, 1958b, p. 23, was proposed by Hayami (1958b), with *Cyrena elliptica* Yokoyama, 1904, as type species and doubtfully assigned to the family Arcticidae. Subsequently, Cox and others (1969) included it in the family Corbiculidae, also with doubts. Hayami (1975) reexamined the type species and concluded that *Yokoyamaina* was a cardiid and considered it to be a subgenus of *Integricardium*, an arrangement followed here (see discussion for *Yokoyamaina* in Genera not Included, p. 172).

Stratigraphic range.—Lower Jurassic (Hettangian)–Paleocene (Danian) (Hayami, 1975; Marincovich, 1993). Cox and others (1969) assigned it a Middle Jurassic–Upper Cretaceous range. With the inclusion of *Yokoyamaina* as subgenus of *Integricardium* (Hayami, 1975), the genus ranges back to the Hettangian. The record of *Integricardium* from the Danian (Marincovich, 1993) extend its range to the Paleocene.

Paleogeographic distribution.—Circumpacific (Fig. 54).

Circumpacific domain: Early Jurassic: Hettangian of Japan (Hayami, 1958d, 1975; Sugawara & Kondo, 2004; Kondo & others, 2006).

Paleoautoecology .- B, Is, S, MS; Sb. Similar to Protocardia.

Mineralogy.—Aragonitic (J. D. Taylor, Kennedy, & Hall, 1973). There are no data about *Integricardium* shell microstructure. See previous discussion in *Tulongocardium*.

Superfamily TELLINOIDEA Blainville, 1814 Family SOWERBYIDAE Cox, 1929 Genus RHAETIDIA Bittner, 1895, p. 222

Type species.—Rhaetidia zittelii Bittner, 1895, p. 223.

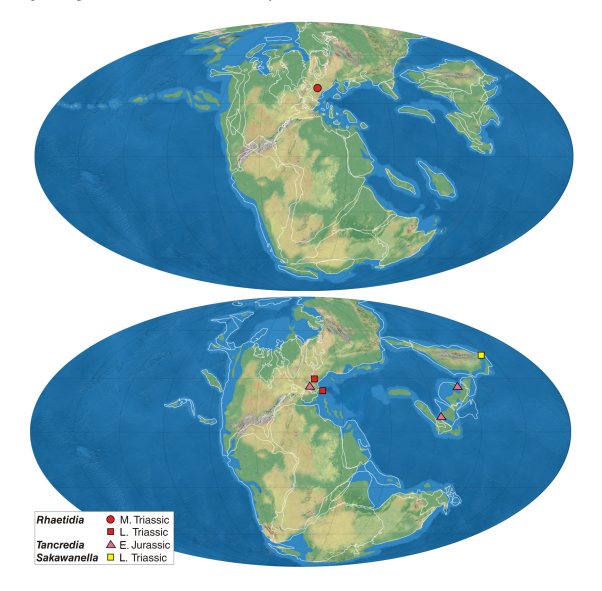


Figure 55. Paleogeographical distribution of Sowerbyidae (Rhaetidia) and Tancrediidae (Tancredia, Sakawanella). 1, Middle Triassic; 2, Late Triassic-Early Jurassic.

Remarks.—Cox and others (1969) doubtfully included *Rhaetidia* in the family Sowerbyidae due to its external morphology being similar to *Sowerbya* d'Orbigny, 1850, but *Rhaetidia* differs from *Sowerbya* by the lack of cardinal teeth and pallial sinus (Jablonski & Bottjer, 1990). Furthermore, these authors noticed a time span of 50 myr between the genera, yet another reason to doubt the relation of this genus to Sowerbyidae. We treat it in this family, pending further studies.

Stratigraphic range.—Middle Triassic (Ladinian)–Upper Triassic (Norian) (Bittner, 1895; S. Conti, 1954). Cox and others (1969) assigned it a Triassic range. According to the included species, its range was Ladinian–Norian. The type species, *R. zitteli* Bittner, 1895, was reported from the Carnian (Bittner, 1895). Other species attributed to this genus are: *R. praenuntia* (Stoppani, 1859 in 1858–1860) and *R. salomoni* Bittner, 1895, both from the Ladinian (Bittner, 1895; Rossi Ronchetti, 1959). Moreover, *Rhaetidia* was also mentioned from the Upper Triassic of Timor (Krumbeck, 1924), with *R. timorensis*

Krumbeck, 1924, and from the Norian of China, with *R. aff. zitteli* Bittner, 1895 (Cowper-Reed, 1927). These last two records were rejected by Jablonski and Bottjer (1990). In the first, they argued that Krumbeck's specimens did not belong to the genus and probably not even to the superfamily Tellinoidea. In the second, they referred to the poor preservation of the specimens studied by Cowper-Reed (1927). Jablonski and Bottjer (1990) recorded *Rhaetidia* from the upper Anisian, but this could not be corroborated in the published literature. The oldest systematic record is Ladinian.

Paleogeographic distribution.—western Tethys (Fig. 55).

Tethys domain: Middle Triassic: Ladinian of the Alps (Bittner, 1895; Rossi Ronchetti, 1959); Late Triassic: Carnian of the Alps (Bittner, 1895); Norian of Italy (S. Conti, 1954).

Paleoautoecology.—B, Is, ?S, SM; Sb. The external form of the shell indicates it probably lived infaunally; perhaps it was a shallow, nonsiphonate (or with short siphons) burrower,

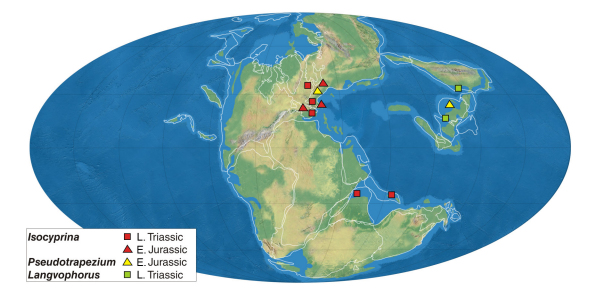


Figure 56. Paleogeographical distribution of Arcticidae (Isocyprina, Pseudotrapezium) and Trapeziidae (Langvophorus). Late Triassic-Early Jurassic.

contrary to living Tellinoidea, which often have long siphons and burrow deeply.

Mineralogy.—Aragonitic (J. D. Taylor, Kennedy, & Hall, 1973). There are no data about *Rhaetidia* shell microstructure. In living tellinoid species studied by J. D. Taylor, Kennedy, and Hall (1973), the shell was entirely aragonitic. Some species have a prismatic outer shell layer, a cross-lamellar middle layer, and a complex cross-lamellar or homogeneous inner layer; in other species, only the last two layers are present.

Family TANCREDIIDAE Meek, 1864 Genus TANCREDIA Lycett, 1850, p. 407

Type species.—Tancredia donaciformis Lycett, 1850, p. 424.

Stratigraphic range.—Lower Jurassic (Sinemurian)–Paleocene (Danian) (Marincovich, 1993; Stiller, 2006). Cox and others (1969) assigned an Upper Triassic–Upper Cretaceous range to *Tancredia*, but, curiously, all the subgenera included in it had their origin in the Jurassic. Except for some dubious records provided by Hallam (1981) from the Norian of North America and by Polubotko and Repin (1990) and Kurushin (1990) from the Upper Triassic of north-eastern Asia, *Tancredia* was not reported from the Triassic. None of the three mentioned papers figured or described the specimens, nor did they indicate the original data source. The oldest sure record is Sinemurian, and the youngest is Danian.

Paleogeographic distribution.—Tethys (Fig. 55). Although we can only confirm its record from the Sinemurian of the Tethys domain, *Tancredia* was also recorded from the Sinemurian of South America (Damborenea, 1996a) and Greenland (Liu, 1995).

Tethys domain: Early Jurassic: Sinemurian of China (J. Chen, 1988; Stiller, 2006), Vietnam (Counillon, 1908), France (Martin, 1860).

Paleoautoecology.—B, Id, S, FM; Db. The superfamily Tellinoidea includes suspensivorous and detritivorous bivalves; it is accepted that

the oldest were suspensivorous and that the detritivorous did not appear until the Early Cretaceous (Jablonski & Bottjer, 1990). S. M. Stanley (1977) considered the members of the family Tancrediidae to be fast burrowers. *Tancredia* had a pallial sinus, indicating long siphons, and therefore it was probably a deep burrower (Fürsich, 1982; Stiller, 2006).

Mineralogy.—Aragonitic (J. D. Taylor, Kennedy, & Hall, 1973). There are no data about *Tancredia* shell microstructure. See discussion in *Rhaetidia* (p. 144).

Genus SAKAWANELLA Ichikawa, 1950, p. 245

Type species.—Sakawanella triadica Ichikawa, 1950, p. 246. *Stratigraphic range.*—Upper Triassic (Carnian) (Hayami, 1975).

Cox and others (1969) and Ichikawa (1950) assigned it the same range: Upper Triassic. Later, Hayami (1975) noticed that *Sakawanella* had only been recorded from the Carnian.

Paleogeographic distribution.—Circumpacific (Fig. 55). Sakawanella was an endemic genus of Japan (Kobayashi & Tamura, 1983b).

Circumpacific domain: Late Triassic: Carnian of Japan (Ichikawa, 1950; Hayami, 1975).

Paleoautoecology.-B, Id, S, FM; Db. Similar to Trancredia.

Mineralogy.—Aragonitic (J. D. Taylor, Kennedy, & Hall, 1973). There are no data about *Sakawanella* shell microstructure. See discussion in *Rhaetidia* (p. 144).

Superfamily ARCTICOIDEA Newton, 1891 Family ARCTICIDAE Newton, 1891 Genus ISOCYPRINA Röder, 1882, p. 90

Type species.—Cardium cyreniforme Buvignier, 1852, p. 15.

Remarks.—Ivimey-Cook and others (1999) noticed that, in most specimens attributed to *Isocyprina*, the hinge is not known and that they were assigned to the genus on the basis of the external shell shape. We follow these authors in their included species list, and

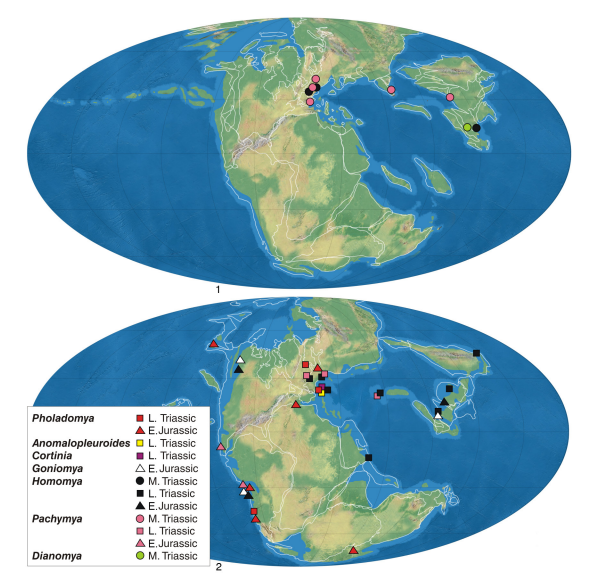


Figure 57. Paleogeographical distribution of Pholadomyidae (*Pholadomya, Anomalopleuroidea, Cortinia, Goniomya, Homomya, Pachymya, Dianomya*). 1, Middle Triassic; 2, Late Triassic–Early Jurassic.

regard *Eotrapezium* Douvillé, 1913, as a subgenus of *Isocyprina*, as Cox and others (1969) did.

Stratigraphic range.—Upper Triassic (Rhaetian)–Lower Cretaceous (Aptian) (Healey, 1908; Hayami, 1975). Cox and others (1969) assigned it an Upper Triassic–Upper Jurassic range, as did Gardner (2005), who proposed the new family Isocyprinidae. However, Hayami (1975) mentioned *I. aliquantula* (Amano, 1957) from the Aptian of Japan. The oldest record of the genus is Rhaetian (see paleogeographic distribution).

Paleogeographic distribution.—Tethys (Fig. 56). Vu Khuc and Huyen (1998) reported the species *I. ewaldi* (Bornemann, 1854) from the Rhaetian of Vietnam. It is the only record from that area, and the specimens were not figured. Damborenea (1996a) recorded *I. ancatruzi* (A. F. Leanza, 1942) from the Sinemurian of South America, but the only figured specimens (A. F. Leanza, 1942) are Pliensbachian in age. This same species is reported from the Pliensbachian of Sonora (Mexico) (Scholz, Aberhan, & González-León, 2008). However, Hallam (1981) listed *Isocyprina* from the Upper Triassic of South America and Europe. Note that Hallam mainly followed Cox and others (1969) and, particularly from South America, Körner (1937), Cox (1949), and Hayami, Maeda, and Ruiz-Fuller (1977). However, none of these three papers discussed *Isocyprina* or any species that was later assigned to it.

Tethys domain: Late Triassic: Rhaetian of the eastern Alps (Austria) (Tomašových, 2006a, 2006b), Burma (Healey, 1908), England (Wright & Benton, 1987; Ivimey-Cook & others, 1999), Tibet ("Lasha block") (J. Yin & Grant-Mackie, 2005), Spain (Gómez, Goy, & Márquez-Aliaga, 2005; Márquez & others, 2010); Early Jurassic: Hettangian of Sweden (Troedsson, 1951), northwestern Europe (Hallam, 1987), Italy (S. Conti, 1954), Portugal (J. C. Fischer & Palain, 1971), Spain (Márquez & others, 2010); Hettangian–Sinemurian of Portugal (Liu, 1995).

Paleoautoecology.—B, Is, S, FM; Sb. Most likely, *Isocyprina* was a moderately fast burrower. Since there is no pallial sinus, it was probably a shallow burrower.

Mineralogy.—Aragonitic (J. D. Taylor, Kennedy, & Hall, 1973). There are no data about the shell microstructure of *Isocyprina*. In living arcticoid species studied by J. D. Taylor, Kennedy, and Hall (1973), the shell is entirely aragonitic. Some species have a prismatic outer shell layer, a cross-lamellar middle layer, and a complex crosslamellar or homogeneous inner layer; in other species, only the last two layers are observed.

Genus PSEUDOTRAPEZIUM Fischer, 1887 in 1880–1887, p. 1075

Type species.—Cypricardia bathonica d'Orbigny, 1850, p. 308.

Stratigraphic range.—Lower Jurassic (Hettangian)–Upper Jurassic (Oxfordian) (Z. Huang, 1986; Hallam, 1987). Cox and others (1969) assigned it a Jurassic (Lias.–Portland.) range. Later, Z. Huang (1986) reported two species from the Hettangian of China that he attributed to *Pseudotrapezium*. These species are *Cypricardia triangularis* Terquem, 1855, and *Cypricardia praelonga* Terquem & Piette, 1865, both recorded from Hettangian beds of Luxemburg and France. It is difficult to know which is the youngest record of this genus, because the information at hand is rather limited. We indicate Oxfordian, following Hallam (1976, 1977, 1987).

Paleogeographic distribution.—Tethys (Fig. 56). Pseudotrapezium was recorded from the Hettangian of Morocco and the Sinemurian of France (Liu, 1995), but we did not locate information about these data.

Tethys domain: Early Jurassic: Hettangian of China and Europe (Z. Huang, 1986).

Paleoautoecology.-B, Is, S, FM; Sb. Similar to Isocyprina.

Mineralogy.—Aragonitic (J. D. Taylor, Kennedy, & Hall, 1973). There are no data about *Pseudotrapezium* shell microstructure. See discussion in *Isocyprina* (p. 146).

Family TRAPEZIIDAE Lamy, 1920 Genus LANGVOPHORUS Vu Khuc, 1977a, p. 677

Type species.—Tancredia (Hettangia) garandi Mansuy, 1912, p. 65. *Remarks.*—Vu Khuc (1977a) introduced this new genus in the

family Trapeziidae; if this is correct, *Langvophorus* extends the range of the family back to the Triassic, but it was formerly regarded as a family with Cretaceous origins (Cox & others, 1969).

Stratigraphic range.—Upper Triassic (Norian–Rhaetian) (Vu Khuc, 1977a). Vu Khuc (1977a) proposed the genus *Langvophorus*, based on the species *T. garandi* Mansuy, 1912, and including *T. chovoensis* Mansuy, 1912, both reported from the Rhaetian of Tonkin (today part of southern China and Vietnam). He assigned it a Norian–Rhaetian range.

Paleogeographic distribution.—Eastern Tethys (Fig. 56). Langvophorus is endemic to southeastern Asia (Vu Khuc & Huyen, 1998).

Tethys domain: Late Triassic: Norian–Rhaetian of China (Cowper-Reed, 1927; Vu Khuc, 1977a); Rhaetian of Vietnam (Vu Khuc, 1977a; Vu Khuc & Huyen, 1998), Tonkin (Vietnam and southern China) (Mansuy, 1912).

Paleoautoecology.—Unknown. It is difficult to interpret how Langvophorus lived, because there is not much information on its morphology nor on the sediments in which it was recorded. It had an elongated shell with a continuous pallial line; if it had siphons, these were very short. The living species of the family Trapeziidae are usually byssate, and some are nestlers (Beesley, Ross, & Wells, 1998).

Mineralogy.—Aragonitic (J. D. Taylor, Kennedy, & Hall, 1973). There are no data about *Langvophorus* shell microstructure. See discussion in *Isocyprina* (p. 146).

Superfamily PHOLADOMYOIDEA King, 1844 Family PHOLADOMYIDAE King, 1844 Genus PHOLADOMYA G. B. Sowerby I, 1823 in 1821–1825, p. unnumbered

Type species.—Pholadomya candida G. B. Sowerby I, 1823 in 1821–1825, p. unnumbered.

Stratigraphic range.—Upper Triassic (Rhaetian)–Holocene (Ivimey-Cook & others, 1999; Harper, Dreyer, & Steiner, 2005). Cox and others (1969) assigned it an Upper Triassic–Holocene range. The oldest record is from the European Rhaetian (Ivimey-Cook & others, 1999). Nowadays, this genus is represented only by its type species and has a very restricted distribution (Harper, Dreyer, & Steiner, 2005).

Paleogeographic distribution.—Tethys, Circumpacific, and Austral (Fig. 57).

Tethys domain: Late Triassic: Rhaetian of England (Ivimey-Cook & others, 1999), Lombardy (Italy) (Chiesa, 1949); Early Jurassic: Hettangian of Europe (Ivimey-Cook & others, 1999), Luxemburg (Terquem, 1855); Hettangian–Sinemurian of Europe and Morocco (Liu, 1995).

Circumpacific domain: Early Jurassic: late Hettangian of Chile (Aberhan, 2004); Sinemurian of Yukon (Canada) (Poulton, 1991), Chile (Aberhan, 1993, 2004).

Austral domain: Late Triassic: Rhaetian of Argentina (Damborenea & Manceñido, 2012). Early Jurassic: Hettangian–Sinemurian of New Zealand (H. J. Campbell & Grant-Mackie, 1995); Sinemurian of Argentina (Damborenea, 1996a; Damborenea & Manceñido, 2005b; Damborenea & Lanés, 2007).

Paleoautoecology.-B, Id, S, SM; Db. The only living species, P. candida G. B. Sowerby I, 1823 in 1821-1825, is a deep burrower with long siphons, especially slow or almost sedentary (Checa & Jiménez-Jiménez, 2003a). It has never been observed alive, and the only information available was provided by two specimens found in the 19th century (Harper, Dreyer, & Steiner, 2005). They have elongated shells with a posterior gape and a deep pallial sinus, which supports the interpretation of a deep burrower. Mehl and Rehfeld (1992) found some specimens of the Bajocian P. fidicula G. B. Sowerby, 1823 in 1821-1825, in which the proximal part of the siphon was preserved by pyritization. In addition, Fürsich (1980) found specimens of several species attributed to Pholadomya in life position, included vertically in the sediment with the posterior part up. Most likely, they had an almost sedentary mode of life, and once they were buried at their living depth, it was unlikely they could reburrow if removed from the sediment (Checa & Jiménez-Jiménez, 2003a). While the living species normally inhabits deep water, Mesozoic species lived in shallow water (Cox in Cox & others, 1969). A similar mode of life is assumed for the family Pholadomyidae.

Mineralogy.—Aragonitic (J. D. Taylor, Kennedy, & Hall, 1973). There are no studies on the shell microstructure of *Pholadomya*; however, J. D. Taylor, Kennedy, and Hall (1973) studied a fragment of the shell of *P. candida*, and it is aragonitic. In other living species of the superfamily Pholadomyoidea, they observed that the shells were always aragonitic and frequently consisted of three layers: an outer prismatic one and nacreous middle and inner layers.

Genus ANOMALOPLEUROIDEA Cox, 1964, p. 45

nom. nov. pro Anomalopleura Leonardi, 1943, p. 62, non Kleine, 1916, p. 70.

Type species.—Anomalopleura elisae Leonardi, 1943, p. 62.

Stratigraphic range.—Upper Triassic (Carnian) (Leonardi, 1943). Cox and others (1969) reported it from the Upper Triassic of Austria and mentioned *Anomalopleura* was proposed in 1948. However, Leonardi proposed this new genus in 1943 from the Carnian of Costalares in Cortina d'Ampezzo (Italy).

Paleogeographic distribution.—western Tethys (Fig. 57).

Tethys domain: Late Triassic: Carnian of Italy (Leonardi, 1943).

Paleoautoecology.—B, ?Id, S, SM; ?Db The internal characters of *Anomalopleuroidea* are not known, so it is difficult to assign, with certainty, a mode of life. It had an elongated shell and a large posterior gape, which might indicate it was a deep burrower. Moreover, the strong inflation of the shell suggests a poor burrower.

Mineralogy.—Aragonitic (J. D. Taylor, Kennedy, & Hall, 1973). There are no data about *Anomalopleuroidea* shell microstructure. See discussion in *Pholadomya* (p. 148).

Genus CORTINIA Leonardi, 1943, p. 63

Type species.—Cortinia catharinae Leonardi, 1943, p. 63.

Stratigraphic range.—Upper Triassic (Carnian) (Leonardi, 1943). Cox and others (1969) reported it from the Upper Triassic of Austria and mentioned it was proposed in 1948. However, Leonardi proposed this new genus in 1943 from the Carnian of Costalares in Cortina d'Ampezzo (Italy).

Paleogeographic distribution.—western Tethys (Fig. 57). Gou (1993) proposed a new species from the Upper Triassic of China, which he doubtfully attributed to *Cortinia*, *C.? elongata* Gou, 1993.

Tethys domain: Late Triassic: Carnian of Italy (Leonardi, 1943).

Paleoautoecology.—B, ?Is, S, SM; ?Sb. Cortinia internal characters are unknown, so it is difficult to know if it was a shallow or deep burrower. The shell is small and has no gape, thus it was probably a shallow burrower.

Mineralogy.—Aragonitic (J. D. Taylor, Kennedy, & Hall, 1973). There are no data about the shell microstructure of *Cortinia*. See discussion in *Pholadomya* (p. 148).

Genus GONIOMYA Agassiz, 1841 in 1840-1845, p. 1

Type species.—Mya angulifera J. Sowerby, 1819, p. 46.

Stratigraphic range.—Lower Jurassic (Hettangian)–Upper Cretaceous (Maastrichtian) (Hayami, 1964; Abdel-Gawad, 1986). Cox and others (1969) assigned it a Jurassic–Upper Cretaceous range and also recorded it from the Eocene. The earliest records are from Upper Cretaceous (Maastrichtian) (e.g., Abdel-Gawad, 1986), but we did not find any record from the Eocene. *Goniomya* is especially reported from the Middle Jurassic and Cretaceous, but it originated in the lower Jurassic (Hayami, 1964). According to Septfontaine (1995), the syntypes of the species *G. renevieri* Jeannet, 1913, deposited in the Cantonal Museum of Geology of Lausanne, are from the Hettangian of Switzerland. In the Lower Jurassic, it was also recorded from the Pliensbachian and Toarcian (Riegraf, 1977; Monari, 1994; Liu, 1995; Harries & Little, 1999; Fürsich and others, 2001; Gahr, 2002; Scholz, Aberhan, & González-León, 2008).

Paleogeographic distribution.—Circumpacific (Fig. 57). In the time interval analyzed here, the genus was recorded from Vietnam and also from the Sinemurian of the Canadian Arctic islands (Aberhan, Hrudka, & Poulton, 1998), with *G. rhombifera* (Goldfuss), but the specimens were not figured or discussed systematically. It was also recorded from the same stage from South America (Damborenea, 1996a; Damborenea & Manceñido, 2005b; Damborenea & Lanés, 2007) and from England (Liu, 1995).

Circumpacific domain: Early Jurassic: middle–late Hettangian of Chile (Aberhan, 2004); Hettangian–Sinemurian of Vietnam (Hayami, 1964; Vu Khuc & Huyen in Sato & Westermann, 1991); Sinemurian of Yukon (Canada) (Poulton, 1991).

Paleoautoecology.—B, Id, S, SM; Db. *Goniomya*, like most members of the family Pholadomyidae, was probably a deep burrower. It had a very elongated shell, inequilateral, moderately inflated, with anterior and posterior gapes and a pallial sinus, characteristics that indicate this mode of life unequivocally. Fürsich (1980, 1982) found specimens belonging to *Goniomya* in life position, with the shell upright in the sediment with the posterior part upward.

Mineralogy.—Aragonitic (J. D. Taylor, Kennedy, & Hall, 1973). There are no data about the shell microstructure of *Goniomya*. See discussion in *Pholadomya* (p. 148).

Genus HOMOMYA Agassiz, 1843 in 1840-1845, p. 154

Type species.—Mactra gibbosa J. Sowerby, 1813, p. 91.

Stratigraphic range.—Middle Triassic (Anisian)–Lower Cretaceous (Valanginian) (Mader, 1982; Lazo, 2007b). Cox and others (1969, p. 832) pointed out that Muschelkalk species attributed to Homomya fit better into Pachymya (Arcomya). However, they assigned it a Middle Triassic (Muschelkalk)–Upper Jurassic (Portland.) range. It is difficult to refer specimens to these and other similar genera (such as Pleuromya Agassiz, 1842 in 1840-1845), because frequently the key internal characters are not observed (Neri & Posenato, 1985). Although these authors were aware of the indications given by Cox and others (1969) regarding the Muschelkalk species, they decided to refer their specimens to "Homomya," following the original conception of the genus rather than Cox's review, because it is not clear which species were transferred to Pachymya (Arcomya). There are no proper discussions of this question. Most Muschelkalk species traditionally attributed to Homomya were relocated in other genera, but there are two species, Homomya albertii (Goldfuss, 1841 in 1833-1841) and Homomya impressa (Alberti, 1864), which are still considered in the literature as belonging to this genus (see e.g., Gall, 1971; Busnardo, 1975; Mader, 1982).

According to Mader (1982), the oldest record of *Homomya* is from the Anisian. Regarding the youngest one, Lazo (2007b) considered that *Pholadomya sanctaecrucis* Pictet & Campiche, 1864 in 1864–1867, and *Pholadomya valangiensis* Pictet & Campiche, 1864 in 1864–1867, from the Valanginian of various localities, are synonyms and belong to *Homomya*, thus the range was extended here.

Paleogeographic distribution.—Tethys and Circumpacific (Fig. 57).

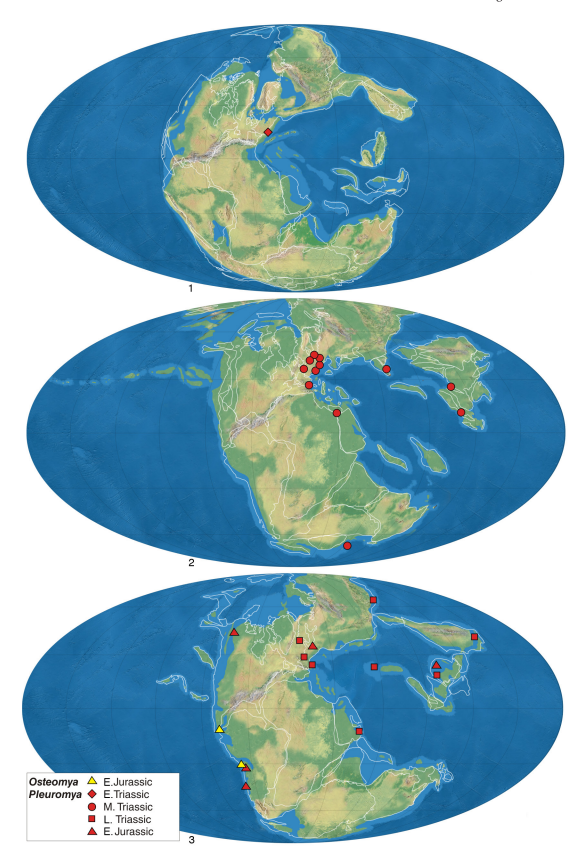


Figure 58. Paleogeographical distribution of Chaenomyidae (Osteomya) Pleuromyidae (Pleuromya). 1, Early Triassic; 2, Middle Triassic; 3, Late Triassic-Early Jurassic.

Tethys domain: Middle Triassic: Anisian of Germany (Mader, 1982), France (Gall, 1971), southestern China (Z. Fang & others, 2009); Late Triassic: Yunnan province (China) (Gou, 1993; Hautmann, 2001b); Carnian of Lombardy (Italy) (Allasinaz, 1962, 1966); Norian of Oman (R. Hudson & Jefferies, 1961; Hautmann, 2001b), Indochina (Hautmann, 2001b); Norian–Rhaetian of Iran (Hautmann, 2001b); Rhaetian of the eastern Alps (Austria) (Tomašových, 2006a, 2006b; Siblík & others, 2010), Hungary (Vörös, 1981), Italy (Sirna, 1968); Early Jurassic: Hettangian of southern China (Z. Huang, 1986).

Circumpacific domain: Late Triassic: Carnian of Japan (Nakazawa, 1956; Hayami, 1975); Early Jurassic: ?Hettangian of Yukon (Canada) (Poulton, 1991); late Sinemurian of Chile (Aberhan, 2004).

Paleoautoecology.—B, Id, S, SM; Db. *Homomya* is externally very similar to *Pholadomya* and has a deep pallial sinus, so it most probably was a deep burrower (Runnegar, 1974). Fürsich (1980) reported *Homomya* sp. from the Bathonian of Poland in life position, buried with the posterior part up.

Mineralogy.—Aragonitic (J. D. Taylor, Kennedy, & Hall, 1973). There are no data about *Homomya* shell microstructure. See discussion in *Pholadomya* (p. 148).

Genus PACHYMYA J. de C. Sowerby, 1826, p. 1

Type species.—Pachymya gigas J. de C. Sowerby, 1826, p. 2.

Stratigraphic range.—Middle Triassic (Anisian)–Upper Cretaceous (Turonian) (Cox & others, 1969; Sha, Chen, & Qi, 1990). Cox and others (1969) assigned it a Middle Triassic–Upper Cretaceous (Turonian) range. They included two subgenera, *P. (Pachymya)* and *P. (Arcomya)* Roemer, 1839, from the Cenomanian and Turonian respectively. The first record for *Pachymya* is from the Anisian of China (Sha, Chen, & Qi, 1990). There are no records after the Turonian, so we follow Cox and others (1969) for the top of the stratigraphic range.

Paleogeographic distribution.—Tethys and Circumpacific (Fig. 57). Cox and others (1969) considered it to be cosmopolitan, but during our study interval, it was only distributed in the Tethys and Circumpacific domains.

Tethys domain: Middle Triassic: Anisian of China (Sha, Chen, & Qi, 1990; Komatsu, Chen, & others, 2004); Ladinian of Spain (Márquez-Aliaga, 1985), Afghanistan (Farsan, 1975), Germany (Kutassy, 1931), southern Alps (Galdieri, 1908; Diener, 1923), Austria (Salomon, 1895); Late Triassic: ?Iran (Hautmann, 2001b); Carnian of the southern Alps (Galdieri, 1908); Norian of Hungary (Kutassy, 1931).

Circumpacific domain: Early Jurassic: late Sinemurian of Chile (Aberhan, 1993, 2004); Sinemurian of ?Sonora (Mexico) (Damborenea in Damborenea & González-León, 1997; Scholz, Aberhan, & González-León, 2008).

Paleoautoecology.—B, Id, S, SM; Db. *Pachymya* was most likely a deep burrower, indicated by its elongated shell and posterior gape (see Cox & others, 1969, p. 836, fig. 2b), although the pallial sinus is shallow or absent (Cox & others, 1969, p. 834).

Mineralogy.—Aragonitic (J. D. Taylor, Kennedy, & Hall, 1973). There are no data about the shell microstructure of *Pachymya*. See discussion in *Pholadomya* (p. 148).

Genus DIANOMYA Guo, 1985, p. 229, 272

Type species.—Dianomya lirulata Guo, 1985, p. 229.

Remarks.—Dianomya was originally included in the family Pholadomyidae by Guo (1985), although Z. Fang and others (2009) suggested it may be better placed in Laternulidae.

Stratigraphic range.—Middle Triassic (Anisian) (Guo, 1985). Guo (1985) proposed *Dianomya* from the Anisian beds (Baifeng Formation) of China.

Paleogeographic distribution.—Eastern Tethys (Fig. 57).

Tethys domain: Middle Triassic: Anisian of southwestern China (Yunnan province) (Guo, 1985).

Paleoautoecology.—B, Id, S, SM; Db. *Dianomya*, like most Pholadomyidae, was probably a deep burrower. It had a very elongated shell, inequilateral, moderately inflated, with anterior and posterior gapes, characteristics that indicate this mode of life. In the diagnosis of the genus, the presence of a pallial sinus was not mentioned, and it is not seen in the figures offered by Guo (1985). *Dianomya* is the first Mesozoic pholadomyoid with oblique posterior costae; this type of ornamentation (Hautmann, Aghababalou, & Krystyn, 2011) could aid during burrowing.

Mineralogy.—Aragonitic (J. D. Taylor, Kennedy, & Hall, 1973). There are no data about the shell microstructure of *Dianomya*. See discussion in *Pholadomya* (p. 148).

Family CHAENOMYIDAE Waterhouse, 1966 Genus OSTEOMYA Moesch, 1874 in 1874–1875, p. 19

Type species.—Mya dilata Phillips, 1829, p. 155.

Remarks.—Cox and others (1969) included this genus in the family Pholadomyidae. Following Runnegar (1974), we consider *Osteomya* to be in the family Chaenomyidae.

Stratigraphic range.—Lower Jurassic (Sinemurian)–Middle Jurassic (Callovian) (Cox & others, 1969; Scholz, Aberhan, & Gónzalez-León, 2008). Cox and others (1969) assigned it a Toarcian–Callovian range. Later, Aberhan (2004) and Scholz, Aberhan, and González-León (2008), respectively, reported *O. dilata* (Phillips, 1829) from the Sinemurian of Chile and Mexico. Guo (1985) proposed a new subgenus, *Osteomya (Yunnanomya)*, from the Anisian of China. This is not taken into account, because Z. Fang and others (2009) regarded it as a junior synonym of *Homomya* Agassiz, 1843 in 1840–1845.

Paleogeographic distribution.—Circumpacific (Fig. 58). During the study interval, the genus was only mentioned from the Circumpacific domain, but, later in the Jurassic, Osteomya was also distributed in the Tethys domain (Jaitly, 1986; Fürsich & others, 2001; Gahr, 2002). Liu (1995) and Damborenea (1996a) reported it from the Hettangian and Sinemurian of Greenland and South America, respectively, but there are no published records to corroborate these data.

Circumpacific domain: Early Jurassic: late Sinemurian of Chile (Aberhan, 2004); Sinemurian of Sonora (Mexico) (Scholz, Aberhan, & González-León, 2008).

Paleoautoecology.—B, Id, S, SM; Db. The characteristics of the *Osteomya* shell suggest that, like other members of this superfamily, it was a deep burrower. It had an elongated shell, strongly inequilateral and a very broad posterior gape (Cox & others, 1969, p. 833). In the specimens figured by Runnegar (1974, pl. 3,17–20), a shallow

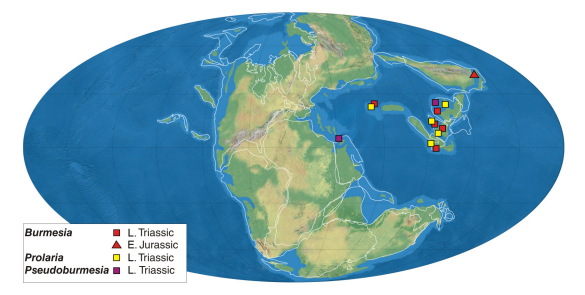


Figure 59. Paleogeographical distribution of Burmesiidae (Burmesia, Prolaria, Pseudoburmesia). Late Triassic-Early Jurassic.

pallial sinus is observed, which indicates that it probably was an intermediate burrower when compared to *Pholadomya*.

Mineralogy.—Aragonitic (J. D. Taylor, Kennedy, & Hall, 1973). There are no data about the shell microstructure of *Osteomya*. See discussion in *Pholadomya* (p. 148).

Family PLEUROMYIDAE Dall, 1900 in 1896–1900 Genus PLEUROMYA Agassiz, 1845 in 1840–1845, p. 231

Type species.—Mya gibbosa J. de C. Sowerby, 1823, p. 19.

Stratigraphic range.-Lower Triassic (Olenekian)-Lower Cretaceous (Valanginian) (Sha & Fürsich, 1994; Posenato, 2008a). Cox and others (1969) assigned it a Triassic-Lower Cretaceous range. We did not find reliable records after the Lower Cretaceous; the youngest record of Pleuromya is Valanginian (Sha & Fürsich, 1994), although the specimens found by these authors are preserved as internal molds. There is some uncertainty concerning the oldest record of this genus. The members of the Lower Triassic "Myacites" group were transferred to various genera, including Pleuromya (see Neri & Posenato, 1985, for a discussion of this issue), but many of the specimens were assigned to this genus based on internal molds that did not show enough detail of characters to make a good generic assignment. Moreover, it is frequently difficult to distinguish between Pleuromya and Homomya (Neri & Posenato, 1985). According to these authors, the attribution of P. elongata (Schlotheim, 1820) to Pleuromya is correct if one considers the original concept of the genus. This same species is also recorded from the Triassic by Leonardi (1935) and Broglio-Loriga and others (1990), but the specimens in all cases are poorly preserved. Fraiser and Bottjer (2007a) and Posenato (2008a) reported Pleuromya from the "Gastropod Oolite" member of the Werfen Formation in Italy, but they did not figure or discuss the material systematically. Therefore, we considered the oldest Triassic record with certain reservations. From the Middle Triassic onward, Pleuromya was well represented (see paleogeographic distribution).

Paleogeographic distribution.—Tethys, Circumpacific, and Austral (Fig. 58). The genus was also mentioned in the Boreal domain, but not figured or described, from the Middle Triassic (Dagys & Kurushin, 1985) and Late Triassic (Kurushin, 1990; Polubotko & Repin, 1990).

Tethys domain: Early Triassic: Olenekian of ?Italy (Neri & Posenato, 1985; Broglio-Loriga & others, 1990; Fraiser & Bottjer, 2007a; Posenato, 2008a); Middle Triassic: Anisian of China (Gu & others, 1976; Sha, Chen, & Qi, 1990), Italy (Posenato, 2008b), Bulgaria (Encheva, 1969), northern Vietnam (Komatsu, Huyen, & Huu, 2010); Buntsandstein of France (Gall, 1971); Muschelkalk of France (Márquez-Aliaga & others, 2002), Spain (Márquez-Aliaga, 1983, 1985; Márquez-Aliaga, Hirsch, & López-Garrido, 1986; Budurov & others, 1991; López-Gómez & others, 1994; Márquez-Aliaga & Martínez, 1996; Márquez-Aliaga & others, 2002), Hungary (Szente, 1997), Poland (Senkowiczowa, 1985), Germany (Fuchs & Mader, 1980; Hagdorn, 1982); Ladinian of Italy (Posenato, 2002), China (Sha, Chen, & Qi, 1990), Afghanistan (Farsan, 1975), Israel (Lerman, 1960), northern Vietnam (Komatsu, Huyen, & Huu, 2010); Late Triassic: China (Gu & others, 1976; J. Chen, 1988); Carnian of China (Gu & others, 1976; Sha, Chen, & Qi, 1990), Italy (Desio, Rossi Ronchetti, & Vigano, 1960); Norian of Iran (Hautmann, 2001b), Oman (R. Hudson & Jefferies, 1961); Rhaetian of the Alps (Austria) (Tomašových, 2006a), Yunnan (China) (Guo, 1985), ?England (Ivimey-Cook & others, 1999), Italy (S. Conti, 1954; Sirna, 1968); Early Jurassic: Sinemurian of Sweden (Troedsson, 1951); ?Hettangian, Sinemurian of China (J. Chen, 1988; Stiller, 2006).

Circumpacific domain: Late Triassic: Carnian of Japan (Nakazawa, 1956; Hayami, 1975); Early Jurassic: early and late Hettangian of Chile (Aberhan, 2004); ?Hettangian, Sinemurian of Yukon (Canada) (Poulton 1991); Sinemurian of Chile (Aberhan, 1993).

Austral domain: Middle Triassic: Ladinian of New Zealand (Marwick, 1953); Early Jurassic: Sinemurian of Argentina (Damborenea & Manceñido, 2005b). Boreal domain: Late Triassic: Carnian of Primorie (Kiparisova, 1972).

Paleoautoecology.—B, IP, S, SM; Db. *Pleuromya* is characterized by an elongated shell, with siphonal and pedal gapes and a deep pallial sinus. These features indicate it had long siphons and burrowed deep in the sediment, like most members of the superfamily Pholadomyoidea. Its shell is moderately to strongly inflated; it would have been a slow burrower. Fürsich (1982) found *Pleuromya* in life position, oriented vertically with the posterior part toward the sediment surface. He compared the mode of life of *Pleuromya* with the living species of *Mya* Linnaeus, 1758, which live deeply buried in the adult stage and are unable to return to their life position if unearthed.

Mineralogy.—Aragonitic (J. D. Taylor, Kennedy, & Hall, 1973). There are no data about the shell microstructure of *Pleuromya*. See discussion in *Pholadomya* (p. 148).

Family BURMESIIDAE Healey, 1908 Genus BURMESIA Healey, 1908, p. 58

Type species.—Burmesia latouchii Healey, 1908, p. 58.

Stratigraphic range.—Upper Triassic (Carnian)–Lower Jurassic (Hettangian) (Hayami, 1975; J. Chen, 1985). Cox and others (1969) assigned it an Upper Triassic–Lower Jurassic range. *Burmesia* was reported from the Norian, Rhaetian, and Hettangian (see paleogeographic distribution). However, Huyen and Vu Khuc (in Sato & Westermann, 1991) mentioned it from the Pliensbachian. The specimens were neither figured nor discussed systematically, and we did not locate any publication to confirm these records.

Paleogeographic distribution.—Eastern Tethys and Circumpacific (Fig. 59).

Tethys domain: Late Triassic: Carnian of China (J. Chen, 1985); Norian of Indonesia (Diener, 1923), China (Gu & others, 1976; Wen & others, 1976; J. Chen, 1985; Gou, 1993; Y. Li, 1994), Iran (Hautmann, 2001b), Vietnam (Vu Khuc & Huyen, 1998; Hautmann, 2001b); Rhaetian of Burma (Healey, 1908), Iran (Hautmann, 2001b).

Circumpacific domain: Early Jurassic: Hettangian of Japan (Hayami, 1959, 1975; Hayami in Sato & Westermann, 1991).

Paleoautoecology.—B, Id, S, SM; Db. Although neither pallial line nor adductor muscles are observed, we assume *Burmesia* had a mode of life similar to other members of the superfamily Pholadomyoidea.

Mineralogy.—Aragonitic (J. D. Taylor, Kennedy, & Hall, 1973). There are no data about *Burmesia* shell microstructure. See discussion in *Pholadomya* (p. 148).

Genus PROLARIA Healey, 1908, p. 60

Type species.—Prolaria sollasi Healey, 1908, p. 60.

Stratigraphic range.—Upper Triassic (Norian–Rhaetian) (Hautmann, 2001b). Cox and others (1969) assigned it a Rhaetian range, as the specimens described by Krumbeck (1914) from the Norian were not considered. Later, *Prolaria* was repeatedly recorded as being from the Norian (see paleo-geographic distribution).

Paleogeographic distribution.—Eastern Tethys (Fig. 59).

Tethys domain: Late Triassic: Norian of China (Gu & others, 1976; Hautmann, 2001b), Sumatra (Indonesia) (Krumbeck, 1914),

Vietnam (Vu Khuc & Huyen, 1998; Hautmann, 2001b); Rhaetian of Burma (Healey, 1908), Iran (Hautmann, 2001b), Indochina (Diener, 1923).

Paleoautoecology.-B, Id, S, SM; Db. Similar to Burmesia.

Mineralogy.—Aragonitic (J. D. Taylor, Kennedy, & Hall, 1973). There are no data about *Prolaria* shell microstructure. See discussion in *Pholadomya* (p. 148).

Genus PSEUDOBURMESIA Gou, 1993, p. 20, 26

Type species.—Pseudoburmesia maantangensis Gou, 1993, p. 21. Remarks.—According to Sha, Chen, and Qi (1990), Pseudoburmesia and its type species were proposed by Gou in 1980, but they do not list the reference. Gou described the genus and its type species as being new in 1993. Sha, Chen, and Qi (1990) proposed a new species, P. yushuensis Sha, Chen, & Qi, 1990, and also included other species: P. posteroradiata (Cox, 1924) and P. qinghaiensis (Lu, 1981). Z. Fang and others (2009, p. 142) indicated: "It may be better to regard this genus as a synonym of Anomalopleuroidea Cox," but no further explanation was given.

Stratigraphic range.—Upper Triassic (Carnian) (Sha, Chen, & Qi, 1990). According to Gou (1993), *Pseudoburmesia* had an Upper Triassic range. Lu (1981) reported *Burmesia? qinghaiensis* Lu, 1981, from the Norian, and Cox (1924) reported *Burmesia? posteroradiata* from the Carnian. Sha, Chen, and Qi (1990), taking into account all the species included in *Pseudoburmesia*, assigned it a Carnian range. *Paleogeographic distribution.*—Tethys (Fig. 59).

Tethys domain: Late Triassic: Carnian of China (Sha, Chen, & Qi, 1990), Jordan (Cox, 1924; Sha, Chen, & Qi, 1990).

Paleoautoecology.-B, Id, S, SM; Db. Similar to Burmesia.

Mineralogy.—Aragonitic (J. D. Taylor, Kennedy, & Hall, 1973). There are no data about *Pseudoburmesia* shell microstructure. See discussion in *Pholadomya* (p. 148).

Superfamily CERATOMYOIDEA Arkell, 1934 in 1929–1937 Family CERATOMYIDAE Arkell, 1934 in 1929–1937 Genus PTEROMYA Moore, 1861, p. 505

Type species.—Pteromya crowcombeia Moore, 1861, p. 506.

Stratigraphic range.—Upper Triassic (Rhaetian)–Lower Jurassic (Hettangian) (Cox, 1963). Cox and others (1969) assigned it a Rhaetian–Hettangian range, following his comprehensive review of *Pteromya* (Cox, 1963).

Paleogeographic distribution.—Western Tethys (Fig. 60). Damborenea (1996a) mentioned *Pteromya* from the southeastern Pacific margin, but it is not a systematic paper. As far as we know, it has only been recorded from Europe.

Tethys domain: Late Triassic: Rhaetian of England (Vokes, 1945; Cox, 1963; Ivimey-Cook & others, 1999), Spain (López-Gómez, Goy, & Márquez-Aliaga, 2005; Márquez-Aliaga & others, 2010); Early Jurassic: Hettangian of Spain (López-Gómez, Goy, & Márquez-Aliaga, 2005; Márquez-Aliaga & others, 2010), England (Cox, 1963; Ivimey-Cook & others, 1999), France (Freneix & Cubaynes, 1984).

Paleoautoecology.—B, Is, S, SM; Sb. According to Runnegar (1974), *Pteromya* was an intermediate burrower. The members of this family bury at different depths, for example, *Ceratomya* Sandberger, 1864, is a shallow burrower, and it is characterized by a robust shell, without siphonal gapes, with subequal adduc-

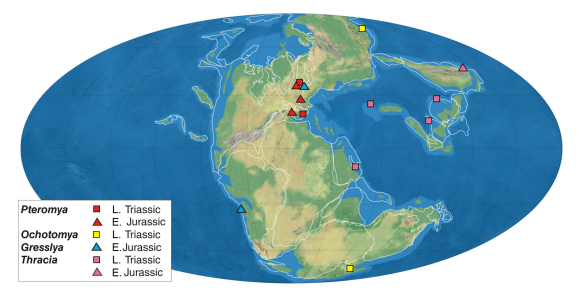


Figure 60. Paleogeographical distribution of Ceratomyidae (Pteromya, Ochotomya, Gresslya) and Thraciidae (Thracia). Late Triassic-Early Jurassic.

tor scars, deep pedal retractors insertions, and with a shallow pallial sinus (Runnegar, 1974). However, *Gresslya* Agassiz, 1843 in 1840–1845, had a deep pallial sinus and posterior gape; it is interpreted as a deep burrower. The shell of *Pteromya* was not as robust as that of *Ceratomya*, and it is more elongated and less inflated, but muscle scars or a pallial sinus are not observed. Since the siphons would have been short, we assign it a shallow burrower mode of life.

Mineralogy.—Aragonitic (J. D. Taylor, Kennedy, & Hall, 1973). There are no data about *Pteromya* shell microstructure. See discussion in *Pholadomya* (p. 148).

Genus OCHOTOMYA Polubotko, 1966, p. 13

Type species.—Ochotomya anyuensis Polubotko, 1966, p. 16.

Stratigraphic range.—Upper Triassic (Norian–Rhaetian) (Polubotko, 1966). Polubotko (1966) proposed *Ochotomya* and included three species: *O. anyuensis* Polubotko, *O. anmandykanensis* (Tuchkov), and *O. terechovae* Polubotko. He reported *Ochotomya* from the Norian–Rhaetian beds of northeastern Russia and noticed that it was probably distributed through Russia, Japan, New Zealand, and North America, and perhaps Italy during the Lower and Middle Triassic. We did not find any record from Lower or Middle Triassic; all the records date from the Upper Triassic (Grant-Mackie, 1981; Okuneva, 1985; Kurushin, 1990; Polubotko & Repin, 1990; Mac-Farlan, 1998; Polubotko, 2010).

Paleogeographic distribution.—Austral and Boreal (Fig. 60).

Austral domain: Late Triassic: Norian–Rhaetian of New Zealand (Grant-Mackie, 1981; MacFarlan, 1998).

Boreal domain: Late Triassic: Norian–Rhaetian of northeastern Russia (Polubotko, 1966, 2010; Okuneva, 1985; Kurushin, 1990; Polubotko & Repin, 1990).

Paleoautoecology.—B, Is, S, SM; Sb. Ochotomya is very similar in external form to Ceratomya Sandberger, 1864 (see mode of life for Pteromya, p. 153). Ochotomya probably was a shallow burrower; it had a robust and smooth globular shell and is usually preserved articulated (Grant-Mackie, 1981).

Mineralogy.—Aragonitic (J. D. Taylor, Kennedy, & Hall, 1973). There are no data about *Ochotomya* shell microstructure. See discussion in *Pholadomya* (p. 148).

Genus GRESSLYA Agassiz, 1843 in 1840-1845, p. 202

Type species.—Lutraria gregaria von Zieten, 1833 in 1830–1833, p. 85.

Stratigraphic range.-Lower Jurassic (upper Hettangian)-Upper Jurassic (Kimeridgian) (Tate & Blake, 1876; Arkell, 1933 in 1929-1937). Cox and others (1969) assigned it a Lower-Upper Jurassic range, and Sepkoski (2002) did the same, but he was more precise: upper Hettangian-upper Tithonian, refering to Hallam (1987) and Kelly (1984). Gresslya was reported from Hettangian-Sinemurian beds in several localities (Hallam, 1987; Liu, 1995; Damborenea, 1996a; Damborenea & Manceñido, 2005b), but none of these papers included a systematic treatment of the specimens; the only paper with figures is Tate and Blake (1876), who recorded G. galathea Agassiz from angulata to oxynotum zones (upper Hettangian-upper Sinemurian). Aberhan (2004) recorded and figured Gresslya sp. A from Sinemurian beds of Chile. Damborenea (in Damborenea & González-León, 1997) recorded Gresslya rotundata (Phillips) from Sinemurian-Toarcian beds of Europe and Canada, but no figures of the material were provided. Regarding the top of the Gresslya range, Kelly (1984) did not consider it in his study, but the youngest record is Kimmeridgian (Arkell, 1933 in 1929-1937). Some authors (e.g., Sha & others, 2009) mentioned it from the Lower Cretaceous, but we did not find any systematic study to corroborate this record.

Paleogeographic distribution.—Eastern Tethys and Circumpacific (Fig. 60). Gresslya was reported from Canada (Poulton, 1991), but he attributed the specimens to this genus only doubtfully.

Tethys domain: Early Jurassic: Hettangian–Sinemurian of England (Tate & Blake, 1876).

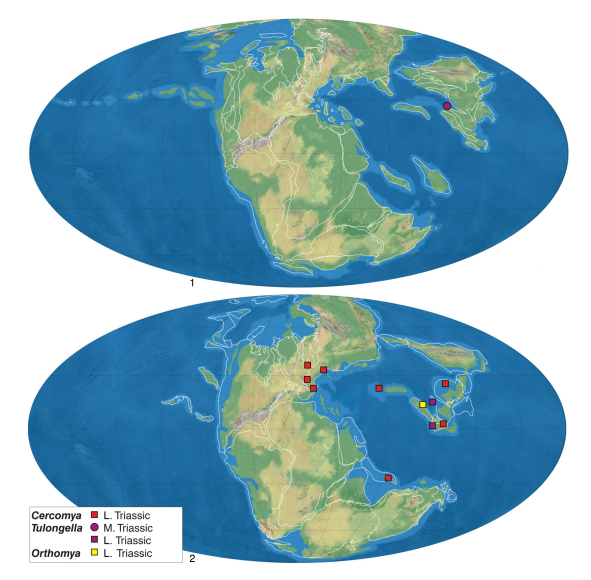


Figure 61. Paleogeographical distribution of Laternulidae (Cercomya, Tulongella, Orthomya). 1, Middle Triassic; 2, Late Triassic.

Circumpacific domain: Early Jurassic: Sinemurian of Chile (Aberhan, 2004).

Paleoautoecology.—B, Id, S, SM; Db. *Gresslya* had a deep pallial sinus and a posterior gape; it is interpreted as a deep burrower (Runnegar, 1974). A deep pallial sinus indicates it had long siphons, although its high inflation might influence its ability to penetrate into the sediment (Pugaczewska, 1986).

Mineralogy.—Aragonitic (J. D. Taylor, Kennedy, & Hall, 1973). There are no data about *Gresslya* shell microstructure. See discussion in *Pholadomya* (p. 148).

Superfamily PANDOROIDEA Rafinesque, 1815 Family THRACIIDAE Stoliczka, 1870 in 1870–1871 Genus THRACIA Leach in J. de C. Sowerby, 1823, p. 20

Type species.—*Mya pubescens* Pulteney, 1799, p. 27.

Stratigraphic range.—Upper Triassic (Norian)–Recent. Although Cox and others (1969) assigned it a Jurassic–Holocene range, *Thracia* was also recorded from the Upper Triassic. There are three species recorded from the Upper Triassic: *T. prisca* Healey, 1908, *T. proavita* R. Hudson & Jefferies, 1961, and *T. applanata* Krumbeck, 1913 (R. Hudson & Jefferies, 1961). Healey (1908) proposed *T. prisca* Healey, 1908, from the Rhaetian of Burma. Later, Gu and others (1976) quoted it from contemporary sediments of China. *Thracia* was also mentioned from the Upper Triassic of Iran (Hautmann, 2001b), the Norian of Oman (R. Hudson & Jefferies, 1961), Indochina (Kutassy, 1931), and Vietnam (Vu Khuc & Huyen, 1998).

Paleogeographic distribution.—Tethys and Circumpacific (Fig. 60). Tethys domain: Late Triassic: Burma (Healey, 1908), China (Gu & others, 1976; Gou, 1993), Iran (Hautmann, 2001b), Oman (R. Hudson & Jefferies, 1961), Indochina (Kutassy, 1931).

Circumpacific domain: Early Jurassic: Hettangian of Japan (Hayami, 1958c, 1975; Kondo & others, 2006).

Paleoautoecology.—B, Id, S, SM; Db. Living species of *Thracia* are deep burrowers with long siphons, and they produce mucus chan-

nels around the siphons, to avoid exposure to the surface (Beesley, Ross, & Wells, 1998). In many of the fossil species, a deep pallial sinus and siphonal gapes are observed; so they probably lived in the same manner. Fürsich (1980) recorded *Thracia* specimens in life position, and they were included in the substrate vertically with the posterior part up.

Mineralogy.—Aragonitic (J. D. Taylor, Kennedy, & Hall, 1973). The shell of living species of *Thracia* is completely aragonitic and is made of two layers, both of homogeneous microstructure (J. D. Taylor, Kennedy, & Hall, 1973). The shell of the Mesozoic specimens contained nacreous structure (Runnegar, 1974).

Family LATERNULIDAE Hedley, 1918 Genus CERCOMYA Agassiz, 1843 in 1840–1845, p. 143

Type species.—Cercomya pinguis Agassiz, 1843 in 1840–1845, p. 145.

Stratigraphic range.—Upper Triassic (Carnian)–Upper Cretaceous (Maastrichtian). Cox and others (1969) assigned it an Upper Triassic–Cretaceous range. The oldest mention of *Cercomya* dates from the Carnian (Sha, Chen, & Qi, 1990) and the youngest from the Maastrichtian (Abdel-Gawad, 1986).

Paleogeographic distribution.—Tethys (Fig. 61). Damborenea (1996a) mentioned it from the Hettangian–Sinemurian of the southeastern Pacific margin, but the first figured specimens are Pliensbachian (Aberhan, 1994a; Scholz, Aberhan, & González-León, 2008).

Tethys domain: Late Triassic: Carnian of China (Sha, Chen, & Qi, 1990), ?Italy (Allasinaz, 1966); Norian–Rhaetian of Iran, the Alps, Burma, Vietnam, and Yunnan (China) (Hautmann, 2001b); Rhaetian of Tibet (so-called Lhasa block) (J. Yin & Grant-Mackie, 2005), England (Ivimey-Cook & others, 1999), ?Hungary (Vörös, 1981).

Paleoautoecology.—B, Id, S, SM; Db. The mode of life of Cercomya was probably similar to Laternula species, the only living genus of this family. Laternula is a deep burrower that has long siphons and anterior and posterior gapes. In juvenile stages, it burrows and buries, but as an adult, it cannot rebury because the foot is atrophied (Beesley, Ross, & Wells, 1998). A pallial line is not observed in Cercomya, so we cannot estimate the size of its siphons, but a posterior siphonal gape is present.

Mineralogy.—Aragonitc (J. D. Taylor, Kennedy, & Hall, 1973). There are no data about the shell microstructure of *Cercomya*. J. D. Taylor, Kennedy, and Hall (1973) studied the shells of various current species of the superfamily Pandoroidea, and, in all except in those belonging to the family Thraciidae, there are three shell layers: a prismatic outer layer and nacreous middle and inner layers.

Genus TULONGELLA Chen & J. Chen in Wen & others, 1976, p. 68

Type species.—Tulongella xizangensis Chen & J. Chen in Wen & others, 1976, p. 69.

Remarks.—Enosolen Guo, 1988, is considered to be a junior synonym of *Tulongella* (Z. Fang & others, 2009).

Stratigraphic range.—Middle Triassic (Anisian)–Upper Triassic (Rhaetian) (Lu & Chen, 1986; Chen & Chen in Wen & others, 1976). Chen and J. Chen (in Wen & others, 1976) erected *Tulongella* and included in it the type species from the Norian of China, *Cuspi*-

daria sp. indet. in Healey (1908) from the Rhaetian of Burma, and *Cuspidaria? problematica* Chen from Upper Triassic of China. Later, Lu and Chen (1986) proposed a new species, *Tulongella qinghaiensis*, from the Anisian of Qinghai (Naocangjiangou Formation). *Enosolen ensatus* Guo, 1988 (type species of *Enosolen*) was also reported from the Anisian (Guo, 1988; Z. Fang & others, 2009).

Paleogeographic distribution.—Eastern Tethys (Fig. 61).

Tethys domain: Middle Triassic: Anisian of Qinghai (China) (Lu & Chen, 1986), western Yunnan (China) (Guo, 1988; Z. Fang & others, 2009); Late Triassic: Norian of China (Xizang, Tibet) (Wen & others, 1976); Rhaetian of Burma (Healey, 1908).

Paleoautoecology.—B, Id, S, SM; Db. See previous discussion of mode of life for *Cercomya*.

Mineralogy.—Aragonitic (J. D. Taylor, Kennedy, & Hall, 1973). There are no data about the shell microstructure of *Tulongella*. J. D. Taylor, Kennedy, and Hall (1973) studied the shells of various current species of the superfamily Pandoroidea, and, in all except in those belonging to the family Thraciidae, there are three shell layers: a prismatic outer layer and nacreous middle and inner layers.

Genus ORTHOMYA Guo, 1985, p. 234, 273

Type species .- Orthomya puerensis Guo, 1985, p. 234.

Stratigraphic range.—Upper Triassic (Carnian) (Guo, 1985). Guo (1985) proposed *Orthomya* and reported it from Carnian of China. No more records of *Orthomya* were located.

Paleogeographic distribution.—Eastern Tethys (Fig. 61).

Tethys domain: Late Triassic: Carnian of southwestern China (Yunnan province) (Guo, 1985).

Paleoautoecology.-B, Id, S, SM; Db. See mode of life for Cercomya.

Mineralogy.—Aragonitic (J. D. Taylor, Kennedy, & Hall, 1973). There are no data about the shell microstructure of *Orthomya*. J. D. Taylor, Kennedy, and Hall (1973) studied the shells of various current species of the superfamily Pandoroidea, and, in all except in those belonging to the family Thraciidae, there are three shell layers: a prismatic outer layer and nacreous middle and inner layers.

GENERA NOT INCLUDED

ACHARAX Dall, 1908a, p. 2

No data are available for the study interval, although Recent Acharax is regarded as being of Paleozoic origin. Cox and others (1969) listed the stratigraphic range of this genus as Miocene to Recent, but they pointed out that probably some fossil species that are similar to Solemya Lamarck, 1818, with external ligament and without internal ribs, known since the Devonian, could be included in Acharax. Pojeta (1988) emended its diagnosis and assigned it a continuous range from lower Permian (Leonardian) to Recent and even accepted a doubtful extension of the range into the Middle Devonian (Eifelian), and Carboniferous (Middle Pennsylvanian). Carter (1990a) emended the diagnosis again and proposed a new subgenus, Acharax (Nacrosolemya), from Carboniferous (Pennsylvanian) beds of Kentucky, and he argued for the need of a good revision of this genus, since there are Devonian and Carboniferous poorly known species attributed to Solemya, which probably would fit better in Acharax. It is interesting to point out that although there are many species attributed to Acharax from Paleozoic deposits, they seem to be far less common in Triassic or Jurassic beds,

because no records were found in deposits of those ages. Although their activity traces are known from Mesozoic beds (Seilacher, 1990), there is no direct relation with actual specimens. It is even possible that none of them were present during our study interval, and that the apparently discontinuous stratigraphic range is in fact due to morphologic convergence.

AGUILERIA White, 1887, p. 34

Aguileria was relatively frequent during the Late Jurassic (Fürsich & Werner, 1988; Liu, 1995) and more abundant during Cretaceous times (Muster, 1995). However, Muster (1995) assigned it an Upper Triassic–Upper Cretaceous range. The Carnian record originated from the reference of *Bakevellia (Bakevelloides) hekiensis* (Kobayashi & Ichikawa, 1952) figured by Hayami (1975, pl. 2,4) as *Aguileria renauxiana* (Mathéron, 1842 in 1842–1843) by Muster (1995), but she did not study Hayami's specimens (deposited in the University of Tokyo Museum). We follow Hayami (1975), since we do not agree that a doubtful single specimen is enough to expand the genus range by more than 50 m.y.

AMERINUMOPECTEN Kasum-Zadeh, 2003, p. 44

This monospecific genus was described by Kasum-Zadeh (2003) from the Upper Triassic of Italy, with the type species *Entolium? amerinum* Sirna, 1968, p. 771. We regard this nomination as unjustified, since the author did not adequately compare it or justify it. Although the ornamentation of this species is unique when compared with other species of *Entolium*, other features (absence of byssal sinus, shape of the auricles, cardinal margin) are perfectly consistent with its attribution to *Entolium*, and the difference in ornamentation may be due to ecological factors (Allasinaz, 1972).

ANGUSTELLA Waagen, 1907, p. 98

Angustella was considered as a junior synonym of Gervillia (Cultriopsis) Cossmann, 1904 (Cox & others, 1969, p. 308), but some authors considered it to be a distinct genus, without justification (Sha, Chen, & Qi, 1990; X. Wang & others, 2008), or as a subgenus of Gervillia (Wen & others, 1976). The name Angustella was later used for a gastropod and a hemipteran (see Nomenclator Zoologicus online: http://uio.mbl.edu/NomenclatorZoologicus/).

ANOMIA Linnaeus, 1758, p. 700

No occurrence from Induan–Sinemurian interval. Cox and others (1969) regarded it as Cretaceous–Holocene and also indicated its doubtful occurrence from the Permian. The origin of the family Anomiidae is now regarded as Jurassic (Todd & Palmer, 2002; Malchus, 2004). Hölder (1990) mentioned *Anomia alpina* Winkler from the Rhaetian. This species was relocated first into *Placunopsis* (Ivimey-Cook & others, 1999) and then into *Pseudoplacunopsis* in Included Genera, p. 70).

ANRADULONECTITES Shurygin & Lutikov, 1991, p. 64

No occurrence from Induan–Sinemurian interval. According to the *Zoological Record*: "*Anradulonectites* (Pectinacea): Gen nov, of Pectinidae, Type species *A. intricatus*, p. 64, Lower Jurassic (Shurygin, B. N. & Lutikov, O.A. 1991. Rossiiskaya Akademiya Nauk Sibirskoe Otdelenie Trudy Instituta Geologii I Geofiziki, 769: 47–78)." Its oldest record is from Pliensbachian times (Zakharov & others, 2006).

ANSHUNOPECTEN (no author)

Anshunopecten was mentioned by Xu (in Z. Yao & others, 1980, table 8, p. 34), without description, figures, or other information accompanying the name. The genus was quoted from the upper Permian of China (H. Yin, 1985; M. Lin & Yin, 1991; L. Li, 1995) and taken into account in Sepkoski (2002) and PBDB (online), but *Anshunopecten* was not formally described, and it is thus invalid (*no-men nudum*) (Z. Fang, personal communication, 2008).

APHANAIA de Koninck, 1877, p. 164

Following Kauffman and Runnegar (1975), *Aphanaia* is regarded as a subgenus of *Atomodesma*. Other authors such as Waterhouse (1979a, 1983, 1987), Astafieva (1991), and Biakov (1992) considered it to be at genus level but none of them justified it. Waterhouse (1958, 1959) included it as a subgenus of *Atomodesma* (the same position was given in Cox & others [1969]), but, later, Waterhouse (1979a) considered it to be at generic level without discussion, even though he mentioned Kauffman and Runnegar (1975).

ARCOMYTILUS Agassiz in J. Sowerby, 1842 in 1842–1844, p. 318

No occurrence from Induan–Sinemurian interval, although it was quoted quite frequently from the Lower Jurassic (Winkler, 1886; Hayami, 1958a, 1975; Cox & others, 1969; Liu, 1999; Fürsich & others, 2001; Gahr, 2002), the oldest record appears to be Pliensbachian.

ARCTOTIS Bodylevsky, 1960, p. 44

No occurrences from the Induan–Sinemurian interval, although Cox and others (1969) assigned it a range from the Lower Jurassic, but all consulted references consider it to be from the Pliensbachian (Kelly, 1984; Jeletzky & Poulton, 1987; Liu, 1995; Sha, 1996; Aberhan, 1998a, 1998b, 2001; Damborenea, 2001; Zakharov & others, 2006). Sepkoski (2002) mentioned its oldest record as Sinemurian, data provided by Hallam (1977), who listed *Arctotis* from Sinemurian of the Arctic region, but we could not confirm it.

ASTARTELLA Hall, 1858, p. 715

Astartella was considered to be a typical Paleozoic genus (e.g., Boyd & Newell, 1968; Cox & others, 1969; Sepkoski, 2002) with a Visean (Carboniferous) origin (Hoare, Heaney, & Mapes, 1989), and its last records were Permian (Ciriacks, 1963; Nakazawa & Newell, 1968; Hayami & Kase, 1977; Biakov, 2002; Clapham & Bottjer, 2007). Recently, Hautmann and others (2011) reported Astartella from the Lower Triassic (Induan) of southern China. The specimens reported by the authors were four valves without internal shell characters; they were attributed to Astartella by the external shape and ornamentation pattern. They resemble astartids but are not clearly referable in particular to Astartella. More available material is necessary to maintain the survivorship of Astartella in the Lower Triassic.

ASTARTOPIS Wöhrmann, 1889, p. 222

Chavan in Cox and others (1969) considered *Astartopis* to be a distinct genus, indicating that Cox regarded it as a subjective synonym

of *Myophoriopis* Wöhrmann, 1889, and he referred it to the family Myophoriopidae. In the *Treatise*, there is no family with this name, and under *Myophoriopis*, Cox does not refer to *Astartopis*. However, *Astartopis* (type: *Myophoria richthofeni* Stur, 1868) was considered to be a synonym of *Myophoriopis* by many authors (e.g., Bittner, 1895; Diener, 1923; Kutassy, 1931; Leonardi, 1943; Allasinaz, 1966), and this is the position followed here, since no substantial differences were noted between the genera.

BARBATIA Gray, 1842, p. 81

No occurrences from the Induan–Sinemurian interval are considered, as records from the Triassic are doubtful. Cox and others (1969) assigned it a Jurassic–Holocene range and mentioned its occurrence from the Triassic with doubts. Sepkoski (2002) assigned it a Triassic (Norian)–Holocene range, referring to Hallam (1981), but, in this paper, the genus was not considered, although in other papers by the same author (Hallam, 1972, 1976, 1977), the oldest recognized record of the genus is Pliensbachian. It was widely distributed during the Middle and Late Jurassic (Kelly, 1984; Fürsich, Palmer, & Goodyear, 1994; Liu, 1995; Delvene, 2000, 2003).

BOSNIOPECTEN Kasum-Zadeh, 2003, p. 55

This genus was proposed by Kasum-Zadeh (2003) from the Carnian of Bosnia, with *Pecten volaris* Bittner, 1903, p. 634, as type species, and he included two other species: *Pecten inaequicostatus* Reis, 1926, and *Pecten subaequicostatus* Bittner, 1895. The author neither compared species nor justified the proposal, and the three species were assigned to *Chlamys (Chlamys)* by Allasinaz (1972), which is followed here. In addition, Kasum-Zadeh (2003) considered his new genus as being present only in the Carnian of Bosnia, but these species were also recorded from Ladinian and Rhaetian beds of Austria and Italy (Allasinaz, 1972).

BRACHIDONTES Swainson, 1840, p. 384

No occurrences from Induan–Sinemurian interval are considered, although Cox and others (1969) and Sepkoski (2002) assigned it a Jurassic–Holocene and Lower Jurassic (?)–Holocene range, respectively. However, the oldest record of *Brachidontes* is from the Pliensbachian of the Iberian Peninsula (Hallam, 1972).

BUPECTEN Guo, 1988, p. 118

Junior synonym of *Entolium* Meek, 1865 (Z. Fang & others, 2009). Guo (1988, p. 118) mentioned it has two oxhorn-shaped auricles in the diagnosis of the genus, and differed from *Entolium* by "two acuminated and high, shooting [projecting] auricles" [translation of the diagnosis offered by Z. Fang & others, 2009, p. 48]. In the opinion of these last authors: "The so-called oxhornlike auricles seem to be artificially processed and polished up," they also noted that, although in Guo's (1988) material there are no right valves of the type species, he included a decription of them in his text. We follow Z. Fang and others (2009).

BUREIOMYA Voronetz, 1938, p. 58

Bureiomya was considered as a junior synonym of *Pholadomya* (*Bucardiomya*) Rollier in Cossmann, 1912 (Cox & others, 1969, p. 829). According to Kobayashi and Tamura (1983b, p. 209), the

genus was proposed from the Triassic of northeastern Siberia. We accept the synonymy, and, therefore, we disregard other records of this genus (e.g., Bychkov & others, 1976; Kurushin, 1990; Polubotko & Repin, 1990).

CALVAENTOLIUM Romanov, 1985, p. 35

Romanov (1985) gave generic names to the four groups established by Staesche (1926) within *Entolium s. l.* (see discussion in Damborenea, 2002a, p. 42–44). Staesche's *Entolium hehlii* d'Orbigny group was included by Romanov (1985) in the new genus *Calvaentolium* (type species: *Pecten magneauritus* Kittl, 1904, from the Anisian of Yugoslavia), plus several Triassic species [*C. pseudodiscites* (Guembel), *C. tribevicianum, C. magneauritum* (Kittl), *C. tridentini* (Bittner), *C. inornatum* (Stoppani), *C. saccoi* (Parona), *C. cainalloi* (Stoppani), *C. contemptibile* (Stoppani)] and Jurassic species [*C. calvum* (Goldfuss, 1935 in 1833–1841), *C. hehlii* (d'Orbigny)]. He assigned it a Triassic–Jurassic range in Europe and Asia. In the absence of a good review on the *Entolium* group, we follow a conservative view and consider this group to be within *Entolium*, following Staesche (1926).

Waller (2006) regarded it as a valid genus and included it in his new family Entoliolidae Waller, 2006, along with other genera, and he even considered *Crenamussium* Newton in Newton & others, 1987, to be a junior synonym of *Calvaentolium* (see discussion for the family Entoliidae in Included Genera, p. 103).

CARDIOMORPHA de Koninck, 1841 in 1841-1844, p. 101

There are no occurrences from the Induan-Sinemurian interval. Cox and others (1969, p. 818) assigned it a Carboniferous range, but they also indicated (p. 115) that some species were reported from the Triassic. Several Triassic species were attributed to Cardiomorpha and described prior to 1969 (Trechmann, 1918; Diener, 1923, p. 229; Krumbeck, 1924; J. P. Smith, 1927; Kutassy, 1931, p. 415; Marwick, 1953) and also after that date (Wen & others, 1976; Sha, Chen, & Qi, 1990). Most of these species were doubtfully referred to Cardiomorpha. Sepkoski (2002) assigned it a Carboniferous-Upper Triassic (Carnian) range, based on Morris (1967). In line with Pojeta and others (1971), we should be careful with studies at lower taxonomic levels of poorly known taxa, such as *Cardiomorpha*, because the evolutionary history of major lineages can give us a better overview of a particular group. They especially referred to Cardiomorpha records from the Ordovician-Devonian and the Triassic. Furthermore, its patchy distribution is very suspicious. We do not include the Triassic records, and we consider Cardiomorpha to be a Carboniferous (see discussion in Pojeta & others, 1971, p. 146; Simões & others, 1997; Waller & Stanley, 2005) and lower Permian (Runnegar, 1965; Runnegar & Newell, 1974) genus.

CARDITA Bruguière, 1792 in 1789-1792, p. 401

No occurrences from Induan–Sinemurian interval are considered; according to Chavan in Cox and others (1969), *Cardita* has a Paleocene–Holocene range. Although it was frequently quoted from the Triassic in papers prior to Cox and others (1969), after the *Treatise*, it was mentioned from the Triassic by Kollarova & Kochánova (1973) and from the Jurassic by Accorsi-Benini (1981). Cox and others (1969) did not comment on all the species attributed to *Cardita* in classical papers (e.g. Stoppani, 1860–1865; Winkler, 1861;

Goldfuss, 1863; Laube, 1865, among others). It is beyond the scope of this paper to review each of these species to determine in what other genera should they be included. This situation is repeated for many Recent genera, such as *Mytilus, Nucula*, and *Corbula*, which had traditionally been used as a mixed bag where all similar forms were included.

CERATOMYA Sandberger, 1864, p. 16

Cox and others (1969) assigned it a Lower–Upper Jurassic range. *Ceratomya* was reported from the Sinemurian of Europe (Etheridge, 1864; Hallam, 1976, 1977, 1987; Liu, 1995). The probable Sinemurian species are: *C. petricosa* (Simpson, 1855) (Hallam, 1976) and *C. gibossa* (Etheridge, 1864) (Damborenea in Damborenea & González-León, 1997), but, although Etheridge (1864) figured the specimens, he did not specify the age of the sediments. On the other hand, there are several papers where the genus is discussed systematically and figured from Pliensbachian beds (e.g., Damborenea in Damborenea & González-León, 1997; Delvene, 2003; Aberhan, 2004; Scholz, Aberhan, & González-León, 2008).

CHAENOCARDIA Meek & Worthen, 1869, p. 170

No occurrences in the study interval are considered; although it was mentioned from the upper Permian of southern China (L. Li, 1995), this will not be taken into account, because it is not a systematic treatment and does not mention the original data source. *Chaenocardia* is regarded as a typical Carboniferous genus (Cox & others, 1969; Newell & Boyd, 1995).

CHIRON Astafieva, 1997, p. 27

Chiron was replaced by *Chironopecten* Astafieva, 2001. Astafieva (1997) proposed *Chiron*, but this name had already been used for an insect, *Chiron* MacLeay, 1819, p. 107 (Astafieva, 2001).

CHULUARIA Waterhouse, 2000, p. 175

Chuluaria is not considered here, because the reasons to separate this genus from *Claraia* are insufficiently justified. Waterhouse (2000) proposed this new genus to accommodate specimens similar to *Claraia* but with differences in form and size of the right anterior auricles, being inequivalve, and the form of byssal notch. These differences are considered by most authors as species-level features (e.g., F. Yang, Peng, & Gao, 2001; He, Feng, & others, 2007).

CINGENTOLIUM Yamani, 1983, p. 6

Cingentolium Yamani, 1983 is a junior objective synonym of *Costentolium* Freneix, 1980, as they are based on the same type species, *Pecten cingulatum* Goldfuss, 1835 in 1833–1841 (see discussion for *Costentolium* and *Neoentolium* Romanov, 1985, below).

CLARAIOIDES Fang, 1993, p. 653, 660

Claraioides is regarded as a junior synonym of *Claraia* Bittner, 1901a (Newell & Boyd, 1995; F. Yang, Peng, & Gao, 2001; He, Feng, & others, 2007). Z. Fang (1993) proposed *Claraioides* to accommodate the upper Permian specimens that were assigned to *Claraia primitiva* Yin, 1983, and *Claraia diana* Guo, 1985, and he also proposed a new species, *Claraioides guizhouensis*. The proposal of this new genus was based on: "... the existence of a subcircular byssal embayment in the right valve (lower valve), by which it may be readily distinguished from the comparable Claraia and Pseudoclaraia" (Z. Fang, 1993, p. 660). Newell and Boyd (1995) and F. Yang, Peng, and Gao (2001) considered it to be a junior synonym of Claraia, based on the lack of a quantitative, comparative study of local populations before proposing the new genus (Newell & Boyd, 1995) and the use of species-level features to discriminate at genus level, in this case the size, shape, and direction of the byssal sinus and the size and shape of the auricles (see discussion in F. Yang, Peng, & Gao, 2001, p. 800-801). Z. Fang (2003) disagreed with F. Yang, Peng, and Gao (2001) and again reaffirmed the validity of *Claraioides*; however, he did not provide further arguments. Kotlyar, Zakharov, and Polubotko (2004), according to Z. Fang (1993), decided to keep Claraioides separate from Claraia, and they referred the upper Permian specimens similar to Claraia to Claraioides (see discussion in Kotlyar, Zakharov, & Polubotko, 2004, p. 524). Delving into the discussion about the supposed synonymy, He, Feng, and others (2007) analyzed the variation in the ornamentation and shape of the byssal sinus in all species included in Claraia with their stratigraphic and paleogeographic distribution (He, Feng, & others, 2007, table 1, p. 1016 and discussion), showing that there is a progressive morphological change in *Claraia* from Changhsingian to Induan, indicating that the shape, size, and orientation within the byssal sinus are not suitable criteria for discrimination at generic level. Z. Fang and others (2009, p. 32) did not agree and provided a list of species included in Claraioides.

COMATAHALOBIA Polubotko in Polubotko, Payevskaya, & Repin, 2001, p. 100

Polubotko, Payevskaya, and Repin (2001), pursuing their revolutionary concept of the "*Halobia*" group (see discussion for the family Halobiidae in Included Genera, p. 91), used once again criteria such as shape and position of the anterior auricle and the ornamentation type to propose a new genus similar to, but separate from, *Halobia*. We follow McRoberts's (1993) criteria and do not take into account this genus.

COSTENTOLIUM Freneix, 1980, p. 89

Entolium (Costentolium) Freneix, 1980, Cingentolium Yamani, 1983, and Neoentolium Romanov, 1985, were proposed for the same species group, which Staesche (1926) called the group of E. cingulatum (Goldfuss, 1835 in 1833–1841) (see Damborenea, 2002a, p. 44). The type species of the three taxa is the same, Pecten cingulatum Goldfuss, so even if it is regarded as a separate genus, it should be called *Costentolium* Freneix, 1980, by priority. Freneix (1980) also included in E. (Costentolium) the following species: E. discites (Schlotheim), E. frontalis (Dumortier), E. proeteum (d'Orbigny), E. renievieri (Oppel), E. spathulatum (Roemer), E. partitum (Sowerby), (?) E. lackeyi Quilty, and (?) E. territorianum Skwarko. Waller (2006) indicated that Neoentolium is a junior synonym of *Cingentolium*, but he did not mention the existence of Costentolium. In this paper, we follow a conservative view and regard all these species as belonging to Entolium in its original sense (see discussion for the family Entoliidae in Included Genera, p. 103).

COSTIGERVILLIA Cox & Arkell, 1948 in 1948-1950, p. 9

No occurrences are recorded in the study interval, although Muster (1995) indicated the possible record of *Costigervillia* from the Upper Triassic. The oldest reliable occurrences of *Costigervillia* are from the Upper Jurassic, with two species, *C. crassicosta* (Morris & Lycett, 1853 in 1851–1855, p. 23) and *C. quincarinata* (Fischer, 1969) (Fürsich & Werner, 1988). Guo (1985) proposed *C. guibaoensis* Guo, 1985, from the Upper Triassic of China, but we were unable to locate information about this species. Apart from this mention, *Costigervillia* was only reported from the Upper Jurassic.

CULTRIOPSIS Cossmann, 1904, p. 510

Following Cox and others (1969, p. 308), *Cultriopsis* is regarded as a subgenus of *Gervillia*, although some authors assigned it a genus rank without justification (e.g., Komatsu, Chen, & others, 2004; Komatsu, Akasaki, & others, 2004).

CUSPIDARIA Nardo, 1840, p. 50

No occurrences from Induan-Sinemurian interval are considered, although there are several mentions of the genus from the Triassic and Jurassic. Prior to Cox and others (1969), Cuspidaria was reported from the Triassic (Diener, 1923, p. 243; Kutassy, 1931, p. 425; Leonardi, 1943; Rossi Ronchetti, 1959; Allasinaz, 1964) and from the Jurassic (Hayami, 1958c, 1959). Cox and others (1969, p. 854) assigned it a Cretaceous-Holocene range, disregarding those mentions. Subsequently, not all authors followed these guidelines when assigning a stratigraphic range to Cuspidaria; Skelton and Benton (1993, p. 260) and Sepkoski (2002) considered Cuspidaria triassica (Stoppani, 1865 in 1860-1865), from the Ladinian of Austria, as the oldest occurrence of the family Cuspidaridae; the original source is Morris (1967). Also, Runnegar (1974) suggested a possible Triassic origin for Cuspidaria; other authors assigned to this genus both Triassic species (Hayami, 1975; Lu & Chen, 1986) and Jurassic species (Hallam, 1976, 1977; Palmer, 1979; Fozy, Kázmér & Szente, 1994; J. D. Hudson & others, 1995; Liu, 1995). These are not taken into account here, since most were based only on general morphology and the presence of a typical, elongated so-called rostrum, as in cuspidarids (also observed in other bivalve families) and, furthermore, in none of them could the microstructure, musculature, and/or hinge be proven to be actually attributable to this family (Harper, Palmer, & Hudson, 2002). In addition, Allasinaz (1966, p. 641) indicated that, from the Triassic species referred to Cuspidaria, the only one that bears some resemblance to the living Cuspidaria is C. alpiscivicae Bittner, 1895. Allasinaz (1966) transferred all Triassic species attributed to Cuspidaria to Solenomorpha, but he did not sufficiently justify this decision (see discussion for Solenomorpha in this section, p. 170). Guo (1985) proposed the subgenus Cuspidaria (Dianocuspidaria) from the Carnian (lower Upper Triassic), based on only two specimens (Z. Fang & others, 2009). For all these reasons, we agree with Harper, Palmer, and Hudson (2002, p. 766-767) that the fossil record of the family Cuspidariidae was wrongly extended back by the inclusion of doubtful species.

CYCLOPELLATIA Cossmann in Cossmann & Pellat, 1907, p. 32

Cyclopellatia is not considered in the study interval because the stratigraphic range assigned by Cox and others (1969) [L. Jurassic (Barr.)] is a mistake, the Barremian being a Lower Cretaceous stage.

DESIDERINECTES Kasum-Zadeh, 2003, p. 50

This genus was proposed by Kasum-Zadeh (2003) from the Upper Triassic of Europe with *Pecten (Chlamys?) desideri* Bittner, 1901b as type species, and including other species that the author did not name. Furthermore, he did not compare or justify the genus proposal. The type species was assigned to *Camptonectes (Annulinectes)* by Allasinaz (1972), and this is followed here.

DIETRICHIA Reck, 1921, p. 434

No occurrences from Induan–Sinemurian interval are considered, although both Cox and others (1969) and Sepkoski (2002) assigned it to have a Jurassic range. *Dietrichia* was only reported from the Oxfordian of Europe and Callovian–Oxfordian of Tunisia (Holzapfel, 1998), so its stratigraphic range is Middle–Upper Jurassic. The same name was used for a Linyphiidae spider genus by Crosby and Bishop (1933) but was recently replaced by *Neodietrichia* (Özdikmen, 2008).

DIMYODON Munier-Chalmas in Fischer, 1886 in 1880–1887, p. 937

Dimyodon is considered a junior synonym of *Atreta* Etallon, 1862 (Fürsich & Werner, 1988, p. 143; Malchus, 2000; Hautmann, 2001a, 2001b). According to Fürsich and Werner (1988), *Dimyodon* and *Atreta* were only distinguished because *Dimyodon* had two muscle scars and internal radial ribs, which are lacking in *Atreta*. However, in their study of the fauna from the Upper Jurassic of Portugal, they reported some specimens attributed to *Atreta* that have these features and concluded that those differences are due to taphonomic processes: specimens with aragonitic inner shell layer replaced by calcite retained these internal structures (muscle and internal radial ribs) and were assigned to *Dimyodon*, while other specimens in which the inner shell layer was dissolved, removing such structures, were assigned to *Atreta*.

DIOTIS Simonelli, 1884, p. 125

It is a junior homonym of *Diotis* Schmarda, 1859 (flatworm Turbellaria). *Diotis* Simonelli, 1884, was replaced by *Caenodiotis* Monari, 1994 (Monari, 1994).

ENANTIOSTREON Bittner, 1901c, p. 70

Checa and others (2003) and Márquez-Aliaga and others (2005) studied the syntypes of *Enantiostreon* and considered it to be an invalid genus. *Enantiostreon* was proposed by Bittner (1901c) to include Triassic ostreid bivalves (see Giebel, 1856) that were attached by their right valves, as observed by Noetling (1880). The type species is *Enantiostreon hungaricum* Bittner, 1912, from the Carnian of Hungary, with unknown hinge, the lower valve is clearly more convex than the upper valve, and specimens attached themselves by their right valves. Cox and others (1969) based their reference on a specimen of *Terquemia difformis* Goldfuss. According to the authors, *E. hungaricum* should be considered as a possible plicatulid

of uncertain generic relations. Some species traditionally included in *Enantiostreon*, such as *E. cristadifformis* Schlotheim, 1823 in 1822–1823, and *E. spondyloides* Schlotheim, 1823 in 1822–1823, clearly belong to Ostreidae, since they attached by the left valve and had a hinge without diagnostic structures (Checa & Jiménez-Jiménez, 2003b); they are included in *Umbrostrea* Hautmann, 2001a (Márquez-Aliaga & others, 2005).

ENOSOLEN Guo, 1988, p. 128

Junior synonym of *Tulongella* Chen & J. Chen in Wen & others, 1976 (Z. Fang & others, 2009, p. 143).

EOMONOTIS Grant-Mackie, 1978a, p. 102

Eomonotis is considered to be a subgenus of *Monotis* (Grant-Mackie, 1978a, 1980a; Grant-Mackie & Silberling, 1990; Silberling, Grant-Mackie, & Nichols, 1997), although some authors, without justification, regarded it as an independent genus (e.g., Kurushin, 1990; Klets, 2006; Wignall & others, 2007).

EOSCHIZODUS Cox, 1951, p. 369

No occurrences from the Induan–Sinemurian interval are considered. Cox (1951) proposed *Eoschizodus* to accommodate Paleozoic myophoriids (Devonian, Carboniferous, and Permian), and the same range was assigned in Cox and others (1969, p. 473): Devonian– Permian. Sepkoski (2002) assigned it a Devonian (Givetian)–Triassic (?lower Anisian) range, based on another database: Skelton and Benton (1993). The only information that Skelton and Benton (1993, p. 254) offered about this genus is "*Eoschizodus truncatus* (Goldfuss, 1837 in 1833–1841), *Strigocephalus* zone, Rhineland, Germany (Newell & Boyd, 1975)" and "Permian specimens are reported (Cox & others, 1969, p. 473)." The *Strigocephalus* zone corresponds to upper Middle Devonian (Newell & Boyd, 1975). According to these authors, *Eoschizodus* was rare and limited to its type species and only recorded from the *Strigocephalus* zone in Germany, so no Permian and Triassic records are here considered.

EPICLARAIA Gavrilova, 1995, p. 132

Epiclaraia is not considered here, because the reasons to separate this genus from *Claraia* are insufficiently justified. Gavrilova (1995) proposed this new genus to accommodate specimens similar to *Claraia* but with a differently shaped right auricle and byssal sinus. These characters were used at the species level within *Claraia* and are not enough to distinguish generic taxa in the group (F. Yang, Peng, & Gao, 2001; He, Feng, & others, 2007) (see discussion for *Claraioides* in this section, p. 159).

FENGJIACHONIA Wu, 1981, p. 377

Junior synonym of *Kija* Lebedev, 1959 (Ma, 1989, p. 611). S. Wu (1981) proposed *Fengjiachonia* from the Jurassic of China, and it was subsequently placed in synonymy by Ma (1989), because he considered both genera as identical in form, external ornamentation, and internal characters.

FILAMUSSIUM Waller, 2006, p. 342

Filamussium is not considered here, because we regard its proposal as unnecessary. The type species, Pecten schafhäutli Winkler, 1859, is best accommodated within *Parvamussium* Sacco, 1897, following its original concept (see discussion for *Parvamussium* in Included Genera, p. 95). Waller (2006) separated *Filamussium* from other propeamussiids by "having a filosus structure on its left valve and in having internal ribs that were probably originally aragonitic rather than calcitic." However, although the author compared his new genus with *Propeamussium*, he did not do a direct comparison with any *Parvamussium* species. Moreover, the presence or absence of the filosus structure may be due to diagenetic processes, and it is observed in other bivalve groups as well (see Hautmann, 2001b, p. 62). Within the Propeamussiidae, genera are recognized by other characters, such as the presence or absence of lateral aperture and byssal notch in the adult stage.

FIMBRIA Megerle von Mühlfeld, 1811, p. 52

Although Hallam (1977, 1990) mentioned *Fimbria* from the Hettangian, we did not find any record until the Pliensbachian. Monari (2003) considered that *Fimbria* had its origin during lower Pliensbachian times in the western Tethys, showing a diversity maximum during the Late Jurassic.

GEMMELLARODUS Di Stefano, 1912, p. 81

Junior synonym of Neomegalodon Gümbel, 1862 (H. Yao & others, 2007). Allasinaz (1965) included Gemmellarodus and Rossiodus Allasinaz, 1965, as subgenera of Neomegalodon, and, although in Cox and others (1969), Neomegalodon was considered as a subgenus of Megalodon, both Rossiodus and Gemmellarodus continued to be used in different ways, as subgenera of Neomegalodon (Tichy, 1975, 1980a; Végh-Neubrandt & others, 1976; Sha, Chen, & Qi, 1990) or as independent genera (Allasinaz & Zardini, 1977; Végh-Neubrandt, 1982; Yancey & Stanley, 1999; H. Yao & others, 2003). H. Yao and others (2007) argued that neither Gemmellarodus nor Rossiodus were sufficiently well known to be useful concepts. Also, they stressed that the hinges of Neomegalodon, Rossiodus, and Gemmellarodus are similar, and, frequently, small-sized megalodontids were attributed to Rossiodus and very inequivalve ones to Gemmellarodus, using characters that are not systematically significant (see discussion in H. Yao & others, 2007, p. 1337).

GIBBOCONCHA De Gregorio, 1930a, p. 30

No occurrences from the Induan–Sinemurian interval are considered, although Cox and others (1969) mentioned *Gibboconcha* from the Lower Jurassic of Sicily, indicating that the description of the genus was made by De Gregorio based on a small fragment possibly belonging to a juvenile specimen of *Cardinia* or *Astarte*, and its internal characters were unknown. No more information about *Gibboconcha* was found, and, taking into account the doubts raised, we did not include it.

GLYPTOLEDA Fletcher, 1945, p. 293, 298

Glyptoleda is considered a subgenus of *Veteranella* Patte, 1926. Puri in Cox and others (1969) regarded *Glyptoleda* as a subgenus of *Veteranella*, and distinguished *V. (Veteranella)* and *V. (Glyptoleda)* by their posterior part not being constricted and constricted, respectively; the first from the Upper Triassic and the second from the Permian. Other authors considered *Glyptoleda* as a junior synonym of *Veteranella* (see Waterhouse, 1980a), but Waterhouse (1980a, 1983, 1987) disagreed and regarded them as valid and independent genera, based primarily on differences in ornamentation. Subsequently, some authors followed Waterhouse in this regard (e.g., J. Chen, Lui, & Lan, 1983; Biakov, 1998, 2006; Zakharov & others, 2006; Z. Fang & others 2009). However, Z. Fang and Cope (2004) warned that the type of ornamentation is of low taxonomic significance, and it may even be variable at the intraspecific level. Biakov (1998, p. 132) mentioned that the ornamentation of *Glyptoleda* is an adaptation to the environment, and therefore is not useful as a taxonomic character. Following Puri in Cox and others (1969) and Z. Fang & Cope (2004), *Glyptoleda* is included as a subgenus of *Veteranella* (see discussion for *Nucundata* in this section, p. 166).

GONILIA Stoliczka, 1871 in 1870-1871, p. 278

No occurrences from the Induan–Sinemurian interval are considered, although Cox and others (1969) assigned it a Jurassic–Holocene range. The Jurassic record was referred to *Gonilia (Ensio)* Cox, 1962. The genus *Ensio* was proposed by Cox (1962), including three species: *Ptychomya agassizii* (holotype) from the Inferior Oolite of England (with a Toarcian–Bajocian range [West, 2007]), *Astarte divaricata* (junior synonym of *Ptychomya agassizii*) from the Bajocian of Lincolnshire Limestone of Santon, and *Astarte eastonii* from the Upper Jurassic of Borneo. Therefore the oldest record of *Gonilia* dates from the Toarcian. Sepkoski (2002) assigned it a Pliocene–Holocene range (see table in Jablonski & others, 2003).

GRYPHELLINA Newell, 1940, p. 289

Newell (1999, p. 4) rejected it because its type species, *Capulus sellardsi* Beede, 1907, is a gastropod.

GUICHIELLA Li & Ding, 1981, p. 329

According to J. Chen and Komatsu (2002), *Guichiella* is considered a subgenus of *Claraia* Bittner, 1901a.

HABONUCULA Singh & Kanjilal, 1977, p. 189

No occurrences from Induan–Sinemurian interval are considered. The Zoological Record assigned it a Jurassic range and those data were incorporated into Sepkoski's database (2002). The original source is Singh and Kanjilal (1977), who proposed the genus from the lower Callovian. Therefore *Habonucula* is not included here. In addition, Jaitly, Fürsich, and Heinze (1995) regarded *Habonucula* as a junior synonym of *Nuculoma* Cossmann in Cossmann & Thièry, 1907.

HEMIMENION Guo, 1988, p. 117

Junior synonym of *Aviculomyalina* Assmann, 1916 (Z. Fang & others, 2009). Guo (1988) proposed *Hemimenion* (family Myalinidae) and included two new species: *H. cuneatum* (type species) and *H. triangulare* from the Anisian of southwestern China. Z. Fang and others (2009) included it as a synonym under *Aviculomyalina*, and we follow this decision.

HUNANONECTES Fang, 1978, p. 465

Junior synonym of *Radulonectites* Hayami, 1957c (Damborenea, 2002a). Z. Fang (1978) proposed *Hunanonectes* as a subgenus of

Camptonectes from the Jurassic of China and included three new species: C. (H.) sanduensis, C. (H.) parachlamys, and C. (H.) yizhangensis). He indicated that it might also be a subgenus of Chlamys, because it was intermediate between Camptonectes and Chlamys. He interpreted the so-called Camptonectes striations as being radial ribs, and thus included it in Camptonectes. Subsequently, Damborenea (2002a) considered it to be a junior synonym of Radulonectites, since the species described by Z. Fang (1978) "have small faintly ornamented shells but are otherwise similar in general shape and other morphological aspects to both North and South Pacific species of Radulonectites." Stiller (2006), although taking into account Damborenea's (2002a) research, decided to keep the two genera separate, but he considered that the three species described by Z. Fang (1978) were intraspecific variations and retained only the type species (H. sanduensis Fang, 1978). This was also done by Z. Fang and others (2009), but no mention of Stiller's paper was made. Stiller compared the type species with other species of Radulonectites (R. japonicus Hayami, 1957c; R. exsertus J. Chen, 1982b) from the Lower Jurassic of China (see discussion and synonym list in Stiller, 2006, p. 23–31), and the only difference he found was that the radial ornamentation was more pronounced in the latter species. We follow Damborenea (2002a) and regard Hunanonectes and Radulonectites as synonyms.

IMPOSIDONIA Waterhouse, 2008, p. 66

Waterhouse (2000, p. 181) described a new species, *Posidonia elegantula*, and, later, Waterhouse (2008) proposed a new genus based on this species. According to Waller and Stanley (2005, p. 19), "species of '*Posidonia*' described by Waterhouse (2000, p. 181) from the Lower Triassic of the Himalayas also appear to be claraiids."

INDIGIROHALOBIA Polubotko, 1984, p. 42

Polubotko (1984) distinguished *Indigirohalobia* by the shape and position of the anterior auricle, by the ornamentation, and the presence of a strong ligament (H. J. Campbell, 1994), features considered by McRoberts (1993) and most authors as being diagnostic at the specific level (see H. J. Campbell, 1994, for a review on the discussion of this topic). Although many authors considered *Indigirohalobia* to be a valid genus (Okuneva, 1985, 1987; Kurushin, 1990; Polubotko, Payevskaya, & Repin, 2001, among others), we regard it as a junior synonym of *Halobia*, following McRoberts (1993) (see discussion of family Halobiidae in Included Genera, p. 91).

INOCERAMUS J. Sowerby in Anonymous, 1814, p. 448

Inoceramus was reported from the Early Jurassic on many occasions (e.g., A. F. Leanza, 1942; Cox & others, 1969; Escobar, 1980). Most of these specimens were attributed to *Inoceramus sensu lato*, and they would be better accommodated in other genera like *Parainoceramus*. According to Harries and Crampton (1998), the true inoceramids did not appear until the Late Jurassic.

IRANOPECTEN Repin in Polubotko, Payevskaya, & Repin, 2001, p. 118

Repin (in Polubotko, Payevskaya, & Repin, 2001) erected *Iranopecten* based on the type species *Indopecten glaber* Douglas, 1929. We are not taking into account this new genus, because we follow the original assignation (Douglas, 1929; and Hautmann, 2001b) and refer this species to *Indopecten* Douglas, 1929.

ISOCARDIOIDES Fan, 1963, p. 523 [540]

Probably a junior synonym of *Schafhaeutlia* Cossmann, 1897 (Gu & others, 1976; Z. Fang & others, 2009).

JURASSICARDIUM Cossmann, 1906, p. 294

There are some reports from the Sinemurian of South America (Damborenea, 1996a; Damborenea & Manceñido, 2005b), but no systematic treatment of specimens was made. *Jurassicardium* was reported from the Upper Jurassic (Schneider, 1995). He noted that the specimens of the type species are lost and he excluded the genus from his analysis.

KRUMBECKIA Diener, 1915, p. 131

Junior synonym of *Schafhaeutlia* Cossmann, 1897 (see Hautmann, 2001b, p. 124). Diener (1915, p. 131) distinguished *Krumbeckia* from *Schafhaeutlia* by "oblique-oval shape of the shell and because the tooth of the hinge was more isolated;" according to Hautmann (2001b), these criteria do not allow separation of the two genera.

LAEVITRIGONIA Lebküchner, 1932, p. 68

No occurrences from the Induan–Sinemurian interval are considered. Cox and others (1969) mentioned it had its origin in the Lower Jurassic. Venzo (1942) noted that the oldest species assigned to this genus is *Trigonia (Laevitrigonia) oviedensis* Lycett, 1881, from the Hettangian of Oviedo (Asturias, Spain). Later, Dubar and others (1971) considered this species as a *Trigonia* and from Kimmeridgian age in Asturias, so we have serious doubts about the affinities of this record, whether it can be attributed to *Laevitrigonia* and even if the sediments from Oviedo referred by Venzo (1942) can be assigned to a Hettangian age. Several authors (Hallam, 1976, 1977; Francis & Hallam, 2003) indicated its origin in the Upper Jurassic of Europe.

LATERNULA Röding in Bolten, 1798, p. 155

Cox and others (1969) assigned it an Upper Cretaceous-Holocene range. Anatina Lamarck, 1818 is considered to be a junior synonym of Laternula, and this is the source of the problem about Triassic species attributed to Laternula. Anatina was widely reported from the Triassic, and, with the establishment of the synonymy discussed, most of the species were referred to other genera, but some remained in an indeterminate state. Some of these species, especially those reported from the Upper Triassic (e.g., Allasinaz, 1962; Sirna, 1968; Linck, 1972; Márquez-Aliaga, Plasencia, & Ros, 2005; Damborenea & Manceñido, 2012), are L. rhaetica (Gümbel, 1861), L. zannonii (Stoppani, 1860–1865), L. amicii (Stopanni, 1863 in 1860–1865), or L. suessi (Oppel, 1857 in 1856-1858). Most of them were based on specimens preserved as internal molds in which key characters were not observed. According to Runnegar (1974), the first laternulid could be Cercomya Agassiz, 1843 in 1840-1845, regarded originally as a subgenus of Anatina (Diener, 1923). A review of these Triassic species is necessary to determine if the range of Laternula should be extended or if they can be accommodated better in other genera. Following Cox and others (1969), we do not consider occurrences of Laternula from the Induan-Sinemurian interval.

LECOMPTEUS Poel, 1959, p. 13

According to Cox and others (1969), the type species designated by Poel (1959) is *Mytilus ornatus* Münster, 1837, in Goldfuss, 1833–1841, and it had a Jurassic range, as well as Cretaceous according to Sepkoski (2002). We could not see the original paper in which the genus was proposed, but all subsequent records reported it from the Lower Cretaceous of Japan (Hayami, 1975, among others). According to Kauffman and H. A. Leanza (2004), Münster (1837, in Goldfuss, 1833–1841) proposed the type species based on specimens from the Upper Cretaceous (Campanian) of Europe. The available information suggests that the presence of *Lecompteus* in the Lowest Jurassic is improbable. Kauffman and H. A. Leanza (2004, p. 1190) proposed to include *Mytilus ornatus* in their new genus *Nodomytilus*, but they did not mention *Lecompteus*, unaware of the position of this species (H. A. Leanza, personal communication, 2007).

LEVICONCHA Waagen, 1907, p. 149

Although Cox and others (1969) considered *Leviconcha* to be a junior synonym of *Neoschizodus* Giebel, 1855, several authors considered it to be a valid genus with different systematic relations: subgenus of *Neoschizodus* (Kobayashi & Tamura, 1968b), subgenus of *Myophoria* (Wen & others, 1976; C. Chen, 1982), or at genus level (Ling, 1988; Sha, Chen, & Qi, 1990). In this paper, we follow Kobayashi and Tamura (1968b), in the absence of a modern revision, because it adequately justifies the position of *Leviconcha*.

LUPHERELLA Imlay, 1967, p. 8

Lupherella was erected as genus by Imlay (1967). Following Damborenea (1987b), Lupherella is considered to be a subgenus of Otapiria Marwick, 1935. Damborenea found that adults of Lupherella boechiformis (Hyatt, 1894) (type species of Lupherella) were extremely similar to juveniles of Otapiria originalis Kiparisova, 1960, and Otapiria neuquensis Damborenea, 1987b (see discussion in Damborenea, 1987b, p. 156). J. Chen (1988) also noticed that the separation between Otapiria and Lupherella at generic level was not well founded. However, other authors (e.g. Aberhan, 1998a; J. Yin, Yao & Sha, 2004) considered Lupherella to be a separate genus.

LYRIOMYOPHORIA Kobayashi, 1954, p. 66

Junior synonym of *Elegantinia* Waagen, 1907 (Kobayashi & Tamura, 1968b; Boyd & Newell, 1999). Cox and others (1969) considered *Lyriomyophoria* to be a valid genus and *Elegantinia* to be a junior synonym of *Gruenewaldia* Wöhrmann, 1889. However, shortly before the publication of Cox and others (1969), Kobayashi and Tamura (1968a, p. 91) decided, following McLearn (1942), to keep both genera as valid, and, subsequently, they considered *Lyriomyophoria* to be a junior objective synonym of *Elegantinia*, because they share the same type species, *Lyriodon elegans* Dunker, 1851, p. 300; this was followed years later by Boyd and Newell (1999, p. 547).

MACTROMYOPSIS Chavan, 1959, p. 506

No occurrences from the Induan–Sinemurian interval are considered. Chavan (1959) proposed *Mactromyopsis* and assigned it a Bajocian–Callovian range. Cox and others (1969, p. 511) assigned it a Jurassic (Charmouthian–Callovian) range in Europe [Charmouthian is an old name that corresponds to Pliensbachian (Morris, 1967; Vera, 1994)]. Interestingly, Sepkoski (2002) assigned it a Jurassic (Hettangian–Callovian) range, based on data provided by Skelton and Benton (1993). However, these last authors assigned a Hettangian–Holocene range to the family Mactromyidae, pointing out that the oldest record for the family was *Mactromyopsis (Mactromyella) inflata* (Thevenin, 1909), from the Charmouthian of France and England. Monari (2003) quoted the same species from the Bajocian of northwestern France. We consider the first record of *Mactromyopsis* to be Pliensbachian.

MAGNOLOBIA Kurushin & Truschelev, 2001, p. 244

Kurushin and Truschelev (2001) reviewed the taxonomy of the genus *Daonella* and proposed a new genus, *Magnolobia* (type species: *Halobia premium* Kiparisova & Popov, 1946, *fide* Kurushin & Truschelev, 2001), a genus close to *Daonella* that differs from it mainly because "ribs are recurved, the anterior elevation is well-developed in the presence of the triangular posterior filed from the valve, in the large number of umbonal crurae, and in the presence of the intercalating ribs in the interrib spaces and in the accessory ribbing." Following McRoberts (1993), these differences are not considered to be sufficient to separate two genera (see discussion for the family Halobiidae in Included Genera, p. 91).

MALLETIA Des Moulins, 1832, p. 85

Although *Malletia* is a genus with an accepted range from the Cretaceous to the Holocene (Sepkoski, 2002), there are some Triassic and Jurassic records. Dagys and Kurushin (1985) reported Malletia pseudopraecursor Kurushin (in Dagys & Kurushin, 1985, pl. V,5-8) and Malletia sp. (pl. V,9 and pl. VI,1) from the Anisian and Ladinian of northern Siberia, but in our opinion, these specimens, especially on the basis of their hinge teeth, belong to Palaeoneilo elliptica var. praecursor (Frech, 1904). This last species was compared with M. pseudopraecursor by Kurushin (in Dagys & Kurushin, 1985), and the only difference between them is that the latter is more elongated than the former. Frech (1904, p. 12, fig. 9) figured three specimens from different ages and localities, showing the species variability in shell elongation. The oldest mention from the Jurassic is ? Malletia sp. from the Pliensbachian of South America (Damborenea, 1987a). Zhakarov and others (2006) also mentioned Malletia from the Pliensbachian of the Boreal region, but they did not treat the genus systematically and did not indicate the original source of data. Therefore Malletia is not considered here because it is not recorded with certainty before the Pliensbachian.

MARTESIA G. B. Sowerby, 1824 in 1821-1825, pl. 23

No occurrences from the Induan–Sinemurian interval are considered. Cox and others (1969) assigned it a ?Carboniferous, Jurassic–Holocene range, and Sepkoski (2002) assigned a Cretaceous (Cenomanian)–Holocene range, based on data offered by Kelly (1988). However, Kelly (1988) considered *Martesia* as a valid genus from the Paleocene to the Recent, since the Mesozoic records were not confirmed and the Carboniferous ones were unfounded (see discussion in Kelly, 1988, p. 366–367).

MCLEARNIA Crickmay, 1930b, p. 45

Mclearnia is not included here, because all quotes from the Lower Jurassic are doubtful, and it is considered to be a subgenus of *Camptonectes* (Kelly, 1984; Kelly, Dhondt, & Zakharov, 1984; Fürsich & Thomsen, 2005). Cox and others (1969) mentioned it as a doubtful genus from the Lower Cretaceous and indicated the need for further investigation. The genus was not widely mentioned, as the type specimens are poorly preserved and it is difficult to compare them with others. Therefore, most species were included in *Boreionectes* Zakharov, 1965, now considered to be a junior synonym of *Mclearnia* (Kelly, Dhondt, & Zakharov, 1984). *Mclearnia* was reported from the Hettangian of northeastern Asia (Kurushin, 1990), but nevertheless, Zakharov and others (2006) reported that its oldest record is from the Aalenian of northern Siberia and the Arctic region. The mention of Kurushin (1990) is not taken into account and considered to be unjustified.

MEGALODON J. de C. Sowerby, 1827, p. 131

Although there are numerous records of *Megalodon* from the Triassic (e.g., Parona, 1888, 1889; Tommasi, 1890; Bittner, 1895; Wurm, 1913; Trechmann, 1918; Schmidt, 1935; Dechaseaux, 1940; Leonardi, 1943; Marwick, 1953; Kiparisova, 1954; Terranini, 1958; Virgili, 1958; R. Hudson & Jefferies, 1961; Allasinaz, 1962, 1964, 1965; Brinkmann, 1966; Encheva, 1972; Fürsich & Wendt, 1977; Hallam, 1981; Lu & Chen, 1986) and from the Jurassic (e.g., Fraser, Bottjer, & Fischer, 2004); following Végh-Neubrandt (1982), we consider *Megalodon s.s.* as a typically Paleozoic genus, specifically restricted to the Devonian. This genus is closely related to *Neomegalodon* and *Triadomegalodon* (Végh-Neubrandt, 1982). The first was originally proposed as subgenus of *Megalodon*, and this is the origin of many of the Triassic quotes. Cox and others (1969), Hallam (2002), and Sepkoski (2002) considered it to be extinguished in the Rhaetian.

MESOMILTHA Chavan, 1938, p. 231

In recent years, *Mesomiltha* was reported from the Upper Triassic (e.g., Ivimey-Cook & others, 1999; Guex & others, 2003, 2004; Lucas & Tanner, 2004), although we could not confirm these records nor those from the lower Jurassic. Ivimey-Cook and others (1999) included specimens from the Rhaetian of England with some doubts, since the diagnostic features were not observed. In the other three papers mentioned above, *Mesomiltha* is listed from the Rhaetian of Nevada, but no systematic treatment was offered. Reports from the Hettangian of Japan (Z. Huang, 1986) are also questionable, because the species were only doubtfully attributed to *Mesomiltha*.

MOLUKKANA Krumbeck, 1923a, p. 219

Cox and others (1969) treated *Molukkana* as doubtful and noticed that it was poorly known and possibly indistinguishable from *Pachymya* (*Pachymya*). It is a monospecific genus; the type species, *M. seranensis* Krumbeck, 1923a, was reported by Krumbeck (1923a) and Cox and others (1969) from the Norian of Indonesia. No reports after 1969 were found, and everything suggests that all the old records originated with Krumbeck (1923a).

MUSCULUS Röding in Bolten, 1798, p. 156

No occurrences from the Induan–Sinemurian interval are considered. Cox and others (1969) assigned it a Jurassic–Holocene range; Sepkoski (2002) specified its origin in the upper Sinemurian [data from Hallam (1977), who mentioned the species *Musculus subcancellata* (Buvignier) from the Sinemurian of Europe, North America, and South America]. We are not taking this quote into account, because it is impossible to corroborate it and we have doubts about its validity. No reports were located in any paper after or before Cox and others (1969). In reviews by Damborenea (1987a, 1987b, 2002a) and Aberhan (1994a) of South American Jurassic bivalves of the Andean Cordillera, *Musculus* was not mentioned, but they included *Modiolus*, a genus with similar external morphology, which can be confused if specimens are not well preserved.

MYTILOIDES Brongniart, 1822, p. 320

Cox and others (1969) regarded *Mytiloides* as a subgenus of *Inoceramus* and assigned it a Jurassic–Upper Cretaceous range. Sepkoski (2002), without indication of the source, considered its origin in the Sinemurian. No mention of *Mytiloides* from the Lower Jurassic was located. Kauffman and Powell (1977) emended the diagnosis of *Mytiloides* and considered it to be a separate genus from *Inoceramus*. Harries and others (1996) noticed that *Mytiloides* had a Cretaceous origin and that all Jurassic species assigned to *Mytiloides* belong to other genera.

MYTILUS Linnaeus, 1758, p. 704

Although Cox and others (1969) assigned it an Upper Jurassic–Holocene range, *Mytilus* was mentioned repeatedly from the Triassic. The species most frequently mentioned for this period is *Mytilus eduliformis* Schlotheim, 1820 (e.g., Zorn, 1971; Busse 1972; Z. Yang & Yin, 1979; Dagys & Kurushin, 1985; Warth, 1990; Budurov & others, 1991; Gou, 1993). Márquez-Aliaga (1983, 1985) mentioned "*Mytilus*" from the Hispanic Muschelkalk, with the quotation marks indicating that it was not a Triassic genus. Waller (in Waller & Stanley, 2005) proposed a new genus, *Promysidiella* (type species: *Mysidiella cordillerana* Newton in Newton & others, 1987), and included several species, among them *Mytilus eduliformis* and *Mytilus otiosus* McLearn, 1947; the author also indicated that "some species described as '*Mytilus*' from the European Muschelkalk may prove to be members of this genus." Therefore we consider that the *Mytilus* range started in the Jurassic.

NEOENTOLIUM Romanov, 1985, p. 37

Romanov (1985) gave generic names to the four groups recognized by Staesche (1926) within *Entolium s.l.* (see discussion in Damborenea, 2002a, p. 42–44). The *Entolium cingulatum* Goldfuss group was referred to the new genus *Neoentolium* (type species: *Pecten cingulatus* Goldfuss, 1835 in 1833–1841), and he added other species [*N. cingulatum* (Goldfuss), *N. renevieri* (Oppel), *N. masticonense* (Lissajons), *N. partitum* (Cox), *N. radiatum* (Andreeva)]. Romanov (1985) assigned it a Hettangian–Kimmeridgian range in Europe and Asia. Two other authors, Freneix (1980) and Yamani (1983), proposed two new genera for Staesche's group, *Costentolium* Freneix, 1980, and *Cingentolium* Yamani, 1983. Waller (2006) considered *Neoentolium* to be a junior synonym of *Cingentolium* and questioned the validity of the latter, as Yamani confused the right and left valves, and several diagnostic features described by him were actually diagenetic marks. But Waller (2006) did not take into account Freneix (1980), who described *Entolium (Costentolium)* Freneix, 1980, based on the same type species as the other two genera, and thus has priority over them.

In the absence of a recent review on the *Entolium* group, we follow a conservative attitude and consider *Neoentolium* to be within *Entolium*, following Staesche (1926) (see discussion for the family Entoliidae in Included Genera, p. 103).

NEOPECTEN Bychkov, 1985, p. 11

Neopecten is not considered here, because we lack information about it, and none of the databases consulted mention it. The only information was provided by the Zoological Record: "Neopecten Gen nov of Aviculopectinidae, Type species N. oxytomaeformis, p. 11, Upper Triassic, Neopecten damesi (Böhm, 1903) Comb nov Transferred from, Pecten, p. 11 Neopecten oxytomaeformis sp. nov., Russia, Upper Triassic, p. 11 (Bychkov, N. 1985. [Upper Triassic molluscs of the Kenkeren Range (Koryak Plateau).]. Pokhialainen, V.P. [Eds]. [Mesozoic Bivalvia and Cephalopoda from northeastern Russia. Collected scientific articles.] Akademiya Nauk SSSR, Magadan. 1985: 1–153. Chapter pagination: 5–24)." It was mentioned from the Norian by Polubotko and Repin (1990) and Zakharov, Kurushin, and Pokhialainen (1996).

NEPTUNELLA Astafieva, 1997, p. 24

Neptunella was replaced by *Neptunopecten* by Astafieva (2001). Astafieva (1997) proposed *Neptunella*, unaware that the name had already been used three times for different mollusks: *Neptunella* Gray, 1854, *Neptunella* Meek, 1864, p. 38, and *Neptunella* Verrill, 1873, p. 639 (Astafieva, 2001).

NUCULA Lamarck, 1799, p. 87

No occurrences from the Induan–Sinemurian interval are considered, following Cox and others (1969), who assigned it a Cretaceous–Holocene range, although many later authors still used the name for Triassic and Jurassic specimens (e.g., Encheva, 1972; Bachmann, 1973; Quintero & others, 1977; Harper, Forsythe, & Palmer, 1998).

NUCULOPSIS Girty, 1911, p. 133

Although *Nuculopsis* was reported from the Triassic (e.g., Hayami, 1975), it was always reported as *Nuculopsis* (*Palaeonucula*). These mentions are not taken into account, because *Palaeonucula* is considered here as a separate genus from *Nuculopsis*, although, prior to Cox and others (1969), *Palaeonucula* was regarded as a subgenus of *Nuculopsis* and even its junior synonym (Nakazawa & Newell, 1968). Carter (1990a, p. 149–150) stated that they were different and distinct genera: *Nuculopsis* possessed a nacreous inner shell layer, while the inner shell layer of *Palaeonucula* was homogeneous, but when internal characters are observed, they can be confused. The range of *Nuculopsis* is emended with respect to Sepkoski (2002), who considered its extinction to have been in the Early Triassic (early Induan). Presumably this datum was taken from Cox and others (1969), although a Carboniferous–Permian range was assigned there.

NUCUNDATA Waterhouse, 1964, p. 641

Nucundata is not taken into account here, because it is considered to be a subgenus of Veteranella Patte, 1926. Waterhouse (1964, p. 641) described Nucundata based on its ornamentation and distinguished it from Glyptoleda by: "A minor change in ornament occurred in a New Zealand stock of Phestiinae, comparable to but less drastic than that shown by the Australasian genus Glyptoleda, and probably equally short-lived. The change simply emphasized tangential instead of concentric ornament." Subsequently, Puri in Cox and others (1969) considered it to be a subgenus of Veteranella, together with Glyptoleda (see discussion for Glyptoleda in this section, p. 161). This author distinguished V. (Glyptoleda) from V. (Nucundata), because the latter had less prominent ornamentation and V. (Veteranella) was distinguished from V. (Nucundata) by the non-constricted or constricted posterior part, respectively. The differences in ornamentation between the three subgenera of Veteranella (Veteranella, Glyptoleda, and Nucundata) were considered to be variations within the genus. However, Waterhouse (1980a, p. 102) noticed that "externally Nucundata and Veteranella are very similar and unaware of the internal characters of Veteranella, for example, the presence or absence of ribs on the escutcheon, or ribs on the umbonal internal part ... [characters to which Waterhouse (1964) gave considerable importance when describing Nucundata and several species of *Glyptoleda*] . . . the name *Nucundata* can be used." This was followed by Waterhouse (1987) and J. Chen, Liu, and Lan (1983), who also proposed a new subfamily, Veteranellinae, using the variation in the ornamentation as criteria. Z. Fang and Cope (2004, p. 1125) disagreed with this last decision, because they considered: "Clearly a factor that may be of lower taxonomic significance at specific level can hardly be significant at subfamilial level. For this reason we reject the subfamily Veteranellinae . . ." We follow Puri in Cox and others (1969) and consider Nucundata to be a subgenus of Veteranella, because variation in the ornamentation is a problematic criterion even at the specific level (Z. Fang & Cope, 2004).

OPISOMA Stoliczka, 1871 in 1870-1871, p. 276

No occurrences from the Induan–Sinemurian interval are considered. Cox and others (1969) assigned it a Jurassic (Infraliassic–Rauracian [equivalent to middle Oxfordian]) range; Sepkoski (2002), a Jurassic (upper Pliensbachian–upper Oxfordian) range based on Hallam (1981), but in this last paper, there is no mention of *Opisoma*. Although the genus was widely reported from the Pliensbachian (especially in "Lithiotis" facies) (Hallam, 1969, 1972, 1976, 1977; Accorsi Benini, 1981; Buser & Debeljak, 1994; Liu, 1995; Aberhan & Hillebrandt, 1999; Damborenea, 2002b; Saadi & others, 2003; Fraser, Bottjer, & Fischer, 2004), there is no evidence from the Sinemurian.

PACHYMYONIA Dun, 1932, p. 411

Pachymyonia is not considered here, because it is regarded as a subgenus of *Myonia*, following Cox and others (1969). Waterhouse (1969) argued that the differences in position and morphology of the adductor and retractor muscles between *Myonia* and *Pachymyonia* perfectly supported the separation of the two genera. This was followed in other papers (Waterhouse, 1987, 2002). However, other

authors preferred to keep it as a subgenus of *Myonia* (Astafieva-Urbaytis, 1976; Scarlato & Starobogatov, 1979).

PACIFIHALOBIA Polubotko, 1990, p. 131

According to McRoberts (1993), *Pacifihalobia* is considered to be a junior synonym of *Halobia* Bronn, 1830a, since the features used for proposing this genus are of specific level (see discussion for the family Halobiidae in Included Genera, p. 91). Polubotko, Payevskaya, and Repin (2001) considered it to be a valid genus.

PALAEOENTOLIUM Romanov, 1985, p. 35

Palaeoentolium is not considered here, because it was based on Pleuronectites discites Schlotheim, 1820, a species with some problems disregarded by Romanov (Waller, 2006, p. 325). Romanov (1985) included the following species: *P. microtis* (Wittenburg), *P. discites* (Schlotheim), *P. marginiplicatum* (Kittl), *P. liscaviense* (Giebel), *P. weissenbachense*, and *P. hallense* (Wohermann). We consider Pleuronectites discites Schlotheim, 1820, as an Entolium (see discussion for the family Entoliidae in Included Genera, p. 103).

PALAEOLOPHA Malchus, 1990, p. 102

Junior synonym of Actinostreon Bayle, 1878 (Hautmann, 2001a; Checa & Jiménez, 2003b; Márquez-Aliaga & others, 2005). The genus proposed by Malchus (1990) was placed in synonymy by Hautmann (2001a), basically for two reasons: first, the type species designated by Malchus, Ostrea haidingeriana Emmrich, 1853, has a shell microstructure that suggests it should be allocated to Actinostreon; second, Malchus (1990) figured the type species of Actinostreon, Ostrea solitaria J. de C. Sowerby, 1825, as Palaeolopha solitaria (Hautmann, 2001a). We follow Hautmann in this case, because the synonymy is more than justified.

PARAHALOBIA Yin & Hsu, 1938 in Chen, 1976, p. 224

Parahalobia was proposed as a subgenus of *Halobia* to accommodate *Halobia* forms lacking radial ornamentation from the Anisian and Ladinian of China (H. J. Campbell, 1994). Subsequently, several authors considered it at the genus level, but none of them justified this decision (e.g., C. Chen & Yu, 1976; Sha, Chen, & Qi, 1990). McRoberts (1993, p. 201) considered *Parahalobia* as a junior synonym of *Halobia* (see discussion for the family Halobiidae in Included Genera, p. 91). However, Sepkoski (2002) considered it in his compendium of genera, taking the data from H. Yin (1985), who regarded it as a subgenus of *Halobia*. H. J. Campbell (1994) indicated that *Parahalobia* could be relocated as a subgenus of *Halobia*, because it is similar in all respects except for the lack of radial ornamentation. For all these reasons, we do not regard *Parahalobia* as an independent genus.

PAULLIA (no author)

Schubert (1993) and Schubert and Bottjer (1995) mentioned *Paullia* as a new genus that would accommodate some species attributed to *Pernopecten*, but they did not formally describe it and mentioned it as a personal communication by Boyd in 1991; however, Boyd never published its description and therefore the name is invalid (*nomen nudum*), although it is taken into account in the PBDB (online).

PECTINULA A. F. Leanza, 1943, p. 244

Junior synonym of Posidonotis Losacco, 1942 (Damborenea, 1986, 1987b). Damborenea (1986, 1987b), after careful consideration of the type species of Pectinula (Pectinula cancellata A. F. Leanza, 1943) and Posidonotis (Posidonotis dainelii Losacco, 1942), concluded that the only difference between them was that the former had an anterior auricle, absent in the latter, a fact that was attributed by Damborenea to the poor preservation of Losacco's material, as in many specimens of Pectinula cancellata, the anterior auricle was not observed. Later, Hayami (1988), after studying his own wellpreserved material, restored the validity of Pectinula. The author based his argument on the differences in auricle shape and in the lack of byssal notch in juvenile stages in his specimens. Very few later papers mentioned the name Pectinula (see Monari, 1994; Waller, 2006); most used Posidonotis, accepting the synonym proposed (see Damborenea, 1993, 2001; Aberhan, 1994a, 1998a, 2001; Aberhan & Pálfy, 1996; Fürsich & others, 2001).

PERAMPLIATA Arkell, 1936 in 1929–1937, p. xx

No occurrences from the Induan–Sinemurian interval are considered. Cox and others (1969) assigned it a Jurassic range, but there is no evidence of *Perampliata* until the Oxfordian (Hallam, 1976; Liu, 1995).

PERIBOSITRA C. Chen, 1981, p. 81

Junior synonym of *Claraia* Bittner, 1901a (Nakazawa, 1992; Z. Fang & others, 2009). C. Chen in Zhao and others (1981) proposed *Peribositra*, based on the type species *P. baoqingensis* Chen, 1981, and included it in the family Posidoniidae. Z. Fang and others (2009) placed it in synonymy with *Claraia*, according to, among others, Waller and Stanley (2005, p. 20), but the latter authors only stated that *Peribositra* was a claraiid, not a posidoniid. We follow Z. Fang and others (2009) in this regard (Chuzhen Chen was the author of *Peribositra* and is a coauthor of Z. Fang and others [2009]).

PERIBOSITRIA Kurushin & Trushchelev, 1989, p. 59

Junior synonym of *Bositra* De Gregorio, 1886 (Waller & Stanley, 2005). *Peribositria* was proposed by Kurushin and Trushchelev (1989) from the Triassic of Siberia and is still used by Russian authors, particularly in biostratigraphic syntheses (e.g., Polubotko, Payevskaya, & Repin, 2001; Klets, 2006; Konstantinov, Sobolev, & Yadernkin, 2007). Waller and Stanley (2005) rejected all differences with *Bositra* mentioned by Kurushin and Trushchelev (1989) to separate the genera, since most of these were due to taphonomic issues (see discussion in Waller & Stanley, 2005, p. 20). However, McRoberts (2010) decided provisionally to retain the genus, because the ligamental system of the Triassic forms is not well known.

PERIHALOBIA Gruber, 1976, p. 192

Although *Perihalobia* was originally proposed as a subgenus of *Halobia*, later it was considered at the generic level, which is totally unjustified, according to several authors (McRoberts, 1993, 2000; H. J. Campbell, 1994), because they are indistinguishable (see discussion for Halobiidae in Included Genera, p. 91).

PHOLADOMYOCARDIA Szajnocha, 1889, p. 88

No occurrences from the Induan–Sinemurian interval are considered. Cox and others (1969) assigned it a Jurassic range in South America (Peru), based on the original description of the genus, but there is no more information about it. Moreover, *Pholadomyocardia* was considered to be a doubtful genus, because no internal characters are known (Cox & others, 1969).

PHYMODONUCULA Guo, 1988, p. 112

Phymodonucula was erected by Guo (1988) from the Carnian of southwestern China and included in the family Nuculidae. Z. Fang and others (2009, p. 6) considered *Phymodonucula* a dubious genus, because the ligament is not known, as none of the specimens have the ligament properly preserved. This is followed here.

PLACUNOPSIS Morris & Lycett, 1853 in 1851-1855, p. 5

Following Todd and Palmer (2002), we consider the origin of *Placunopsis* to be in the Middle Jurassic. There are several Triassic species attributed to *Placunopsis* (see discussion for the genus *Pseudoplacunopsis* in Included Genera, p. 70).

PLATYMYOIDEA Cox, 1964, p. 42

Cox and others (1969) assigned it a Lower Jurassic–Lower Cretaceous range. The oldest report dates from the Pliensbachian (Damborenea in Damborenea & González-León, 1997; Aberhan, 2004). Liu (1995) quoted the genus from the Sinemurian of Morocco, but we could not corroborate this information. Furthermore, Hautmann (2001b) attributed a single, fragmentary specimen from the Norian of Iran to *Platymyoidea* sp. We are not taking this into account, because it is necessary to check it with new and better material. For all of these reasons, we will not consider occurrences from the Induan–Sinemurian interval until reliable information is available.

PLESIOCYPRINA Fischer, 1887 in 1880–1887, p. 1072

Although several sources (Cox & others, 1969; Sepkoski, 2002; Gardner, 2005; J. Yin & Grant-Mackie, 2005) assigned an Upper Triassic (Rhaetian)–Upper Jurassic range, it seems that all data originated from Cox and others (1969). Nevertheless, *Plesiocyprina* was not reported for our study interval. Neither Diener (1923) nor Kutassy (1931) referred to it in their treatises. The oldest record is Toarcian (Gahr, 2002).

PLEUROMYSIDIA Ichikawa, 1954, p. 52

Junior synonym of *Otapiria* Marwick, 1935 (Zakharov, 1962). According to Ando (1988), Zakharov established the synonymy of *Pleuromysidia* with *Otapiria*, because he did not observe significant morphological differences between them, and this was followed by most authors (e.g., Hayami, 1975; Ando, 1983, 1988; Kobayashi & Tamura, 1983b; Okuneva, 1986; Damborenea, 1987b).

PLICATOSTYLUS Lupher & Packard, 1930, p. 204

Junior synonym of *Lithiotis* Gümbel, 1871 (Broglio-Loriga & Neri, 1976; Accorsi-Benini & Broglio-Loriga, 1977). This synonymy was accepted by most subsequent authors (e.g., P. L. Smith & Tip-

PLICATULA Lamarck, 1801, p. 132

Cox and others (1969) assigned Plicatula a Middle Triassic-Holocene range. Subsequently, it was consistently mentioned from both the Triassic and the Jurassic. The problem with this genus and its range is that many species attributed to Plicatula are now regarded as belonging to other genera, such as Harpax Parkinson, 1811, Eoplicatula Carter, 1990a, and Pseudoplacunopsis Bittner, 1895. Hautmann (2001a) did not consider Plicatula from the Triassic. Regarding the Jurassic specimens, in most of them, the hinge is not preserved, and it is difficult to know which species should be assigned to Harpax and which to Plicatula. Damborenea (2002a) believed that the genera coexisted in the early history of the group during the Upper Triassic, but no specific records were found. Following Hautmann (2001a), we do not consider *Plicatula* to be from the Triassic, and, although it may have been present in the lower Jurassic, we do not have enough evidence to decide.

POLIDEVCIA Chernyshev, 1951, p. 25

Since its proposal over 50 years ago, there has been an open debate about the validity of *Polidevcia*. According to different authors, it was considered to be a junior synonym of *Phestia* (McAlester in Cox & others, 1969), a junior synonym of *Culunana* Lintz, 1958, a subgenus of *Phestia* (Nakazawa & Newell, 1968; González, 1969; Carter, 1990a), or a valid and independent genus (Ciriacks, 1963; Waterhouse, 1964) (see discussions in Carter, 1990a, p. 153; Bradshaw, 1999, p. 70; and Anelli, Rocha-Campos, & Simões, 2002, p. 170–171). Here we follow the approach adopted by Carter (1990a), treating *Polidevcia* as a subgenus of *Phestia*.

POSIDONIA Bronn, 1828, p. 268

According to Waller and Stanley (2005), *Posidonia* is a Paleozoic genus. Most species described under the genus *Posidonia* from the Lower and Middle Triassic belong to *Bositra* De Gregorio, 1886 (as emended in Waller & Stanley, 2005).

PRAECHLAMYS Allasinaz, 1972, p. 340

Although some authors (Waller & Marincovich, 1992; Monari, 1994; Damborenea, 2002a) considered *Praechlamys* Allasinaz, 1972, at the generic level to be separate from *Chlamys*, Allasinaz (1972) described it as a subgenus of *Chlamys*. Taking into account the problems with the so-called *Chlamys*-like forms (see discussion for *Chlamys* in Included Genera, p. 97), we follow a conservative attitude and follow Allasinaz's original concept, even knowing the problems involved (see Damborenea, 2002a). Other authors, such as Posenato (2008b), followed this same position, and Hautmann (2001b) even questioned the validity of *Praechlamys*, since, in his opinion, the type of ornamentation is not an important taxonomic character at subgeneric level, and Allasinaz (1972) used this criterion when separating the subgenera *C. (Chlamys)*, *C. (Praechlamys)*, and *C. (Granulochlamys)*.

PRIMAHALOBIA Polubotko, 1988, p. 98

Polubotko recognized *Primahalobia* as a new subgenus of *Indigirohalobia* Polubotko, 1984, based on the angular relationship of the anterior auricle (H. J. Campbell, 1994). Subsequently, Polubotko, Payevskaya, and Repin (2001) raised it to generic level. Following McRoberts (1993), we do not consider it to be valid, because the diagnostic criteria for *Primahalobia* are used by most authors at the specific level (see discussion for the family Halobiidae in Included Genera, p. 91).

PRONOELLA Fischer, 1887 in 1880–1887, p. 1087

No occurrences from Induan–Sinemurian interval are considered. Cox and others (1969) included two subgenera in *Pronoella*: *P. (Pronoella*) and *P. (Gythemon*); only the first one interests us because of its Lower Jurassic range. Nevertheless, no records were located from Hettangian or Sinemurian beds, although it was widely mentioned from the Pliensbachian (Hallam, 1972, 1976, 1977, 1987; Fürsich, 1982; Liu, 1995; Holzapfel, 1998; Gahr, 2002).

PROPEAMUSSIUM De Gregorio, 1884, p. 119

Following Damborenea (1998), we consider that *Propeamussium* ranges from the Upper Cretaceous to the Recent. Following the original concept of *Parvamussium* and *Propeamussium*, Triassic and Jurassic *Propeamussium* records are better accommodated in *Parvamussium* Sacco, 1897 (see discussion for *Parvamussium* in Included Genera, p. 95).

PROSPONDYLUS Zimmermann, 1886, p. 105

Prospondylus was considered to be a junior synonym of *Pseudo-monotis* von Beyrich, 1862, by Newell (1937), and this was followed in Cox and others (1969). Newell and Boyd (1970) reestablished the validity of *Prospondylus*. Although *Prospondylus* was reported from the Triassic, those records are not considered, since it is regarded as a Paleozoic genus (Newell & Boyd, 1970; Márquez-Aliaga & others, 2005). Hautmann (2001a, 2001b) pointed that it may be present in Triassic sediments ("*Hinnites*" comptus Bittner, 1898), but this is not proven.

PROTODICERAS G. Böhm, 1892, p. 51

Although *Protodiceras* was reported from the Sinemurian, no occurrences from the Induan–Sinemurian interval are considered. Cox and others (1969) reported it from the lower Lower Jurassic. Hallam (1976) quoted *Protodiceras pumilum* (Guembel) from the Sinemurian and Pliensbachian of Europe, data incorporated by Sepkoski (2002) in his database. However, Hallam (1977) only reported it from the Pliensbachian of Europe. No undoubtful Sinemurian records were found.

PSEUDOCLARAIA Z. Zhang, 1980, p. 443

Junior synonym of *Claraia* Bittner, 1901a (Newell & Boyd, 1995; F. Yang, Peng, & Gao, 2001; He, Feng, & others, 2007; Z. Fang & others, 2009). Z. Zhang (1980) proposed *Pseudoclaraia* and designated *Claraia wangi* (Patte, 1935) as type species. He considered *Pseudoclaraia* to be very close to *Claraia* and argued that it differed from it by the larger right anterior auricle and by differences in shape and orientation of the byssal sinus. Nakazawa

(1996) included it as a subgenus of *Claraia*, since differences were not great and often their differentiation was very difficult. Newell and Boyd (1995) considered Pseudoclaraia to be a junior synonym of Claraia, because no quantitative and comparative studies of local populations were made before the proposal of the new genus. F. Yang, Peng, and Gao (2001) reached the same conclusion about the synonymy after carrying out a comparative study between Claraioides, Claraia, and Pseudoclaraia; they concluded that the only differences between these three genera were: 1) right anterior auricle shape, and 2) byssal sinus shape. These differences, in their opinion, were not reliable to separate the genera, because the byssal sinus shape is due to the adaptation of Claraia to different environments, and, on the other hand, the diagnosis of Claraia states nothing about the size and shape of the byssal sinus; therefore, this character can only be used as diagnostic at the specific level. For all these reasons, F. Yang, Peng, and Gao (2001) considered Pseudoclaraia to be a junior synonym of Claraia. Kotlyar, Zakharov, and Polubotko (2004) argued for the validity of Pseudoclaraia but did not provide new arguments. He, Feng, and others (2007), following Boyd and Newell (1995) and F. Yang, Peng, and Gao (2001), reached the same conclusion regarding the synonymy of Pseudoclaraia after studying the variation in ornamentation and byssal sinus shape of all species included in Claraia in relation to their stratigraphic ranges and paleogeographic distribution (see He, Feng, & others, 2007, table 1, p. 1016 and discussion). The authors showed there was a gradual morphological change from the early Changhsingian to Induan, indicating that the shape, size, and orientation of the byssal sinus was not a valid criterion for discrimination at the generic level in this group.

PSEUDOMONOTIS von Beyrich, 1862, p. 10

Although *Pseudomonotis* is frequently mentioned from the Triassic and Jurassic (see section PBDB age range for example, online), we follow Cox and others (1969), Newell and Boyd (1970, 1989, 1995), and Hautmann (2001a, 2001b) in the idea that it was an exclusively Paleozoic genus. Newell and Boyd (1970, 1989, 1995) showed the external morphological similarity between *Pseudomonotis* and *Gryphaea* (see Newell & Boyd, 1995, fig. 47). In many cases, these genera can only be distinguished by the shell microstructure and the attachment valve (right in *Pseudomonotis* and left in *Gryphaea*) (Newell & Boyd, 1995). In the words of Newell and Boyd (1989, p. 2): "*Pseudomonotis* is not an oyster. It is an aberrant scallop, an aviculopectinacean with an oysterlike habit of fixation." Because of this external resemblance, *Pseudomonotis* is commonly confused with Triassic and Jurassic ostreids.

PSEUDOPACHYMYTILUS Krumbeck, 1923b, p. 87

No occurrences from the Induan–Sinemurian interval are considered, although both Cox and others (1969) and Sepkoski (2002) assigned it a Lower Jurassic range, because only Pliensbachian records were reported (e.g., Hallam, 1972; Liu, 1995, 1999; Posenato & Avanzini, 2004).

PSEUDOPIS Cox, 1946, p. 44

No occurrences from Induan–Sinemurian interval are considered, although Cox and others (1969) assigned it a Lower Jurassic range, the same age indicated by Cox (1946) when proposing *Pseudopis*. The oldest record is Pliensbachian (Hallam, 1976, 1977, 1987; Liu, 1995).

PTEROCLARAIA Guo, 1985, p. 266

Waterhouse (2000, p. 167) raised *Pteroclaraia* to genus level (Guo [1985] had proposed it as a subgenus of *Claraia*), arguing that "the size and definition of the posterior wing are distinctive and easily recognised features." Nevertheless, together with J. Chen and Komatsu (2002) and Z. Fang and others (2009), we consider *Pteroclaraia* to be a subgenus of *Claraia*, since this character is not regarded as diagnostic at the genus level (He, Feng, & others, 2007).

PTEROHALOBIA Guo, 1985, p. 266

Z. Fang and others (2009) placed *Pterohalobia* in synonymy with *Daonella* Mojsisovics, 1874. Their reasons were: "The anterior auricle of the figures of the holotype were not appropriately clipped off by Guo (1985, pl. 16,4b) and its ornamentation is very similar to that of *Daonella boeckhi* Mojsisovics." Guo (1985) erected *Pterohalobia* based on *P. productalata* Guo, 1985, from the Anisian of southwestern China. The material figured by Guo (1985) is very poor.

PTEROPERNA Lycett, 1850, p. 421

Fürsich and Werner (1988, p. 106) considered *Pteroperna* as a junior synonym of *Pteria* Scopoli, 1777, because, in their view, "the main diagnostic feature of *Pteroperna*, the presence of 1 to 4 external longitudinal grooves on the exterior of the posterior wing, does not justify separation at the generic level. Such grooves occasionally are also present in individuals of Recent species of *Pteria*. Furthermore, populations from the Kimmeridgian of Consolaçao show that the number and distinctness of such grooves varies greatly." Hautmann (2001b) provisionally considered it to be a subgenus of *Pteria*, as his specimens are similar to the type species of *Pteroperna*. We judge it to be more appropriate to keep it as a subgenus of *Pteria*, because, although they are very similar, *Pteroperna* is a useful concept and easily recognized in the Upper Triassic and Jurassic.

PTEROPIRIA Waterhouse, 2008, p. 169

This genus was proposed by Waterhouse (2008, p. 169) with *Otapiria tailleuri* Imlay, 1967, p. 3, as type species. It was diagnosed as "characterized by presence of well developed posterior wing on left and right valve." Within the range of morphological variability accepted within *Otapiria, O. taillieuri* Imlay is extremely similar to the type species of *Otapiria, O. marshalli* (Trechmann), see Imlay (1967, pl. 1) for figures of both species.

PTYCHOSTOLIS Tullberg, 1881, p. 14

No occurrences from the Induan–Sinemurian interval are considered, although both Cox and others (1969) and Sepkoski (2002) assigned it a Jurassic range. Even though it was impossible to see Tullberg (1881), where *Ptychostolis* was proposed and which is also the only record located, we know the author assigned it a Tithonian range. However, it is difficult to know the exact age of the specimens studied by Tullberg (1881), because they come from two different localities and four different lithological units; the author reported the species *Ptychostolis nordenskioeldii* from the Kalkstein of Skodde Bituminoeser Bay, where only *Ammonites okensis* (d'Orbigny) (now

PULVINITES Defrance in Blainville, 1824, p. 316

Palmer (1984) considered that *Pulvinites* had its origin in the Middle Jurassic, but Damborenea (1987b) assigned some Pliensbachian South American specimens to this genus and noted that some Hettangian specimens of France could be included in *Pulvinites*. A review of these specimens is needed in order to include *Pulvinites* in our study interval.

QUADRATIA Yin, 1974, p. 25

It is a junior homonym of *Quadratia* Muir-Wood & Cooper, 1960, p. 161 (Brachiopoda; family Productellidae) (Z. Fang, 2009). The name was replaced by *Guizhoumyophoria*, new name, based on the same type species, *Quadratia quadrata* Yin (Z. Fang, 2009).

RETROCERAMUS Koschelkina, 1957, p. 29

No occurrences from the Induan–Sinemurian interval are considered, although a doubtful mention from the Sinemurian in PBDB (online) was located. This is from Aberhan's (1995) unpublished material, and it is only doubtfully attributed to this genus (*Retroceramus* sp). In addition, J. Chen (1982a) proposed a new species, *Retroceramus? xiaoshuiensis* Chen from the Upper Triassic of southern China; this uncertain mention is not taken into account. Throughout the consulted literature, the origin of *Retroceramus* is regarded as Middle Jurassic (Hallam, 1976, 1977; Crampton, 1988; Damborenea, 1996b; N. Hudson, 2003; Damborenea & Manceñido, 2005a; Zakharov & others, 2006) or Toarcian at the earliest (Hallam, 1987; J. Yin & Grant-Mackie, 2005).

RHYNCHOPTERUS Gabb, 1864, p. 31

Junior synonym of *Pteria* Scopoli, 1777 (see Silberling & Nichols, 1982, p. 66). *Rhynchopterus* Gabb, 1864, is a junior homonym of *Rynchopterus* Shrank, 1798, a bettle genus. The name proposed by Shrank was a mistake, because surely he meant *Rhychophorus* (Silberling & Nichols, 1982), but even so, it is a valid name and therefore *Rhynchopterus* Gabb should not be used for a bivalve. Instead of proposing a new name for the poorly known bivalve genus, it is more appropriate to leave it as a probable synonym of *Pteria s.l.*

ROSSIODUS Allasinaz, 1965, p. 120

Junior synonym of *Neomegalodon* Gümbel, 1862 (H. Yao & others, 2007). See discussion for *Gemmellarodus* in this section (p. 161).

RUGICLARAIA Waterhouse, 2000, p. 179

Waterhouse (2000) proposed *Rugiclaraia* with *Claraia aurita* (Hauer, 1850) as type species. Waterhouse (2000, p. 179) indicated that *R. aurita* differs from the type species of *Claraia, Claraia clarae* (Hauer, 1850) by the absence of radial ornament, but He, Feng, and others (2007) attributed a concentric and radial ornamentation to both species. Another feature mentioned by Waterhouse (2000) for his new genus is the auricle size and shape, characters used to discriminate species by most authors (F. Yang, Peng, & Gao, 2001).

For all these reasons, we do not consider *Rugiclaraia* to be a valid genus and include its type species in *Claraia*.

SATURNELLA Astafieva, 1994, p. 16

Name replaced by *Saturnopecten* Astafieva, 2001. Astafieva (1994) proposed *Saturnella* without knowing that the name was already being used for a Jurassic foraminiferan, *Saturnella* Hedinger, 1993, p. 33 (Astafieva, 2001).

SATURNOPECTEN Astafieva, 2001, p. 557

According to Newell and Boyd (1995, p. 85), *Saturnopecten* was not taken into account, because it was based on very few specimens, mostly broken or poorly preserved, and no studies of microstructure and/or hinges were carried out.

SEPTIFER Recluz, 1848 in 1848-1849, p. 275

Although both Cox and others (1969) and Sepkoski (2002) assigned it a Triassic-Holocene range; Carter (1990a) argued that the Triassic specimens attributed to Septifer were not related to the type species (Mytilus bilocularis Linnaeus, 1758, p. 705), and they differed in shell microstructure, mineralogy, and other features, such as the position of the ligament. The Triassic species attributed to Septifer are: S. eduliformis (Schlotheim, 1820) (Ürlichs, 1992; Posenato, 2002; Posenato & others, 2002); S. rugulosus (Bittner, 1895) (Zardini, 1981); Mytilus (Septifer) praeacutus Klipstein, 1843 in 1843–1845 (Diener, 1923); and Mytilus (Septifer) praeacutiformis Wilckens, 1909 (Diener, 1923); all are mentioned from Ladinian and Carnian. No more records of the genus are known until the Cretaceous. The species eduliformis was included in Promysidiella Waller & Stanley, 2005, based on the study of collections in various museums (Waller & Stanley, 2005). Carter (1990a) showed that rugulosus did not match Septifer due to substantial differences with its type species. And praeacutus and praeacutiformis were originally described in the genus Mytilus, being included in the subgenus of Septifer by Diener (1923). Szente and Vörös (in Budai & others, 2003) mentioned Septifer? sp. from the Middle Triassic, but this record is not taken into account, because its generic attribution is doubtful. Although this topic requires a major review, it seems that Septifer has not been recorded from the Triassic.

SICHUANTRIGONIA Gou, 1993, p. 24

Sichuantrigonia was placed in synonymy with *Acanomyphoria* Guo, 1985, by Z. Fang and others (2009), because the ornament is very similar in both genera and that is practically the only difference.

SOLENOMORPHA Cockerell, 1903, p. 559

Solenomorpha is a typically Paleozoic genus. Cox and others (1969) assigned it a Lower Devonian–upper Permian range. However, Allasinaz (1966) mentioned it from the Carnian when he transferred the species *Cuspidaria gladius* Laube, 1865, to *Solenomorpha*. Allasinaz (1966) also considered that all Triassic species (without mentioning them) attributed to *Cuspidaria* should be included in *"Solenopsis"* (junior synonym of *Solenomorpha*). We have no opinion on the subject. The only species mentioned by Allasinaz (1966) was *C. gladius*, and in the figures (pl. 41,1–2), the characters he discussed in the text are not observed. If we compare these figures with those

offered by Cox and others (1969, p. 821, fig. F3, 2) or by some others on Paleozoic specimens (e.g., LaRocque, 1950, pl. 17,8–11; Hoare, Heaney, & Mapes, 1989, fig. 7.9), they do not seem to have much in common. *Solenomorpha gladius* was subsequently mentioned from the Carnian following Allasinaz (1966) (Jelen, 1988; Jurkovsek, 1993). We follow Cox and others (1969) in the range considered for this genus.

SOMAPTERIA Tamura, 1960, p. 224

No occurrences from the Induan–Sinemurian interval are considered. According to Cox and others (1969), *Somapteria* was reported from the Upper Jurassic of Japan. *Somapteria* is a monospecific genus limited to the Kimmeridgian of Japan (Hayami, 1975). Tëmkin (2006) assigned it a Middle Triassic range, but this must be a mistake, because none of the sources mentioned in his table offered this information.

SPONDYLUS Linnaeus, 1758, p. 690

No occurrences from the Induan-Sinemurian interval are considered. The oldest specimen attributable to Spondylidae was reported from the Bajocian of France (Harper in Waller, 2006); according to Skelton and Benton (1993), the first species attributable to this family should be Spondylus consobrinus Deslongchamps. Malchus (2004) extended the family range back to the upper Permian, but he did not justify it. Spondylus is rarely mentioned from the Jurassic, and it is often mistaken for *Eopecten* Douvillé, 1897 (Waller, 2006). Cox and others (1969) assigned it a Jurassic-Holocene range, but in previous papers, it was widely quoted from the Triassic. Frequently, these specimens could be referred to the Triassic Newaagia Hertlein, 1952 (Waller, 2006). Moreover, there is another similar genus with which it may be mistaken: Spondylopecten Röder, 1882. Johnson (1984) stated that Spondylopecten ranged from the Middle Jurassic, but Waller (2006) reported it from the Upper Triassic, referring to some specimens in the Mesozoic Stratigraphy collections of the Smithsonian Institution from Nevada, attributed to Spondylopecten. However, Waller (2006) did not describe or figure the material, and we are not taking this record into account.

STREBLOCHONDRIA Newell, 1938, p. 80

Although Streblochondria is considered to be a Paleozoic genus (Newell, 1938; Newell & Boyd, 1995), several Triassic records were located (e.g., Nakazawa, 1971; Tamura, 1973; Hayami, 1975; Tamura & others, 1975; Fraiser & Bottjer, 2007a), but none of them proved its occurrence after the upper Permian. Nakazawa (1971) tentatively included his specimens from the Triassic of Japan in "Streblopteria" matsushitai Nakazawa, 1971, because they were referable to the subfamily Streblochondriniinae Newell, 1938, and he preferred this genus over others of this subfamily. Hayami (1975) simply repeated Nakazawa's data (1971). Tamura (1973) tentatively referred his specimens from the Middle Triassic of Malaysia to this genus (Streblochondria? sp.), because, although they showed the typical ornamentation of Streblochondria, the ligament was not observed in the three deformed specimens (two external molds and the upper part of internal mold). Tamura and others (1975) repeated Tamura's data (1973). Finally, Fraiser and Bottjer (2007a) assigned four valves from the Triassic of Japan to Streblochondria, but they

neither figured nor described the specimens, and we are not taking this record into account.

STREBLOPTERINELLA [Kurushin, 1998]

The generic name *Streblopterinella* was introduced by N. I. Kurushin (1998) in his unpublished doctoral thesis. The genus *Streblopterinella* Kurushin comprised seven species: *Streblopteria newelli* Kurushin, 1982; *Streblopteria jakutica* Kurushin in Dagys & Kurushin, 1985; *Streblopteria egorovi* Kurushin, 1984, as well as four further species which are *nomina nuda* (described in Kurushin, 1998). Unfortunately, the generic name *Streblopterinella* has never been officially published (in accordance with ICZN rules, 1999) by the author, and it is considered *nomen nudum* (Konstantinov, personal communication, 2011).

TAIMYRODON Sanin, 1973, p. 92

Taimyrodon was originally described from the Upper Cretaceous of northern Siberia (Sanin, 1973). Subsequently, it was mentioned from Triassic (Dagys & Kurushin, 1985; Kurushin, 1990; Klets, 2006) and Lower Jurassic deposits (Meledina & Shurygin, 2001; Zakharov & others, 2006). Most of these records did not figure the specimens, and specimens in the others do not show the hinge features and therefore could belong to *Palaeoneilo*.

TEINONUCULANA Zhang in Zhang, Wang, & Zhou, 1977, p. 9

R. Zhang in R. Zhang, Wang, and Zhou (1977) proposed *Teinonuculana* (Lower Jurassic of China), based on the type species *T. guangdongensis* Zhang in Zhang, Wang, & Zhou, 1977, with special emphasis on ornamentation. According to J. Yin and McRoberts (2006), the diagnostic features of the type species of *Teinonuculana* (elongate form, taxodont dentition, and characteristic rostrum) fit perfectly with those of *Ryderia* Wilton, 1830, and the external ornamentation of *Teinonuculana* is the same as in English specimens of *Ryderia texturata* (Terquem & Piette, 1865) from the Lower Jurassic (see J. Yin & McRoberts, 2006, p. 106). For those reasons, J. Yin and McRoberts (2006) included *Teinonuculana* as a subjective synonym of *Ryderia*, and this is followed here. These authors also warned about the difficulties of studying the ornamentation of these species, since conmarginal or V-shaped ribs are often destroyed or deformed by taphonomic processes.

TOMMASINA Cox, 1964, p. 44

Cox (1964) proposed the name *Tommasina* to replace *Mytiliconcha* Tommasi, 1911, p. 35 (*non Mytiloconcha* Conrad, 1862, p. 290, as *Mytiliconcha* [erroneously] on p. 579). Almost all authors regarded this name change as valid, but Stiller and Chen (2006) defended the validity of *Mytiliconcha* as an unnecessary replacement, following Vokes (1980). In our opinion, the name is indeed unnecessary, because the name proposed by Conrad was *Mytiloconcha*, and therefore it is not a homonym of *Mytiliconcha* (ICZN Art. 56.2, 1999). Moreover, *Mytiliconcha* is not included here, because it is a junior synonym of *Protopis* Kittl, 1904 (see Waller & Stanley, 2005, p. 9). These authors considered *Tommasina* [=*Mytiliconcha*] to be a junior synonym of *Protopis*, because they have the same diagnostic features, although the hinge structure

of *Protopis* is unknown. Hautmann (2008) believed both genera show significant differences on external morphology, but the poor knowledge of *Tommasina* [=*Mytiliconcha*] makes it difficult to establish its taxonomic relationships.

TRICHITES Voltz in Thurmann, 1833, p. 13

No occurrences from the Induan–Sinemurian interval are considered. Cox and others (1969) assigned it to have a Middle Jurassic (Bajocian)–Lower Cretaceous range, but Sepkoski (2002) considered *Trichites* to be from the Norian, based on Hallam (1981). We could not locate the original quotation nor any other reference that substantiates the datum in the references mentioned by Hallam (1981) for the Norian of North America, where the genus was supposedly reported. *Trichites* was frequently mentioned from the Middle and Upper Jurassic (Fürsich, 1980; Damborenea, 1987a; Aberhan, 1994b; Liu, 1995; Fürsich & Hautmann, 2005; Sano & others, 2010) and also from the Lower Jurassic, but the oldest record is from the Pliensbachian of South America (Pérez, 1982; Damborenea, 1987a; Liu, 1999), and the Toarcian of Europe (Gahr, 2002).

VENTALIUM De Gregorio, 1930a, p. 23

Cox and others (1969) considered it to be a doubtful genus, and no later mention of *Ventalium* was found. The range assigned was Jurassic, but it was impossible to corroborate its occurrence within the range of study. Furthermore, Cox and others (1969, p. 371) stated: "Type imperfect, characters not fully known but considered by De Gregorio to be related to *Pecten*." Damborenea (1987b, p. 198) considered it to be a junior synonym of *Eopecten* Douvillé, 1897: "This genus [*Ventalium*] from the lower Jurassic of Sicily was based on incomplete specimens that are in all aspects comparable with fragments of *Eopecten*, especially in ornamentation pattern."

VIETNAMICARDIUM Vu Khuc, 1977a, p. 678

Junior synonym of *Tulongocardium* Chen, Chen, & Zhang in Wen & others, 1976 (Schneider, 1995). Vu Khuc (1977a) proposed *Vietnamicardium* based on Upper Triassic specimens from Vietnam, with *V. vietnamicum* (Vu Khuc in Vu Khuc & others, 1965) as type species. A year earlier, Chen and others (in Wen & others, 1976) had proposed the subgenus *Cardium* (*Tulongocardium*), and *Cardium vietnamicum* Vu Khuc was one of its listed species. For this reason, *Vietnamicardium* was considered a synonym of *C. (Tulongocardium*) (Schneider, 1995, p. 322). On the other hand, Hautmann (2001b, p. 146), although discussing the possible synonymy, considered *Vietnamicardium* to be a distinct genus and included in it some species previously assigned to *Cardium* (*Tulongocardium*). Sepkoski (2002), by mistake, called it *Tulongcardium*. Since the synonymy proposed by Schneider seems sufficiently justified, we follow him, and we also agree to regard *Tulongocardium* at the generic level.

VOKESELLA Chavan, 1952, p. 97

No occurrences from the Induan–Sinemurian interval are considered. Both Cox and others (1969) and Sepkoski (2002) assigned it a Jurassic range without further discussion. Liu (1995) reported it from the Kimmeridgian.

WALLEROBIA Waterhouse, 2008, p. 176

Junior synonym of *Enteropleura* Kittl, 1912 (J. Chen & Stiller, 2010). Waterhouse (2008) proposed *Wallerobia* based on *Enteropleura jenksi* Hopkin & McRoberts, 2005, p. 797 as type (*=Enteropleura* sp. A of Waller in Waller & Stanley, 2005). J. Chen and Stiller (2010, p. 526) rejected it and regarded it as a junior synonym of *Enteropleura* because "the morphological features emphasized by Waterhouse (2008) to distinguish *Wallerobia* from *Enteropleura* are only of species-level significance, and the character of the morphological differences does not justify the formal erection of a new genus-level taxon." We are not taking into account this new genus, because we follow Hopkin and McRoberts (2005), Waller (in Waller & Stanley, 2005), J. Chen and Stiller (2007, 2010), and McRoberts (2010), and include the type species in the genus *Enteropleura* Kittl, 1912.

XINANOPECTEN Feng, Cui, & Liu, 1992, p. 512

Feng, Cui, and Liu (1992) proposed Xinanopecten from the upper Permian of southern China, based on the type species X. orbicularis n. sp. In its diagnosis, they especially referred to the ornamentation, the only feature that differentiates it from the genus Streblochondria Newell, 1938. According to the diagnosis, this ornament consists of "Radial costae wide and clear, concentric lines very weak in the middle and posterior parts of the shell body; but radial costae narrow and weak, concentric lines clear in the anterior part of the shell body. Costae growing bifurcately on the left and right valves." To begin with, the poor preservation of the figured specimens could be the cause of the observed differences between the anterior and posterior parts; moreover, if we compare the diagnosis of Xinanopecten with the diagnosis given by Newell (1937) for Streblochondria, Xinanopecten could be easily included in Streblochondria, since the variation on ornamentation is used by Newell (1937) and by other authors (Ciriacks, 1963; Newell & Boyd, 1985, 1995) as a diagnostic criterion at specific level; and, finally, the diagnosis of Xinanopecten is based on three poorly preserved specimens, which is not enough to characterize the population variation. For all these reasons, we do not include it here.

YOKOYAMAINA Hayami, 1958b, p. 23

Yokoyamaina is considered to be a subgenus of *Integricardium* Rollier, 1912 in 1911–1918 (Hayami, 1975). Hayami (1958b) proposed it from the Lower Jurassic of Japan. Later, Hayami (1975), after reviewing the type species, decided that *Yokoyamaina* fits better as a subgenus of *Integricardium*, and this is followed by most authors (Hallam, 1977; Matsukawa & Nakada, 2003; Kondo & others, 2006). However, some databases and other authors have not discussed the issue, and they considered it to be at the genus level (Vokes, 1980; Schneider, 1995; Sepkoski, 2002; PBDB, 2005).

ZANDAIA Yin & Nie, 1990, p. 106 [255]

Junior synonym of *Praeotapiria* Kurushin in Dagys & Kurushin, 1985 (Z. Fang & others, 2009). Yin and Nie (1990) proposed *Zandaia*, based on *Z. angusta* Yin & Nie, 1990, from the upper Lower Triassic of the Himalayan region (China), but *Z*. Fang and others (2009) placed it in synonymy with *Praeotapiria*. Kurushin (in Dagys & Kurushin, 1985) proposed *Praeotapiria* as a subgenus of *Otapiria* Marwick, 1935. According to Ando (1988), this new subgenus is unnecessary, since morphological differences with other species of *Otapiria* are very subtle. *Zandaia* is better placed in synonymy with *Otapiria*.

ZITTELIHALOBIA Polubotko, 1984, p. 42

Polubotko distinguished Zittelihalobia from Halobia, based on the shape and position of the anterior auricle and ligament (H. J. Campbell, 1994). Zittelihalobia was considered to be a junior synonym of Halobia by McRoberts (1993), while H. J. Campbell (1994) preferred to keep it as a subgenus of Halobia. However, many authors had a different concept of this group and considered it to be a genus (Okuneva, 1987; Kurushin, 1990, 1991; Vu Khuc & Tran Huyen, 1998; Polubotko, Payevskaya, & Repin, 2001; Klets, 2006; Konstantinov, Sobolev, & Yadernkin, 2007). We follow here a conservative approach, considering that this taxon should be maintained at subgenus level at the most.

Other doubtful genera

There are several genera that were included in *Incertae sedis* in Cox and others (1969), and no more information about them was found. These genera are: *Bleta* De Gregorio, 1930b, p. 17; *Carnidia* Bittner, 1901c, p. 6; *Cruciella* Koken, 1913, p. 35; *Ensia* De Gregorio, 1930b, p. 21; *Gerlus* De Gregorio, 1930b, p. 21; *Gingillum* De Gregorio, 1930a, p. 27; *Psammoconcha* Tommasi, 1896, p. 61; *Rebusum* De Gregorio, 1930a, p. 27; and *Taeniodon* Dunker, 1848, p. 179 (see Vokes, 1945, for some information about *Taeniodon*).

Also, there are several genera considered by Sepkoski (2002) in our study interval that we do not consider, because no records in this interval are found: *Cyrtopinna* Mörch, 1853, p. 51; *Atomodesma* von Beyrich, 1865, p. 71; *Cyrtorostra* Branson, 1930, p. 44; *Etheripecten* Waterhouse, 1963, p. 193; *Lopha* Bolten, 1798, p. 168; *Eoschizodus* Cox, 1951, p. 369; *Astartellopsis* Beurlen, 1954, p. 128; *Cowperesia* Mendes, 1952, p. 86; *Mactromya* Agassiz, 1843 in 1840–1845, p. 187; *Terraia* Cox, 1934, p. 269; *Unicardium* d'Orbigny, 1850, p. 218; and *Panopea* Menard, 1807.

ACKNOWLEDGMENTS

This database was initially developed as part of the Ph.D. thesis for S. Ros (Valencia University), and generous help and encouragement from Miquel De Renzi is very warmly acknowledged. We also thank Andrea Allasinaz, Renato Posenato, David Bottjer, Alberto Riccardi, and Antonio Checa, who provided research facilities and friendly hospitality to S.R. during short stays at Università degli Studi di Torino, Università degli Studi di Ferrara, University of Southern California, Museo de Ciencias Naturales de La Plata, and Universidad de Granada.

Many colleagues from around the world contributed to obtain difficult references and kindly shared their knowledge on Mesozoic Bivalvia. Special thanks to Miguel O. Manceñido, Jingeng Sha, Frank Stiller, Christopher A. McRoberts, Stefano Monari, Renato Posenato, Hamish J. Campbell, Zonjie Fang, Héctor A. Leanza, A. G. Konstantinov, Andrzej Kaim, Evgeny Sobolev, Antonio G. Checa, Andrea Allasinaz, Antonio P. Jiménez, Alistair Crame, Gregori López, Jiarun Yin, Jack Grant-Mackie, Thomas Waller, Michael Amler, Noel Morris, Graciela Delvene, Andrea Sterren, Jinhua Chen, and Dario Lazo. The paleogeographic maps were drawn by Scientific Illustrator Oscar Sanisidro (Valencia), with our instructions; the contribution of Pablo Plasencia in this endeavor is also acknowledged. This paper benefited by the thorough reviews of Hamish Campbell and Martin Aberhan, who are gratefully acknowledged.

Research was funded by a doctoral grant from the Ministerio Español de Ciencia y Tecnología and a post-doctoral grant of the MAEC-AECID (Ministerio de Asuntos Exteriores y de Cooperación y Agencia Española de Cooperación Internacional) to S.R. This paper is a contribution to the Ministry of Science and Innovation of Spain Project "CGL2011-24408".

REFERENCES

(*) References not seen by the authors.

- Abbott, R. T. 1954. American Seashells, with illustrations by F. M. Bayer, 1st edit., 2nd printing. Van Nostrand Reinhold Co. Toronto, Inc. xiv + 541 p., 40 pl.
- Abdel-Gawad, G. I. 1986. Maastrichtian non-cephalopod mollusks (Scaphopoda, Gastropoda and Bivalvia) of the middle Vistula Valley, central Poland. Acta Geologica Polonica 36:69–224.
- Aberhan, Martin. 1993. Faunal replacement in the Early Jurassic of northern Chile: Implications for the evolution in Mesozoic benthic shelf ecosystem. Palaeogeography, Palaeoclimatology, Palaeoecology 103:155–177.
- Aberhan, Martin. 1994a. Early Jurassic Bivalvia of Northern Chile. Part I. Subclasses Palaeotaxodonta, Pteriomorphia, and Isofilibranchia. Beringeria 13:114 p.
- Aberhan, Martin. 1994b. Guild-structure and evolution of Mesozoic benthic shelf communities. PALAIOS 9:516–545.
- Aberhan, Martin. 1998a. Early Jurassic Bivalvia of western Canada. Part I. Subclasses Palaeotaxodonta, Pteriomorphia and Isofilibranchia. Beringeria 21:57–150.
- Aberhan, Martin. 1998b. Paleobiogeographic patterns of pectinoid bivalves and the Early Jurassic tectonic evolution of Western Canadian Terranes. PALAIOS 13:129–148.
- Aberhan, Martin. 1999. Terrane history of the Canadian Cordillera: Estimating amounts of latitudinal displacement and rotation of Wrangelliaand Stikinia. Geological Magazine 136(5):481–492.
- Aberhan, Martin. 2001. Bivalve palaeobiogeography and the Hispanic Corridor: time of opening and effectiveness of a proto-Atlantic seaway. Palaeogeography, Palaeoclimatology, Palaeoecology 165:375–394.
- Aberhan, Martin. 2004. Early Jurassic Bivalvia of northern Chile. Part II. Subclass Anomalodesmata. Beringeria 34:117–154.
- Aberhan, Martin, John Alroy, F. T. Fürsich, Wolfgang Kiessling, Matthew Kosnik, J. S. Madin, M. E. Patzkowky, & P. J. Wagner. 2004. Ecological attributes or marine invertebrates. Unpublished [in Paleobiology Database].
- Aberhan, Martin, & F. T. Fürsich. 1997. Diversity analysis of Lower Jurassic bivalves of the Andean Basin and the Pliensbachian-Toarcian mass extinction. Lethaia 29:181–195.
- Aberhan, Martin, & Axel von Hillebrandt. 1996. Taxonomy, ecology, and palaeobiogeography of *Gervilleioperna (Gervilleiognoma) aurita* n. subgen. n. sp. (Bivalvia) from the Middle Jurassic of northern Chile. Paläontologische Zeitschrift 70:79–96.
- Aberhan, Martin, & Axel von Hillebrandt. 1999. The bivalve *Opisoma* in the Lower Jurassic of northern Chile. Profil 16:149–164.
- Aberhan, Martin, M. Hrudka, & T. P. Poulton. 1998. Lower Jurassic and Aalenian bivalve ranges of Western and Northern Canada. *In* P. A. Johnston & J. W. Haggart, eds., Bivalves: An Eon of Evolution - Paleobiological Studies Honoring Norman D. Newell. University of Calgary Press. Calgary. p. 47–50.
- Aberhan, Martin, Wolfang Kiessling, & F. T. Fürsich. 2006. Testing the role of biological interactions in the evolution of mid-Mesozoic marine benthic ecosystems. Paleobiology 32(2):259–277.

- Aberhan, Martin, & Heike Muster. 1997. Palaeobiology of Lower Jurassic Bakevelliid Bivalves from Western Canada. Palaeontology 40(3):799–815.
- Aberhan, Martin, & Joszef Pálfy. 1996. A low oxygen tolerant East Pacific flat clam (*Posidonotis semiplicata*) from the Lower Jurassic of the Canadian Cordillera. Canadian Journal of Earth Sciences 33(7):993–1006.
- Accorsi-Benini, Claudia. 1981. Opisoma Stoliczka, 1871 lamellibranco eterodonte della facies a "Lithiotis" (Giurassico inf., Liassico); revisione morfologica e tassonomica. Bollettino della Societa Paleontologica Italiana 20(2):197–228.
- Accorsi-Benini, Claudia, & Carmela Broglio-Loriga. 1975. Isognomon (Mytiloperna) v. Ihering 1903 e Gervilleioperna Krumbeck 1923 fra i grossi Lamellibranchi della facies a "Lithiotis" del Veneto. Atti Accademia Nazionale dei Lincei, Classe di Scienze Fisiche Matematiche e Naturali, Rendiconti (Ser 8ª) 57(3–4):233–245.
- Accorsi-Benini, Claudia, & Carmela Broglio-Loriga. 1977. *Lithiotis* Gümbel, 1871, e *Cochlearites* Reis, 1903, I, Revisione morfologica e tassonomica. Bolletino della Societa Paleontologica Italiana 16:15–60.
- Acharyya, S. K. 2000. Break up of Australia-India-Madagascar Block, opening of the Indian Ocean and continental accretion in Southeast Asia with special reference to the characteristics of the Peri-Indian collision zones. Gondwana Research 3(4):425–443.
- Adams, Henry, & A. Adams. 1854–1858. The Genera of Recent Mollusca, Arranged According to their Organization. Van Voorst. London. Vol. 2:1–604; collation: i–xl + 605–661; vol. 3: 138 pl.
- Agassiz, J. L. R. 1840. Mémoire sur les moules de mollusques vivans et fossiles. Mémoire de la Societé Scientifique Naturel Neuchâtel 2(4):1–48 p., 9 pl.
- Agassiz, J. L. R. 1840–1845. Études Critiques zur les Mollusques Fossiles (contenant les Trigonies & les Myes du Jura et de la Craie Suisses). Published by the author. Neuchatel. 4 livr., Livr. 1, Memoire sur les Trigonies: 1–58, 11 pl. Livr. 2–4, Monographie des Myes: 1–287, 94 pl.
- Alberti, Friedich von. 1864. Ueberlick über die Trias, mit Berücksichtigung ihres Vorkommens in den Alpen. Verlag der J-G. Cottaschen Buchhandlung. Stuttgart. 353 p., 7 pl.
- Alencaster de Cserna, Gloria. 1961. Paleontología del Triásico superior de Sonora. Parte III. Fauna fósil de la formación Santa Clara (Cárnico) del Estado de Sonora. Paleontología Mexicana 11(III):44 p.
- Allasinaz, Andrea. 1962. Il Trias in Lombardia (Studi geologici e paleontologici). III. Studio paleontologico e biostratigrafico del Retico dei dintorni di Endine (Bergamo). Rivista Italiana di Paleontologia e Stratigrafia LXVIII(3):307–376.
- Allasinaz, Andrea. 1964. Il Trias in Lombardia (studi geologici e paleontologici). V. I fossili carnici del grupo di cima Camino (Brescia (1)). Rivista Italiana di Paleontologia e Stratigrafia LXX(2):185–262.
- Allasinaz, Andrea. 1965. Il Trias in Lombardia (Studi geologici e paleontologici). IX. Note Tassonomiche sulla familia Megalodontidae. Rivista Italiana di Paleontologia e Stratigrafia LXXI(1):111–152.
- Allasinaz, Andrea. 1966. Il Trias in Lombardia (studi geologici e paleontologici). XVIII La fauna a Lamellibranchi dello Julico (Carnico medio). Rivista Italiana di Paleontologia e Stratigrafia LXXII(3):609–752.
- Allasinaz, Andrea. 1972. Revisione dei Pettinidi Triassici. Rivista Italiana di Paleontologia e Stratigrafia LXXVIII(2):189–428.
- Allasinaz, Andrea, Marcel Gutnic, & André Poisson. 1974. La formation de l'Isparta Çay: Calcaires à Halobies, Grès à plantes et Radiolarites d'âge Carnien (?)-Norien (Taurides-Région d'Isparta-Turquie). *In* Die Stratigraphie der alpin-mediterranen Trias. Schrifteneihe Erdwissenschaftlichen Kommissionen Österreichische Akademie der Wissenschaften 2:11–21.
- Allasinaz, Andrea, & Rinaldo Zardini. 1977. Megalodontidae e Dicerocardiidae del Triassico Superiore di Cortina d'Ampezzo. I, Testo, II, Tavole. Rivista Italiana di Paleontologia e Stratigrafia, Memoria XV:215 p.
- Allen, J. A. 2004. The Recent species of the genera *Limatula* and *Limea* (Bivalvia, Limacea) present in the Atlantic, with particular reference to those in deep water. Journal of Natural History 38:2591–2653.

- Allen, J. A., & F. J. Hannah. 1986. A reclassification of the Recent genera of the Subclass Protobranchia (Mollusca: Bivalvia). Journal of Conchology 32:225–249. (*)
- Allen, J. A., & H. L. Sanders. 1969. Nucinella serrei Lamy (Bivalvia; Protobranchia), a monomyarian solemyid and possible living actinodont. Malacologia 7:381–396.
- Alma, F. H. 1925. Eine Fauna des Wettersteinkalkes bei Innsbruck. Annalen des Naturhistorischen Museums, Wien 40:111–129, 1 pl.
- Amano, Kazutaka, R. G. Jenkins, & Yoshinori Hikida. 2007. A new gigantic Nucinella (Bivalvia: Solemyoida) from the Cretaceous cold-seep deposit in Hokkaido, Northern Japan. The Veliger 49(2):84–90.
- Amano, Masahisa. 1957. The Lower Cretaceous fauna from Hagino in southern Shikoku, Japan. Kumamoto Journal of Sciences (series B) 2(2):77–121.
- Amler, M. R. W. 1989. Die Gattung *Parallelodon* Meek and Worthen 1866 (Bivalvia, Arcoida) im mitteleuropäischen Unterkarbon. Geologica et Paleontologica 23:53–69.
- Amler, M. R. W. 1999. Synoptical classification of fossil and Recent Bivalvia. Geologica et Palaeontologica 33:237–248.
- Amler, M. R. W., Rudolf Fischer, & Nicole Rogalla. 2000. Systematische Einheiten. In H. K. Erben, G. Hillmer & H. Ristedt, eds., Muscheln. Haeckel-Bücherei. Enke. Band 5, p. 70–133.
- Amler, M. R. W., & C. F. Winkler Prins. 1999. Lower Carboniferous marine bivalves from the Cantabrian Mountains (Spain). Scripta Geologica 120:1–45.
- Ando, Hisao. 1983. Paleontological significance of late Triassic Bivalve Monotis. Part I: A review. Fossils 33:13–27.
- Ando, Hisao. 1984. Paleontological significance of late Triassic Bivalve Monotis. Part II: In the case of materials from the Southern Kitakami Mountains, Northeast Japan. Fossils 35:1–15.
- Ando, Hisao. 1986. Evolution and paleobiogeography of late Triassic bivalve *Monotis* from Japan. Proceedings of the International Symposium on Shallow Tehthys 2(2):233–246.
- Ando, Hisao. 1987. Paleobiological study of the Late Triassic bivalve *Monotis* from Japan. The University Museum, The University of Tokyo, Bulletin 30:1–110.
- Ando, Hisao. 1988. Mode of occurrence of *Otapiria dubia* (Bivalvia) from the Upper Triassic of West Kyushu, Southwest Japan. *In* J.A. Grant-Mackie, K. Masuda, K. Mori& K. Ogasawara, eds., Professor Tamio Kotaka Commemorative Volume on Molluscan Paleontology. Saito Ho-on Kai Special Publication. Sendai. p. 265–279.
- Ando, Hisao, Masayuki Noda, & Yuichiro Sato. 1987. Discovery and its significance of the Upper Triassic *Monotis* Bed from the Kurosegawa Terrain in East Kyushu. Journal of Geography, Tokyo 96(3):33–37.
- Anelli, L. E., A. C. Rocha-Campos, & M. G. Simões. 2002. Protobranch bivalves from the Piauí Formation (Middle Pennsylvanian), Parnaíba Basin, Brazil. Revista Española de Paleontología 17(2):165–176.
- Anelli, L. E., A. C. Rocha-Campos, & M. G. Simões. 2006. Pennsylvanian pteriomorphian bivalves from the Piauí Formation, Parnaíba basin, Brazil. Journal of Paleontology 80(6):1125–1141.
- Anonymous. 1814. Linnean Society. Annals of Philosophy 4:448-449.
- Anonymous. 1974. A Handbook of the Stratigraphy and Paleontology in Southwest China. Nanjing Institute of Geology and Paleontology, ed. Science Press, Academia Sinica. 454 p.
- Archiac, Etienne J. A. D. 1843. Description géologique du Département de l'Aisne. Mémoires de la Société Géologique de France, 1° sér., 5(1), Mém. 3:129–424.
- Arkell, W. J. 1929–1937. A Monograph of British Corallian Lamellibranchia. Palaeontographical Society, London [Monographs]. The Palaeontographical Society. Part 1, 81 [1929]:1–72, pl. 1–4. Part 2, 82 [1930]:73–104, pl. 5–8. Part 3, 83 [1931]:105–132, pl. 9–12. Part 4, 84 [1932]:133–180, pl. 13–20. Part 5, 85 [1933]:181–228, pl. 21–28. Part 6, 86 [1934]:229–276, pl. 29–36. Part 7, 87 [1934]:277–324,

pl. 37-44. Part 8, 88 [1935]:325-350, i-xvi, pl. 45-49. Part 9, 89 [1936]:351-376, xvii-xxii, pl. 50-56. Part 10, 90 [1937]:377-392.

- Arkhipov, Yu. V., & A. M. Trushchelev. 1980. A New Genus of Pectinids from the Triassic of the Northeast USSR and Central Siberia. In A. S. Dagys, ed., Palaeontology and Stratigraphy of Trias of the Middle Siberia. Academy of Sciences of the USSR, Siberian Branch, Institute of Geology and Geophysics, Transactions. Nauka. Moscow. 448:10–17 + 109–110. In Russian. (*)
- Arp, Gernot. 2007. Sedimentabfolge und Makrobenthos-Assoziationen der Schwarzjura-Transgression bei Sulzkirchen (Hettangium-Pliensbachium, Südliche Frankenalb). Jahrestagung der Paläontologischen Gesellschaft. Freiberg/Sachsen. Poster 77.
- Arp, Gernot, Friedrich Bielert, V. E. Hoffman, & Thomas Löffler. 2005. Palaeoenvironmental significance of lacustrine stromatolites of the Arnstadt Formation ("Steinmergelkeuper," Upper Triassic, N-Germany). Facies 51:419–441.
- Arthaber, G. V. 1908. Die alpine Trias des Mediterran-Gebietes. Ladinische Stufe. (1903–1908), 3rd part. Lethaea Geognostica 1:272–294.
- Asgar-Deen, Michèle, Russell Hall, Jim Craig, & Cynthia Riediger. 2003. New biostratigraphic data from the Lower Jurassic Fernie Formation in the subsurface of west-central Alberta and their stratigraphic implications. Canadian Journal of Earth Sciences 40:45–63.
- Assmann, P. 1916. Die Brachiopoden und Lamellibranchiaten der Oberschlesischen Trias. Jahrbuch der Königlich Preussischen Geologischen Landesanstalt zu Berlin 36(1)(3):586–638, pl. 30–36. Published for 1915.
- Astafieva, M. M. 1986. The Permian bivalved molluscs *Parainoceramus* and *Kolymia*. Paleontological Journal 20(4):23–31.
- Astafieva, M. M. 1991. The bipolar distribution of Inoceramus-like bivalved mollusks in the Permian. Paleontological Journal 25:33–39.
- Astafieva, M. M. 1993. Permskie inoceramopodobnye dvustvorchatye mollyuski Rossii [Permian Inoceramus-like bivalve molluscs of Russia]. Moskva: Izdatel'stvo "Nauka" [Moscow: Publishing House "Science"]. 128 p. In Russian.
- Astafieva, M. M. 1994. A revision of the Chaenocardiidae (=Streblochondriidae) (Bivalvia, Pectinoidea). Paleontological Journal 28(1A):1–22.
- Astafieva, M. M. 1997. New representatives of the Aviculopectinidae (Bivalvia). Paleontological Journal 31(1):22–27.
- Astafieva, M. M. 1998. The development of Permian Marine Bivalves of the Boreal Region (East-European and the Taimyr-Kolyma Subregions). Paleontological Journal 32(2):241–247.
- Astafieva, M. M. 2001. New names for genera of Late Paleozoic bivalves. Paleontological Journal 35(5):557.
- Astafieva-Urbaytis, K. A. 1976. The genus Myonia (Bivalvia) in the Permian of the northeast of the USSR. Paleontological Journal 10(1):23–36.
- Awad, G. H. 1945. On the occurence of Marine Triassic (Muschelkalk) deposits in Sinai. Bulletin de L'Institut d'Egypte 27:397–429.
- Ayyasami, Krishnan. 2006. Role of oysters in biostratigraphy: A case study from the Cretaceous of the Ariyalur area, southern India. Geosciences Journal 10(3):237–247.
- Bachmann, G. H. 1973. Die Karbonatischen Bestandteile des Oberen Muschelkalkes (Mittlere Trias) in südwest-Deutschland und ihre Diagenese. Arbeiten aus dem Institut für Geologie und Paläontologie der Universität Stuttgart, N.F. 68:1–99.
- Bailey, J. B. 1978. Disjunct distributions in time and space; an occurrence of the European Triassic bivalve *Palaeonucula strigilata* Goldfuss in the Mississippian of Arkansas. Abstracts with Programs, Geological Society of America 10(6):246.
- Bambach, R. K. 1977. Species richness in marine benthic habitats through the Phanerozoic. Paleobiology 3:152–157.
- Bambach, R. K. 1983. Ecospace Utilization and Guilds in Marine Communities Through the Phanerozoic. *In* M. J. S. Tevesz & P. L. McCall, eds., Biotic Interactions in Recent and Fossil Benthic Communities. Plenum. New York. p. 719–746.

- Bambach, R. K., A. M. Bush, & D. H. Erwin. 2007. Autecology and the filling of ecospace: key metazoan radiations. Palaeontology 50(1):1–22.
- Barboza-Gudino, J. R., Margario Tristán-González, & J. R. Torres-Hernández. 1990. Tectonic setting of pre-Oxfordian units from central and northeastern Mexico: A review. Geological Society of America, Special Paper 340:197–210.
- Barthel, K. W. 1958. Eine marine Faunula aus der mittleren Trias von Chile. Neues Jahrbuch f
 ür Geologie und Pal
 äontologie Abhandlungen 106(3):352–382.
- Bartholomä, Alfred. 1983. Ein neuer Fund von *Daonella* sp. aus dem Oberen Muschelkalk (Mittl. Trias) von Crailsheim (Südwestdeutschland). Neues Jahrbuch für Geologie und Paläontologie Monatshefte 1:1–4.
- Baumgarte, Diethelm. 1973. Die Gattung Astarte Sowerby im Oberen Muschelkalk am Südostrand des Eggegebirges. Notizblatt des Hessischen Landesamtes-Amt Bodenforschung 101:166–175.
- Baumgarte, Diethelm. 1975. Die Myophorien der germanischen Trias in Niederhessen Vorkommen und Verbreitung. Notizblatt des Hessischen Landesamtes-Amt Bodenforschung 103:53–70.
- Bayle, Emile. 1878. Explication de la carte Géologique Détaillée de la France, Atlas, Première Partie. Fossiles principaux des terrains, vol. 4. Imprimerie Nationale. Paris. 158 pl.
- Bayle, Emile, & H. Coquand. 1851. Mémoire sur les Fossiles recueillis dans le Chili par M. Ignace Domeyko et sur les terrains auxquels ils appartiennent. Mémoires de la Société Géologique de France (series 2) 4(1):1–47, pl. 1–8.
- Beede, J. W. 1907. Invertebrate paleontology of the upper Permian Red Beds of Oklahoma and the Panhandle of Texas. Kansas University Science Bulletin 4:113–1171.
- Beesley, P. L., G. J. B. Ross, & Alice Wells, eds. 1998. Mollusca: The Southern Synthesis. Fauna of Australia, vol. 5, part A. CSIRO Publishing. Melbourne. 563 p.
- Begg, J. G. 1981. The basement geology and palaeontology of the Wairaki Hills, Southland. Unpublished Ph.D. Thesis. University of Otago Library. Dunedin, New Zealand. 400 p. (*)
- Begg, J. G., & H. J. Campbell. 1985. *Etalia*, a new Middle Triassic (Anisian) bivalve from New Zealand, and its relationship with other pteriomorphs. New Zealand Journal of Geology and Geophysics 28(4):725–741.
- Beltan, Laurence, Suzanne Freneix, Philippe Janvier, & Oscar Lopez-Paulsen. 1987. La faune triasique de la formation de Vitiacua dans la región de Villamontes (Département de Chuquisaca, Bolivie). Neues Jahrbuch für Geologie und Paläontologie Monatshefte 2:99–115.
- Bernard, Félix. 1897. Études comparatives sur la coquille des lamellibranches. II. Les genres *Philobrya* et *Hochstetteria*. Journal de Conchyliologie (Paris) 45(1):1–47, pl. 1.
- Beurlen, K. 1954. As faunas de lamellibranquios do sistema Gonduanico no Parana. In F. W. Lange, ed., Paleontologia do Parana. Comissão de Comemorações do Centenario do Parana. Curitiba. p. 107–136, pl. 1–7.
- Beushausen, H. E. L. 1895. Die Lamellibranchiaten des rheinischen Devon mit Ausschluss der Aviculiden. Abhandlungen der Königlich Preussischen Geologischen Landesanstalt (new series) 17:1–514.
- von Beyrich, E. 1845. *Protocardia*, eine neue Gattung fossilier Muscheln. Zeitschrift für Malakozoologie 1845:17–20.
- von Beyrich, E. 1862. Zwei aus dem deutschen Muschelkalk noch nicht bekannte Avicula-artige Muscheln. Zeitschrift der Deutschen Geologischen Gesellschaft 14:9–10.
- von Beyrich, E. 1865. Über eine Kohlenkalk Fauna von Timor. Abhandlungen der Königlichen Akademie Wissenschaften. Berlin. (1864, Phys.):61–98.
- Biakov, A. S. 1992. New species of Permian Inoceramid bivalves from Northeast Russia. Palaontologicheskii Zhurnal 1:27–37 [Paleontological Journal 26(1):32–46].
- Biakov, A. S. 1998. Permian ctenodont nivalves from Northeastern Russia. Paleontological Journal 32(2):129–132.

- Biakov, A. S. 2002. Permian bivalves from the Transbaikal Region. Paleontologicheskii Zhurnal 5:20–28 [Paleontological Journal 36(5):460–468].
- Biakov, A. S. 2006. Permian bivalve mollusks of Northeast Asia. Journal of Asian Earth Sciences 26:235–242.
- Biakov, A. S. 2007. Permian biostratigraphy of the Northern Okhotsk Region (Northeast Asia). Stratigraphy and Geological Correlation 15(2):161–184.
- Bice, D. M., C. R. Newton, S. McCauley, P. W. Reiners, & C. A. McRoberts. 1992. Shocked Quartz at the Triassic-Jurassic Boundary in Italy. Science 255:443–445.
- Bigot, A. P. D. 1895. Contributions à l'étude de la faune jurassique de Normandie. Deuxième mémoire: sur les Opis. Mémoires de la Société Linnéenne de Normandie 18(2):39 p., 2 pl. (*)
- Bistram, A. F. von 1903. Beiträge zur Kenntnis der Fauna des unteren Lias in der Val Solda. Geologisch-paläontologische Studien in den Comasker Alpen, I. Berichte der Naturforschenden Gesellschaft zu Freiburg I, Br. 13:116–214, pl. 1–8.
- Bittner, Alexander 1890. Brachiopoden der alpinen Trias. Abhandlungen der Kaiserlich-Königlichen Geologischen Reichsanstalt, Wien 14:1–325, Pls. 1–41.
- Bittner, Alexander. 1891. Triaspetrefakten von Balia in Kleinasien. Jahrbuch der Kaiserlich-Königlichen. Geologischen Reichsanstalt, Wien, 41:97–116.
- Bittner, Alexander. 1892. Neue Arten aus der Trias von Balia in Kleinasien. Jahrbuch der Kaiserlich-Königlichen Geologischen Reichsanstalt, Wien 42(1):77–89.
- Bittner, Alexander. 1894. Zur Kenntnis der Nuculiden und Arciden der Fauna von Sct. Cassian. Verh. der Kaiserlich-Königlichen Geologischen Reichsanstalt, Wien VII:186–191.
- Bittner, Alexander. 1895. Lamellibranchiaten der alpinen Trias. I Theil: Revision der Lamellibranchiaten von Sct. Cassian. Abhandlungen der Kaiserlich-Königlichen Geologischen Reichsanstalt, Wien XVIII(1):1–235.
- Bittner, Alexander. 1898. Beiträge zur Paläontologie, insbesondere der triadischen Ablagerungen centralasiatischer Hochgebirge. Jahrbuch der Kaiserlich-Königlichen Geologischen Reichsanstalt, Wien 48(4):689–718.
- Bittner, Alexander. 1899. Versteinerungen aus den Trias-Ablagerungen des Süd-Ussuri-Gebietes in der ostsibirischen Küstenprovinz. Mémoires du Comité Geologiqué de St. Petersbourg 7(4):35 p.
- Bittner, Alexander. 1900. Ueber die triadische Lamellibranchiaten-Gattung Mysidioptera Sal. und deren Beziehungen zu palaeozoischen Gattungen. Jahrbuch der Kaiserlich-Königlichen Geologischen Reichsanstalt, Wien 50(1):59–66.
- Bittner, Alexander. 1901a. Ueber *Pseudomonotis Telleri* und verwandte Arten der unteren Trias. Jahrbuch der Kaiserlich-Königlichen Geologischen Reichsanstalt, Wien 50(4):559–592.
- Bittner, Alexander. 1901b. Lamellibranchiaten aus der Trias von Hudiklanec nächst Loitsch in Krain. Jahrbuch der Kaiserlich-Königlichen Geologischen Reichsanstalt, Wien 51(2):225–234.
- Bittner, Alexander. 1901c. Lamellibranchiaten aus der Trias des Bakonyerwaldes. Resultate der wissenschaftlichen Erforschung des Balatonsees, 1. Paläontologie der Umgebung des Balatonsees 2(3):1–106.
- Bittner, Alexander. 1903. Brachiopoden und Lamellibranchiaten aus der Trias von Bosnien, Dalmatien, und Venetien. Jahrbuch der Kaiserlich-Königlichen Geologischen Reichsanstalt, Wien 52(3–4):495–646.
- Bittner, Alexander. 1912. Lamellibranchiaten aus der Trias des Bakonyer Waldes. Resultate der Wissenschaftlichen Erforschung des Balatonsees, II Band: Paläontologie der Umgebung des Balatonsees. Hölzel. Wien. p. 1–107.
- Blainville, Henri-Marie D. de. 1814. Mémoire sur la classification méthodique des animaux mollusques, et établissement d'une nouvelle considération pour y parvenir. Bulletin des Sciences par la Société Philomatique de Paris, Zoologie 1814:175–180.
- Blainville, Henri-Marie D. de. 1820, 1821, 1824, 1829. Mollusques, vol. 32. In F. Cuvier & "plusieurs Professeurs du Jardin du Roi, et des princi-

pales Écoles de Paris," 1816–1830, Dictionnaire des Sciences Naturelles dans lequel on Traite Méthodiquement des Différens Êtres de la Natur, Considérés soit en Eux-mêmes, d'après l'état Actuel de nos Connoissances, soit Relativement a l'Utilité qu'en Peuvent Retirer la Médecine, l'Agriculture, le Commerce et les Arts. Suivi d'une Biographie des Plus Célèbres Naturalistes. F. G. Levrault. Strasbourg and Paris. p. 1–392.

- Bodylevsky, V. I. 1960. Novyi pozdneyurskii predstavitel' avikulopektinid Taimyra [A new Late Jurassic representative of aviculopectinids from Taimyr]. *In* B. P. Markovskii, ed., Novye vidy drevnikh rastenii i bespozvonochnykh SSSR, 2 [New species of prehistoric plants and invertebrates of the USSR; Part 2]. Ministerstva Geologii i Okhran' Nedr, Vsesoiuznyi Nauchno-Issledovatel'skii. Gosgeoltekhizdat, Moskva: 44–45, pl. 7. In Russian.
- Boettger, Oskar. 1880. Die Conchylien der unteren Tertiarschichten. In R. D. M. Verbeek, O. Boettger & K. von Fritsch, eds., Die Tertiarformation von Sumatra und ihre Tierreste. Palaeontographica Supplement 3(8–9):29–120. (*)
- Böhm, Georg. 1884. Beitrag zur Kenntnis der grauen Kalke von Venetien. Deutsche Geologische Gesellschaft, Zeitschrift 36:737–782, 12 pl.
- Böhm, Georg. 1892. Megalodon, Pachyerisma und Diceras. Berichten der Naturforschender Gesellschaft zu Freiburg (new series) 6:33–55.
- Böhm, Georg. 1893. Coelastarte und Heteropis. Berichten der Naturforschender Gesellschaft zu Freiburg (new series) 7(2):169–178, 1 pl.
- Böhm, Joachim. 1903. Description de la faune des couches de Pereiros. Communications du Service Géologique du Portugal 5(1):1–48, pl. 1–3.
- Böhm, Joachim. 1922. Zur systematischen Stellung der Gattung Neithea Drouet. Jahrbuch der Preussischen Geologischen Landesanstalt zu Berlin 40(2):129–147.
- Bolten, Joachim F. 1798. [Published by Röding]. Museum Boltenianum sive Catalogus Cimeliorum e Tribus Regnis Naturae quae olim Collegerat Joa. Fried. Bolten. Pars Secunda. Continens Conchylia sive Testacea Univalvia, Bivalvia, & Multivalvia. Johan Christi Trappii. Hamburgi. 199 p.
- Boit, Bernardo. 1966. Fauna de la facies occidental del Noriano al Oeste de Colquijirca (Cerro de Pasco). Publicaciones del Museo de Historia Natural Javier Prado (Universidad Nacional Mayor de San Marcos, Lima), serie C Geología 11:1–15.
- Boni, Alfredo. 1943. Revisione della fauna Triassica Bresciana: la fauna del Trias Inferiore. Rivista Italiana di Paleontologia e Stratigrafia 49(2):1–41.
- Borghi, Piero. 1937. Su alcune nuove località fossilifere del Trias Superiore Lombardo. Atti Società Italiana di Science Naturali 77:21–35.
- Born, Ignatius [Ignaz] von. 1778. Index Rerum Naturalium Musei Caesarei Vindobonensis. Pars Ima (Testacea). Verzeichniss der Natürlichen Seltenheiten des K. K. Naturalien Kabinets zu Wien, Erster Theil, Schalthiere. Joannus Paulus Kraus. Vienna. xlii + 458+[78] p., 1 pl.
- Bornemann, Johann G. 1854. Ueber die Liasformation in der Umgegend von Göttingen und ihre organischen Einschlüsse. PhD Thesis, Berlin. 77 p., 4 pl.
- Bosc, L. A. G. 1801. Histoire Naturelle de Coquilles, Contenant leur Description, les Moeurs des Animaux qui les Habitent et Leurs Usages, avec Figures Desinées d'après Nature. *In* G. L. Leclerk de Buffon, ed., Histoire naturelle de Buffon, classée par ordres, genres et espèces, d'après le systême de Linné; avec les caractères génériques et la nomenclature linnéenne; Par René-Richard Castel; auteur du poëme des Plantes Nouvelle édition. Déterville, Paris. Mollusca. Vol. 1:343 p.; vol. 2:330 p., pl. 1–15; vol. 3:292 p., pl. 16–28; vol. 4:280 p., pl. 29–36; vol. 5:255 p., pl. 37–44.
- Bouchet, Philippe, & J. P. Rocroi. 2010. Nomenclator of bivalve families with a classification of bivalve families by R. Bieler, J. G. Carter, & E. V. Coan. Malacologia 52(2):1–184.
- Boué, M. A. 1835. Aperçu sur la constitution géologique des Provinces Illyriennes. Mémoir de la Société Géologique de France (Serie 1) 2(1):41–89.
- Boyd, D. W., & N. D. Newell. 1968. Hinge grades in the evolution of crassatellacean bivalves as revealed by Permian genera. American Museum Novitates 2328:1–52.

- Boyd, D. W., & N. D. Newell. 1976. Diagenetic image reversal in a Triassic Pelecypod. Contributions to Geology 14(2):65–68.
- Boyd, D. W., & N. D. Newell. 1979. Permian pelecypods from Tunisia. American Museum Novitates 2686:1–23.
- Boyd, D. W., & N. D. Newell. 1997. A reappraisal of Trigoniacean families (Bivalvia) and a description of two early Triassic species. American Museum Novitates 3216:1–14.
- Boyd, D. W., & N. D. Newell. 1999. *Lyriomyophoria* Kobayashi, 1954, a junior synomym of *Elegantinia* Waagen, 1907. Journal of Paleontology 73(3):547–548.
- Boyd, D. W., D. E. Nice, & N. D. Newell. 1999. Silt injection as a mode of fossilization: a Triassic example. PALAIOS 14(6):545–554.
- Boyer, D. L., D. J. Bottjer, & M. L. Droser. 2004. Ecological Signature of Lower Triassic Shell Beds of the Western United States. PALAIOS 19:372–380.
- Brack, Peter, & Hans Rieber. 1993. Towards a better definition of the Anisian/Ladinian boundary: New biostratigraphic data and correlations of boundary sections from the southern Alps. Eclogae Geologicae Helvetiae 86(2):415–527.
- Bradshaw, M. A. 1999. Lower Devonian bivalves from the Reefton Group, New Zealand. Memoir of the Association of Australasian Palaeontologists 20:1–171.
- Brand, Uwe, & F. M. G. McCarthy. 2005. The Allerø-Younger Dryas-Holocene sequence in the west-central Champlain Sea, eastern Ontario: a record of glacial, oceanographic, and climatic changes. Quaternary Science Reviews 24:1463–1478.
- Branson, C. C. 1930. Paleontology and stratigraphy of the Phosphoria Formation University of Missouri Studies 5(2):1–99.
- Branson, C. C. 1942. Parallelodon, Grammatodon, and Beushausenia (= Cosmetodon, new name). Journal of Paleontology 16(2):247-249.
- Brasil, L. 1895. Observations sur le Bajocien de Normandie. Bulletin Laboratoire Géologique, Faculté des Sciences, Caen 1895:1–21.
- Brinkmann, Roland. 1966. Triásico. In Roland Brinkmann, Compendio de Geología Histórica. Editorial Labor. Barcelona. p. 153–181.
- Brocard, Christian, & Jean Philip. 1989. Les bioconstructions à *Placunopsis ostracina* v. Schlotheim dans le Ladinien supérieur de Provence (SE de la France). Bulletin de la Société Géologique de France V(6):1201–1206.
- Brocchi, Giambattista B. 1814. Conchiologia Fossile Subapennina con Osservazioni Geologiche sugli Apennini e sul Suolo Adiacente. Dalla Stamperia Reale. Milano. vol. 1:1–240, pl. 1–16; vol. 2:241–712.
- Broglio-Loriga, Carmela, Simonetta Cirilli, Vittorio de Zanche, Donato di Bari, Piero Gianolla, G. F. Laghi, William Lowrie, Stefano Manfrin, Adelaide Mastandrea, Paolo Mietto, Giovanni Muttoni, Claudio Neri, Renato Posenato, M. C. Rechichi, Roberto Rettori, & Guido Roghi. 1999. The Prati di Stuores/Stuores Wiesen section (Dolomites, Italy): a candidate global stratotype sction and point for the base of the Carnian stage. Rivista Italiana di Paleontologia e Stratigrafia 105(1):37–78.
- Broglio-Loriga, Carmela, Ferenc Góczán, Janos Haas, Katalin Lenner, Claudio Neri, Anna Oravecz Scheffer, Renato Posenato, János Szabó, & Agnes Tóth Makk. 1990. The Lower Triassic Sequences of The Dolomites (Italy) and Transdanubian Mid-Mountains (Hungary) and their correlation. Memorie de scienze geologiche XLII:41–103.
- Broglio-Loriga, Carmela, Antonio Ietto, & Renato Posenato. 1993. Banchi a *Cornucardia* nell'unità di San Donato (Triassico superiore, Calabria settentrionale). Atti Ticinensi di Scienze della Terra 36:121–129.
- Broglio-Loriga, Carmela, Daniele Masetti, & Claudio Neri. 1982. La Formazione di Werfen (Scitico) delle Dolomiti occidentali: sedimentologia e biostratigrafia. Rivista Italiana di Paleontologia e Stratigrafia 88(4):501–598.
- Broglio-Loriga, Carmela, & Sabina Mirabella. 1986. Il genere *Eumorphotis* Bittner 1901 nella biostratigrafia dello Scitico, formazione de Werfen (Dolomiti). Memorie di Scienze Geologiche XXXVIII:245–281.

- Broglio-Loriga, Carmela, & Claudio Neri. 1976. Aspetti paleobiologici e paleogeografici della facies a "*Lithiotis*" (Giurese inf.). Rivista Italiana di Paleontologia e Stratigrafia 82:651–706.
- Broglio-Loriga, Carmela, Claudio Neri, M. Pasini, & Renato Posenato. 1988. Marine fossil assemblages from upper Permian to Lowermost Triassic in the Western Dolomites (Italy). Memoria della Società Geologica Italiana 34:5–44.
- Broglio-Loriga, Carmela, Claudio Neri, & Renato Posenato. 1980. La "Lingula zone" dello Scitico (Triassico Inferiore). Stratigrafia e paleoecologia. Annali dell'Università di Ferrara. Serie IX: Scienze Geologiche e Paleontologiche VI(6):93–124.
- Broglio-Loriga, Carmela, Claudio Neri, & Renato Posenato. 1986. The early Macrofaunas of the Werfen Formation and the Permian-Triassic Boundary in the dolomites (Southern Alps, Italy). Studi Trentini di Scienze Naturali (Acta Geologica) 62:3–18.
- Broglio-Loriga, Carmela, & Renato Posenato. 1986. Costatoria (Costatoria?) subrotunda (Bittner, 1901). A Smithian (Lower Triassic) marker from Thethys. Rivista Italiana di Paleontologia e Stratigrafia 92(2):189–200.
- Broglio-Loriga, Carmela, & Renato Posenato. 1996. Adaptive strategies of Lower Jurassic and Eocene multivincular bivalves. In Antonietta Cherchi, ed., Autecology of selected fossil organisms: Achievements and problems. Bolletino della Società Paleontologica Italiana, Special Volume 3:45–61.
- Broili, Ferdinand. 1904. Die Fauna Pachycardientuffe der Seiser Alp (mit Ausschluss der Gastropoden und Cephalopoden). Palaeontographica 50(4–5):145–227.
- Brongniart, A. 1822. Description géologique des couches des environs de Paris parmi lesquelles se trouvent les gypses à ossements. *In* G. Cuvier, ed., Recherches sur les Ossemens Fossiles, vol. 2, no. 2. Dufour et E. d'Ocagne. Paris. p. 220–648.
- Bronn, H. G. 1824. System der Urweltlichen Konchylien durch Diagnose, Analyse und Abbildung der Geschlechter Erläutert zum Gebrauche bei Vorlesungen über Petrefaktenkunde und zur Erleichterung des Selbststudiums Derselben. Academischen Buchhandlung von J. C. B. Mohr. Heilderberg. p. 6–56, 7 pl.
- Bronn, H. G. 1828. *Posidonia becheri*, eine neue fossile Muschel der Uebergangs-Periode. Zeitschrift oder Taschenbuch f
 ür Mineralogie von Leonhard 1:262–269.
- Bronn, H. G. 1830a. Ueber die Muschel-Versteinerungen des süd-deutschen Steinsalzgebirges, welche bisher unter dem Namen *Pectinites salinarius* zusammenbegriffen wurden. Jahrbuch für Mineralogie Geognosie; Geologie und Petrefaktenkunde 1:279–285.
- Bronn, H. G. 1830b. Gaea Heidelbergensis, oder Mineralogische Beschreibung der Gegend von Heidelberg. Mit einer petrographischen Karte. Karl Groos. Heidelberg. 237 p.
- Bronn, H. G. 1831. Italiens Tertiär-Gebilde und deren Organische Einschlüsse. Vier Abhandlungen. Mit einer Steindrucktafel. Neue Akademische Buchandlungen von Karl Groos, Heidelberg. p. 1–176, 7 pl.
- Bronn, H. G. 1834–1838. Lethaea Geognostica. Schweizerbart, Stuttgart. Vol. 1:1–544 p. (1834–1837); Vol. 2:545–1350 (1837–1838); Atlas:1–16, pl. 1–47 (1834–1837).
- Bronn, H. G. 1848–1849. Index Palaeontologicus oder Übersicht der bis jetzt bekannten fossilen Organismen, bearbeitet unter Mitwirkung der Herren Prof. H. R. Göppert und Herm. v. Meyer, bearbeitet von Dr. H. G. Bronn. Erste Abtheilung A. Nomenclator Palaeontologicus, in Alphabetischer Ordnung. Schweizerbart. Stuttgart. 1381 p. [1848.] Zweite Abtheilung B. Enumerator Palaeontologicus: Systematische Zusammenstellung und Geologische Entwicklungs-Gesetze der Organischen Reiche. 980 p. [1849].
- Brookfield, M. E., R. J. Twitchett, & Craig Goodings. 2003. Palaeoenvironments of the Permian-Triassic transition sections in Kashmir, India. Palaeogeography, Palaeoclimatology, Palaeoecology 198:353–371.

- Broom, R. 1927. On a new type of mammal-like reptile from the South African Karroo Beds (*Anningia megalops*). Proceedings of the Zoological Society of London 97(1):227–232.
- Brown, Thomas. 1844. Illustrations of the Recent Conchology of Great Britain and Ireland, with the Description and Localities of all the Species, Marine, Land and Fresh water, Drawn and Coloured from Nature, 2nd edit. London & Edinburgh. 144 p., 59 pl.
- Bruguière, Jean G. 1789–1792. Encyclopédie Méthodique, ou par ordre de matières. Histoire Naturelle des Vers. Tome premier. Panckoucke. Paris. Première partie: i–xviii + 1–344; Deuxième partie: p. 345–585.
- Bruguière, Jean G. 1797. Tableau Encyclopédique et Méthodique des Trois Règnes de la Nature Contenant l'Helminthologie, ou les Vers Infusoires, les Vers Intestins, les Vers Mollusques. H. Agasse. Paris. Plates to Vers, Coquilles, Mollusques et Polypiers, pl. 190–286. (*)
- Bubnoff, S. von. 1921. Die ladinische Fauna von Forno (Mezzovalle) bei Predazzo. Verhandlungen naturhistorisch-medizinischen Vereins zu Heidelberg (Neue Folge) Band XIV:257–636.
- von Buch, L. 1838. Über den zoologischen Character der Secondär-Formationen in Süd-Amerika. Monatsberichte der Deutschen Königlichen Akademie der Wissenschaften 1838:54–67.
- Budai, Tamás, György Lelkes, Sándor Kovács, József Pálfy, Olga Piros, Imre Szabó, István Szente, & Attila Vörös. 2003. The Pelsonian Substage on the Balaton Highland (Middle Triassic, Hungary). Geologica Hungarica, Series Palaeontologica Part I (55):1–50.
- Budurov, Kiril, Fransesc Calvet, Antonio Goy, Ana Márquez-Aliaga, Leopoldo Márquez, Ekaterina Trifonova, & Alfredo Arche. 1991. Middle Triassic Stratigraphy and Correlation in part of the Tethys Realm (Bulgaria and Spain). Muschelkalk Internationale Tagung (Muschelkalkmuseum Hagdorn Ingelfingen), Schöntal (Baden-Württemberg, Alemania). p. 16–17.
- Budurov, Kiril, Francesc Calvet, Antonio Goy, Ana Márquez-Aliaga, Leopoldo Márquez, Ekaterina Trifonova, & Alfredo Arche. 1993. Middle Triassic Stratigraphy and Correlation in part of the Tethys Realm (Bulgaria and Spain). *In* H. Hagdorm & A. Seilacher, eds., Muschelkalk Schöntaler Symposium 1991. Sonderbände der Gesellschaft für Naturkunde in Wüttemberg II. Ed. Goldschneck. Korb. p. 157–164.
- Bujnovsky, A., & Maria Kochanová. 1973. Utesy Hlavného Dolomitu Revúckej Doliny a ich megalodontová fauna [Reefs of Hauptdolomite and the fauna of Megalodonts in the Revúcka Dolina Valley]. Geologické práce 60:169–195. In Slovak.
- Bujnovsky, A., Maria Kochanová, & J. Pevny. 1975. Korytnica Limestones A New litho-stratigraphical unit and its fauna. Geologické práce 63:21–53.
- Burckhardt, Carlos, & Salvador Scalia. 1905. Le faune marine du Trias Supérieur de Zacatecas. Boletín del Instituto Geológico de Mexico 21:44 p.
- Buser, Stanko, & Irena Debeljak. 1994. Lower Jurassic beds with bivalves in south Slovenia. Geologija (Ljubljana) 37:23–62.
- Busnardo, Robert. 1970. Faunules du Trias Subbetique (Andalousie). Documents des laboratoires de Géologie de la Faculté des Sciences de Lyon. Notes et Mémoires 37:55–83.
- Busnardo, Robert. 1975. Prébétique et Subbétique de Jaén à Lucena (Andalousie). Intoduction et Trias. Documents des laboratoires de Géologie de la Faculté des Sciences de Lyon 65:1–183.
- Busse, Erwin. 1972. Fazies und Fauna Oberen Muschelkalkes von Willebadessen. Philippia 1/3:110–126.
- Busse, Erwin., & Manfred Horn. 1978. Neue Fossilfunde im Mittleren Buntsandstein NW-Hessens und ihre Bedeutung fuer die Palaeogeographie. Geologisches Jahrbuch Hessen 106:131–142.
- Buvignier, Amand. 1852. Statistique Géologique, Minéralogique, Minérallurgique et Paléontologique du Département de la Meuse. J.-B. Baillière. Paris. 694 p.; Atlas, 52 p., 32 pl.
- Bychkov, Yu M. 1985. [Late Triassic molluscs of the Kenkeren Range (Koryak Plateau)]. In V. P. Pokhialainen, ed., Mesozoic Bivalvia and Cephalopoda from the North Eastern USSR. Collected Scientific Articles. Akademiya Nauk, SSSR, Magadan, 153:5–24. In Russian.

- Bychkov, Yu. M., A. S. Dagys, A. F. Efimova, & I. V. Polubotko. 1976. Atlas Triasovoj fauny i fory Severo-Vostoka SSSR [Atlas of Triassic fauna and flora from North-East USSR]. 'Nedra'. Moskva. 267 p. In Russian.
- Cafiero, B., & Paola Capoa de Bonardi. 1980. Stratigraphy of the pelagic Triassic in the Budva-Kotor area (Crna-Gora, Montenegro, Yugoslavia). Bolletino della Società Paleontologica Italiana 19(2):179–204.
- Cafiero, B., & Paola Capoa de Bonardi. 1982. Biostratigrafia del Trias pelagico della Sicilia. Bolletino della Società Paleontologica Italiana 21(1):35–71.
- Calvet, Francesc, Xavier Berástegui, R. M. Florensa, Mariona Losantos, César Puig, Leopoldo Márquez, Ana Márquez-Aliaga, Nuria Solé de Porta, & Ekaterina Trifonova. 1994. El Triásico (Muschelkalk) de la zona de Gerri de la Sal (Les Nogueres, Pirineos Catalanes). *In* A. Arche, ed., III Coloquio de Estratigrafía y Paleogeografía del Pérmico y Triásico de España, Cuenca (España). Cuenca. p. 27–28.
- Calzada, Sebastián. 1982. Algunos lamelibranquios liásicos de Alfara (Tarragona). Acta Geológica Hispánica 17(1–2):121–127.
- Campbell, H. J. 1982. *Halobia* (Bivalvia, Triassic) and a gastropod from Torlesse Supergroup rocks of Wellington, New Zealand. New Zealand Journal of Geology and Geophysics 25:487–492.
- Campbell, H. J. 1983. *Monotis* (Bivalvia, Triassic) from Torlesse Supergroup rocks, St Arnaud Range, southeast Nelson. New Zealand Journal of Geology and Geophysics 26:103–108.
- Campbell, H. J. 1984. New records and taxa of Permian and Triassic fossils from New Caledonia and New Zeland. Alcheringa 8:151–167.
- Campbell, H. J. 1987. Triassic records of the genus *Lingula* (Brachiopoda: Inarticulata) in New Zealand. New Zealand Journal of Geology and Geophysics 17(1):9–16.
- Campbell, H. J. 1994. The Triassic Bivalves *Daonella* and *Halobia* in New Zealand, New Caledonia, and Svalbard. Institute of Geological and Nuclear Sciences Monograph 4:1–166.
- Campbell, H. J., & J. A. Grant-Mackie. 1995. Jurassic Pholadomyidae (Bivalvia) from New Zealand and New Caledonia. New Zealand Journal of Geology and Geophysics 38:47–59.
- Campbell, H. J., & J. A. Grant-Mackie. 2000. The Marine Triassic of Australasian and its interregional correlation. *In* H. Yin, J. M. Dickins, G. R. Shi, & J. Tong, eds., Permian-Triassic Evolution of Tethys and Western Circum-Pacific. Developments in Palaeontology and Stratigraphy 18:235–255.
- Campbell, J. D., & E. R. Force. 1973. Kaihikuan stage (Middle Triassic): definition and type locality. New Zealand Journal of Geology and Geophysics 16(2):209–219.
- Campbell, K. A., E. A. Nesbitt, & Joanne Bourgeois. 2006. Signatures of storms, oceanic floods and forearc tectonism in marine shelf strata of the Quinault Formation (Pliocen), Washington, United States. Sedimentology 53:945–969.
- Capoa de Bonardi, Paola. 1970. Le Daonelle e la Halobie della serie calcareo-silico-marnosa della Lucania (Appennino Meridionale). Studio Paleontológico e biostratigrafico. Memoria della Società dei Naturalisti in Napoli 78:1–130.
- Carter, J. G. 1990a. Evolutionary significance of shell microstructure in the Palaeotaxodonta, Pteriomorphia and Isofilibranchia (Bivalvia: Mollusca). *In* J. G. Carter, ed., Skeletal Biomineralization: Patterns, Processes and Evolutionary Trends. Van Nostrand Reinhold. New York. p. 136–296.
- Carter, J. G. 1990b. Shell microstructural data for the Bivalvia. *In* J. G. Carter, ed., Skeletal Biomineralization: Patterns, Processes and Evolutionary Trends. Van Nostrand Reinhold. New York. p. 297–411.
- Carter, J. G. 1990c. Skeletal Biomineralization: Patterns, Processes and Evolutionary Trends. II. Atlas and index. *In J. G. Carter*, ed., Van Nostrand Reinhold. New York. 101 p.
- Carter, J. G., E. Barrera, & M. J. S. Tevesz. 1998. Thermal potentiation and mineralogical evolution in the Bivalvia (Mollusca). Journal of Paleontology 72(6):991–1010.

- Carter, J. G., & Michael Hautmann. 2011. Shell microstructure of the basal Pectinid *Pleuronectites laevigatus*: Implications for Pectinoid phylogehy (Mollusca: Bivalvia: Pteriomorphia). Journal of Paleontology 85(3):464–467.
- Carter, J. G., D. R. Lawrence, & Howard Sanders. 1990. Shell Microstructural data for the Bivalvia. Part II. Orders Nuculoida and Solemyoida. *In J. G. Carter*, ed., Skeletal Biomineralization: Patterns, Processes and Evolutionary Trends. Van Nostrand Reinhold. New York. Vol. 1, p. 303–319.
- Carter, J. G., & G. D. Stanley, Jr. 2004. Late Triassic gastrochaenid and lithophaginid borings (Mollusca: Bivalvia) from Nevada (United States) and Austria. Journal of Paleontology 78(1):230–234.
- Carter, J. G., & M. J. S. Tevesz. 1978. Shell microstructure of a middle Devonian (Hamilton Group) Bivalve fauna from Central New York. Journal of Paleontology 52(4):859–880.
- Castell, C. P., & L. R. Cox. 1975. British Mesozoic fossils. British Museum (Natural History) 703:207 p.
- Caswell, B. A., A. L. Coe, & A. S. Cohen. 2009. New range data for marine invertebrate species across the early Toarcian (Early Jurassic) mass extinction. Journal of the Geological Society, London 166:859–872.
- Cavanaugh, C. M. 1983. Symbiotic chemoautotrophic bacteria in marine invertebrates from sulphide-rich habitats. Nature 302:58–61.
- Chavan, André. 1938. Essai critique de classification des lucines. Journal de Conchyliologie (Paris) 82(3):215–243.
- Chavan, André. 1945. Les lamellibranches hétérodontes des sables astartiens de Cordebugle (Calvados). I. Astartidae, Cyprinidae, Isocardiidae. Journal de Conchyliologie (Paris) 8(2):41–86, pl. 1.
- Chavan, André. 1947. L'évolution des faunes marines de Mollusques dans le Nord-Ouest de l'Europe, de la fin du Crétacé a celle de l'Éocene. Bulletin de la Société Géologique de France (series 5) 16(4–6):193–212. Published for 1946.
- Chavan, André. 1951. Dénominations supraspécifiques de Mollusques modifiées ou nouvelles. Société Géologique de France, Comptes Rendus (1951) 11–12:210–212.
- Chavan, André. 1952. Les pélécypodes des Sables Astartiens de Cordebugle (Calvados). Schweizerische Paläontologische Abhandlungen [Mémoires Suisses de Paléontologie] 69:1–132. (*)
- Chavan, André. 1954. Les *Pleurophorus* et genres voisins. Cahiers Géologiques de Seyssel (Ain) 22:1–200.
- Chavan, André. 1959. Essai critique de classification des Mactromyidae. Cahiers Géologiques 53:505–510.
- Checa, A. G., & A. P. Jiménez-Jiménez. 2003a. Evolutionary morpholgy of oblique ribs of Bivalves. Palaeontology 46(4):709–724.
- Checa, A. G., & A. P. Jiménez-Jiménez. 2003b. Rib fabrication in Ostreoidea and Plicatuloidea (Bivalvia, Pteriomorphia) and its evolutionary significance. Zoomorphology 122:145–159.
- Checa, A. G., A. P. Jiménez-Jiménez, Ana Márquez-Aliaga, & Hans Hagdorn. 2003. Revisión de *Enantiostreon* e implicaciones sobre el origen de Plicatulidae y Dimyidae (Plicatuloidea, Pteriomorphia, Bivalvia). Libro de Resúmenes XIX Jornadas de la Sociedad Española de Paleontología, Morella. p. 51–52.
- Checa, A. G., A. P. Jiménez-Jiménez, Ana Márquez-Aliaga, & Hans Hagdorn. 2006. Further comments on the origin of oysters. Palaeogeography, Palaeoclimatology, Palaeoecology 240:672–674.
- Chemnitz, J. H. 1784. Neues systematisches Conchylien-Cabinet, vol. 7. Bauer & Raspe. Nürnberg. 356 p., pl. 37–69.
- Chen Chu-zhen. 1976. In Gu Zhi-wei, Huang Bao-yu, Chen Chu-zhen, Wen Shi-xuan, Ma Qi-hong, Lan Xiu, Xu Jun-tao, Liu Lu, Wang Shumei, Wang De-you, Qiu Ran-zhong, Huang Zao-qi, Zhang Zao-ming, Chen Jin-hua, & We Pei-li, eds., The Lamellibranch Fossils of China. Nanjing Institute of Geology and Palaeontology, Academia Sinica. Science Press. Beijing. 522 p., 150 pl.

- Chen Chu-zhen. 1981. Lamellibranchiata. In Zhao Jin-ke, Sheng Jin-zhang, Yao Zhao-qi, Liang Xi-luo, Chen Chu-zhen, Rui Lin, & Liao Zhuoting, eds., The Changhsingian and Permian-Triassic boundary of south China. Bulletin Nanjing Institute, Geology and Palaeontology, Academia Sinica 1981(2):81–83.
- Chen Chu-zhen. 1982. Some late Paleozoic and Triassic bivalves from Xizang. In Paleontology of Xizang. *In* Anonymous, ed., The series of the scientific expedition to the Qinghai-Xizang Plateau Vol. IV. Science Press. Beijing, p. 211–224. In Chinese.
- Chen Chun-zhen, Qihong Ma, & Zuoming Zhang. 1974. Triassic Lamellibranchiata. *In* Nanjing Institute of Geology and Palaeontology, Academia Sinica, ed., Handbook of stratigraphy and palaeontology of the south-western region. Science Press. Beijing. p. 326–343, pl. 173–179. In Chinese.
- Chen Chun-zhen, & Yu Jing-shan. 1976. Monotidae. *In* Nanjing Institute of Geology and Palaeontology, Academia Sinica, eds., Palaeontological Atlas of Lamellibranchia from China. Publishing House Science. Beijing. p. 225–227. In Chinese.
- Chen Jin-hua. 1982a. A study on the bivalve fauna of the Upper Triassic in Guangdong, Hunan and Jiangxi provinces, South China. Bulletin of Nanjing Institute of Geology and Paleontology, Academia Sinica 4:279–306. In Chinese.
- Chen Jin-hua. 1982b. Liassic bivalve fossils from Mt. Jinji of Guangdong. Acta Palaeontologica Sinica 21(4):404–416. In Chinese with English abstract.
- Chen Jin-hua. 1982c. Assemblages of *Daonella* (Bivalvia) in south Guizhou Province. Scientia Geologica Sinica 4:235–238. In Chinese with English abstract.
- Chen Jin-hua. 1985. On the new horizons of *Burmesia* (Bivalvia). Kexue Tongbao 30(12):1660–1663.
- Chen Jin-hua. 1988. Early Jurassic marine bivalves from Guangdong-Nanling district, southern China. Bulletin of Nanjing Institute of Geology and Palaeontology 12:23–94. In Chinese with English abstract. (*)
- Chen Jin-hua. 1999. Discussion on the "Aucellina" from the Longzhaogou Group and on the significance of pseudoctenolium in classification. Acta Palaeontologica Sinica 38(4):454–468. In Chinese with English abstract.
- Chen Jin-hua. 2003. Macroevolution of Bivalvia after the end-Permian mass extinction in South China. *In* Rong J. Y. & Fang Z. J., eds., Faunal Recovery After the Palaeozoic Three Mass Extinctions. Science Press. Beijing, p. 647–700. In Chinese with English abstract.
- Chen Jin-hua, & Chen Chu-zhen. 1980. *Jianchuania*, a new Genus of Bivalve from the Upper Triassic in Yunnan. Acta Paleontologica Sinica 19(1):57–60. In Chinese with English abstract.
- Chen Jin-hua, & Toshifumi Komatsu. 2002. So-called middle Triassic "*Claraia*" (Bivalvia) from Guangxi, South China. Acta Palaeontologica Sinica 41(3):434–447. In Chinese with English abstract.
- Chen Jin-hua, & Liu Lu. 1981. Discovery of Early Jurassic *Cardinia* beds in Rucheng, Hunnan. Acta Palaeontologica Sinica 20(4):376–382. In Chinese with English abstract.
- Chen Jin-hua, Liu Lu, & Lan Xiu. 1983. Veteranellinae, a new subfamily of Nuculanidae (Bivalvia), with the classificatory significance of ornamentations in Palaeotaxodonta. Acta Palaeontologica Sinica 22(6):616–627. In Chinese with English abstract.
- Chen Jin-hua, & Frank Stiller. 2007. The halobiid bivalve genus *Enteropleura* and a new species from the Middle Anisian of Guangxi, southern China. Acta Palaeontologica Polonica 52(1):53–61.
- Chen Jin-hua, & Frank Stiller. 2008. Two new late Triassic nuculanoidean bivalve genera from north-west Yunnan, China. Neues Jahrbuch für Geologie und Paläontologie Abhandlungen 249(3):357–370.
- Chen Jin-hua, & Frank Stiller. 2010. An early *Daonella* from the Middle Anisian of Guangxi, southwestern China, and its phylogenetical significance. Swiss Journal of Geosciences 103:523–533.

- Chen Jin-hua, Frank Stiller, & Toshifumi Komatsu. 2006. Protostrea from the Middle Triassic of southern China, the earliest dimyoid bivalve? Neues Jahrbuch für Geologie und Paläontologie Monatshefte 3:148–164.
- Chen Jin-hua, Wang Yu-mao, Wu Quin-rong, Li Lin-pao, Zhao Rhongde, & Chen Hai-chou. 1992. A study of bivalve zonal succession from upper part of Middle Triassic in northwest Guangxi, S. China. Acta Palaeontologica Sinica 31(4):403–419. In Chinese with English abstract.
- Chen Jin-hua, & Yang Sheng-qiu. 1983. *Eleganuculana*, new genus, and some other bivalves of the Upper Triassic of Kangmar, Xizang. Acta Palaeontologica Sinica 22(3):355–358. In Chinese with English abstract.
- Chen Zhong-qiang, Kunio Kaiho, A. D. George, & Tong Jinnan. 2006. Survival brachiopod faunas of the end-Permian mass extinction from the southern Alps (Italy) and South China. Geological Magazine 143(3):301–327.
- Chen Zhong-qiang, & K. J. McNamara. 2006. End-Permian extinction and subsequent recovery of the Ophiuroidea (Echinodermata). Palaeogeography, Palaeoclimatology, Palaeoecology 236:321–344.
- Chen Zhong-qiang, Shi G. R., & Kunio Kaiho. 2004. New ophiuroids form the Permian/Triassic boundary beds of South China. Palaeontology 47(5):1301–1312.
- Chen Zhong-qiang, Shi G. R., Yang Fengqing, Gao Yongquan, Tong Jinnan, & Peng Yuanqiao. 2006. An ecologically mixed brachiopod fauna from Changhsingian deep-water basin of South China: consequence of end-Permian global warming. Lethaia 39:79–90.
- Chernyshev, B. I. 1939. Tip Myagkotelyye-Molluska, Klass Plastinchatozhhabernyye-Lamellibranchiata. *In* I. I. Gorsky, ed., Atlas rukovodyashchikh form iskopayemykh faun SSSR, vol. 5, Sredneiy i verkhniy otdely kamennougol'noy sistemy. Tsentral'nyy Nauchno-Issledovatel'skiy Geologo-Rasveddochnyy Institut. Leningrad & Moscow. p. 113–126, pl. 28–31.
- Chernyshev, B. I. 1943. Do systematyky verkhn'o-paleozoys'kykh Taxodonta. [Contributions to the systematics of the Upper Paleozoic Taxodonta] Akademii Nauk Ukrayins'koyi RSR, Instytut Geologichnikh Nauk. Kiev. 40 p. (*)
- Chernyshev, B. I. 1951. Semeistvo Ledidae iz kamennougolnykh otlozhenii SSSR [= The family Ledidae from the Coal Measures of the U.S.S.R]. Trudy Institut Geologicheskikh Nauk, Seriya Stratigrafii i Paleontologii, Akademiya Nauk, Ukraine S.S.R., Kiev 2:1–40.
- Chiesa, Cesare. 1949. Contributo alla conoscenza del Retico nella Lombardia occidentale. Rivista Italiana di Paleontologia e Stratigrafia 55(1):18–33.
- Chiplonkar, G. W., & R. M. Badve. 1977. Development of multiple resilifers in *Lopha (Actinostreon) diluviana* (Linne) from the Upper Cretaceous of South India. Current Science 46:242.
- Chong, Guillermo, & Axel von Hillebrandt. 1985. El Triásico Preandino de Chile entre los 23°30' y 26°00' de latitud Sur. IV Congreso Geológico Chileno I:162–210.
- Chronic, H. 1952. Molluscan fauna from the Permian Kaibab formation, Walnut Canyon, Arizona. Bulletin of the Geological Society of America 63(2):95–165, pl. 1–11.
- Chumakov, N. M., & M. A. Zharkov. 2003. Climate during the Permian–Triassic Biosphere Reorganizations. Article 2. Climate of the Late Permian and Early Triassic: General Inferences. Stratigraphy and Geological Correlation 11(4):361–375.
- Ciriacks, K. W. 1963. Permian and Eotriassic Bivalves of the Middle Rockies. Bulletin of the American Museum of Natural History 125(1):1–100.
- Cisterna, G. A., & A. F. Sterren. 2003. Variaciones composicionales de la "Fauna de Levipustula" en la Precordillera Argentina. Ameghiniana 40(4), Suplemento Resúmenes, 11–R.
- Clapham, M. E., & D. J. Bottjer. 2007. Permian marine paleoecology and its implications for large-scale decoupling of brachiopod and bivalve abundance and diversity during the Lopingian (Late Permian). Palaeogeography, Palaeoeclimatology, Palaeoecology 249:283–301.

- Coan, Eugene V., Valentich Scott, Paul, & Bernard, Frank R. 2000. Bivalve Shells of Western North America. Marine Bivalve Mollusks from Arctic Alaska to Baja California. Santa Barbara Museum of Natural History Monographs 2:1–764.
- Cockerell, T. D. A. 1903. The name Solenopsis. Nature 67:559.
- Conrad, T. A. 1842. Observations of the Silurian and Devonian systems of the United States, with descriptions of new species of organic remains. Journal of the Academy of Natural Sciences of Philadelphia 8(2):228–280.
- Conrad, T. A. 1858. Observations of a group of Cretaceous fossil shells, found in Tippah County, Mississippi, with descriptions of fifty-six new species. Journal of the Academy of Natural Sciences of Philadelphia (series 2) 3:323–336.
- Conrad, T. A. 1862. Descriptions of new genera, subgenera, and species of Tertiary and recent shells. Proceedings of the Academy of Natural Science of Philadelphia 14:284–291.
- Conrad, T. A. 1867. Descriptions of new genera and species of fossil shells. American Journal of Conchology 2:8–16.
- Conti, M. A., & Stefano Monari. 1991. Bivalve and gastropod fauna from the Liassic Ammonitico Rosso facies in the Bilecik area (Western Pontides, Turkey). Geologica Romana 27:245–301.
- Conti, M. A., & Stefano Monari. 1992. Thin-shelled bivalves from the Jurassic Rosso Ammonitico and Calcari a *Posidonia* Formations of the Umbrian-Marchean Apennine (Central Italy). Paleopelagos 2:193–213.
- Conti, Sergio. 1954. Stratigrafia e Paleontologia della Val Solda (Lago di Lugano). Memorie descrittive della Carta Geologica d'Italia 30:241 p.
- Cooper, R. A., ed. 2004. The Zew Zealand Geological Timescale. Institute of Geological & Nuclear Sciences Monograph 22. 284 p.
- Cope, J. C. W. 1995. The early evolution of the Bivalvia. *In* J. D. Taylor, ed., Origin and evolutionary radiation of the Mollusca. Oxford University Press. Oxford. xiv + 392 p.
- Cope, J. C. W. 1997. The early phylogeny of the class Bivalvia. Palaeontology 40(3):713–746.
- Corazzari, Diana, & A. M. Lucchi-Garavello. 1980. Nuove specie di Bivalvi della formazione di San Cassiano (conca di Cortina d'Ampezzo e Dintorni). Annali della Universita di Ferrara, Sezione IX: Scienze Geologiche e Paleontologiche 7(3):37–57.
- Cossmann, Maurice. 1897. Paléoconchologie et ouvrages généraux. Revue Critique de Paléozoologie. Paris. 1:46–68.
- Cossmann, Maurice. 1904. Note sur l'Infralias de La Vendée et des Deux-Sèvres. II. Pélécypodes. Bulletin de la Société Géologique de France (série 4) 3(5):497–537, pl. 16–18.
- Cossmann, Maurice. 1906. Description de quelques pélécypodes jurassiques de France. Premier article. Comptes Rendus, Association Française pour l'avancement des Sciences 34(2):284–297.
- Cossmann, Maurice. 1912. Quelques pelécypodes jurassiques recueillis en France. Comptes Rendus, Association Française pour l'avancement des Sciences 1912:11 p.
- Cossmann, Maurice. 1920. Rectifications de nomenclature. Revue Critique de Paléozoologie 24(2):81–83.
- Cossmann, Maurice. 1921. Description de quelques pélécypodes jurassiques recueillis en France. 2 Série, 1 Art. Comptes Rendus de l'Association Française pour l'Avancement des Sciences (Congrès de Strasbourg, 1920):1–29. (*)
- Cossmann, Maurice, & Edmond Pellat. 1907. Le Barrémiem supérieur à facies Urgonien de Brouzet-lez-Alais (Gard). Mémoires de la Société Géologique de France 15(37:5–42.
- Cossmann, Maurice, & P. Thièry. 1907. Note sur le Callovien de la Haute-Marne. Bulletin, Société d'Agriculture, Lettres, Sciences et Arts de la Haute-Saône (1907)2:69–147, pl. 1–3.
- Counillon, Henri. 1908. Sur le gisement liasique du Huu-Nien Province de Quang-Nam (Annam). Bulletin de la Société Géologique de France (4 Série) 8:524–532.

- Covacevich, Vladimir, & Fernando Escobar. 1979. La presencia del género Otapiria Marwick, 1935 (Mollusca: Bivalvia) en Chile y su distribución en el ámbito Circumpacífico. Actas del Segundo Congreso Geológico Chileno 3:H165–H187.
- Covacevich, Vladimir, Ernesto Pérez, & Fernando Escobar. 1991. Presencia del género *Kalentera* Marwick, 1953 (Mollusca Bivalvia) en el Sinemuriano al sur de Taltal, Chile. Actas del 6º Congreso Geológico Chileno 1991. Resúmenes expandidos. Servicio Nacional de Geología y Mineria. Santiago. 1:68–71.
- Cowper-Reed, F. R. 1927. Palaezoic and Mesozoic fossils from Yunnan. Memoirs of the Geological Survey of India. Palaeontologia Indica (new series) 10(1):1–291.
- Cox, Leslie R. 1924. A Triassic Fauna from the Jordan Valley. The Annals and Magazine of Natural History (series 9) 14(79):52–96, pl. 1-2.
- Cox, Leslie R. 1925. The Fauna of the Basal Shell-Bed of the Portland Stone, Isle of Portland. Proceedings of the Dorset Natural History Field Club 46:113–172.
- Cox, L. R. 1926. Notes on the Lamellibranchia of the Black Marl of Black Ven and Stonebarrow. Quarterly Journal of the Geological Society of London 82:180–184.
- Cox, Leslie R. 1929. A synopsis of the Lamellibranchia and Gastropoda of Portland beds of England, Part 1. Lamellibranchia. Proceedings of the Dorset Natural History and Archaeology Society 50:131–202, pl. 1–6.
- Cox, Leslie R. 1934. Lamelibranquios de los estratos gondwánicos del Uruguay. Boletín del Instituto de Geología y Perforaciones del Uruguay 21:1–13.
- Cox, Leslie R. 1936. The Gastropoda and Lamellibranchia of the Green Ammonite beds of Dorset. Quarterly Journal of the Geological Society, London 92:456–471.
- Cox, Leslie R. 1937a. Notes on Jurassic Lamellibranchia. I. On the occurrence of the genus *Palaeoneilo* in the Jurassic of Great Britain. Proceedings of the Malacological Society of London 22:190–193.
- Cox, L. R. 1937b. Notes on Jurassic Lamellibranchia. II. On *Indogramma-todon*, a new subgenus from the Jurassic of the Indo-African Province. Proceedings of the Malacological Society of London 22:194–198.
- Cox, Leslie R. 1937c. Notes on Jurassic Lamellibranchia. V. On a new subgenus of *Mytilus* and a new *Mytilus*-like genus. Proceedings of the Malacological Society of London 22:339–348.
- Cox, L. R. 1940. The Jurassic Lamellibranch Fauna of Kachh (Cutch). Memoirs of the Geological Survey of India, Palaeontologia Indica (series 9) 3:1–157.
- Cox, Leslie R. 1943. The English Upper Lias and Inferior Oolite species of *Lima*. Malacological Society of London, Proceedings 25(5–6):151–187, pl. 6–29.
- Cox, Leslie R. 1946. *Tutcheria* and *Pseudopis*, new lamellibranch genera from the Lias. Proceedings of the Malacological Society of London 27:34–48.
- Cox, Leslie R. 1949. Moluscos del Triásico Superior del Perú. Boletín, Instituto Geológico de Perú 12:1–45.
- Cox, Leslie R. 1951. The lamellibranch genus *Schizodus* and other Palaeozoic Myophoriidae. Geological Magazine 88(5):362–371.
- Cox, Leslie R. 1952. Notes on the Trigoniidae, with outlines of a classification of the family. Proceedings of the Malacological Society of London 29:45–70.
- Cox, Leslie R. 1953. Lower Cretaceous Gastropoda, Lamellibranchia and Annelida from Alexander Island (Falkland Islands Dependencies). Scientific Reports of the British Antarctic Survey, Falkland Islands Dependencies Survey 4:1–14, 2 pl.
- Cox, Leslie R. 1954. Taxonomic notes on Isognomonidae and Bakevelliidae. Proceedings of the Malacological Society of London 31(2):46–49.
- Cox, Leslie R. 1958. *Anningella*, nom. nov. for *Anningia* Cox non Broom. Proceedings of the Geological Society of London 1557:44.
- Cox, Leslie R. 1962. New genera and subgenera of Mesozoic bivalvia. Palaeontology 4(4):592–598. Published for 1961.

- Cox, Leslie R. 1963. The Rhaetic-Hettangian bivalve genus *Pteromya* Moore. Palaeontology 6(3):582–95.
- Cox, Leslie R. 1964. Notes concerning the taxonomy and nomenclature of fossil Bivalvia (mainly Mesozoic). Proceedings of the Malacological Society of London 36:39–48.
- Cox, Leslie R., & W. J. Arkell. 1948–1950. A Survey of the Mollusca of the British Great Oolite Series. Primarily a nomenclatorial revision of the monographs by Morris and Lycett (1851–55), Lycett (1863) and Blake (1905–1907). Palaeontographical Society Monographs, London 102: i–xiii + 1–48 (1948); 103: xiv–xxiv + 49–105 (1950).
- Cox, Leslie R., N. D. Newell, C. C. Branson, R. Casey, André Chavan, A. H. Coogan, Colette Dechaseaux, C. A. Fleming, F. Haas, L. G. Hertlein, A. M. Keen, A. LaRocque, A. L. McAlester, B. F. Perkins, H. S. Puri, L. A. Smith, T. Soot-Ryen, H. B. Stenzel, R. D. Turner, & J. Weir. 1969. Systematic Descriptions. *In* R. C. Moore, ed., Treatise on Invertebrate Paleontology, Part N, Mollusca 6, Bivalvia, vol. 1 & 2. The Geological Society of America & the University of Kansas. Boulder, Colorado & Lawrence, Kansas. ii + p. 1–952.
- Crame, J. A. 1982. Late Jurassic inoceramid bivalves from the Antarctic peninsula and their stratigraphic use. Palaeontology 25(3):555–603.
- Crame, J. A. 1995. Occurrence of the bivalve genus *Manticula* in the Early Cretaceous of Antarctica. Palaeontology 38(2):299–312.
- Crame, J. A. 1996. A new oxytomid bivalve from the Upper Jurassic Lower Cretaceous of Antarctica. Palaeontology 39(3):615–628.
- Crampton, J. S. 1988. Comparative taxonomy of the bivalve families Isognomonidae, Inoceramidae and Retroceramidae. Palaeontology 31(4):965–996.
- Credner, H. 1851. Über die Gervillien der Trias-Formation in Thüringen. Neues Jahrbuch für Mineralogie 1851:641–657.
- Crickmay, Colin H. 1930a. The Jurassic rocks of Ashcroft, British Columbia. University of California, Publications in Geological Sciences 19(2):23–74, pl. 2–7.
- Crickmay, Colin H. 1930b. Fossils from Harrison Lake area, British Columbia. Bulletin, National Museum Canada 63:33-66.
- Crickmay, C. H. 1932. Contributions toward a monograph of the Trigoniidae, I. American Journal of Science 24:443–464.
- Crosby, C. R., & S. C. Bishop. 1933. American Spiders: Erigoneae, males with cephalic pits. Annals of the Entomological Society of America 26:105–182.
- Császár, Géza, & Dragica Turnšek. 1996. Vestiges of atoll-like formations in the Lower Cretaceous of the Mecsek Mountains, Hungary. Cretaceous Research 17:419–442.
- Dagys, A. S. 1974. Triasovye brakhiopodi (Morfologiia, sistema, filogeniia, stratigraficheskoe znachenie i biogeografiia) [Triassic brachiopods (Morphology, classification, phylogeny, significance and biogeography)]. Novosibirsk: Izdatel'stvo "Nauka." Sibirskoe Otdelenie. 387 p. In Russian.
- Dagys, A. S., & A. A. Dagys. 1994. Global correlation of the terminal Triassic. Mémoires de Géologie (Lausanne) 22:25–34.
- Dagys, A. S., A. Yu. Egorov, A. M. Kazakov, & N. I. Kurushin. 1989. Stratigraphiya tiasovykh otlozheniy Yogo-Vostochnogo Taymyra [Stratigraphy of Triassic sediments of south-eastern Taymyr]. *In* Verkhniy Paleozoy I trias Sibiri [Upper Paleozoic and Triassic of Siberia]. Trudy Instituta Geologii i Geofiziki [Proceedings of the Institute of Geology and Geophysics] 732:71–91. In Russian. (*)
- Dagys A. S., & N. I. Kurushin. 1985. The Triassic Brachiopods and Bivalves in the North of the Central Siberia. Transactions 633:160 p. In Russian.
- Dall, William H. 1895. Contributions to the Tertiary fauna of Florida, with especial reference to the Miocene silex-beds of Tampa and the Pliocene beds of the Caloosahatchie River, Part III. A new classification of the Pelecypoda. Wagner Free Institute of Science of Philadelphia, Transactions 3(3):483–570.
- Dall, William H. 1896–1900. Pelecypoda. In K. A. von Zittel, ed., Textbook of Palaeontology, vol. 1. McMillan & Co. New York & London. p. 346–429.

- Dall, William H. 1898. Contributions to the Tertiary fauna of Florida with especial reference to the Miocene Silex beds of Tampa and the Pliocene beds of the Caloosahatchie River, including in many cases a complete revision of the generic groups treated and of their American Tertiary Species. Part IV. 1. Prionodesmacea: *Nucula* to *Julia*. 2. Teleodesmacea: *Teredo* to *Ervilia*. Wagner Free Institute of Science, Philadelphia, Transactions 3(4):i–viii + p. 571–947, pl. 23–35.
- Dall, William H. 1901. Synopsis of the Lucinacea and of the American species. U. S. National Museum, Proceedings 23(1237):779–833, pl. 39–42.
- Dall, William H. 1908a. A revision of the Solenomyadidae. The Nautilus 22(1):1–2.
- Dall, William H. 1908b. Reports on the dredging operations off the west coast of Central America to the Galapagos, to the West coast of Mexico, and in the Gulf of California, in charge of Alexander Agassiz, carried on by the U. S. Fish Commission steamer "Albatross," during 1891, Lieut. Commander Z. L. Tanner, U.S.N., commanding. XXXVII. Reports on the scientific results of the expedition to the eastern tropical Pacific, in charge of Alexander Agassiz, by the U. S. Fish Commission steamer "Albatross," from October, 1904, to March, 1905, Lieut. commander L. M. Garrett, U.S.N., commanding. XIV. The Mollusca and the Brachiopoda. Museum of Comparative Zoology, Harvard University, Bulletin 43(6):205–487, pl. 1–22.
- Dall, William H., Bartsch, Paul, & Rehder, HaraldA. 1938. A manual of the recent and fossil marine pelecypod mollusks of the Hawaiian Islands. Bernice P. Bishop Museum Bulletin 153:3–233.
- Damborenea, S. E. 1986. El género *Posidonotis* Losacco (Bivalvia, Jurásico Inferior): su distribución estratigráfica y paleogeográfica. Actas del IV Congreso Argentino de paleontología y bioestratigrafía (Mendoza, 1986) 4:45–51.
- Damborenea, S. E. 1987a. Early Jurassic Bivalvia of Argentina. Part 1: Stratigraphical introduction and superfamilies Nuculanacea, Arcacea, Mytilacea and Pinnacea. Palaeontographica Abteilung A 199(1–3):23–111.
- Damborenea, S. E. 1987b. Early Jurassic Bivalvia of Argentina. Part 2: Superfamilies Pteriacea, Buchiacea and part of Pectinacea. Palaeontographica Abteilung A 199(4–6):113–216.
- Damborenea, S. E. 1993. Early Jurassic South American pectinaceans and circum-Pacific palaeobiogeography. Palaeogeography, Palaeoclimatology, Palaeoecology 100(2):109–123.
- Damborenea, S. E. 1996a. Palaeobiogeography of early Jurassic bivalves along the southeastern Pacific margin. Actas del XIII Congreso Geológico Argentino y III Congreso de Exploración de Hidrocarburos V:151–167.
- Damborenea, S. E. 1996b. Andean Jurassic Inoceramids as Potential Bioevent Markers for the Austral Realm. GeoResearch Forum 1–2:433–442.
- Damborenea, S. E. 1998. The bipolar bivalve *Kolymonectes* in South America and the diversity of Propeamussiidae in Mesozoic Times. *In* P. Johnston & J. W. Haggart, eds., Bivalves: An Eon of Evolution - Paleobiological Studies Honoring Norman D. Newell. University of Calgary Press. Calgary. p. 143–155.
- Damborenea, S. E. 2000. Hispanic Corridor: Its evolution and the biogeography of Bivalve Molluscs. GeoResearch Forum 6:369–380.
- Damborenea, S. E. 2001. Unidades paleobiogeográficas marinas jurásicas basadas sobre moluscos bivalvos: una visión desde el hemisferio sur. Anales Academia Nacional de Ciencias Exactas, Físicas y Naturales, Buenos Aires 53:141–160.
- Damborenea, S. E. 2002a. Early Jurassic bivalves of Argentina. Part 3: Superfamilies Monotoidea, Pectinoidea, Plicatuloidea and Dimyoidea. Palaeontographica Abteilung A265:1–119.
- Damborenea, S. E. 2002b. Jurassic evolution of Southern Hemisphere marine palaeobiogeographic units based on benthonic bivalves. Geobios 35(M. sp. 24):51–71.
- Damborenea, S. E. 2004. Early Jurassic Kalentera (Bivalvia) from Argentina and its palaeobiogeographical significance. Ameghiniana 41(2):185–198.

- Damborenea, S. E., & C. M. González-León. 1997. Late Triassic and Early Jurassic Bivalves from Sonora, Mexico. Revista Mexicana de Ciencias Geológicas 14(2):178–201.
- Damborenea, S. E., & Silvia Lanés. 2007. Early Jurassic shell beds from marginal marine environments in southern Mendoza, Argentina. Palaeogeography, Palaeoclimatology, Palaeoecology 250:68–88.
- Damborenea, S. E., & M. O. Manceñido, 1979. On the palaeogeographical distribution of the pectinid genus *Weyla* (Bivalvia, Lower Jurassic). Palaeogeography, Palaeoclimatology, Palaeoecology 27:85–102.
- Damborenea, S. E., & M. O. Manceñido. 1988. Weyla: Semblanza de un bivalvo jurásico andino. Actas V Congreso Geológico Chileno II:13–25.
- Damborenea, S. E., & M. O. Manceñido. 1992. A comparison of Jurassic marine benthonic faunas from South America and New Zealand. Journal of the Royal Society of New Zealand 22:131–152.
- Damborenea, S. E., & M. O. Manceñido. 2005a. Tafonomía y modo de vida de especies de *Retroceramus* (Bivalvia) del Jurásico medio de Argentina. Ameghiniana 42(4) Suplemento Resúmenes:23R.
- Damborenea, S. E., & M. O. Manceñido. 2005b. Biofacies analysis of Hettangian-Sinemurian bivalve/brachiopod associations from the Neuquén Basin (Argentina). Geologica Acta 3(2):163–178.
- Damborenea, S. E., & M. O. Manceñido. 2012. Late Triassic bivalves and brachiopods from southern Mendoza, Argentina. Revue de Paléobiologie Volume Spécial 11:317–344.
- Damborenea, S. E., I. V. Polubotko, I. I. Sey, & K.V. Paraketsov. 1992. Bivalve zones and assemblages of the circum-Pacific region. *In* G. E. G. Westermann, ed., The Jurassic of the Circum-Pacific. Cambridge University Press. Ontario. p. 300–307.
- Dance, S. P. 1962. The authorship of the Portland Catalogue (1786). Journal of the Society for the Bibliography of Natural History 4(1):30–34.
- Dechaseaux, Colette. 1940. Megalodon, Pachyerisma, Protodiceras, Diceras, Pterocardium et l'origine des Diceras. Bulletin de la Société Géologique de France 9:207–218.
- Defrance, M. J. L. 1816. Avicule. In Dictionnaire des Sciences Naturelles, vol. 3. F. G. Levrault. Strasbourg & Paris. p. 141.
- Defrance, M. J. L. 1820. Gervillie, *Gervillia*. In Dictionnaire des Sciences Naturelles, vol. 18. F. G. Levrault. Strasbourg & Paris. p. 502–503.
- Defrance, M. J. L. 1824a. Gervillia. In Dictionnaire des Sciences Naturelles, vol. 38. F. G. Levrault. Strasbourg & Paris. p. 316.
- Defrance, M. J. L. 1825a. Opis. In Dictionnaire des Sciences Naturelles, vol. 36. F. G. Levrault. Strasbourg & Paris. p. 219.
- Defrance, M. J. L. 1825b. Nucule (Fossiles). In Dictionnaire des Sciences Naturelles, vol. 35. F. G. Levrault. Strasbourg & Paris. p. 216–219.
- De Gregorio, Antonio. 1884. Nota intorno ad alcune nuove conchiglia mioceniche de Sicilia. Naturalista Siciliano 3:119–120.
- De Gregorio, Antonio. 1886. Monographie des fossiles de Valpore (Mont Grappa) du sous-horizon Grappin de Greg. Annales de Géologie et de Paléontologie 2:1–20.
- De Gregorio, Antonio. 1930a. Monografia dei fossili Liassici di Monte San Giuliano conservati nel mio privato gabinetto geologico. Annales de Géologie et de Paléontologie de Palerme 53:1–57.
- De Gregorio, Antonio. 1930b. Fossili triassici delle Cave di Billiemi presso Palermo. Annales de Géologie et de Paléontologie. Palermo 54:1–40, pl. 1–7.
- Delvene, Graciela. 2000. Taxonomie und Palökologie der Bivalven im Mittelund Oberjura der Keltiberischen Ketten (Spanien). PhD Thesis. University of Zaragoza. 214 p.
- Delvene, Graciela. 2001. Middle and Upper Jurassic bivalves from the Iberian Range (Spain). Beringeria 28:43–104.
- Delvene, Graciela. 2003. Las asociaciones de bivalvos del Jurásico Medio y Superior en la Cordillera Ibérica (España). Geobios 36:519–531.
- De Renzi, Miquel, & Ana Márquez-Aliaga. 1980. Primary and diagenetic features in the microstructure of some Triassic bivalves. Revista del Instituto de Investigaciones Geológicas 34:101–116.

- Dercourt, Jean, L. E. Ricou, & Bruno Vrielynck, eds. 1993. Atlas Peri-Tethys Palaeogeographical Maps.CCGM/CGMW. Paris. 307 p.
- Deshayes, G. P. 1830–1832. Encyclopédie méthodique. Agasse. Paris. Histoire Naturelle des vers. Tome 2. Première partie:1–256; Deuxième partie:1–594.
- Desio, Ardito. 1929. Studi geologici sulla regione dell'Albenza (Prealpi Bergamasche). Memorie della Scietà Italiana di Scienze Naturali e del Museo Civico di Storia Naturale di Milano 10(1):81–125.
- Desio, Ardito, Carla Rossi Ronchetti, & P. L. Vigano. 1960. Sulla stratigrafia del Trias in Tripolitania e nel Sud-Tunisino. Rivista Italiana di Paleontologia e Stratigrafia LXVI(3):273–322.
- Des Moulins, C. 1832. Déscription d'un nouveau genre de coquille vivante, bivalve, des mers du Chili. Actes de la Société Linnéenne de Bordeaux 5:83–92.
- De Toni, A. 1913. Illustrazione della fauna triasica di Valdepena (Cadore). Memorie dell' Instituto Geologico della R. Università di Padova 2:1–194.
- Dèzes, Pierre. 1999. Tectonic and metamorphic evolution of the Central Himalayan Domain in Southeast Zanskar (Kashmir, India). Mémoires de Géologie (Lausanne), 32:145 p.
- Dhondt, A. V. 1989. Late Cretaceous *Limea (Pseudolimea)* species of Europe. Bulletin de l'Institut Royal des Sciences Naturelles de Belgique 59:105–125.
- Dickins, J. M. 1957. Lower Permian pelecypods and gastropods from the Carnarvon Basin, western Australia. Australia, Bureau of Mineral Resources, Geology and Geophysics, Bulletin 41:1–75, 10 pl.
- Dickins, J. M. 1963. Permian pelecypods and gastropods from Western Australia. Australia, Bureau of Mineral Resources, Geology and Geophysics, Bulletin 63:1–203.
- Dickins, J. M. 1999. Mid-Permian (Kubergandian-Murgabian) bivalves from the Khuff Formation, Oman: implications for world events and correlation. Rivista Italina di Paleontologia e Stratigrafia 105(1):23–35.
- Dickins, J. M., & R. A. McTavish. 1963. Lower Triassic marine fossils from the Beagle Ridge (BMR 10) Bore, Perth Basin, Western Australia. Australian Journal of Earth Sciences 10(1):123–140.
- Diener, Carl. 1908. Ladinic, Carnic and Noric faunae of Spiti. Palaeontologia Indica, Serie 15, Himalayan Fossils 5(3):1–157.
- Diener, Carl. 1915. Zur systematischen Stellung der Pelecypodengattung *Pomarangina*. Centralblatt f
 ür Mineralogie, Geologie und Pal
 äontologie 1915:129–131. (*)
- Diener, Carl. 1923. Lamellibranchiata Triadica. In C. Diener, ed., Fossilium Catalogus I: Animalia. W. Junk. Berlin. 19:1–257.
- Dillwyn, L. W. 1817. A descriptive catalogue of recent shells, arranged according to the Linnean method; with particular attention to the synonymy. John & Arthur Arch. London. vol. 1:1–580; vol. 2:581–1092 + 29 p. (*)
- Di Stefano, G. 1912. La dolomita principale dei dintorni di Palermo e di Castellammare del Golfo (Trapani). Palaeontographica Italica 18:57–104, tav. 8–17.
- Dittmar, Alphons von. 1864. Die Contorta-Zone (Zone der Avicula contorta Portl.). Ihre Verbreitung und ihre organischen Einschlüsse. H. Manz. München. 217 p.
- Douglas, J. A. 1929. A marine Triassic fauna from eastern Persia. Quarterly Journal of the Geological Society 85:624–648.
- Douglas, J. A., & W. J. Arkell. 1932. The Stratigraphical Distribution of the Cornbrash: II. The North-Eastern Area. Quarterly Journal of the Geological Society, London 88(2):112–170, pl. 10–12.
- Douvillé, H. F. 1897. Essai de classification systématique des pectinidés. Bulletin de la Société Géologique de France (série 3) 25:202–205.
- Douvillé, H. F. 1904. Paléontologie, Mollusques Fossiles. In J. de Morgan, Mission Scientifique en Perse, vol. 3, no. 4. E. Leroux. Paris. p. 191–380.
- Douvillé, H. F. 1913. Classification des Lamellibranches. Bulletin de la Société Géologique de France (série 4) 12(7):419–467.

- Dubar, Gonzague, Renné Mouterde, Carmina Virgili, & L. C. Suárez. 1971. El Jurásico de Asturias (Norte de España). Cuadernos de Geología Ibérica 2:561–580.
- Duff, K. L. 1975. Palaeoecology of a bituminous shale—The Lower Oxford Clay of Central England. Palaeontology 18(3):443–482.
- Duff, K. L. 1978. Bivalvia from the English Lower Oxford Clay (Middle Jurassic). Palaeontographical Society Monographs 132:1–137.
- Dumortier, Vincent E. 1864–1874. Études Paléontologiques sur les Dépots Jurassiques du Bassin du Rhône. F. Savy, Paris. Part I, Infra-Lias, p. 1–187, pl. 1–30 (1864); part II, Lias-Inférieur, p. 1–252, pl. 1–50 (1867); part III, Lias-Moyen, p. 1–348 p., pl. 1–45 (1869); part IV, Lias Supérieur, p. 1–335, pl. 1–62 (1874).
- Dun, W. S. 1932. The lower marine forms of *Myonia*, with notes on a proposed new genus, *Pachymyonia*. Records of the Australian Museum 18(8):411–414.
- Dunbar, C. O. 1924. Kansas Permian insects, part 1. The geologic occurrence and the environment of the insects. American Journal of Science 7:171–208.
- Dunker, W. B. R. H. 1846. Über die in dem Lias bei Halberstadt vorkommenden Versteinerungen, part 1. Palaeontographica 1(1):34–41.
- Dunker, W. B. R. H. 1848. Nachtrag zu der Beschreibung der in dem Lias bei Halberstadt vorkommenden Versteinerungen. Palaeontographica 1(4):176–181.
- Dunker, W. B. R. H. 1851. Über die im Muschelkalk von Oberschlesien bis jetz gefundenen Mollusken. Palaeontographica 1(6):283–310.
- von Eichwald, J. K. E. I. 1862. Die vorweltliche Fauna und Flora des Grünsandes der Umgegend von Moskwa. Bulletin de la Societé Imperiale des Naturalistes de Moscou 35(2):355–410.
- Emel'yantsev, T. M., I. S. Gramberg, A. I. Kratsova, and P. S. Puk. 1960. Geologiya i perspektivy neftegazonosnosti nizov'yev r. Leny. [Geology and prospects of oil and gas bearing of the lower Lena]. Trudy Nauchno-Issledovatel'skogo Instituta, Geologii Arktiki 108:724 p. (In Russian)
- Emmrich, A. 1853. Geognostische Beobachtungen aus den östlichen bayerischen und den angränzenden österreichischen Alpen. Jahrbuch der Kaiserlich-Königlichen Geologischen Reichsanstalt 4(2):326–394.
- Encheva, Milka. 1969. Several bivalvia from Campilian by the village Konare, District of Tolbouhin. Bulletin Geological Institute, Serie Paleontology XVIII:47–58.
- Encheva, Milka. 1972. Les fossiles de Bulgarie. II Le Trias. Academie Bulgare des Sciences. p. 1–152.
- Escobar, Fernando. 1980. Paleontologia y bioestratigrafia del Triásico Superior y Jurásico Inferior en el área de Curepto, Provincia de Talca. Instituto de Investigaciones Geológicas de Chile - Boletin 35:46–48.
- Étallon, Auguste. 1862. Études paléontologiques sur les terrains jurassiques du Haut-Jura. Monographie de l'étage Corallien. 2ième partie. Mémoires de la Société d'Émulation du Département du Doubs (3)6: 53–241. Besançon.
- Etayo Serna, Fernando, Nuria Solé De Porta, Jaime De Porta, & Tatiana Gaona. 2003. The Batá Formation of Colombia is truly Cretaceous, not Jurassic. Journal of South American Earth Sciences 16:113–117.
- Etheridge, Robert. 1864. Description of new species of Mollusca & c. Quarterly Journal of the Geological Society of London 20(2):112-114.
- Etter, Walter. 1996. Pseudoplanktonic and benthic invertebrates in the Middle Jurassic Opalinum Clay, northern Switzerland. Palaeogeography, Palaeoclimatology, Palaeoecology 126(3–4):325–341.
- Eudes-Deslongchamps, J. A. 1860. Essai sur les Plicatules fossiles des terrains du Calvados et sur quelques autres genres voisins ou démembrès de ces coquilles. Mémoires de la Societé Linneénne de Normandie 11:1–164, pl. 7–20.
- Fallahi, Mohammed, Bernhard Gruber, & Gottfried Tichy. 1983. Gastropoden und Bivalven aus dem oberen Teil der Nayband-Formation (Obertrias) von Baqirabad (Isfahan, Iran). *In* Helmuth Zapfe ed., Neue Beiträge zur Biostratigraphie der Tethys-Trias. Österreichische Akademie

der Wissenschaften, Schriftenreihe der Erdwissenschaftlichen Kommissionen 5:57-82.

- Fan Jia-song. 1963. On Lower Jurassic Lamellibranchiata from Guangdong (Kuangtung). Acta Palaeontologica Sinica 11(4):508–543. In Chinese with English summary.
- Fang Lee-shing, & Shen Pouyan. 1988. A living mechanical file: the burrowing mechanism of the coral-boring bivalve *Lithophaga nigra*. Marine Biology 97:349–354.
- Fang Zon-jie. 1978. On a new subgenus of Pectinidae (Lamellibranchia). Acta Palaeontologica Sinica 17(4):461–466. In Chinese with English abstract.
- Fang Zon-jie. 1982. On genus *Permoperna* (Bivalvia). Acta Palaeontologica Sinica 21(5):45–552. In Chinese with English abstract.
- Fang Zon-jie. 1993. On "Claraia" (Bivalvia) of late Permian. Acta Palaeontologica Sinica 32(6):653–661. In Chinese with English abstract.
- Fang Zon-jie. 2003. Discussion on the study of the genera *Claraia* and *Claraioides* (Bivalvia) of late Permian age. Acta Palaeontologica Sinica 42(4):613–619. In Chinese with English abstract.
- Fang Zon-jie. 2009. Guizhoumyophoria, a new name for Quadratia Yin, 1974 (Bivalvia: Trigonioida: Myophoriidae), preoccupied by Quadratia Muir-Wood and Cooper, 1960 (Brachiopoda: Strophomenida: Productellidae). Journal of Paleontology 83(6):998.
- Fang Zon-jie, Arthur Boucot, Vladimir Covacevich, & Francisco Hervé. 1998. Discovery of Late Triassic fossils in the Chonos metamorphic complex, southern Chile. Revista Geológica de Chile 25(2):165–173.
- Fang, Zon-jie, Chen Jin-hua, Chen Chu-zhen, Sha Jin-geng, Lan Xiu, & Wen Shi-xuan. 2009. Supraspecific taxa of the Bivalvia first named, described, and published in China (1927–2007). The University of Kansas Paleontological Contributions (new series) 17:157 p
- Fang, Zon-jie, & J. C. W. Cope. 2004. Early Ordovician bivalves from Dali, West Yunnan, China. Palaeontology 47(5):1121–1158.
- Fang, Zon-jie, & N. J. Morris. 1997. The genus *Pseudosanguinolites* and some modioliform bivalves (mainly Palaeozoic). Palaeoworld 7:49–74.
- Fantini Sestini, Nerina. 1966. Il Trias in Lombardia (Studi geologici e paleontologici). XIX. Studio biometrico di "Myophoria kefersteini" (Muenster). Rivista Italiana di Paleontologia e Stratigrafia 72(4):1023–1082.
- Fantini Sestini, Nerina. 1984. Gastropods and bivalves of the middle Anisian from Kokaeli Peninsula (Turkey). Rivista Italiana di Paleontologia e Stratigrafia 90(3):321–330.
- Farabegoli, Enzo, M. C. Perri, & Renato Posenato. 2007. Environmental and biotic changes across the Permian-Triassic boundary in western Tethys: The Bulla parastratotype, Italy. Global and Planetary Change 55:109–135.
- Farsan, N. M. 1972. Stratigraphische und Paläo-geographische stellung der Khenjan-Serie und deren Pelecypoden (Trias, Afghanistan). Palaeontographica Abteilung A 140(4–6):131–191.
- Farsan, N. M. 1975. Pelecypoden aus der Khenjan-serie von Zentral-Afghanistan (Mittel Trias). Palaeontographica Abteilung A 149(4–6):119–138.
- Feng Qinglai, Chongpan Chonglakmani, Dietrich Helmcke, Rucha Ingavat-Helmcke, & Liu Benpei. 2005. Correlation of Triassic Stratigraphy between the Simao and Lampang-Phrae Basins: implications for the tectonopaleogeography of Southeast Asia. Journal of Asian Earth Sciences 24:777–785.
- Feng, Qinglai, Cui Xinsheng, & Liu Benpei. 1992. Discovery of Late Permian bivalve-fauna from Laochang of Lancang, West Yunnan and its biogeographic characteristics. Earth Science–Journal of China University of Geosciences 17(5):512–520. In Chinese with English abstract.
- de Férussac, André E. J. d'Audebard. 1821–1822. Tableaux Systèmatiques des Animaux Mollusques suivis d'un Prodrome Général pour tous les Mollusques Terrestres ou Fluviatiles Vivants ou Fossiles. Première Partie, Tableaux Systématiques Généraux. Arthus-Bertrand. Paris. J. B. Sowerby. London. 110 p.
- Fischer, A. G., & D. J. Bottjer. 1995. Oxygen-depleted waters: a lost biotope and its role in ammonite and bivalve evolution. Neues Jahrbuch für Geologie und Paläontologie Abhandlungen 195(1–3):133–146.

- Fischer, J. C. 1969. Géologie, paléontologie et paléoécologie du Bathonien au SudOuest du Massif Ardennais. Mémoire, Musée National d'Histoire Naturelle n.s. (series C) Sciences de la Terre 20:1–321, pl. 1–21.
- Fischer, J. C., & C. Palain. 1971. Nouvelles Observations Sédimentologiques et Paléobiologiques sur l'Hettangien du Portugal. Comunicações dos Serviços Geológicos de Portugal LV:105–132.
- Fischer, Paul. 1880–1887. Manuel de Conchyliologie et de Paléontologie Conchyliologique, ou Histoire Naturelle des Mollusques Vivants et Fossiles suivi d'un Appendice sur les Brachiopodes par D. P. Oehlert. Avec 23 planches contenant 600 figures dessinées par S. P. Woodward. Savy, Paris. xxiv + 1369 p., 23 pl., 1138 text fig. (1:1–112 [1880]; 2:113–192 [1881]; 3:193–304 [1881]; 4:305–416 [1882]; 5:417–512 [1883]; 6:513–608 [1883]; 7:609–688 [1884a]; 8:689–784 [29, 1885]; 9:785–896 [1885]; 10:897–1008 [1886]; 11:1009–1369 [1887]).
- Fleming, C. A. 1957. The Triassic Lamellibranch *Torastarte bensoni* Marwick. Proceedings of the Malacological Society of London 32(4):173–174.
- Fleming, C. A. 1962. Two new genera of Triassic Trigoniidae from New Zealand. Proceedings of the Malacological Society of London 35(1):1–4.
- Fleming, C. A. 1963. A new genus and species of Trigoniidae from the Middle Triassic of New Zealand. New Zealand Journal of Geology and Geophysics 6(5):843–846.
- Fleming, C. A. 1964. History of the Bivalve Family Trigoniidae in the South-West Pacific. The Australian Journal of Science 26(7):196–204.
- Fleming, C. A. 1987. New Zealand Mesozoic bivalves of the superfamily Trigoniacea. New Zealand Geological Survey Paleontological Bulletin 53:1–104.
- Fleming, C. A., F. W. Munden, & R. P. Suggate. 1954. An Upper Triassic lamellibranch from the Southern Alps of North Westland, New Zealand (Sheet S53). Transactions of the Royal Society of New Zealand 82:111–114.
- Fleming, John. 1828. A History of British Animals, Exhibiting the Descriptive Characters and Systematical Arrangement of the Genera and Species of Quadrupeds, Birds, Reptiles, Fishes, Mollusca, and Radiata of the United Kingdom; including the Indigenous, Extirpated, and Extinct Kinds, Together with Periodical and Occasional Visitants. Bell & Bradfute, Edinburgh, & James Duncan. London. 565 p.
- Fletcher, Harold O. 1945. A new genus *Glyptoleda* and a revision of the genus *Nuculana* from the Permian of Australia. Records of the Australian Museum 21:293–312.
- Fozy, István, Miklós Kázmér, & István Szente. 1994. A unique Lower Tithonian fauna in the Gerecse Mts, Hungary. Proceedings of the 3rd Pergola International Symposium. Palaeopelagos Special Publication 1:155–165.
- Fraiser, M. L., & D. J. Bottjer. 2005a. Fossil preservation during the aftermath of the end-Permian mass extinction: taphonomic processes and palaeoecologial signals. *In* D. J. Over, J. R. Morrow & P. B Wignall, eds., Understanding Late Devonian and Permian-Triassic Biotic and Climatics Events: Towards an Integrated Approach. Elsevier. Amsterdam. Chapter 11:299–312.
- Fraiser, M. L., & D. J. Bottjer. 2005b. Restructuring in benthic levelbottom shallow marine communities due to prolonged environmental stress following the end-Permian mass extinction. Comptes Rendus Palevol 4:515–523.
- Fraiser, M. L., & D. J. Bottjer. 2007a. When bivalves took over the world. Paleobiology 33(3):397–413.
- Fraiser, M. L., & D. J. Bottjer. 2007b. Elevated atmospheric CO₂ and the delayed biotic recovery from the end-Permian mass extinction. Palaeogeography, Palaeoclimatology, Palaeoecology 252:164–175.
- Francis, A. O., & Anthony Hallam. 2003. Ecology and evolution of Jurassic trigoniid bivalves in Europe. Lethaia 36:287–304.
- Fraser, N. M. 1997. Large epifaunal bivalves from Mesozoic buildups of Western North America. M. S. Thesis. Faculty of the Graduate School University of Southern California. Unpublished. 312 p.

- Fraser, N. M., & D. J. Bottjer. 2001a. The beginning of Mesozoic bivalve reefs: Early Jurassic "Lithiotis" facies bioherms. *In* Abstracts of California paleontology conference 2001. PaleoBios 21(1):3–4.
- Fraser, N. M., & D. J. Bottjer. 2001b. Dissecting the Lithiotis facies; implications for the Early Jurassic reef eclipse. Abstracts with Programs. Geological Society of America 33(6): 378.
- Fraser, N. M., D. J. Bottjer, & A. G. Fischer. 2004. Dissecting "Lithiotis" Bivalves: Implications for the Lower Jurassic reef eclipse. PALAIOS 19:51–67.
- Frebold, Hans, & T. P. Poulton. 1977. Hettangian (Lower Jurassic) rocks and faunas, northern Yukon Territory. Canadian Journal of Earth Sciences 14(1):89–101.
- Frech, F. 1891. Die Devonischen Aviculiden Deutschlands. Ein Beitrag zur Systematik und Stammesgeschichte der Zweischaler. Abhandlungen zur Geologischen Specialkarte von Preussen und den Thüringischen Staaten 9(2):1–261; 9(3):199–459, 18 pl.
- Frech, Fritz. 1904. Neue Zweischaler und Brachiopoden aus der Bakonyer Trias. Resultate der Wissenschaftlichen Erforschung des Balatonsees 1(1):1–137.
- Frech, F. 1905. Nachträge zur den Cephalopoden und Zweischalern der Bakonyer Trias (Werfener und Cassianer Estherienschichten). Resultate der Wissenschaftlichen Erforschung des Balatonsees, Wien 1(1):1–30.
- Frech, Fritz. 1907. Ueber Aviculiden von palaeozoischem Habitus aus der Trias von Zacatecas. Comptes Rendus du 10° Congrès Géologique International, Mexico 10:327–335.
- Frech, F. 1909. Die Leitfossilien der Werfener Schichten und Nachträge zur Fauna des Muschelkalkes, der Cassianer und Raibler Schichten, sowie der Rhaet und des Dachsteindolomites (Hauptdolomit). Resultate der Wissenschaftlichen Erforschung des Balatonsees 1(1), Anhang. Palaeontologie der Umgebung des Balatonsees 2(6):1–96, pl. 1–16.
- Freitas, T. A., F. Brunton, & T. Bernecker. 1993. Silurian megalodont bivalves of the Canadian Arctic and Australia: paleoecology and evolutionary significance. PALAIOS 8:450–464.
- Freneix, Suzanne. 1965. Les bivalves du Jurassique Moyen et Supérieur du Sahara Tunisien. Annales de Paléontologie (Invertébrés) 51(1):65 p.
- Freneix, Suzanne. 1980. Bivalves Néocrétacés de Nouvelle-Calédonie. Signification biogéographique, biostratigraphique, paléoécologique. Annales de Paléontologie (Invertébrés) 66(2):67–134.
- Freneix, Suzanne. 1999. Bivalves du Trias des Rochers de los Pastores (Ouest D'Algeciras, Andalousie, Espagne). Geodiversitas 21(2):137–146.
- Freneix, Suzanne, & Jacques Avias. 1977. Caledogonia, a new smooth trigoniacean bivalve from the Triassic of New Caledonia. Alcheringa 1:279–291.
- Freneix, Suzanne, & René Cubaynes. 1984. Biofaciès à *Cuneigervillia—Pte-romya* et biofaciès à *Eomiodon* dans l'Hettangien du Quercy (Aquitaine orientale). Geobios 17(1):5–17.
- Fuchs, Günter, & Dotlef Mader. 1980. Fossiliferous marine Muschelkalk in the Oberbettinger Triasgebiet (Western Eifel, Germany). Neues Jahrbuch für Geologie und Paläontologie Monatshefte 11:651–672.
- Fürsich, F. T. 1980. Preserved life positions of some Jurassic bivalves. Palaeontologische Zeitschrift 54(3–4):289–300.
- Fürsich, F. T. 1981. Salinity-controlled benthic associations from the Upper Jurassic of Portugal. Lethaia 14(3):203–223.
- Fürsich, F. T. 1982. Upper Jurassic bivalves from Milne Land, East Greenland. Grønlands Geologiske Undersøgelse Bulletin 144:1–126.
- Fürsich, F. T., Roman Berndt, Thomas Scheuer, & Matthias Gahr. 2001. Comparative ecological analysis of Toarcian (Lower Jurassic) benthic faunas from southern France and east-central Spain. Lethaia 34:169–199.
- Fürsich, F. T., Susanne Freytag, Jochen Röhl, & Annette Schmid. 1995. Palaeoecology of benthic associations in salinity-controlled marginal marine environments: Examples from the Lower Bathonian (Jurassic) of the Causses (southern France). Palaeogeography, Palaeoclimatology, Palaeoecology 113:135–172.

- Fürsich, F. T., & Michael Hautmann. 2005. Bivalve reefs from the Upper Triassic of Iran. Annali dell'Università degli Studi di Ferrara, sezione Museologia Scientifica e Naturalistica, Special volume in honour of Carmela Loriga. p. 13–23.
- Fürsich, F. T., Wolfgang Oschmann, A. K. Jaitly, & I. B. Singh. 1991. Faunal response to transgressive-regressive cycles: example from the Jurassic of western India. Palaeogeography, Palaeoclimatology, Palaeoecology 85:149–159.
- Fürsich, F. T., T. J. Palmer, & K. L. Goodyear. 1994. Growth and disintegration of bivalve-dominated patch reefs in the Upper Jurassic of southern England. Palaeontology 37(1):131–171.
- Fürsich, F. T., & Elsebeth Thomsen. 2005. Jurassic biota and biofacies in erratics from the Sortland area, Vesterålen, northern Norway. Norges Geologiske Undersøkelse Bulletin 443:37–53.
- Fürsich, F. T., & Jobst Wendt. 1977. Biostratinomy and Palaeoecology of the Cassian formation (Triassic) of the southern Alps. Palaeogeography, Palaeoeclimatology, Palaeoecology 22:257–323.
- Fürsich, F. T., & Winfried Werner. 1988. The Upper Jurassic Bivalvia of Portugal. Part II. Pteriomorphia (Pterioida exclusive Ostreina). Comunicações dos Serviços Geológicos de Portugal 74:105–164.
- Fürsich, F. T., Markus Wilmsen, Kazem Seyed-Emami, Fabrizio Cecca, & M. R. Majidifard. 2005. The upper Shemshak Formation (Toarcian-Aalenian) of the Eastern Alborz (Iran): Biota and palaeoenvironments during a transgressive-regressive cycle. Facies 51:365–384.
- Gabb, William M. 1864. Description of the Triassic fossils of California and the adjacent territories. California Geological Survey, Paleontology 1:17–35.
- Gabb, William M. 1870. Descriptions of some Secondary fossils from the Pacific States. American Journal of Conchology vol. 5–18. 5 pl.
- Gaetani, Maurizio. 1970. Faune Hettangiane della parte orientale della provincia di Bergamo. Rivista Italiana di Paleontologia e Stratigrafia 76(3):355–442.
- Gaetani, Maurizio., V. Lozowski, J. Szulc, Alfredo Arche, Francesc Calvet, & José López-Gómez. 2000a. Olenekian (245–243 Ma). *In* J. Dercourt, M. Gaetani, B. Vrielynck, E. Barrier, B. Biju-Duval, M. F. Brunet, J. P. Cadet, S. Crasquin & M. Sandulescu, eds., Atlas Peri-Tethys, Palaeogeographical maps. JCCGM/ CGMW. Paris. Explanatory notes. p. 27–32.
- Gaetani, Maurizio., V. Lozowski, J. Szulc, Alfredo Arche, Francesc Calvet, & José López-Gómez. 2000b. Early Ladinian (238–235 Ma). *In* J. Dercourt, M. Gaetani, B. Vrielynck, E. Barrier, B. Biju-Duval, M. F. Brunet, J. P. Cadet, S. Crasquin & M. Sandulescu, eds., Atlas Peri-Tethys, Palaeogeographical maps. JCCGM/ CGMW. Paris. Explanatory notes. p. 33–39.
- Gaetani, Maurizio., V. Lozowski, J. Szulc, Alfredo Arche, Francesc Calvet, & José López-Gómez. 2000c. Late Norian (215–212 Ma). In J. Dercourt, M. Gaetani, B. Vrielynck, E. Barrier, B. Biju-Duval, M. F. Brunet, J. P. Cadet, S. Crasquin & M. Sandulescu, eds., Atlas Peri-Tethys, Palaeogeographical maps. JCCGM/ CGMW. Paris. Explanatory notes. p. 41–48.
- Gahr, M. E. 2002. Palökologie des Makrobenthos aus dem Unter-Toarc SW-Europas. Beringeria 31:3–204.
- Gair, H. S., D. R. Gregg, & I. G. Speden. 1962. Triassic fossils from Corbies Creek, North Otago. New Zealand Journal of Geology and Geophysics 5(1):92–113.
- Galdieri, Agostino. 1908. Sul Trias dei dintorni di Giffoni. Atti della Accademia Pontaniana 38(8):123 p.
- Gall, J. C. 1971. Faunes et paysages du grès à Voltzia du Nord des Vosges. Essai paléoécologique sur le Buntsandstein supérieur. Mémoires Service de la Carta Géologique d'Alsace et de Lorraine. Université Louis Pasteur de Strasbourg 34:1–318.
- Gan Xiuming, & Yin Hongfu. 1978. Lamellibranchiata. *In* Stratigraphy and Palaeontology Working Team of Guizhou Province, ed., Paleontological Atlas of SW China, Volume of Guizhou Province, pt. 2. Geological Publishing House. Beijing. p. 305–393. In Chinese. (*)

- Ganelin, V. G., & A. S. Biakov. 2006. The Permian biostratigraphy of the Kolyma-Omolon region, Northeast Asia. Journal of Asian Earth Sciences 26:225–234.
- Gang Li. 2001. Palaeontology and biostratigraphy of the Lower Cretaceous Qihulin Formation in eastern Heilongjiang, northeastern China. PhD Thesis. University of Heidelberg. 128 p., https://heidi.ub.uni-heidelberg. de/volltextserver/volltexte/ 2001/1819/pdf/GangLi.pdf. Checked August 2012.
- Gao Yong-qun, Yang Feng-qing, & Peng Yuan-qiao. 2004. Significance of *Claraia* from the upper Permian of South Guizhou, China. Alcheringa 28(2):469–476.
- García-March, J. R. 2005. Aportaciones al conocimiento de la Biología de *Pinna nobilis* Linneo, 1758 (Mollusca: Bivalvia) en el litoral mediterráneo Ibérico. PhD Thesis. Servicio de Publicaciones de la Universidad de Valencia. 332 p.
- García-March, J. R., Ana Márquez-Aliaga, & J. G. Carter. 2008. The duplivincular ligament of recent *Pinna nobilis* L., 1758: further evidence for pterineid ancestry of the Pinnoidea. Journal of Paleontology 82(3):621–627.
- Gardner, R. N. 2005. Middle-Upper Jurassic bivalves of the superfamily Veneroidea from New Zealand and New Caledonia. New Zealand Journal of Geology and Geophysics 48:325–376.
- Gardner, R. N., & H. J. Campbell. 1997. The bivalve genus Grammatodon from the Middle Jurassic of the Catlins District, South Otago, New Zealand. New Zealand Journal of Geology and Geophysics 40(4):487–498.
- Gardner, R. N., & H. J. Campbell. 2002. Middle to Upper Jurassic bivalves of the genera *Neocrassina* and *Trigonopis* from New Zealand. New Zealand Journal of Geology and Geophysics 45:323–347.
- Gavrilova, V. A. 1995. A new Pterinopectinid genus (Bivalvia) from the Lower Triassic of the Mangyshlak. Paleontologicheskii Zhurnal 29(1): 132–137. Translationin Paleontological Journal 29(1):154–160.
- Gavrilova, V. A. 1996. On the systematics of Triassic Pterinopectinidae (Bivalvia). Paleontological Journal 30(5):497–505.
- Gelati, Romano, & Andrea Allasinaz. 1964. Nuovo contributo alla conoscencza del Trias superiore dell'Alta Valtellina. Rivista Italiana di Paleontologia LXX(1):15–64.
- Geyer, O. F. 1973. Das präkretazische Mesozoikum von Kolumbien. Geologisches Jahrbuch B 5:1–155.
- Giebel, C. G. A. 1852. Allgemeine Palaeontologie: Entwuf einer Systematischen Darstellung der Fauna und Flora der Vorwelt, zum Gebrauche bei Vorlesungen und zum Selbtsunterrichte. Ambrosius Abel. Leipzig. 413 p.
- Giebel, C. G. A. 1855. Kritisches über die Myophorien des Muschelkalkes. Zeitschrift für die Gesammten Naturwissenschaften, Berlin 5:34–36.
- Giebel, C. G. 1856. Die Versteinerungen in Muschelkalk von Lieskau bei Halle. Abhandlungen Naturwissenschaftlichen Vereines I:53–126.
- Gillet, Suzette. 1924. Études sur les lamellibranches néocomiens. Mémoires de la Société Géologique de France (nouvelle série) 3:1–224.
- Girty, G. H. 1911. On some new genera and species of Pennsylvanian fossils from the Wewaka Formation of Oklahoma. Annals of the New York Academy of Science 21:119–156.
- Gmelin, J. F. 1791. Caroli a Linné, systema naturae. Tom. I. Pars VI. Lipsiae. p. 3021–3910.
- Goldfuss, G. A. 1833–1841. Petrefacta Germaniae tam ea, quae in Museo Universitatis Regiae Borussicae Fridericiae Wilhelmiae Rhenanae Servantur quam alia Quaecunquae in Museis Hoenighusiano Muensteriano Aliisque Extant, Iconibus et Descriptionibus Illustrata. Abbildungen und Beschreibungen der Petrefacten Deutschlands und der Angrenzender Länder unter Mitwirkung des Herrn Grafen Georg zu Münster. Arnz & Co. Düsseldorf. Zweiter Theil [Bivalvia], part 1, p. 1–68, with Atlas, pl. 72–96 [1833]; part 2, p. 69–140, with Atlas. pl. 97–121 [1835]; part 3, p. 141–224, with Atlas, pl. 122–146 [1837]; part 4, p. 225–312, with Atlas, pl. 147–165 [1841].

- Goldfuss, G. A. 1863. Abbildungen und Beschreibungen der Petrefacten Deutschlands und der angrenzenden Länder. Divisio Quarta: Molluscorum acephalicorum reliquiae. Muschelthiere der Vorwelt. 1 Bivalvia. Petrefacta Germaniae IV(1):1–273.
- Golebiowski, R. 1990. Facial and faunistic changes from Triassic to Jurassic in the Northern Calcareous Alps (Austria). Les Cahiers de l'Université Catholique de Lyon, Série Sciences 3:175–184.
- Golonka, Jan. 2004. Plate tectonic evolution of the southern margin of Eurasia in the Mesozoic and Cenozoic. Tectonophysics 381:235–273.
- Golonka, Jan. 2007. Late Triassic and Early Jurassic palaeogeography of the world. Palaeogeography, Palaeoclimatology, Palaeoecology 244:297–307.
- Golonka, Jan, & David Ford. 2000. Pangean (Late Carboniferous–Middle Jurassic) paleoenvironment and lithofacies. Palaeogeography, Palaeoclimatology, Palaeoecology 161:1–34.
- Gómez, J. J., Antonio Goy, & Ana Márquez-Aliaga. 2005. Bioestratigrafía del tránsito Triásico-Jurásico en Asturias: ammonoideos y bivalvos entre Avilés y Caravia (España). Geo-Temas 8:183–187.
- González, C. R. 1969. Nuevas especies de Bivalvia del Paleozoico Superior del sistema de Tepuel, provincia de Chubut, Argentina. Ameghiniana 6(3):236–350.
- González, C. R. 1992. La fauna de Cerro Veladero, Carbónico de la cuenca del Rio Blanco, La Rioja. Acta Geologica Lilloana 17(2):101–113.
- Goodwin, D. H. 1997. The Importance of paleoecology in assessing paleogeographic relationships of the Antimonio Formation. *In C. M.* González-León & G. D. Stanley, Jr., eds., US - Mexico Cooperative Research: International Workshop on the Geology of Sonora Memoir. Publicaciones Ocasionales 1: 36–38.
- Gordillo, Sandra, & A. E. Aitken. 2000. Palaeoenvironmental interpretation of Upper Quaternary marine molluscan assemblages, Canadian Arctic Archipelago. Géographie physique et Quaternaire 54(3):301–315.
- Gortani, Michele. 1902. Nuovi Fossili raibliani della Carnia. Rivista Italiana di Paleontologia 8:76–94.
- Goto, Michiharu. 1983. Some bivalves from the Lower Jurassic Kuruma Group of Central Japan. Transactions and Proceedings of the Palaeontological Society of Japan (new series) 130:79–84.
- Gou Zong-hai. 1985. Inoceramidae fossils in Xizang (Tibet). Acta Palaeontologica Sinica 24(5): 485–491. In Chinese with English abstract.
- Gou Zong-hai. 1993. Bivalve fauna of Upper Triassic in Maantang area, Jiangyou, Sichuan. Acta Palaeontologica Sinica 32(1):13–30. In Chinese with English abstract.
- Gou Zong-hai. 2003. A bivalve fauna of Early and Middle Jurassic epoch in Nieniexiongla region, Nyalam County, Tibet. Journal of Chengdu University of Technology 30(2):147–154. In Chinese.
- Goy, Antonio, & Ana Márquez-Aliaga. 1998. Bivalvos del Triásico Superior en la Formación Imón (Cordillera Ibérica, España). Boletín de la Real Sociedad Española de Historia Natural (Sec. Geol.) 94(1–2):77–91.
- Gradstein, F. M., & J. G. Ogg. 2004. Geologic Time Scale 2004–why, how and where next! Lethaia 37:175–181.
- Grant-Mackie, J. A. 1960. On a new *Kalentera* (Pelecypoda: Cypricardiacea) from the Upper Triassic of New Zealand. New Zealand Journal of Geology and Geophysics 3:74–80.
- Grant-Mackie, J. A. 1976. The Upper Triassic bivalve *Monotis* in the southwest Pacific. Pacific Geology 11:47–56.
- Grant-Mackie, J. A. 1978a. Subgenera of the Upper Triassic bivalve *Monotis*. New Zealand Journal of Geology and Geophysics 21(1):97–111.
- Grant-Mackie, J. A. 1978b. Status and identity of the New Zealand Upper Triassic bivalve *Monotis salinaria* var. *richmondiana* Zittel 1864. New Zealand Journal of Geology and Geophysics 21(3):375–402.
- Grant-Mackie, J. A. 1978c. Systematics of New Zealand *Monotis* (Upper Triassic Bivalvia) - Subgenus *Entomonotis*. New Zealand Journal of Geology and Geophysics 21(4):483–502.

- Grant-Mackie, J. A. 1978d. Systematics of New Zealand *Monotis* (Upper Triassic Bivalvia): Subgenus *Maorimonotis*. Journal of the Royal Society of New Zealand 3(3):293–322.
- Grant-Mackie, J. A. 1980a. Systematics of New Zealand *Monotis* (Upper Triassic Bivalvia): subgenus *Eomonotis*. New Zealand Journal of Geology and Geophysics 23:639–663.
- Grant-Mackie, J. A. 1980b. Systematics of New Zealand Monotis (Upper Triassic Bivalvia): Subgenus Inflatomonotis. New Zealand Journal of Geology and Geophysics 23:629–637.
- Grant-Mackie, J. A. 1981. New Zealand Warepan (Upper Triassic) sequences: Murihiku Supergroup of the North Island. Journal of the Royal Society of New Zealand 11(3):231–256.
- Grant-Mackie, J. A. 1984. New Zealand Warepan (Upper Triassic) sequences: Murihiku Supergroup of the South Island. Journal of the Royal Society of New Zealand 14(2):175–206.
- Grant-Mackie, J. A. 1994. Mesozoic Bivalvia from Clerke and Mermaid canyons, Northwest Australian continental slope. *In* N. Exon, ed., Geology of the outer North West Shelf, Australia. AGSO Journal of Australian Geology and Geophysics 15(1):119–125.
- Grant-Mackie, J. A., & N. J. Silberling. 1990. New data on the Upper Triassic bivalve *Monotis* in North America, and the new subgenus *Pacimonotis*. Journal of Paleontology 64(2):240–254.
- Grant-Mackie, J. A., & von Helmuth Zapfe. 1973. Otapiria (Monotidae, Bivalvia) aus den Zlambach-Schichten des Salzkammergutes, OÖ. Anzeiger der Österreichischen Akademie der Wissenschaften, Mathematisch-Naturwissenschaftlichen Klasse, Jahrgang 7:45–49.
- Gray, J. E. 1824. A Supplement to the Appendix of Captain Perry's Voyage for the Discovery of a North West Passage, in the Years 1819–1820, Containing an Account of the Subjects of Natural History. Appendix X. Natural History, Shells: ccxl–ccxlvi. *In* W. E. Parry, Journal of a Second Voyage for the Discovery of a Northwest Passage from the Atlantic to the Pacific, Performed in the Years 1821–22–23, in His Majesty's Ships Fury and Hecla, under the Orders of Captain William Edward Parry, R. N., F. R. S. J. Murray, London.
- Gray, J. E. 1840. Shells of molluscous animals. *In* Synopsis of the contents of the British Museum, 42nd ed. G. Woodfall & Son. London. p. 105–152.
- Gray, J. E. 1842. Molluscs. In [multiauthored] Eastern Zoological Gallery. In [multiauthored], Synopsis of the Contents of the British Museum, 44nd ed. G. Woodfall and Son. London. p. 48–92.
- Gray, J. E. 1847. A list of the genera of Recent Mollusca, their synonyma and types. Zoological Society of London, Proceedings 15(178):129–219.
- Gray, J. E. 1854. A revision of the arrangement of the families of bivalve shells (Conchifera). Annals and Magazine of Natural History (series 2) 13(77):408–418; 14(79):21–28.
- Gray, J. E. 1858. On Carpenteria and Dujardinia, two genera of a new form of Protozoa with attached multilocular shells filled with sponge, apparently intermediate between Rhizopoda and Porifera. Proceedings of the Zoological Society of London 26(1):266–271.
- Griffin, Miguel. 1991. Eocene bivalves from the Río Turbio Formation, Southwestern Patagonia (Argentina). Journal of Paleontology 65(1):119–146.
- Griffin, Miguel, & S. N. Nielsen. 2008. A revision of the type specimens of Tertiary molluscs from Chile and Argentina described by d'Orbigny (1842), Sowerby (1846) and Hupé (1854). Journal of Systematic Palaeontology, Published online by Cambridge University Press 06 Feb 2008 doi:10.1017/S1477201907002374.
- Groeber, Pablo. 1924. Descubrimiento del Triásico marino en la República Argentina. Comunicaciones del Museo de Historia Natural de Buenos Aires 2(9):87–94.
- Gruber, Bernhard. 1976. Neue Ergebnisse auf dem Gebiete der Ökologie, Stratigraphie und Phylogenie der Halobien (Bivalvia). Mitteilungen der Gesellschaft der Geologie und Bergbaustudenten in Österreich, Wien 23:181–198.

- Gruber, Bernhard. 1984. Bemerkungen zur Morphologie and Ökologie von Otapiria marshalli alpine (Zapfe) aus den Zlambachmergeln (Obertrias) von Osterreich. Mitteilungen der Gesellschaft der Geologie und Bergbaustudenten Österreich, Wien 29:207–216. (*)
- Gu Zhiwei, Chen Chuzhen, Li Jinhua, & Cheng Zhengxiu. 1980. Waagenoperna (Waagenoperna) and its distribution in the "Lower Jurassic" of South China. Acta Palaeontologica Sinica 19(6):445–460. In Chinese with English abstract.
- Gu Zhiwei, Huang Baoyu, Chen Chuzhen, Wen Shixuan, Ma Qihong, Lan Xiu, Xu Juntao, Liu Lu, Wang Shumei, Wang Deyou, Qiu Ranzhong, Huang Zhaoqi, Zhang Zuoming, Chen Jinhua, & Wu Peili. 1976. The Fossil Lamellibranchiata of China. Nanjing Institute of Geology and Palaeontology, Academia Sinica, Science Press. Beijing. 580 p. In Chinese.
- Gu Zhiwei, Li Zia-hun, & Yu Xi-han. 1997. Lower Cretaceous bivalves from the eastern Heilongjiang Province of China. Science Press. Beijing. 301 p.
- Guérin-Franiatte, Simone. 1990. Exemples du changement caractéristique des faunes vers la fin du Trias et le début du Jurassique. Cahiers de l'Université Catholique de Lyon, Série Sciences 3:17–25.
- Guex, Jean, Annachiara Bartolini, Viorel Atudorei, & David Taylor. 2003. Two negative ¹³C_{org} excursions near the Triassic - Jurassic boundary in the New York Canyon area (Gabbs Valley Range, Nevada). Bulletin de Géologie, Lausanne 360:1–4.
- Guex, Jean, Annachiara Bartolini, Viorel Atudorei, & David Taylor. 2004. High-resolution ammonite and carbon isotope stratigraphy across the Triassic–Jurassic boundary at New York Canyon (Nevada). Earth and Planetary Science Letters 225:29–41.
- Gugenberger, Odomar. 1935. Die *Cardita*-Schichten von Launsdorf in Mittelkärnten und ihre Fauna. V, Lamellibranchiaten. Theil 1. Sitzungsberichte der Mathematisch-Naturwissenschaftliche Klasse der Akademie der Wissenschaften, Wien 144(1):231–252, pl. 1.
- Guillaume, Louis. 1928. Revision des Posidonomyes Jurassiques. Bulletin de la Société Géologique de France (4ème série) 27(3–5):217–234.
- Gümbel, C. W. 1861. Geognostische Beschreibung des bayerischen Alpengebirges und seines Vorlandes. Verlag von Justus Perthes. Gotha. 950 p., 42 pl.
- Gümbel, C. W. 1862. Die Dachsteinbivalve (*Megalodon triqueter*) und ihre alpinen Verwandten. Sitzungsberichte, Akademie der Wissenstchaften, mathematischen und naturwissenchaftlichen Klasse 45:326–377, Taf. 1–7.
- Gümbel, C. W. 1871. Die sogenannten Nulliporen (Lithothamnium und Dactylopora) und ihre Betheiligung an der Zusammensetzung der Kalkgesteine, pt 1: Die Nulliporen des Pflanzenreichs (Lithothamnium). Abhandlungen der Mathematisch-Physikalischen Classe der Königlich Bayerischen Akademie der Wissenschaften 11:13–52.
- Gümbel C. W. 1880. Geognostische Mitteilungen aus den Alpen. VI: Ein geognostischer Streifzug durch die Bergamasker Alpen. Sitzungsberichte der königliche Akademie der Wissenschaften Mathematisch-Naturwissenschaftliche Klasse 10, 164 p., München.
- Guo Fu-xiang, 1985. Fossil Bivalves of Yunnan. Yunnan Science and Technology Publishing House. Kunming. 319 p. In Chinese with English abstract.
- Guo Fu-xian. 1988. New genera of fossil bivalves from Yunnan. Yunnan Geology 7:112–144. In Chinese with English abstract. (*)
- Haas, O. 1909. Bericht über neue Aufsammlungen in den Zlambachmergeln der Fischerwiese bei Alt-Aussee. Beiträge zur Paläontologie und Geologie, Österreich-Ungarns und des Orient 22:143–167.
- Hagdorn, Hans. 1982. The "Bank der kleinen Terebrateln" (Upper Muschelkalk, Triassic) near Schwäbisch Hall (SW-Germany) - a tempestite condensation horizon. *In* G. Einsele & A. Seilacher, eds., Cyclic and Event Stratification. Springer. Berlin. p. 263–285.
- Hagdorn, Hans. 1991. The Muschelkalk in Germany-an introduction. In H. Hagdorn, ed., Muschelkalk, A Field Guide. Goldschneck-Verlag Werner K. Weidert. p. 7–21.

- Hagdorn, Hans. 1995. Farbmuster und Pseudoskulptur bei Muschelkalkfossilien. [Colour Patterns Pseudosculture in Muschelkalk Fossils]. Neues Jahrbuch für Geologie und Paläontologie Monatshefte 1995(1–3):85–108.
- Hagdorn, Hans, & Theo Simon. 1983. Ein Hartgrund im Unteren Muschelkalk von Göttingen. Aufschluss 34:255–263.
- Hagdorn, Hans, & Theo Simon. 1985. Geologie und Landschaft des Hohenloher Landes. Herausgegeben vom Historischen Verein für Württembergisch Franken, dem Stadtarchiv Schwäbisch Hall und dem Hohenlohe-Zentralarchiv Neuenstein 28:186 p.
- Hagdorn, Hans, & Theo Simon. 1991. Ökostratigraphische Leitbänke im Oberen Muschelkalk. *In* H. Hagdorm & A. Seilacher, eds., Muschelkalk Schöntaler Symposium 1991. Sonderbände der Gesellschaft für Naturkunde in Württemberg 2. Korb (Goldschneck). Stuttgart. 2:193–208.
- von Hagenow, F. 1842. Monographie der Rügen'schen Kreideversteinerungen. III. Abteilung: Mollusken. Neues Jahrbuch für Mineralogie, Geognosie, Geologie und Petrefaktenkunde 9:528–575.
- Hall, James. 1852. Natural History of New York. Paleontology of New York, vol. 2. Containing Descriptions of the Organic remains of the Lower Division of the New-York System (Equivalent in part to the Middle Silurian Rocks of Europe). Van Benthuysen. Albany. 362 p., pl. 1–85.
- Hall, James. 1858. Palaeontology of Iowa. Report of the Geological Survey of the State of Iowa 1(2):1–724.
- Hall, J. 1859–1861. Natural History of New York. Palaeontology of New York. Palaeontology. State of New York. Albany. Part 1 (text):i–xii, 1–533 (1859). Part 2 (plates):pl. 1–120 (1861).
- Hall, James, & R. P. Whitfield. 1869. Preliminary notice of the lamellibranchiata shells of the upper Helderberg, Hamilton and Chemung Groups with others from the Waverly sandstones, part 2. New York State Museum. Albany. p. 1–96.
- Hall, James, & R. P. Whitfield. 1877. Part II. Palaeontology. In C. King, ed., Annual Report to the Secretary of War on the U. S. Geological Exploration of the Fortieth Parallel. Government Printing Office. Washington, D. C. p. 198–302.
- Hallam, Anthony. 1969. Faunal realms and facies in the Jurassic. Palaeontology 12(1):1–18.
- Hallam, Anthony. 1972. Diversity and density characteristics of Pliensbachian - Toarcian molluscan and brachiopod faunas of the North Atlantic margins. Lethaia 5:389–412.
- Hallam, Anthony. 1976. Stratigraphic distribution and ecology of European Jurassic Bivalves. Lethaia 9:245–259.
- Hallam, Anthony. 1977. Jurassic bivalve biogeography. Paleobiology 3:58–73.
- Hallam, Anthony. 1981. The end-Triassic bivalve extinction event. Palaeogeography, Palaeoclimatology, Palaeoecology 35:1–44.
- Hallam, Anthony. 1987. Radiations and extinctions in relation to environmental change in the marine Lower Jurassic of northwest Europe. Paleobiology 13(2):152–168.
- Hallam, Anthony. 1990. Bivalvia. Les Cahiers de l'Université Catholique de Lyon, Série Sciences 3:55–60.
- Hallam, Anthony 1996. Recovery of the marine fauna in Europe after the end-Triassic and early Toarcian mass extinctions. *In* M. B. Hart, ed., Biotic Recovery from Mass Extinction Events. Geological Society, Special Publication 102:231–236.
- Hallam, Anthony. 2002. How catastrophic was the end-Triassic mass extinction? Lethaia 35:147–157.
- Hallam, Anthony, & Zain El Shaarawy. 1982. Salinity reduction of the end-Triassic sea form the Alpine region into northwestern Europe. Lethaia 15(2):105–198.
- Hallam, Anthony, & P. B. Wignall. 1997. Extinctions within and at the close of the Triassic. *In* Mass Extinctions and their Aftermath. Oxford University Press. New York. p. 142–160.

- Hallam, Anthony, & P. B. Wignall. 2000. Facies changes across the Triassic-Jurassic boundary in Nevada, United States. Journal of the Geological Society (London) 157(1):49–54.
- Hallam, Anthony, P. B. Wignall, Yin Jiarun, & J. B. Riding. 2000. An investigation into possible facies changes across the Triassic - Jurassic boundary in southern Tibet. Sedimentary Geology 137:101–106.
- Hansson, H. G. 1998. NEAT (North East Atlantic Taxa): Scandinavian marine Mollusca Check-List. Internet Edition, 128 p., http://www.tmbl. gu.se/libdb/taxon/neat_pdf/NEAT*Mollusca.pdf. Checked August 2012.
- Harper, E. M., Hermann Dreyer, & Gerhard Steiner. 2005. Reconstructing the Anomalodesmata (Mollusca: Bivalvia): morphology and molecules. *In* Rüdiger Bieler FLS, ed., Bivalvia—A look at the Branches. Zoological Journal of the Linnean Society 148:395–420.
- Harper, E. M., G. T. W. Forsythe, & T. J. Palmer. 1998. Taphonomy and the Mesozoic marine revolution; preservation state masks the importance of boring predators. PALAIOS 13(4):352–360.
- Harper, E. M., & T. J. Palmer. 1993. Middle Jurassic cemented Pectinids and the missing rigth valves of *Eopecten*. Journal of Molluscan Studies 59:63–72.
- Harper, E. M., T. J. Palmer, & J. D. Hudson. 2002. The Middle Jurassic bivalve "*Cuspidaria*" *ibbetsoni*; a corbulid not a septibranch. Palaeontology 45(4):759–769.
- Harries, P. J., & J. S. Crampton. 1998. The Inoceramids. American Paleontologist 6(4):2–6.
- Harries, P. J., E. G. Kauffman, J. S. Crampton (Redacteurs), P. Bengtson, S. Cech, J. A. Crame, A. V. Dhondt, G. Ernst, H. Hilbrecht, G. R. López Mortimore, K. A. Tröger, I. Walaszcyk, & C. J. Wood. 1996. Lower Turonian Euramerican Inoceramidae: A morphologic, taxonomic, and biostratigraphic overview. A report from the first Workshop on Early Turonian Inoceramids (Oct. 5–8, 1992) in Hamburg, Germany. Mitteilungen aus dem Geologisch - Paläontologischen Museum der Universität Hamburg 77:641–671.
- Harries, P. J., & C. T. S. Little. 1999. The early Toarcian (Early Jurassic) and the Cenomanian-Turonian (Late Cretaceous) mass extinctions: similarities and contrasts. Palaeogeography, Palaeoclimatology, Palaeoecology 154:39–66.
- Hauer, Franz. 1850. Über die von Herrn Bergrath W. Fuchs in der Venetianer Alpen gesammelten Fossilien. Denkschriften der Kaiserlichen Akademie der Wissenschaften in Wien, Mathematisch-naturwissenschaftlichen Klasse 2(1):109–126, pl. 18–21.
- Hauer, Franz. 1853. Über die Gliederung der Trias-Lias und Juragebilde in den nordöstlichen Alpen. Jahrbuch, Geologisches Reichsanstalt 4:715–784.
- Hauer, Franz. 1857. Ein Beitrag zur Kenntnis der Fauna der Raibler Schichten. Sitzungsberichte der Kaiserlichen Akademie der Wissenschaften, Wien, mathematisch-naturwissenschaftliche Classe 24:537–566, pl. 1–6.
- Hautmann, Michael. 2001a. Taxonomy and phylogeny of cementing Triassic bivalves (Families Prospondylidae, Plicatulidae, Dimyidae and Ostreidae). Palaeontology 44(2):339–373.
- Hautmann, Michael. 2001b. Die Muschelfauna der Nayband Formation (Obertrias, Nor - Rhät) des östlichen Zentraliran. Beringeria 29:3–181.
- Hautmann, Michael. 2003. Notes on *Gruenewaldia* Wöhrmann, 1889 (Bivalvia, Trigonioida, Triassic). Neues Jahrbuch für Geologie und Paläontologie Monatshefte 3:145–152.
- Hautmann, Michael. 2004. Early Mesozoic evolution of alivincular bivalve ligaments and its implications for the timing of the "Mesozoic marine revolution." Lethaia 37:165–172.
- Hautmann, Michael. 2006a. Shell mineralogical trends in epifaunal Mesozoic bivalves and their relationship to seawater chemistry and atmospheric carbon dioxide concentration. Facies 52:417–433.
- Hautmann, Michael. 2006b. Shell morphology and phylogenetic origin of oysters. Palaeogeography, Palaeoclimatology, Palaeoecology 240: 668-671.

- Hautmann, Michael. 2008. Taxonomy and Phylogeny of the Triassic bivalve families Mysidiellidae Cox, 1964 and Healeyidae new family. Journal of Paleontology 82(3):555–564.
- Hautmann, Michael. 2010. The first scallop. Paläontologische Zeitschrift 84(2):317–322.
- Hautmann, Michael, Babak Aghababalou, & Leo Krystyn. 2011. An unusual Late Triassic nuculid bivalve with divaricate shell ornamentation and the evolutionary history of divaricate ribs in Triassic bivalves. Journal of Paleontology 85(1):22-28.
- Hautmann, Michael, Hugo Bucher, Thomas Brühwiler, Nicolas Goudemand, Andrzej Kaim, & Alexander Nützel. 2011. An unusually diverse mollusc fauna from the earliest Triassic of South China and its implications for benthic recovery after the end-Permian biotic crisis. Geobios 44:71–85.
- Hautmann, Michael, & M. Golej. 2004. Terquemia (Dentiterquemia) eudesdeslongchampsi new subgenus and species, an interesting cementing bivalve from the Lower Jurassic of the Western Carpathians (Slovakia). Journal of Paleontology 78(6):1086–1090.
- Hautmann, Michael, & Alexander Nützel. 2005. First record of a heterodont bivalve (Mollusca) from the Early Triassic: palaeoecological significance and implications for the "Lazarus problem." Palaeontology 48(6):1131-1138.
- Hautmann, Michael, Frank Stiller, Huawei Cai, & Sha Jingeng. 2005. An unusual post-extinction bivalve fauna from the Triassic-Jurassic transition near Germig, southern Tibet–a preliminary report. *In* Sha J. & Wang I., eds., Abstracts of International Symposium on the Jurassic boundary events. Nanjing. p. 19–21.
- Hayami, Itaru. 1957a. Liassic *Gervillia & Isognomon* in Japan (Studies on the Liassic pelecypods in Japan, 2). Japanese Journal of Geology and Geography 28(1-3):95–106.
- Hayami, Itaru. 1957b. Liassic *Chlamys*, "*Camptonectes*" and other pectinids from the Kuruma group in central Japan (Studies on the Liassic pelecypods in Japan, 5). Transactions and Proceedings of the Palaeontological Society of Japan (new series) 28:119–127.
- Hayami, Itaru. 1957c. Radulonectites, a new Pectinid genus, from the Liassic Kuruma group in Central Japan. Transactions and Proceedings of the Palaeontological Society of Japan (new series) 27:89–93.
- Hayami, Itaru. 1958a. Liassic Volsella, Mytilus and some other dysodont species in Japan (Studies on the Liassic pelecypods in Japan, 6). Transactions and Proceedings of the Palaeontological Society of Japan (new series) 29:155–165.
- Hayami, Itaru. 1958b. A review of the so-called Liassic "cyrenoids" in Japan (Studies on the Liassic pelecypods in Japan, 7). Recent Progress of Natural Sciences in Japan 29(1–3):11–27.
- Hayami, Itaru. 1958c. Supplementary descriptions of the Liassic pelecypods from the Kuruma and Shizukawa groups in Japan (Studies on the Liassic pelecypods in Japan, 9). Transactions and Proceedings of the Palaeontological Society of Japan (new series) 30:193–199.
- Hayami, Itaru. 1958d. Some Hettangian Pelecypods from the "*Trigonia*sandstone" of Shizukawa Group in Northeast Japan. Japanese Journal of Geology and Geography 29(1–3):99–110.
- Hayami, Itaru. 1959. Pelecypods of the Mizunuma Jurassic in Miyagi prefecture with some stratigraphical remarks (Studies on the Liassic Pelecypods in Japan. 10). Transactions and Proceedings of the Palaeontological Society of Japan (new series) 34:66–77.
- Hayami, Itaru. 1960. Jurassic inoceramids in Japan. Journal of the Faculty of Science, University of Tokyo, Section 2: Geology, Mineralogy, Geography, Geophysics 12(2):277–328.
- Hayami, Itaru. 1961. Pelecypods from the Liassic Yamaoku Formation in west Japan. (Studies on the Liassic pelecypods in Japan, 11). Transactions and Proceedings of the Palaeontological Society of Japan (new series) 43:113–116.
- Hayami, Itaru. 1964. Some Lower Jurassic Pelecypods from South Viet-Nam, collected by Dr. H. Fontaine. Contributions to the Geology and

Palaeontology of Southeast Asia, XV. Japanese Journal of Geology and Geography 35(2–4):163–174.

- Hayami, Itaru. 1965. Lower Cretaceous marine pelecypods of Japan, part I. Memoirs of the Faculty of Science, Kyushu University (series D, Geology) 15:221–349.
- Hayami, Itaru. 1969a. Notes on Mesozoic 'planktonic' bivalves. Journal of the Geological Society of Japan 75(7):375–385.
- Hayami, Itaru. 1969b. The occurrence of *Tutcheria* from the Lower Jurassic of West Japan. Transactions and Proceedings of the Palaeontological Society of Japan, New Series 73:26–31.
- Hayami, Itaru. 1975. A systematic survey of the Mesozoic Bivalvia from Japan. The University Museum, The University of Tokyo, Bulletin 10:249 p.
- Hayami, Itaru. 1985. Range chart of selected bivalve species of Japanese Jurassic. Circum-Pacific Jurassic Field Conference, Tsukuba III:44–50
- Hayami, Itaru. 1988. A Tethyan bivalve, *Posidonotis dainellii*, from the Lower Jurassic of west Japan. Transactions and Proceedings of the Palaeontological Society of Japan (new series) 151:564–569.
- Hayami, Itaru, & Tomoki Kase. 1977. A systematic survey of the Paleozoic and Mesozoic gastropoda and Paleozoic bivalvia from Japan. The University Museum, The University of Tokyo Bulletin 13:154 p.
- Hayami, Itaru, Shiro Maeda, & Carlos Ruiz-Fuller. 1977. Some Late Triassic Bivalvia and Gastropoda from the Domeyko Range of North Chile. Transactions and Proceedings of the Palaeontological Society of Japan (new series) 108:202–221.
- Hayasaka, Ichiro. 1967. Some Permian fossils from Southern Kitakami. IX Two Pelecypod Genera: *Allorisma* and *Myoconcha*. Proceedings of the Japan Academy (series B: Physical and Biological Sciences) 43:517–521.
- He Weihong, Feng Qinglai, E. A. Weldon, Gu Songzhu, Meng Youyan, Zhang Fan, & Wu Shunbao. 2007. A late Permian to Early Triassic bivalve fauna from the Dongpan section, Southern Guangxi, South China. Journal of Paleontology 81(5):1009–1019.
- He Weihong, Shi G. R., Feng Qinglai, & Peng Yuanqiao. 2007. Discovery of late Changhsingian (latest Permian) brachiopod Attenuatella species from South China. Alcheringa 31:271–284.
- Healey, Maud. 1908. The fauna of the Napeng Beds or the Rhaetic Beds of Upper Burma. Memoirs of the Geological Survey of India, Palaeontologia Indica (new series) 2(4):89 p.
- Hedinger, A. S. 1993. Upper Jurassic (Oxfordian-Volgian) foraminifera from the Husky Formation, Aklavik Range, District of MacKenzie, Northwest Territories. Geological Survey of Canada Bulletin 439:173 p.
- Hedley, Charles. 1918. A check-list of the marine fauna of New South Wales. Part 1, Mollusca. Journal of the Royal Society of New South Wales 51(supplement):M1–M120.
- Heilprin, Angelo. 1887. The Geographical and Geological distribution of animals. D. Appleton & Co. New York. 435 p.
- Heinberg, Claus. 1978. Bivalves from the White Chalk (Maastrichtian) of Denmark; II Arcoida. Bulletin of the Geological Society of Denmark 27(3–4):105–116.
- Heinberg, Claus. 1999. Lower Danian bivalves, Stevns Klint, Denmark: continuity across the K/T boundary. Palaeogeography, Palaeoclimatology, Palaeoecology 154(1–2):87–106.
- Hermann, J. 1781. Brief über einige Petrefacten. Der Naturforscher, Halle 15:115–134.
- Hertlein, L. G. 1952. *Newaagia*, new name for *Philippiella* Waagen. Journal of Paleontology 26(2):275.
- Hillebrandt, Axel von. 1981. Kontinentalferschiebung und die paläozoogeographischen Beziehungen des südamerikanischen Lias. Geologische Rundschau 70(2):570–582.
- Hillebrandt, Axel von. 1990. The Triassic/Jurassic boundary in Northern Chile. Les Cahiers de l'Université Catholique de Lyon (série Sciences) 3:27–53.
- Hind, Wheelton. 1896–1905. A monograph of the British Carboniferous Lamellibranchiata. Palaeontographical Society Monographs. Palaeonto-

graphical Society, London. Vol. 1: p. 1–80, pl. 1–2 [1896]; p. 81–208, pl. 3–15 [1897]; p. 209–276, pl. 16–25 [1898]; p. 277–360, pl. 26–39 [1899]; p. 361–476, pl. 40–54 [1900]; title page and index [1901]; vol. 2: p. 1–222 [1901?–1905].

- Hoare, R. D. 1993. Mississipian (Chesterian) Bivalves from the Pennsylvanian Stratotype Area in West Virginia and Virginia. Journal of Paleontology 67(3):374–396.
- Hoare, R. D., M. J. Heaney III, & R. H. Mapes. 1989. Bivalves (Mollusca) from the Imo Formation (Mississipian, Chesterian) of north-central Arkansas. Journal of Paleontology 63(5):582–603.
- Hodges, Peter. 1991. The relationship of the Mesozoic bivalve *Atreta* to the Dimyidae. Palaeontology 34(4):963–970.
- Hodges, Peter. 2000. The Early Jurassic Bivalvia from the Hettangian and Lower Sinemurian of south-west Britain; Part I. Monograph of the Palaeontographical Society of London, United Kingdom 154(614):1–64.
- Hoernes, R. 1880. Materialen zu einer Monographie der gattung Megalodus mit besonderer Berücksichtigung mesozoischer Formen. Denkschriften Akademie Wissenschaften Wien 42:91–126, pl. 1–7.
- Hölder, Helmut. 1990. Über die Muschelgattung *Placunopsis* (Pectinacea, Placunopsidae) in Trias und Jura. Stuttgarter Beiträge zur Naturkunde Series B (Geologie und Palaeontologie) 165:1–63.
- Hölder, Helmut. 1995. On the genus *Tutcheria* Cox, 1946 (Lamellibranchiata, Heterodonta) in the German Liassic. Neues Jahrbuch für Geologie und Paläontologie Abhandlungen 198(1–2):69–83.
- Holland, S. M. 1995. The stratigraphic distribution of fossils. Paleobiology 21(1):92–109.
- Holzapfel, Susanne. 1998. Palökologie benthischer Faunen-gemeinschaften und Taxonomie der Bivalven im Jura von Südtunesien. Beringeria 22:199 p.
- Hopkin, E. K., & C. A. McRoberts. 2005. A new Middle Triassic flat clam (Pterioida: Halobiidae) from the Middle Anisian of North-Central Nevada, United States. Journal of Paleontology 79(4):796–800.
- Hsu Te-you. 1936–1937. Contribution to the marine Lower Triassic Fauna of Southern China. Bulletin of the Geological Society of China 16:303–324. In Chinese.
- Hsu T. Y. & Chen K. 1943. Revision of the Chingyen Triassic fauna from Kweichou. Bulletin of the Geological Society of China 23(3–4):129–138.
- Hu Xiumian, Luba Jansa, & Wang Chengshan. 2008. Upper Jurassic–Lower Cretaceous stratigraphy in south-eastern Tibet: a comparison with the western Himalayas. Cretaceous Research 29(2):301–315.
- Huang Kainian, & N. D. Opdyke. 2000. Magnetostratigraphic investigations of the Middle Triassic Badong Formation in South China. Geophysical Journal International 142:74–82.
- Huang Zhi-hui. 1986. Discovery of early Jurassic bivalves from Shenzen area. Acta Palaeontologica Sinica 25(5):577–584. In Chinese with English abstract.
- Hudson, J. D., R. G. Clements, J. B. Riding, M. I. Wakefield, & Willian Walton. 1995. Jurassic paleosalinities and brackish-water communities - a case study. PALAIOS 10:392–407.
- Hudson, Neville. 2003. Stratigraphy and correlation of the Ururoan and Temaikan Stage (Lower–Middle Jurassic, ?Sinemurian–Callovian) sequences, New Zealand. Journal of the Royal Society of New Zealand 33(1):109–147.
- Hudson, R. G. S., & R. P. S. Jefferies. 1961. Upper Triassic Brachiopods and lamellibranchs from the Oman peninsula, Arabia. Palaeontology 4(1):1–41.
- Hyatt, Alpheus. 1894. Trias and Jura in the western States. Geological Society of America, Bulletin 5:395–434.
- Ichikawa, Koichiro. 1949. *Trigonucula* (nov.) and other taxodont pelecypods from the Upper Triassic of the Sakawa basin in Shikoku, Japan. Japanese Journal of Geology and Geography 21(1–4):267–272, pl. 10.
- Ichikawa, Koichiro. 1950. Sakawanella, new genus, and some other pelecypods from the upper Triassic Kochigatani group in the Sakawa Basin,

Shikoku, Japan. Journal of the Faculty of Science, University of Tokyo, Section 2: Geology, Mineralogy, Geography, Geophysics 7(3–5):245–256.

- Ichikawa, Koichiro. 1954. Late Triassic pelecypods from the Kochigatani group in the Sakuradani and Kito areas, Tokushima Prefecture, Shikoku, Japan, part 1. Journal of the Institute of Polytechnics, Osaka City University [G] 1:35–57, pl. 1–2.
- Ichikawa, Koichiro. 1958. Zur Taxonomie und Phylogenie der Triadischen "Pteriidae" (Lamellibranch.) mit besonderer Berücksichtigung der Gattungen *Claraia, Eumorphotis, Oxytoma* und *Monotis*. Palaeontographica A 111:131–212.
- Ichikawa, Koichiro, & Yin E. H. 1966. Discovery of Early Triassic Bivalves from Kelantan, Malaya. Journal of Geosciences, Osaka City University 9:101–108.
- ICZN (International Commission on Zoological Nomenclature). 1954. Opinion no. 292. Opinion, International Commission on Zoological Nomenclature 8:117–130.
- ICZN (International Commission on Zoological Nomenclature). 1999. International Code of Zoological Nomenclature. Fourth edition. International Trust for Zoological Nomenclature. London. 306 p.
- Ihering, H. von. 1903. Notes sur quelques Mollusques fossils du Chile. Revista Chilena de Historia Natural VII(2):120-127.
- Imhoff, J. F., Heiko Sahling, Jörg Süling, & Thomas Kath. 2003. 16S rDNA-based phylogeny of sulfphur-oxidising bacterial endosymbionts in marine bivalves from cold-seep habitats. Marine Ecology Progress Series 249:39–51.
- Imlay, R. W. 1967. The Mesozoic Pelecypods Otapiria Marwick and Lupherella Imlay, new genus, in the United States. Contributions to Paleontology, US Geological Survey Professional Paper 573–B:11 p.
- Iredale, Tom. 1931. Australian molluscan notes. No. 1. Records of the Australian Museum 18:201–235.
- Ishikawa, Hideo, Shiro Maeda, Tetsuya Kawabe, & César Rangel. 1983. Some Jurassic Trigoniids from Peru. The Bulletin of the Faculty of Education, Chiba University 32(2):37–44.
- Ivimey-Cook, H. C., Peter Hodges, Andrew Swift, & J. D. Radley. 1999. Bivalves. *In* Andrew Swift & D. M. Martill, eds., Fossils of the Rhaetian Penarth Group. The Palaeontological Association Field Guides to Fossils, London 9(8):83–127.
- Jablonski, David, & D. J. Bottjer. 1990. Onshore-offshore trends in marine invertebrate evolution. In R. M. Ross & W. D. Allmon, eds., Causes of Evolution: A Paleontological Perspective. University of Chicago Press. Chicago. p. 21–75.
- Jablonski, David, Kaustuv Roy, J. W. Valentine, R. M. Price, & P. S. Anderson. 2003. The impact of the pull of the recent on the history of marine diversity. Science 300:133–1135.
- Jagt, W. M., Christian Neumann, & A. S. Schulp. 2007. Bioimmuring Upper Cretaceous and recent oysters: "A view from within." Geologica Belgica 10(1–2):121–126.
- Jaitly, A. K. 1986. Revised morphotaxonomic description of six Middle Jurassic Pholadomyoid clams from Kala Dongar, Pachchham Island, District Kachchh, Gujarat. Indian Minerals 40(1):39–46.
- Jaitly, A. K. 1988. Some Middle Jurassic clams from Kala Dongar, Pachchham Island, Kachchh, Gujarat. Indian Minerals 42(2):117–125.
- Jaitly, A. K., F. T. Fürsich, & Michael Heinze. 1995. Contributions to the Jurassic of Kaschchh, Western India IV. The bivalve fauna. Subclasses Palaeotaxodonta, Pteriomorpha, and Isofilibrachia. Beringeria 16:147–257.
- Jaworski, Erich. 1914. Beiträge zur Kenntnis der Lias-Volen Südamerikas und der Stammesgeschichte der Gattung Vola. Palaeontologische Zeitschrift 1:273–320.
- Jaworski, Erich. 1915. Beiträge zur Kenntnis des Jura in Süd-Amerika. Teil II. Spezieller, paläontologischer Teil. Neues Jahrbuch für Mineralogie, Geologie und Paläontologie 40(2):364–456, pl. 5–8.
- Jaworski, Erich. 1922. Die marine Trias in Süd-Amerika. Neues Jahrbuch für Mineralogie, Geologie, und Paläontologie 47(Supplement):93–200.

- Jeannet, Alphonse. 1912–1913. Monographie géologique des Tours d'Aï et des régions avoisinantes (Préalpes vaudoises), 2 vol. Beiträge zur Geologischen Karte der Schweiz (new series) 34(1-2):465 p. (*)
- Jeannet, Alphonse. 1951. Stratigraphie und Paläontologie des oolithischen Eisenerzlagers von Herznach und seiner Umgebund. Beiträge zur Geologie, Schweizerisches Geotechnisches Serie 13(5):240 p.
- Jefferies, R. P. S., & P. Minton. 1965. The mode of life of two Jurassic species of "*Posidonia*" (Bivalvia). Palaeontology 8(1):156–185.
- Jelen, Bogomir. 1988. Karnijska školj na favna na Lesnem brdu in njen paleobiološki pomen [The Karnian bivalves (Mollusca) from Lesno brdo, Slovenia, NW Yugoslavia and their paleobiological significance]. Geologija 31:11–127. In Slovak with English abstract.
- Jeletzky, J. A., & T. P. Poulton. 1987. A new genus and subgenus and two new species of latest Jurassic oxytomid bivalves from Arctic Canada. Canadian Journal of Earth Sciences 24(4):711–722.
- Jiang Baoyu, Sha Jingeng, & Pan P. 2008. Some Early Jurassic nonmarine bivalves from the Southern Junggar Basin, NW China. In Abstracts Volume of Fifth International Symposium of IGCP506: Marine and non marine Jurassic: Global correlation and major geological events. March, Tunisia. p. 71.
- Jiménez de Cisneros, Daniel. 1923. La fauna de los estratos de "Pygope Aspasia" Menegh. del Liásico medio del Rincón de Egea en el NW de la provincia de Murcia. Trabajos del Museo Nacional de Ciencias Naturales de Madrid (serie Geología) 30:1–55.
- Johnson, A. L. A. 1984. The palaeobiology of the bivalve families Pectinidae and Propeamussiidae in the Jurassic of Europe. Zitteliana 11:3–235.
- Johnson, A. L. A., & M. J. Simms. 1989. The timing and cause of late Triassic marine invertebrate extinctions: evidence from scallops and crinoids. *In* E. Donovan, ed., Mass extinction: processes and evidence. Belhaven Press. London. p. 174–194.
- Jones, D. S., & D. K. Jacobs. 1992. Photosymbiosis in *Clinocardium nuttalli*: Implications for tests of photosymbiosis in fossil mollusks. PALAIOS 7:86–95.
- Jurkovsek, Bogdan. 1978. Biostratigrafija karnijske stopnje v okolici Mežice [Biostratigraphy of the Carnian beds in the Mežica area]. Geologija 21:173–208. In Slovak with English abstract.
- Jurkovsek, Bogdan. 1982a. Halobiidae in Monotidae v Sloveniji in Njihov biostratigrafski Pomen [Halobiidae and Monotidae of Slovenia and their biostratigraphical significance]. Zbornik Radova, X jubilarni kongres, Knjiga 1. Budva. p. 147–154. In Slovak.
- Jurkovsek, Bogdan. 1982b. Zgornjetriasne Monotidae v Sloveniji [Upper Triassic Monotidae in Slovenia]. Mining and Metallurgy Quaterly 29(2–3):122–131. In Slovak with English abstract.
- Jurkovsek, Bogdan. 1983. Fassanske plasti z daonelami v Sloveniji [Fassanian beds with daonellas in Slovenia]. Geologija 26:29–70. In Slovak with English abstract.
- Jurkovsek, Bogdan. 1984. Langobardske plasti z daonelami in pozidonijami v Sloveniji [Langobardian beds with daonellas and posidonias in Slovenia]. Geologija 27:41–95. In Slovak with English abstract.
- Jurkovsek, Bogdan. 1993. Karnijske plasti s koljkami južno od Rovt [Carnian beds with pelecypods south of Rovte]. Geologija 36:195–205. In Slovak with English abstract.
- Jurkovsek, Bogdan, & Bogomir Jelen. 1990. Fosilne školjke iz karnijskih plasti pri Orlah [Fossil lamellibranchs from the Carnian beds at Orle]. Geologija 31/32:129–132. In Slovak with English abstract.
- Jurkovsek, Bogdan, & Tea Kolar-Jurkovsek. 1986. A Upper Triassic (Carnian) fish skeleton (family Birgeriidae) from Slovenia, NW Yugoslavia. Neues Jahrbuch für Geologie und Paläontologie Monatshefte 8:475–478.
- Kaim, Andrzej. 1997. Brachiopod-bivalve assemblages of the Middle Triassic Terebratula Beds, Upper Silesia, Poland. Acta Palaeontologica Polonica 42(2):333–359.

- Kaim, Andrzej. 2001. Faunal dynamics of juvenile gastropods and associated organisms across the Valanginian transgression-regression cycle in central Poland. Cretaceous Research 22:333–351.
- Kambe, Nobukazu. 1957. On the myophorians from the Miharaiyama group in Hyogo Prefecture. Reports of the Geological Survey of Japan 173:1–19.
- Kashiyama, Yuichiro, & Tatsuo Oji. 2004. Low-diversity shallow marine benthic fauna from the Smithian of northeast Japan: paleoecologic and paleobiogeographic implications. Paleontological Research 8(3):199–218.
- Kasum-Zadeh, A. A. 2003. Reviziyi systematika Mezozoyskikh predstaviteley otryada Pectinida Azerbaydzhaia. [= Advance in research of Mesozoic Bivalve Mollusks in Azerbaijan (Order Pectinoida: Revision and Systematics)]. "El-ALliance." Baku. 112 p. Translation by Rosanne D'Aprile Johnson, VIARC, Smithsonian Institution.
- Kauffman, E. G. 1969. Form, Function, and Evolution. In R. C. Moore, ed., Treatise on Invertebrate Paleontology, Part N, Mollusca 6, Bivalvia, vol. 1. The Geological Society of America & The University of Kansas. Boulder & Lawrence. p. 129–205.
- Kauffman, E. G., & H. A. Leanza, 2004. A remarkable new genus of Mytilidae (Bivalvia) from the Lower Cretaceous of Southwestern Gondwanaland. Journal of Paleontology 78(6):1187–1191.
- Kauffman, E. G., & J. D. Powell. 1977. Part 2: Paleontology. In E. G. Kauffman, D. E. Hattin, & J. D. Powell, eds., Stratigraphic, paleontologic, and paleo-environmental analysis of the Upper Cretaceous rocks of Cimarron County, northwestern Oklahoma. Geological Society of America Memoir 149:47–150.
- Kauffman, E. G., & Bruce Runnegar. 1975. Atomodesma (Bivalvia), and Permian species of the United States. Journal of Paleontology 49(1):23–41.
- Kay, E. Alison. 1965. The Reverend John Lightfoot, Daniel Solander and the Portland Catalogue. Nautilus 79(1):10–19.
- Kaye, W. J. 1919. New Species and Genera of Nymphalidce, Syntomidce, and Sphingidce in the Joicey Collection. The Annals and Magazine of Natural History 4(20):84–93.
- Kear, David. 1961. Ururoan beds of the Hetherington inlier, West Auckland. New Zealand of Geology and Geophysics 4:231–238.
- Kegel, W. & M.T. da Costa. 1951. Espécies neopaleozóicas do Brasil, da familia Aviculopectinidae, ornamentadas com costelas fasciculadas. Boletim do Departamento Nacional de Producão Mineral, Divisão de Geologia e Mineralogia 137:1–48.
- Kelly, S. R. A. 1980. *Hiatella*-a Jurassic bivalve squatter? Palaeontology 23(4):769–781.
- Kelly, S. R. A. 1984. Bivalvia of the Spilsby Sandstone and Sandringham Sands (late Jurassic-early Cretaceous) of eastern England, Part I. Monograph of the Palaeontographical Society of London 137(566):94 p.
- Kelly, S. R. A. 1988. Cretaceous wood-boring bivalves from western Antarctica with a review of the Mesozoic Pholadidae. Palaeontology 31(2):341–372.
- Kelly, S. R. A. 1995a. New Trigonioid bivalves from the early Jurassic to earliest Cretaceous of the Antarctic Peninsula region: systematics and austral paleobiogeography. Journal of Paleontology 69(1):66–84.
- Kelly, S. R. A. 1995b. New trigonioid bivalves from the Albian (Early Cretaceous) of Alexander Island, Antarctic Peninsula: systematics, paleoecology, and austral Cretaceous paleobiogeography. Journal of Paleontology 69(2):264–279.
- Kelly, S. R. A., A. V. Dhondt, & V. A. Zakharov. 1984. *Boreionectes* Zakharov 1965 (Bivalvia: Pectinidae) - a synonym of *Maclearnia* (Crickmay 1930). Journal of Paleontology 58(1):109–114.
- Kenig, F., J. D. Hudson, J. S. Sinninghe Camsté, & B. N. Popp. 2004. Intermittent euxinia: Reconciliation of a Jurassic black shale with its biofacies. Geology 32(5):421–424.
- Kennedy, W. J., N. J. Morris, & J. D. Taylor. 1970. The shell structure, mineralogy and relationships of the Chamacea (Bivalvia). Palaeontology 13(3):379–413.

- Kerr, M. B. 1875. Outlines of the geology of North Carolina. Report of the Geological Survey of North Carolina (Raleigh) 1:1–141.
- King, William. 1844. On a new genus of Palaeozoic shells. Annals and Magazine of Natural History 14(92):313–317.
- King, William. 1848. A Catalogue of the Organic Remains of the Permian Rocks of Northumberland and Durham. Published by the author. Newcastle upon Tyne. p. iv + 16.
- King, William. 1850. A Monograph of the Permian Fossils of England. Palaeontographical Society Monograph, London. 1–258 p., 28 pl.
- Kiparisova, L. D. 1938. The Lower Triassic Pelecypoda of the Ussuriland. Travaux de l'Institut Géologique de l'Académie des Sciences de l'Russie 7:197–311.
- Kiparisova, L. D. 1954. Polevoj Atlas kharakternykh kompleksov fauny i flory triasovykh otlozhenij Primorskogo Kraya [A Field manual for the identification of Triassic fossils (Mainly Mollusks) and plant remains of the Maritime Territory, USSR]. Vse-soyuznyi Nauchno-Issledovatelskii Geologi-cheskii Institut, Trudy. Moscow. 52 p. In Russian.
- Kiparisova, L. D. 1972. Paleontologischeskoe Obosnovanie Stratigrafii Triasovykh Otlozhenij Primorskoto Kraya. Chast' 2. Pozdnetriasovye Dvustvorchatye Mollyuski i Obshchaya Stratigrafia [Palaeontological basis of Triassic stratigraphy of the Primorye region. 2. Late Triassic molluscs and general stratigraphy]. Trudy Vsesoyuznyi Nauchno-issledovatelyskij Geologicheskij Institut (VSEGEI), Novaya Seriya [Proceedings of VSEGEI, New Series] 181:246 p. In Russian.
- Kiparisova, L. D., Y. M. Bychkov, & I. V. Polubotko. 1966. Pozdnetriasovyye dvustvorchatyye molluyuski Severovostoka SSSR [Late Triassic Bivalvie Mollusca from Northeastern USSR]. Ministry of Geology of the USSR, All-Union Geological Research Institute. Magadan. 230 p. In Russian.
- Kittl, Ernst. 1904. Geologie der Umgebung von Sarajevo. Jahrbuch der kaiserlich-königlichen Geologischen Reichsanstalt, Wien 53:515–748, 3 pl.
- Kittl, Ernst. 1907. Die Triasfossilien vom Heureka Sund. Report of the Second Norwegian Arctic Expedition in the "Fram" 1898–1902. Kristiania 7:1–44.
- Kittl, Ernst. 1912. Materialien zu einer Monographie der Halobiidae und Monotidae der Trias. Resultate der Wissenschaftlichen Erforschung des Balantonsees, I Band, I teil. Paläontologie. 229 p.
- Kleemann, Karl. 1990. Evolution of the chemically boring Mytilidae (Bivalvia). *In* B. Morton, ed., The Bivalvia. Proceedings of a Memorial Symposium in Honour of Sir Charles Maurice Yonge, Edinburgh, 1986. Hong Kong University Press. Hong Kong. p. 111–124.
- Kleemann, Karl. 1994. Mytilid bivalve *Lithophaga* in Upper Triassic coral *Pamiroseris* from Zlambach beds compared with Cretaceous *Lithophaga alpina*. Facies 30:151–154.
- Kleine, R. 1916. Neue Taphroderini (Brenthidae, Col.). Entomologische Mitteilungen 5(1/4):1–92.
- Klets, A. G., I. V. Budnikov, R. V. Kutygin, A. S. Biakov, & V. S. Grinenko. 2006. The Permian of the Verkhoyansk-Okhotsk region, northeastern Rusia. Journal of Asian Earth Sciences 26:258–268.
- Klets, T. V. 2005. Palaeobiogeographic zoning of Triassic seas of Northeastern Asia based on Conodontophoridae. Albertiana 32:40–56.
- Klets, T. V. 2006. Diversity and stages in evolution of Conodontophorids and Bivalve Mollusks in Triassic seas of Siberia and the Far East. Stratigraphy and Geological Correlation 14(2):174–184.
- von Klipstein, August. 1843–1845. Beiträge zur geologischen Kenntniss der östlichen Alpen. Mitteilungen aus dem Gebiete der Geologie und Paläontologie, vol. 1. Georg Friedrich Heyer. Giessen. 312 p., 20 pl.
- Klug, Christian, Hans Hagdorn, & Michael Montenari. 2005. Phosphatized soft-tissue in Triassic bivalves. Palaeontology 48(4):833–852.
- Kobayashi, T. 1931. Notes on a new occurrence of Ladino-Carnic limestone at Sambosan, Tosa Province, Japan. Japanese Journal of Geology and Geography 8:251–258.

- Kobayashi, Teiichi. 1954. Studies on the Triassic trigonians in Japan, Part 1. Preliminary notes. Japanese Journal of Geology and Geography 25(1–2):61–80.
- Kobayashi, Teiichi. 1957. Some Trigonians from the Hida Plateau Region, Central Japan. Japanese Journal of Geology and Geography 28(1-3):43–46.
- Kobayashi, Teiichi, & Koichiro Ichikawa. 1949a. *Myophoria* and other Upper Triassic pelecypods from the Sakawa Basin in Shikoku, Japan. Japanese Journal of Geology and Geography Transactions 21(1–4):177–192.
- Kobayashi, Teiichi, & Koichiro Ichikawa. 1949b. *Tosapecten* nov. and other Upper Triassic Pectinidae from the Sakawa Basin, in Shikoku, Japan. Japanese Journal of Geology and Geography Transactions 21(1–4):163–176.
- Kobayashi, Teiichi, & Koichiro Ichikawa. 1950. Late Triassic Mytilus, Volsella, Pleurophorus and Myoconcha from the Sakawa Basin in Shikoku, Japan. Journal of the Faculty of Science, University of Tokyo, Section 2: Geology, Mineralogy, Geography, Geophysics 7(3–5):207–215.
- Kobayashi, Teiichi, & Koichiro Ichikawa. 1952. The Triassic fauna of the Heki formation in the Province of Tamba (Kyoto Prefecture), Japan. Japanese Journal of Geology and Geography 22:55–84, pl. 1–3.
- Kobayashi, Teiichi, & Yoshio Kaseno. 1947. A new Liassic species of *Trigonia* s. str. (i. e. *Lyriodon*) from the Kitakami mountains, Nippon. Japanese Journal of Geology and Geography 20(2–4):41–43.
- Kobayashi, Teiichi, & Masaru Katayama. 1938. Further evidence as to the chronological determination of so-called Rhaeto-Liassic floras with a description of *Minetrigonia*, a new subgenus of *Trigonia*. Proceedings of the Imperial Academy, Tokyo 14(5):184–189.
- Kobayashi, Teiichi, & Kazuo Mori. 1954a. Studies on the Jurassic trigonians in Japan. Part II. *Prosogyrotrigonia* and the Trigoniinae. Japanese Journal of Geology and Geography 25(3–4):155–175.
- Kobayashi, Teiichi, & Kazuo Mori. 1954b. The Vaugoniinae from the Kitakami Mountains in North Japan. On the Jurassic Trigonians in Japan, Part III. Japanese Journal of Geology and Geography 26(1-2):73-88.
- Kobayashi, Teiichi, & Minoru Tamura. 1968a. *Myophoria* (s. l.) in Malaya with a note on the Triassic Trigoniacea. Contributions to the Geology and Palaeontology of Southeast Asia, LX. Geology and Palaeontology of Southeast Asia 5:88–137.
- Kobayashi, Teiichi, & Minoru Tamura. 1968b. Upper Triassic Pelecypods from Singapore. Contributions to the Geology and Palaeontology of Southeast Asia LXI. Geology and Palaeontology of Southeast Asia 5:138–150.
- Kobayashi, Teiichi, & Minoru Tamura. 1983a. On the Oriental Province of the Tethyan Realm in the Triassic Period. Proceedings of the Japan Academy (series B) 59(7):203–206.
- Kobayashi, Teiichi, & Minoru Tamura. 1983b. The Arcto-Pacific Realm and the Trigoniidae in the Triassic Period. Proceedings of the Japan Academy (series B) 59(7):207–210.
- Kochanová, Maria. 1967. The occurrence of *Variamussium schafhaeutli* (Winkler) in the Norian of West Carpathians (East Slovakia). Rivista Italiana di Paleontologia e Stratigrafia 73(3):713–738.
- Kochanová, Maria. 1985. Middle Triassic Bivalvia from Area of Gemerská Hôrka in Slovak Karst (West Carpathians). Západné Karpaty (série paleontológia) 10:55–73.
- Kochanová, Maria. 1987. Obertriassische Muscheln von Ostré V šky (Slowakischer Karst). Západné Karpaty (série paleontológia) 12:31–46.
- Kochanová, Maria., J. Mello, & Miloš Siblík. 1975. Fosílie wettersteinskych vápencov slovenského krasu (lokalita Silicka) [Fossils from the Wetterstein limestones of the Slovakian karst (Silicka locality)]. Geologické práce 63:55–66. In Slovak.
- Koken, E. 1913. Beiträge zur Kenntnis der Schichten von Heiligenkreuz (Abteital, Südtirol). Abhandlungen der kaiserlich-königlichen Geologischen Reichsanstalt, Wien 16:1–43, 6 pl.

- Kollarova, Vanda, & Maria Kochanová. 1973. Molluskenfauna des Bleskový prame bei Drnava "Nor, Westkarpaten." Verlag der Slowakischen Akademie der Wissenschaften. 213 p. In Slovak with French abstract.
- Komatsu, Toshifumi, Masahiko Akasaki, Chen Jinhua, Cao Meizhen, & Frank Stiller. 2004. Benthic fossil assemblages and depositional facies of the Middle Triassic (Anisian) Yuqing Member of the Qingyan Formation, southern China. Paleontological Research 8(1):43–52.
- Komatsu, Toshifumi, Chen Jinhua, Cao Meizhen, Frank Stiller, & Hajime Naruse. 2004. Middle Triassic (Anisian) diversified bivalves: Depositional environments and bivalve assemblages in the Leidapo Member of the Qingyan Formation, southern China. Palaeogeography, Palaeoclimatology, Palaeoecology 208:207–223.
- Komatsu, Toshifumi, & Maeda Haruyoshi. 2005. Stratigraphy and fossil bivalve assemblages of the mid-Cretaceous Goshoura Group, southwest Japan. Paleontological Research 9(2):119–142.
- Komatsu, Toshifumi, & D. T. Huyen. 2006. Lower Triassic bivalve fossils from the Song Da and An Chau Basins, North Vietnam. Paleontological Research 11(2):135–144.
- Komatsu, Toshifumi, D. T. Huyen, & Chen Jinhua. 2006. Depositional environments and fossil bivalves in the lowermost parts of the Triassic System in North Vietnam and South China. Journal of Geography 115(4):470–483.
- Komatsu, Toshifumi, D. T. Huyen, & Chen Jinhua. 2007. Bivalve assemblages in North Vietnam and South China following the End-Permian crisis. *In* S. G. Lucas & J. A. Spielmann, eds., The Global Triassic. New Mexico Museum of Natural History and Science Bulletin 41:134–136.
- Komatsu, Toshifumi, D. T. Huyen & N. D. Huu. 2010. Radiation of Middle Triassic bivalve: Bivalve assemblages characterized by infaunal and semi-infaunal burrowers in a storm- and wave-dominated shelf, An Chau Basin, North Vietnam. Palaeogeography, Palaeoclimatology, Palaeoecology 291:190–204.
- Komatsu, Toshifumi, Ryo Saito, & F. T. Fürsich. 1993. Mode of occurrence and composition of bivalves of the Middle Jurassic Mitarai Formation, Tetori Group, Japan. Paleontological Research 5(2):121–129.
- Kondo, Yasuo, Takeshi Kozai, Naoki Kikuchi, & Kazuhiro Sugawara. 2006. Ecologic and taxonomic diversification in the Mesozoic brackish-water bivalve faunas in Japan, with emphasis on infaunalization of heterodonts. Gondwana Research 10:316–327.
- de Koninck, L. G. 1841–1844. Description des animaux fossiles, qui se trouvent dans le terrain carbonifère de Belgique. Liège, 650 p., 69 pl. (1–96[1841]; 97–240[1842]; 241–480[1843]; 481–650[1844]).
- de Koninck, L. G. 1877. Recherches sur les Fossiles Paléozoïques de la Nouvelle-Galles du Sud (Australie). Mémoires de la Societé Sciences Naturelles, Liége (serie 2) 6(7):1–373.
- Konstantinov, A. G., E. S. Sobolev, & A. V. Yadernkin. 2007. Detailed biostratigraphy of Triassic deposits in the Lena lower reaches (northern Yakutia). Russian Geology and Geophysics 48:721–736.
- Körner, Karl. 1937. Marine (Cassianer-Raibler) Trias am Nevado de Acrotambo (Nord-Peru). Palaeontographica Abteilung A 86:145–237.
- Koschelkina, Z. W. 1957. Palaeontological basis of the distribution of strata in the marine Jurassic deposits of the Vilyusk syncline and Verkhoyansk depression. *In* Transactions of the International Conference on the Development of the Unified Stratigraphical Schemes in Siberia. Leningrad. p. 27–31.
- Koschelkina, Z. V. 1963. Jurassic stratigraphy and Bivalvia of the Vilyuysk syncline and Verkhoyansk depression. Trudy Severo-vostoka Kompleks. Nauchno-issled. Intituta, Sibirskoe Otdelenie, Akademija Nauk SSSR 5:1–220. In Russian.
- Kotlyar, G. V., Y. D. Zakharov, & I. V. Polubotko. 2004. Upper Changhsingian fauna of the Northwestern Caucasus Mountains, Russia. Journal of Paleontology 78(3):513–527.

- Kozai, Takeshi, Keisuke Ishida, & Yasuo Kondo. 2006. Radiolarians, bivalves and the J/K boundary in the Birafu Formation, southern Kurosegawa Belt, Central Shikoku, SW Japan. Geosciences Journal 10(3):217–235.
- Kristan-Tollmann, Edith, Simon Barkham, & Bernhard Gruber. 1987. Pötschenschichten, Zlambachmergel (Hallstätter Obertrias) und Liasfleckenmergel in Zentraltimor, nebst ihren Faunenelementen. Mitteilungen der Österrieches Geologisches Gesellschaft 80:229–285.
- Krumbeck, L. 1913. Obere Trias von Buru und Misól (Die Fogi-Schichten und Asphaltschiefer West-Burus und der Athyridenkalk des Misól-Archipels). *In* G. Böhm, ed.,Beiträge zur Geologie von Niederländisch-Indien. II. Abteilung. 1. Abschnitt. Palaeontographica Suppl. 4:1–161. Taf. 1–11. Stuttgart.
- Krumbeck, Lothar. 1914. Obere Trias von Sumatra (Die Padang-Schichten von West-Sumatra nebst Anhang). Palaeontographica Supplement IV(II):197–266.
- Krumbeck, Lothar. 1923a. Brachiopoden, Lamellibranchiaten und Gastropoden aus der oberen Trias der Insel Seran (Mittel-Seran). Palaeontographica Supplement 4(3):185–246.
- Krumbeck, Lothar. 1923b. Zur Kenntnis des Jura der Insel Timor sowie des Aucellen-Horizontes von Seran und Buru. Paläontologie von Timor 12(20):1–120, pl. 172–177.
- Krumbeck, Lothar. 1924. Die Brachiopoden, Lamellibranchiaten und Gastropoden der Trias von Timor II. Paläontologischer Teil. Paläontologie von Timor 13(22):144–417.
- Krystyn, Leo, Sylvain Richoz, Aymond Baud, & R. J. Twitchett. 2003. A unique Permian–Triassic boundary section from the Neotethyan Hawasina Basin, Central Oman Mountains. Palaeogeography, Palaeoclimatology, Palaeoecology 191:329–344.
- Kues, B. S. 2004. Marine invertebrate assemblages from the Late Pennsylvanian (Virgilian) Holder Formation, Dry Canyon, Sacramento Mountais, south-central New Mexico. New Mexico Geology 26(2):43–53.
- Kumpera, O., F. Prantl, & B. R ži ka. 1960. Revision of the Nuculanidae from the Ostrava-Karviná District (Pelecypoda). Acta Musei Nationalis Pragae (serie B) Historia Naturalis 16(1–2):17–80, pl. 1–4.
- Kurushin, N. I. 1982. The first occurrence of the genus *Streblopteria* in Triassic of Siberia. Transactions of the Institute of Geology and Geophysics, Siberian branch of the Academy of Sciences 462:60–63.
- Kurushin, N. I. 1984. Novye karnijskie Paleotaksodonty Vostochogo Tajmya [New Carnian Palaeotaxodonta of Eastern Taimyr]. In V. S. Surkov, ed., Novye vidy drevnikh bespozvonochnykh i rastenii neftegazonosnykh provintsii Sibiri [New species of fossil invertebrates and plants from oil and gas bearing provinces of Siberia]. Siberian Geological Research Institute, Novosibirsk. p. 61–66. In Russian. (*)
- Kurushin, N. I. 1987. Drevnejshie triasovye dvustvorchatye mollyuski Yakutii [The oldest Triassic bivalves from Yakutia]. Trudy Instituta Geologii i Geofiziki [Proceedings of the Institute of Geology and Geophysics] (Novosibirsk) 689:99–110. In Russian.
- Kurushin, N. I. 1990. Bivalve changes at the Triassic/Jurassic boundary of the Northeastern Asia. Les Cahiers de l'Université Catholique de Lyon (série Sciences) 3:185–190.
- Kurushin, N. I. 1991. Otkrytie zony Zittelihalogia zitteli (Bivalvia) v osnovanii Verkhnego Triasa na vostochnom Tajmyre [Discovery of the Zittelihalobia zitteli (Bivalvia) zone at the Upper Triassic base in eastern Taimyr]. Geologii i Geofiziki (Novosibirsk) 10:54–58. In Russian with English abstract.
- Kurushin, N. I. 1998. Triassic Bivalves of Northeastern Asia (Morphology, System, Biostratigraphy, Paleoecology, and Paleobiogeography).ScD Thesis. OIGGM, Novosibirsk. 45 p. In Russian. (*)
- Kurushin, N. I., & A. M. Trushchelev. 1989. Posidoniids from Triassic deposits in Siberia and the Far East. *In* A. S. Dagys & V. N. Dubatolov, eds., Upper Paleozoic and Triassic of Siberia. Academy of Sciences of the USSR, Siberian Department, Transactions of the Institute of Geology and Geophysics 172:57–71. In Russian. (*)

- Kurushin, N. I., & A. M. Truschelev. 2001. *Magnolobia*: A new bivalve genus from the Boreal Ladinian. Paleontological Journal 35(3):243–248.
- Kutassy, Andreas. 1931. Lamellibranchiata Triadica II. In W. Quenstedt, ed., Fossilium Catalogus I: Animalia. W. Junk. Berlin. 51:261–477.
- Kutassy, Andreas. 1934. Pachyodonta mesozoica. Fossilium Catalogus I. Gravenhage. 68:1–202.
- Kuznetzov, A. P., & A. A. Schileyko. 1984. On the gutless Protobranchia (Bivalvia). Nauchnye Doklady Vysshei Shkoly Biologicheskie Nauki 2:39–49. In Russian with English abstract.
- de Lamarck, J. B. P. A. de Monet. 1799. Prodrome d'une nouvelle classification des coquilles, comprenant une rédaction appropriée des caractères génériques, et l'établissement d'un grand nombre de genres nouveaux. Société d'Histoire Naturelle de Paris, Mémoires 1:63–91.
- de Lamarck, J. B. P. A de Monet. 1801. Système des Animaux sans Vertèbres, ou Tableau Général des Classes, des Ordres, et des Genres de ces Animaux. Présentant leurs Caractères Essentiels et leur Distribution, d'apres la Considération de leurs Rapports Naturels et de leur Organisation, et Suivant l'Arrangement établi dans les Galeries du Muséum National d'Histoire Naturelle, Parmi leurs Dépouilles Conservées, Précedée du Discours d'Ouverture du Cours de Zoologie, Donné dans le Muséum National d'Histoire Naturelle. L'an 8 de la Republique [= 1801]. Deterville [or chez l'auteur], Paris. viii + 432 p., tables.
- de Lamarck, J. B. P. A. de Monet. 1809. Philosophie Zoologique, ou Exposition des Considérations Relative à l'Histoire Naturelle des Animaux; à la Diversité de Leur Organisation et des Facultés qu'ils en Obtiennent; aux Causes Physiques qui Maintiennent en Eux la Vie et Donnent lieu aux Mouvements qui'ils Exécutent; enfin, á celles qui Produisent, les Unes le Sentiment, et les Autres l'Intelligence de Ceux qui eu sont Doués. Chez Dentu [et] L'Auteur, Paris, 2 vols: Vol. 1, xxv + 422 p., + Table des Matières, p. 423–428; Vol. 2, 466 p., + Table des Matières, p. 467–475.
- de Lamarck, J. B. P. A. de Monet. 1818. Histoire Naturelle des Animaux sans Vertèbres, vol. 5. Paris. 612 p.
- de Lamarck, J. B. P. A. de Monet. 1819. Histoire Naturelle des Animaux sans Vertèbres, vol. 6, part 1. Paris. 232 p.
- Lamy, Édouard. 1920. Révision des Cypricardiacea et des Isocardiacea vivants du Muséum d'Histoire Naturelle de Paris. Journal de Conchyliologie (Paris) 64[(4)18](4):259–307, pl. 8.
- LaRocque, A. 1950. Pre-traverse Devonian pelecypods of Michigan. Contributions from the Museum of Paleontology, University of Michigan VII(10):271–366.
- Laube, G. C. 1865. Die Fauna der Schichten von St. Cassian. Ein Beitrag zur Palaontologie der Alpinen Trias. II Abtheilung. (Brachiopoden und Bivalven). Denkschriften der kaiserlich-königlichen Akademie der Wissenschaften, Mathematisch-Naturwissenschaftliche Classe 25:32–76.
- Laube, G. C. 1866. Die Fauna der Schichten von St. Cassian. II. Abtheilung. Brachiopoden und Bivalven. Denkschriften der kaiserliche-königlichen Akademie der Wissenschaften, Mathematisch-Naturwissenschaftliche Classe 25:1–76, 9 pl.
- Laws, R. A. 1982. Late Triassic depositional environments and molluscan associations from west-central Nevada. Palaeogeography, Palaeoclimatology, Palaeoecology 37:131–148.
- Lazo, D. G. 2003. Taxonomy, facies relationships and palaeobiology of bakevelliid bivalves from the Lower Cretaceous of west-central Argentina. Cretaceous Research 24:765–788.
- Lazo, D. G. 2007a. Early Cretaceous bivalves of the Neuquén Basin, west-central Argentina: notes on taxonomy, palaeobiogeography and palaeoecology. Geological Journal 42:127–142.
- Lazo, D. G. 2007b. The bivalve *Pholadomya gigantea* in the Cretaceous of Argentina: Taxonomy, taphonomy and paleogeographic implications. Acta Palaeontologica Polonica 52(2):375–390.
- Leach, W. E. 1819. Descriptions des nouvelles espèces d'animaux découvertes par le vasseau Isabelle dans un voyage au pôle boréal. Journal de Physique, de Chimie, d'Histoire Naturelle et des Arts 88(6):462–467.

- Leanza, A. F. 1940. Dos nuevas especies del Género *Cucullaea* Lam. del Lias de Piedra Pintada. Notas del Museo de La Plata, Paleontología 5(23):207–215.
- Leanza, A. F. 1942. Los pelecípodos del Lias de Piedra Pintada en el Neuquén. Revista del Museo de La Plata (n.s.), Paleontología 2(10):206 p.
- Leanza, A. F. 1943. *Pectinula*, nuevo género de Pelecípodos en el Lías de Neuquén. Notas del Museo de La Plata, Paleontología 8(53):241–249.
- Leanza, H. A. 1993. Jurassic and Cretaceous Trigoniid Bivalves from West-Central Argentina. Bulletins of American Paleontology 105(343):95 p.
- Leanza, H. A. 1996. Jurassic trigoniacenas from Argentina: a review. Georesearch Forum 1–2:67–78.
- Leanza, H. A., & J. I. Garate Zubillaga. 1987. Faunas de trigonias (Bivalvia) del Jurásico y Cretácico Inferior de la Provincia del Neuquen Argentina, conservadas en el Museo Juan Olsacher de Zapala. *In* W. Volkheimer, ed., Bioestratigrafía de los sistemas regionales del Jurásico y Cretácico de America del sur 1:201–255.
- Lebküchner, R. 1932. Die Trigonien des süddeutscher Jura. Palaeontographica 77:1–119, pl. 1–16.
- Lebedev, I. V. 1959. Mezozojskie Peletsipody iz Chulymo-Enisejskoj Vprodiny. Trudy Tomsk Polytechnic Institute. 164:60–63.
- Leonardi, Piero. 1935. Il Trias inferiore delle Venezie. Memorie dell'Istituto Geologico della R. Università di Padova XI:1–136.
- Leonardi, Piero. 1943. La fauna Cassiana di Cortina d'Ampezzo. 1ª parte. Introduzione geologica e lamellibranchi. Memorie degli Instituti di Geologia e Mineralogia dell'Università di Padova XV:1–78.
- Leonardi, Piero. 1960. Studio statistico-sedimentologico di alcune faune werfeniane della Valle di Fiemme nel Trentino. Studi Trentini di Scienze Naturali XXXVII(1):17–29.
- Lerman, A. 1960. Triassic pelecypods from southern Israel and Sinai. Bulletin Research Council of Israel 9(1):1–60.
- Lewy, Zeev. 1976. Morphology of the shell in Gryphaeidae. Israel Journal of Earth-Sciences 25(2):45–50.
- Li Jin-hua, & Ding Bao-liang. 1981. Two new lamellibranch genera from the Lower Triassic of Anhui. Acta Palaeontologica Sinica 20(4):325–330. In Chinese with English abstract.
- Li Ling. 1995. Evolutionary change of bivalves from Changxingian to Griesbachian in South China. Acta Palaeontologica Sinica 34(3):350–369. In Chinese with English abstract.
- Li Xiaochi. 1990. The Marine Jurassic and Lower Cretaceous of Southern Xizang (Tibet): Bivalve Assemblages, Correlation, Paleoenvironments and Paleogeography. PhD thesis. University of Auckland. 280 p.
- Li Xiaochi, & J. A. Grant-Mackie. 1994. New Middle Jurassic-Lower Cretaceous bivalves from southern Tibet. Journal of Southeast Asian Earth Sciences 9(3):263–276.
- Li Xubing, Meng Fansong, & Wang Chuanshang. 2005. Paleoecological characteristics of fossil bivalves from the Guanling biota in Guizhou. Geology in China 32(1):41–47. In Chinese with English abstract.
- Li Yun. 1994. A Late Triassic *Burmesia mojiangensis* sp. nov. of Mojiang, Yunnan. Acta Palaeontologica Sinica 33(2):259–261. In Chinese with English abstract.
- Li Zishun, & Yu Xihan. 1982. The Middle and Late Jurassic Bivalvia from eastern Heilongjiang Province. Bulletin of the Shenyang Institute of Geology and Mineral Resources 5:73–200. In Chinese with English abstract.
- Li Zishun, Zhan Lipei, Zhu Xiufang, Zhang Jinghua, Jin Ruogu, Liu Guifang, Sheng Huaibien, Shen Guimei, Dai Jinye, Huang Hengquan, Xie Longchun, Yan Zheng, & Yao Jianxin. 1986. Mass extinction and geological events between Paleozoic and Mesozoic era. Acta Geologica Sinica 1:1–15. In Chinese.
- Licharew, B., ed. 1939. The Atlas of the Leading Forms of the Fossil Fauna of the U.S.S.R., vol. 6, Permian. Central Geological and Prospecting Institute. Leningrad. 228 p. In Russian.
- Licharew, B. K., & O. L. Einor. 1941. On the problem of the age of the Upper Paleozoic deposits of the south-east part of the Kolymia Basin.

Doklady Akademii Nauk SSSR [Comptes Rendus (Doklady) of the Academy of Sciences of URSS (new series)] 31(2):150–152. In Russian. (*)

- [Lightfoot, John]. 1786. A Catalogue of the Portland Museum, Lately the Property of the Duchess Dowager of Portland, Deceased. London. 194 p. Published as anonymous.
- Liljedahl, Louis. 1984. *Janeia silurica*, a link between nuculoids and solemyoids (Bivalvia). Palaeontology 27(4):693–698.
- Lin Li, Zhu Lidong, Pang Yancun, & Sha Jingeng. 2007. A new genus of Pergamidiidae (Bivalvia) from the Late Triassic of the Changtai-Gacun area, eastern Qinghai-Xizang Plateau, China. Journal of Asian Earth Sciences 30(1):108–112.
- Lin Mingyue, & Yin Hongfu. 1991. Preliminary research on Changxingian ecostratigraphy of Southwest Guizhou. Earth Science - Journal of China University of Geosciences 18(2):127–134.
- Linck, Otto. 1972. Die marine Fauna des süddeutschen oberen Gipskeupers, insbesondere der sogenannten Anatinenbank (Trias, Karn, Mittl. Keuper, km1) und deren Bedeutung-Evertebraten I. Jahreshefte des Geologischen Landesamt Baden-Württemberg 14:145–253.
- Ling Quixian. 1988. Triassic bivalve fauna in Zhen'an area of the eastern Qinling Mts. Acta Palaeontologica Sinica 27(5):615–628. In Chinese with English abstract.
- Link, H. F. 1807. Beschreibung der Naturalien-Sammlung der Universität zu Rostock. Adlers Erben, Rostock. Abtheilung 3:101–165.
- Linnaeus, C. A. 1758. Systema Naturae per Regna Tria Naturae, Secundum Classes, Ordines, Genera, Species, cum Characteribus, Differentiis, Synonymis, Locis. Editio Decima, reformata. Tomus I, Regnum Animale. p. 1–823.
- Linnaeus, C. A. 1764. Museum S:ae R:ae M:tis Ludovicae Ulricae Reginae Svecorum, Gothorum, Vandalorumque . . . In quo animalia rariora, exotica, imprimis insecta & conchilia describuntur & determinantur. Literis & impensis Direct. Laur. Salvii. 8 + 720 p. Stockholm. (*) Please spell out highlighted words
- Lintz, Joseph. 1958. The fauna of the Ames and Brush Creek shales of the Conemaugh Formation of western Maryland. Journal of Paleontology 32(1):97–112.
- Lissajous, Marcel. 1923. Étude sur la faune du Bathonien des environs de Mâcon. Travaux du Laboratoire de Géologie de la Faculté des Sciences de Lyon, Fasc. V, Memorie 3:282 p.
- Little, C. T. S., & R. C. Vrijenhoek, 2003. Are hydrothermal vent animals living fossils? Trends in Ecology and Evolution 18(11):582–588.
- Liu Chun-lian. 1995. Jurassic bivalve palaeobiogeography of the Proto-Atlantic and the application of multivariate analysis methods to palaeobiogeography. Beringeria 16:3–123.
- Liu Chun-lian. 1999. Distribution of Jurassic bivalve faunas in relation to palaeogeography and palaeoclimate. Acta Palaeontologica Sinica 38(4):544–550. In Chinese with English abstract.
- Llompart, Carmen, Juan Rosell, Ana Márquez-Aliaga, & Antonio Goy. 1987. El Muschelkalk de la isla de Menorca. Cuadernos de Geología Ibérica 11:323–339.
- Llopis LLadó, Noel. 1952. Sobre el área de dispersión de Daonella cf. lommeli v. Mojs., en el NE de España. Memorias y Comunicaciones, Instituto Geológico, Consejo Superior de Investigaciones Científicas, Barcelona IX:39–47.
- López-Gómez, José, Antonio Goy, & Ana Márquez-Aliaga. 2005. Bioestratigrafía del tránsito Triásico-Jurásico en Asturias: Ammonoideos y bivalvos entre Avilés y Caravia (España). Geo-Temas 8:183–187.
- López-Gómez, José, Ana Márquez-Aliaga, Alfredo Arche, & Antonio Goy. 1987. Las facies Muschelkalk de Henarejos (Cuenca): Sedimentología y fauna del tramo superior. Cuadernos de Geología Ibérica 11:665–676.
- López-Gómez, José, Ana Márquez-Aliaga, Francesc Calvet, Leopoldo Márquez, & Alfredo Arche. 1994. Aportaciones a la estratigrafía y paleontología del Triásico Medio de los alrededores de Agost, Alicante.

Boletín de la Real Sociedad Española de Historia Natural (Sección Geología) 89(1–4):109–120.

- de Loriol, Perceval. 1891. Études sur les mollusques des couches coralligènes inférieurs du Jura Bernois, accompagnées d'une Notice Stratigraphique par E. Koby, Prof. Troisième Partie. Mémoires de la Société Paléontologique Suisse 18:175–258.
- de Loriol, Perceval. 1900. Étude sur les mollusques et brachiopodes de l'Oxfordien inférieur ou Zone à Ammonites renggeri du Jura Lédonien. Mémoires de la Société Paléontologique Suisse 27:1–143, pl. 1–6.
- de Loriol, Perceval. 1901. Étude sur les mollusques et brachiopodes de l'Oxfordien supérieur et moyen du Jura Bernois. Premier Supplément. Mémoires de la Societé Paléontologique Suisse 28:1–119, pl. 1–7.
- de Loriol, Perceval. 1904. Étude sur les Mollusques et Brachiopodes de l'Oxfordien supérieur et moyen du Jura Lédonien. Mémoires de la Societé Paléontologique Suisse 31:1–161, pl. 20–27.
- Losacco, U. 1942. Un nuovo genere aaleniano di Aviculidae della Sabina. Rivista Italiana di Paleontologia e Stratigrafia 48(1):9–16. Published for 1940.
- Lu Yi-ju. 1981. Late Triassic lamellibranchs from Datong of Qinghai, NW China. Acta Palaeontologica Sinica 20(6):577–585. In Chinese with English abstract.
- Lu Yi-ju, & Chen Chu-zhen. 1986. Triassic bivalves from Mt Burhan Budai, Qinghai. *In* Academia Sinica, ed., Carboniferous and Triassic Systems and fossils from the southern slope of Mt. Barhan Budai, Quinghai, China. Anhai Science and Technology Publishing House. Wuhu. p. 141–169. In Chinese with English abstract.
- Lucas, S. G., & J. W. Estep. 1997. The Liassic bivalve Weyla from Sonora, Mexico. In C. M. González-León & G. D. Stanley, Jr., eds., US–Mexico Cooperative Research. International Workshop on the Geology of Sonora Memoir, Publicaciones Ocasionales 1:45.
- Lucas, S. G., & C. M. González-León. 1994. Marine Upper Triassic strata at Sierra la Flojera, Sonora, México. Neues Jahrbuch für Geologie und Paläontologie Monatshefte 1:34–40.
- Lucas, S. G., & L. H. Tanner. 2004. Late Triassic extinction events. Albertiana 31:31–40.
- Lucas, S. G., D. G. Taylor, Jean Guex, L. H. Tanner, & Karl Krainer. 2007. Updated proposal for Global Stratotype Section and Point for the base of the Jurassic System in the New York Canyon area, Nevada, United States. ISJS Newsletter 34(1):34–42.
- Lupher, A. W., & E. L. Packard. 1930. The Jurassic and Cretaceous rudistids of Oregon. University of Oregon Publication (Geology series) 1(3):203–225, pl. 1–6.
- Lycett, J. 1850. Tabular view of fossil shells from the middle division of the Inferior Oolite in Gloucestershire. Annals and Magazine of Natural History, London 2(6):401–425, pl. 11.
- Lycett, J. 1857. The Cotteswold Hills. Piper Stevenson & Spencer. London. 170 p., 7 pl.
- Lycett, J. 1863. Supplementary monograph on the Mollusca from the Stonesfield Slate, Great Oolite, Forest Marble and Cornbrash. Paleontographical Society. London. 129 p.
- Lycett, J. 1881. The Fossil Trigoniae (Supplement number 1). Monograph of the Paleontographical Society of London 35(164):1–3.
- M'Coy, Frederick. 1844. A synopsis of the characters of the Carboniferous limestone fossils of Ireland. Dublin. 207 p., 29 pl.
- M'Coy, Frederick. 1851. Descriptions of some new Mountain Limestone fossils. Annals and Magazine of Natural History (serie 2) 7:167–175.
- Ma Qihong. 1989. A new recognition of "Pseudocardiniidae" (Lamellibranchia). Acta Palaeontologica Sinica 28(5):604–616. In Chinese with English abstract.
- Ma Qihong, Chen Jinhua, Lan Xiu, Gu Zhiwei, Chen Chuzhen, & Lin Minji. 1976. Mesozoic lamellibranchs from Yunnan. *In* Nanjing Institute of Geology and Palaeontology, Academia Sinica, ed., Mesozoic fossils from Yunnan, Vol. 1. Science Press. Beijing. p. 161–386. In Chinese. (*)

- MacFarlan, D. A. B. 1998. Mesozoic stratigraphy of the Marokopa area, southwest Auckland, New Zealand. New Zealand Journal of Geology and Geophysics 41:297–310.
- Machalski, Marcin. 1998. Oyster life positions and shell beds from the Upper Jurassic of Poland. Acta Palaeontologica Polonica 43(4):609-634.
- Mack, G. H., Mike Leeder, Marta Pérez-Arlucea, & B. D. J. Bailey. 2003. Sedimentology, Paleontology, and Sequence Stratigraphy of Early Permian Estuarine Deposits, South-Central New Mexico, United States. PALAIOS 18:403–420.
- MacLeay, W. S. 1819. Horæ Entomologicæ: or Essays on the Annulose Animals, vol. 1, part I. S. Bagster. London. 160 p.
- Mader, Detlef. 1982. Genese des mitteleuropäischen Buntsandsteins. Entwicklungsgeschichte einer kontinentalen Rotformation. Naturwissenschaften 69:311–325
- Maeda, Shiro, Hideo Ishikawa, Tetsuya Kawabe, & L. G. Morales. 1983. Some Triassic Trigoniids from Peru. Bulletin of the Faculty of Education, Chiba University 32(2):47–57.
- Mahler, Horst, & Jürgen Sell. 1991. Die "vulgaris/costata-Bank" (Oberer Buntsandstein, Mitteltrias) - ein lithostratigraphisch verwertbarer biostratigraphischer Leithorizont mit chronostratigraphischer Bedeutung. *In* H. Hagdorm & A. Seilacher, eds., Muschelkalk Schöntaler Symposium 1991. Sonderbände der Gesellschaft für Naturkunde in Wüttemberg 2. Stuttgart, Korb (Goldschneck). 2:187–192.
- Malchus, Nikolaus. 1990. Revision der Kreide-Austern (Bivalvia: Pteriomorphia) Ägyptens (Biostratigraphie, Systematik). Berliner Geowissenschaftliche Abhandlungen A125:1–231.
- Malchus, Nikolaus. 1998. Multiple parallel evolution and phylogenetic significance of shell chambers and chomata in the Ostreoidea (Bivalvia). *In* P. A. Johnston & J. W. Haggart, eds., Bivalves: An Eon of Evolution Paleobiological Studies Honoring Norman D. Newell. University of Calgary Press. Calgary. p. 393–407.
- Malchus, Nikolaus. 2000. Early shell stages of the Middle Jurassic bivalves *Camptochlamys* (Pectinidae) and *Atreta* (Dimyidae) from Poland. Journal of Molluscan Studies 66:577–581.
- Malchus, Nikolaus. 2004. Constraints in the ligament ontogeny and evolution of Pteriomorphian Bivalvia. Palaeontology 47(6):1539–1574.
- Malchus, Nikolaus. 2008. Problems concerning early oyster evolution: A reply to Márquez-Aliaga and Hautmann. Palaeogeography, Palaeoclimatology, Palaeoecology 258:130–134.
- Malchus, Nikolaus, & Martin Aberhan. 1998. Transitional gryphaeate/ exogyrate oysters (Bivalvia: Gryphaeidae) from the Lower Jurassic of Northern Chile. Journal of Paleontology 72(4):619–631.
- Malinowskiej, Lidii. 1979. Budowa geologiczna Polski. Atlas skamieniato ci Przewodnich i Charakterystycznych. Mezozoik, Trias [Geology of Poland. Atlas of Guide and Characteristic Fossils. Mesozoic, Triassic]. Ed. Wydawnictwa Geologiczne. Tomo III, 286 p. In Polish.
- Mallada, Lucas. 1880. Sinopsis de las especies fósiles que se han encontrado en España. Terreno Mesozoico (Sistemas Triásico y Jurásico). Boletín Comunicaciones Mapa Geológico de España. Tomo II. Imprenta y fundición de Manuel Tello 7:241–256.
- Manceñido, M. O., C. R. González, & S. E. Damborenea. 1976. La fauna de la Formación del Salto (Paleozoico Superior de la Provincia de San Juan); Parte III, Bivalvia I. Ameghiniana 13(1):65–84.
- Mander, Luke, Richard Twitchett, & M. J. Benton. 2008. Palaeoecology of the Upper Triassic extinction event in the SW UK. Journal of the Geological Society of London 165(1):319–332.
- Mángano, M. G., L. A. Buatois, R. R. West, & C. G. Maples. 1998. Contrasting behavioral and feeding strategies recorded by tidal-falt bivalve trace fossils from the Upper Carboniferous of Eastern Kansas. PALAIOS 13:335–351.
- Mansuy, Henri. 1912. Contribution à la géologie du Tonkin. Paléontologie. Mémoires du Service Géologique de l'Indochine 1(4):55–71, pl. 10–13.

- Mansuy, Henri. 1914. I, Nouvelle contribution à la Paléontologie du Yunnan. II, Description d'espèces nouvelles des terrains paléozoïques et triasiques du Tonkin. III, Contribution a la Paléontologie du Laos. IV, Gisement Liasique des schistes de Trian (Cochinchine). V, Étude des faunes paléozoïques et mésozoïques des feuilles de Phú-nho-quan et de Son-tây (Tonkin). Mémoire Service géologique de l'Indochine 3(2):91 p., 10 pl. .
- Mansuy, Henri. 1919. Faunes Triasiques et Liasiques de Na Cham, Tonkin. Mémoire Service géologique de l'Indochine 6(1): 39 p., 5 pl .
- Mariani, Ernesto. 1919. Sulla fauna retica lombarda. Atti della Società Italiana di Science Naturali 58:104–146.
- Marincovich, Louie, Jr. 1993. Danian Mollusks from the Prince Creek Formation, Northern Alaska, and Implications for Arctic Ocean Paleogeography. The Paleontological Society Memoir 35, Journal of Paleontology 67(5):1–35.
- Marincovich, Louie Jr., K. B. Barinov, & A. E. Oleinik. 2002. The Astarte (Bivalvia: Astartidae) that document the earliest opening of Bering Strait. Journal of Paleontology 76(2):239–245.
- Marinov, V. A., S. V. Meledina, O. S. Dzyuba, O. S. Urman, O. V. Yazikova, V. A. Luchinina, A. G. Zamirailova, & A. N. Fomin. 2006. Biofacies of Upper Jurassic and Lower Cretaceous Sediments of Central West Siberia. Stratigraphy and Geological Correlation 14(4):418–432.
- Márquez-Aliaga, Ana. 1983. Aportación a sistemática de los bivalvos triásicos. COL-PA (Coloquios de Paleontología), Publicaciones del Departamento de Paleontología, Universidad Complutense 38:33–47.
- Márquez-Aliaga, Ana. 1985. Bivalvos del Triásico Medio del Sector Meridional de la Cordillera Ibérica y de los Catalánides. Doctoral Thesis. Colección Tesis Doctorales. Editorial de la Universidad Complutense de Madrid 40:429 p.
- Márquez-Aliaga, Ana, Kiril Budurov, & Vicente Martínez. 1996. Registro de conodontos del Triásico Medio en un nautiloideo de Espejeras (Alicante, España). Coloquios de Paleontología 48:76–83.
- Márquez-Aliaga, Ana, S. E. Damborenea, José Gómez, & Antonio Goy. 2010. Bivalves from the Triassic-Jurassic transition in northern Spain (Asturias and western Basque-Cantabrian Basin). Ameghiniana 47(2):185–205.
- Márquez-Aliaga, Ana, S. E. Damborenea, & Antonio Goy. 2008a. A preliminary study of Hettangian Bivalves ftom the Triassic-Jurassic passage beds in northern Spain. *In* Abstract Volume, Fifth International Symposium of IGCP506: Marine and non marine Jurassic: Global correlation and major geological events. Hammamet. p. 67–68.
- Márquez-Aliaga, Ana, S. E. Damborenea, & Antonio Goy. 2008b. Rhaetian and Hettangian Bivalves from the Triassic-Jurassic boundary beds in Asturias and Palencia. *In* Ruiz-Omeñaca, J. I., Piñuela, L. & García-Ramos, J. C., Libro de Resúmenes de las XXIV Jornadas de la Sociedad Española de Paleontología. Museo del Jurásico de Asturias, Colunga. p. 267–268.
- Márquez-Aliaga, Ana, Graciela Delvene, Anna García-Forner, & Sonia Ros. 2002. Catálogo de los bivalvos del Triásico depositados en el Museo Geominero (IGME, Madrid). Boletín Geológico y Minero 113(4):429–444.
- Márquez-Aliaga, Ana, Anna Gandin, Antonio Goy, & Pablo Plasencia. 2000. Nuevas aportaciones paleontológicas del Triásico Medio de Cerdeña (Italia). *In* J. B. Díez & A. C. Balbino, eds., Resúmenes I Congreso Ibérico de Paleontología. Ed. Universidad de Évora. p. 103–105.
- Márquez-Aliaga, Ana, Anna García-Forner, Graciela Delvene, & Sonia Ros. 2001. La colección de bivalvos del Triásico de Serra, área de Sagunto (Valencia), depositada en el Museo Geominero (IGME) Madrid. Publicaciones del Seminario de Paleontología de Zaragoza (SEPAZ) 5(2):614–620.
- Márquez-Aliaga, Ana, Anna García-Forner, & Pablo Plasencia. 2002. "Colección Triásico de Teruel" en el Museo de Geología de la Universitat de València. *In* G. Meléndez & E. Peñalver, eds., Patrimonio paleontológico de Teruel. Instituto de estudios turolenses. Teruel. p. 385–398.
- Márquez-Aliaga, Ana, & Soledad García-Gil. 1991. Paleontología y ambientes del Triásico Medio en el sector noroccidental de la Cordillera

Ibérica (Provs. de Soria y Guadalajara, España). Estudios Geológicos 47(1–2):85–95.

- Márquez-Aliaga, Ana, Francis Hirsch, & A. C. López-Garrido. 1986. Middle Triassic Bivalves from the Hornos-Siles formation (Sephardic province, Spain). Neues Jahrbuch für Geologie und Paläontologie Monatshefte 173(2):201–227.
- Márquez-Aliaga, Ana, A. P. Jiménez-Jiménez, A. G. Checa, & Hans Hagdorn. 2005. Lower oysters and their supposed Permian ancestors. Palaeogeography, Palaeoclimatology, Palaeoecology 229(1–2):127–136.
- Márquez-Aliaga, Ana, & José López-Gómez. 1989. Paleontología y ambientes sedimentarios del Triásico Medio, Muschelkalk, de la Cordillera Ibérica I: Cuenca y Valencia, España. Estudios Geológicos 45:387–398.
- Márquez-Aliaga, Ana, & Leopoldo Márquez. 2000. Fosildiagénesis de bivalvos del Triásico Medio del Prebético (Murcia, España): una aproximación. Boletín Geológico y Minero 111(5):33–46.
- Márquez-Aliaga, Ana, Leopoldo Márquez, Marcelo March, Antonio Goy, & J. M. Brito. 1987. Aspectos paleontológicos del Muschelkalk de la zona de Calanda (provincia de Teruel). Cuadernos de Geología Ibérica 11:677–691.
- Márquez-Aliaga, Ana, & Vicente Martínez. 1990a. Aspectos Microestructurales de bivalvos triásicos prismáticos. Iberus. Revista Española de Malacología 9(1–2):69–74.
- Márquez-Aliaga, Ana, & Vicente Martínez. 1990b. Aspectos Microestructurales de bivalvos triásicos foliares. Iberus. Revista Española de Malacología 9(1–2):75–80.
- Márquez-Aliaga, Ana, & Vicente Martínez. 1994. Procesos de colonización y su problemática, ejemplificada por bivalvos del Triásico español. Boletín de la Real Sociedad Española de Historia Natural (Sección Geología) 89(1–4):121–130.
- Márquez-Aliaga, Ana, & Vicente Martínez. 1996. Asociaciones de bivalvos del Triásico en España. Revista Española de Paleontología Nº Extraordinario, p. 101–111.
- Márquez-Aliaga, Ana, & Plinio Montoya. 1991. El Triásico de Alicante: Un efecto Lázaro en los estudios paleontológicos. Revista Española de Paleontología Nº Extraordinario, p. 115–125.
- Márquez-Aliaga, Ana, Pablo Plasencia, & Sonia Ros. 2005. Bivalvos del Retiense de la Formación Imón (Tarragona, España): en homenaje a Francesc Calvet. Geo-Temas 8:193–196.
- Márquez-Aliaga, Ana, & Sonia Ros. 2002. Taphonomy Richness of the Scarce Spanish Bivalves Triassic Record: Some Examples to be Discussed. *In* M. De Renzi, M. V. Pardo, M. Belinchón, E. Peñalver, P. Montoya & A. Márquez-Aliaga, eds., Current Topics on Taphonomy and Fossilization, 2002. Editorial Ilmo. Ayuntamiento de Valencia. Valencia. p. 199–206.
- Márquez-Aliaga, Ana, & Sonia Ros. 2003. Associations of bivalves of Iberian Peninsula (SPAIN): Ladinian. Albertiana 28:85–88.
- Márquez-Aliaga, Ana, José I. Valenzuela-Ríos, Pablo Plasencia, & Sonia Ros. 2004. Los fósiles del Muschelkalk (Triásico Medio) en el sector oriental de la Península Ibérica. *In* E. Baquedano & S. Rubio, eds., Miscelánea en Homenaje a Emiliano Aguirre. II: Palentología. Museo Arqueológico Regional de Alcalá de Henares. Madrid. p. 276–291.
- Martens, Eduard, & Georg Pfeffer. 1886. Die Mollusken von Süd-Georgien nach der Ausbeute der Deutschen Station 1882–3. Jahrbuch der Hamburgischen Wissenschaftlichen Anstalten III:65–135.
- Martin, M. J. 1860. Paléontologie stratigraphique de l'infra-Lias du département de la Cote-d'or. Mémoire de la Société géologique de France (2 serie) 7(1):1–100.
- Martín-Algarra, Agustín, Ana Márquez-Aliaga, Nuria Solé de Porta, & J. M. Valenzuela. 1993. La serie Triásica de los Pastores (Algeciras). Estudios Geológicos 49(1–2):1–20.
- Martín-Algarra, Agustín, Nuria Solé de Porta, & Ana Márquez-Aliaga. 1995. Nuevos datos sobre la estratigrafía y procedencia paleogeográfica del Triásico de las escamas del Corredor del Boyar (Cordillera Bética occidental). Cuadernos de Geología Ibérica 19: 279–307.

- Martínez, Vicente, & Ana Márquez-Aliaga. 1992. Aspectos fosildiagenéticos de *Placunopsis* del Muschelkalk Español. *In* A. Alemany, ed., Historia Natural '91. Universitat Illes Balears. p. 563–569.
- Martínez, Vicente, & Ana Márquez-Aliaga. 1994. Fosildiagénesis de Gervillia joleaudi (SCHMIDT, 1935): Estudio metodológico previo. Boletín de la Real Sociedad española de Historia Natural (Sección Geología) 89(1–4):199–208.
- Marwick, J. 1935. Some new genera of the Myalinidae and Pteridae of New Zealand. Transactions & Proceedings of the Royal Society of New Zealand 65(3):295–303, pl. 34–36.
- Marwick, John. 1953. Divisions and Faunas of the Hokonui System. New Zealand Geological Survey, Palaeontological Bulletin 21:141 p.
- Marwick, John. 1956. Three fossil Mollusca from the Hokonui system (Triassic and Jurassic). Transactions of the Royal Society of New Zealand 83(3):489–491.
- von Martens, E. C., & G. Pfeffer. 1886. Mollusken, Krebse und Echinodermen von Cumberland-Sund nach der Ausbeute der deutschen Nordexpedition 1882 und 1883. Jahrbuch der Hamburgischen Wissenschaftlichen Anstalten 3:23–50, 1 pl.
- Mathéron, Philippe [E.]. 1842–1843. Catalogue méthodique et descriptif des corps organisés fossiles du Départment des Bouches-du-Rhône et Lieux circonvoisins; précedé d'un mémoire sur les terrains supérieurs au Grés Bigarré du S.E. de La France. 269 p., 41 pl.
- Maton, W. G., & T. Rackett. 1807. A descriptive catalogue of the British Testacea. Transactions of the Linnean Society of London 8:17–250, 6 pl.
- Matsukawa, Masaki, & Kosuke Nakada. 2003. Adaptive strategy and evolution of corbiculoids based on the Japanese Mesozoic fossils. Bulletin Tokyo Gakugei University Section 4, Mathematics and Natural Sciences 55:161–189. In Japanese with English abstract.
- McAlester, A. L. 1968. Type species of Paleozoic Nuculoid Bivalve Genera. Memoir, Geological Society of America 105:1–143.
- McLearn, F. H. 1941. Preliminary descriptions of some new Triassic pelecypods from the Peace River foothills, B. C. The Canadian Field-Naturalist 55:31–33. (*)
- McLearn, F. H. 1942. The Neo-triassic *Cassianella* fauna of Tyaughton Creek Valley, B. C. The Canadian Field-Naturalist 56:99–103.
- McLearn, F. H. 1946. Upper Triassic faunas in Halfway, Sikanni Chief, and Prophet River Basins, Northeastern British Columbia. Geological Survey of Canada Paper 46–25:6–9.
- McLearn, F. H. 1947. Upper Triassic faunas of Pardonet Hill, Peace River Foothills, British Columbia. Geological Survey of Canada, Paper 47–24:16 p.
- M'Coy, F. 1844. A Synopsis of the Characters of the Carboniferous Limestone Fossils of Ireland. Dublin University Press. Dublin. 207 p.
- M'Coy, F. 1851. Description of some new Mountain Limestone fossils. Annals and Magazine of Natural History 2(7):167–175.
- McRoberts, C. A. 1992. Systematics and paleobiogeography of Late Triassic *Gryphaea* (Bivalvia) from the North American Cordillera. Journal of Paleontology 66(1):28–39.
- McRoberts, C. A. 1993. Systematics and biostratigraphy of halobiid bivalves from the Martin Bridge Formation (Upper Triassic), northeast Oregon. Journal of Paleontology 67(2):198–210.
- McRoberts, C. A. 1994. The Triassic-Jurassic ecoestratigraphic transition in the Lombardian Alps, Italy. Palaeogeography, Palaeoclimatology, Palaeoecology 110:145–166.
- McRoberts, C. A. 1997a. Late Triassic (Norian, Rhaetian) Bivalves from the Antimonio Formation, Northwestern Sonora, Mexico. Revista Mexicana de Ciencias Geológicas 14(2):167–171.
- McRoberts, C. A. 1997b. Late Triassic North American Halobiid Bivalves: diversity trends and circum-Pacific correlations. *In* J. M. Dickins, Z. Yang, H. Yin, S. G. Lucas, & S. K. Acharyya, eds., Late Paleozoic and Early Mesozoic Circum-Pacific Events and their Global Correlation. Cambridge University Press. Cambridge. Chapter 22. p. 198–208.

- McRoberts, C. A. 2000. A primitive *Halobia* (Bivalvia: Halobioidea) from the Triassic of Northeast British Columbia. Journal of Paleontology 74(4):599–603.
- McRoberts, C. A. 2005. Extinction and survival of Permian to Early Triassic marine Myalinidae (Bivalvia: Pterioida). Albertiana 33:60–61.
- McRoberts, C. A. 2010. Biochronology of Triassic bivalves. In S. G. Lucas, ed., The Triassic Timescale. Geological Society, London, Special Publications 334:201–219.
- McRoberts, C. A., & R. B. Blodgett. 2000. Late Triassic (Norian) Mollusks from Taylor Mountains Quadrangle, Soutwestern Alaska. U. S. Geological Survey Professional Paper 1662:55–67.
- McRoberts C. A., & J. G. Carter. 1994. Nacre in an early gryphaeid bivalve (Mollusca). Journal of Paleontology 68(6):1405–1408.
- McRoberts, C. A., Leo Krystyn, & Adriel Shea. 2008. Rhaetian (Late Triassic) *Monotis* (Bivalvia: Pectinoida) from the Eastern Northern Calcareous Alps (Austria) and the End-Norian crisis in pelagic faunas. Palaeontology 51(3):721–735.
- McRoberts, C. A., & N. D. Newell, 2005. Marine Myalinidae (Bivalvia: Pterioida) from the Permian of West Texas. American Museum Novitates 3469:1–15.
- McRoberts, C. A., C. R. Newton, & Andrea Allasinaz. 1995. End-Triassic bivalve extinction: Lombardian Alps, Italy. Historical Biology 9:297–317.
- Meek, F. B. 1864. Checklist of the invertebrate fossils of North America. Cretaceous and Jurassic. Smithsonian Miscellaneous Collections 7(177):1–40.
- Meek, F. B. 1865. Description of fossils from the auriferous slates of California. Geological Survey of California, Geology 1:477–482.
- Meek, F. B. 1877. Part I Palaeontology. Professional Papers of the Engineer Department, U.S. Army. No. 18. Report of the Geological Exploration of the Fortieth Parallel, Secretary of War 4:1–197.
- Meek, F. B., & F. V. Hayden. 1859. Description of new organic remains collected in Nebraska Territory in the year 1857, by Dr. F. V. Hayden, geologist to the Exploring Expedition under the command of Lieut. G. K. Warren, Top. Engr. U.S. Army, together with some remarks on the geology of the Black Hills and portions of the surrounding country. Proceedings of the Academy of Natural Sciences, Philadelphia 10:41–59.
- Meek, F. B., & F. V. Hayden. 1860. Systematic catalogue, with synonyma, & c., of Jurassic, Cretaceous and Tertiary fossils collected in Nebraska, by the Exploring Expeditions under the command of Lieut. G. K. Warren, of U.S. Topographical Engineers. The Academy of Natural Sciences of Philadelphia, Proceedings 12:417–432.
- Meek, F. B., & F. V. Hayden. 1864. Palaeontology of the Upper Missouri. Smithsonian Contributions to Knowledge 14(172):1–135.
- Meek, F. B., & A. H. Worthen. 1866. Descriptions of Paleozoic fossils from the Silurian, Devonian, and Carboniferous rocks of Illinois and other western states. Chicago Academy of Natural Science, Proceedings 1(1865–1866):11–23.
- Meek, F. B., & A. H. Worthen. 1869. Descriptions of new Carboniferous fossils from the western states. Academy of Natural Sciences of Philadelphia, Proceedings 21:136–172.
- Megerle von Mühlfeld, J. K. 1811. Entwurf eines neuen Systems der Schalthiergehäuse, Erste Abtheilung, Die Muscheln. Der Gesellschaft Naturforschender Freunde zu Berlin, Magazin für die Neuesten Entdeckungen in der Gesammten Naturkunde. Fünfter [V] Jahrgang, Berlin. In der Realschulbuchhandlung 5:38–72, fig. 1–3, pl. 3.
- Mehl, Johannes, & K. U. Rehfeld. 1992. Sipho-Erhaltung bei *Pholadomya fidicula* (Pelecypoda) aus dem Mittleren Jura (Bajocium), ein Vergleich mit der rezenten *Pholadomya candida*. Senckenbergiana Lethaea 71(5–6):367–375.
- Meledina, S. V., & B. N. Shurygin. 2001. Ammonoidei i Dvustvorchatye mollyuski iz verkhnego perkhnego Plinsbakha srednej Sibiri [Upper Pliensbachian ammonoids and bivalve mollusks from Central Siberia]. Novosti paleontologii i stratigrafii (Prilogenie k zhurnalu Geologiya I Geofizika t. 42) [News on Paleontology and Stratigraphy (Geology and

Geophysics, Supplement to vol. 42)] 4:35–47. In Russian with English abstract.

- Melnikova, G. K. 2006. The Early Jurassic Fauna from the Gurumdy and Mynhajir Zones of the East Pamirs. Doklady Earth Science 407(2):172–174.
- Melville, R. V. 1956. Stratigraphical paleontology, ammonites excluded, of the SDL Park Borehole. Bulletin of the Geological Survey of Great Britain 11:67–139.
- Menard de La Groye, F. J. B. 1807. Mémoire sur un nouveau genre de coquille de la famille des Solénoides. Annales du Muséum d'Histoire Naturelle (Paris) 9(50–51):131–139.
- Mendes, J. C. 1944. Lamelibranquios triassicos de Rio Claro (estado de Sao Paulo). Boletim da Faculdade de Filosofia, Ciências e Letras da Universidade de São Paulo, Geología 45(1):41–75.
- Mendes, J. C. 1945. Considerações sobre a estratigrafia e idade da formação Estrada Nova. Boletim da Faculdade de Filosofia, Ciências e Letras da universidade de São Paulo, Geologia 50(2):27–34.
- Mendes, J. C. 1952. A Formação Corumbataí na região do Rio Corumbataí (estratigrafia e descrição dos lamellibrânquios). Boletim da Faculdade de Filosofia, Ciências e Letras, Universidade São Paulo, Geologia 145(8):1–119.
- Meneghini, J. 1853. Nuovi fossili toscani illustrati dal Prof. G. Meneghini. In Appendice alle considerazioni sulla geologia estratigrafica toscana dei Professori Cavalieri P. Savi e G. Meneghini. Annali Universitá Toscana 3:1–40.
- Mensink, Hans, & Gottfried Tichy. 1977. Megalodontiden aus dem Keuper der Südpyrenäen südlich Seo de Urgel (Ein Beitrag zur Stratigraphie der Trias in Nordspanien). Geologie Paläontologie Mitteilungen Innsbruck 7(1):1–19.
- Metcalfe, Ian. 1998. Palaeozoic and Mesozoic geological evolution of SE Asian Region: multidisciplinary constraints and implications for biogeography. *In* R. Hall & J. D. Holloway, eds., Biogeography and Geological Evolution of SE Asia. Backhuys Publishers. Leiden, The Netherlands. p. 25–41.
- Metcalfe, Ian. 1999. The ancient Tethys oceans of Asia: How many? How old? How deep? How wide? UNEAC Asia Papers 1:1–9.
- Metwally, M. H. N. 1993. An analysis of the ranges of pectinid genera (Bivalvia) and their paleoecological interpretation. Neues Jahrbuch für Geologie und Paläontologie Monatshefte 4:193–208.
- Michalík, Jozef, Otília Lintnerová, Andrzej Gazdzicki, & Ján Soták. 2007. Record of environmental changes in the Triassic–Jurassic boundary interval in the Zliechov Basin, Western Carpathians. Palaeogeography, Palaeoclimatology, Palaeoecology 244:71–88.
- Miller, S. A. 1877. The American Palaeozoic Fossils, a Catalog of the Genera and Species, with names of authors, dates, places of publication, groups of rock in which found, and the etymology and signification of the words, and an introduction devoted to the stratigraphical geology of the Palaeozoic rocks. Published by the author. Cincinnati. 253 p.
- Milova, L. V. 1969. Novye ranneyurskie dvustvorchatye molluski bassejna reki Bol'shogo Anyuya. *In* N. A. Shilo, ed., Inotseramy yury i mela Severo-Vostoka SSSR. Akamemiya Nauk SSSR, Sibirskoye Otdeleniye, Severo Vostochenyj Kompleksnoyj, Trudy 32:177–189, pl. 1–3. Magadan.
- Milova, L. V. 1976. Stratigrafiya i dvustvorchatye mollyuski triasovo-yurskikh otlozhenii Severnogo Priokhot'ya [Triassic-Jurassic Stratigraphy and Bivalvia of northern Priokhotye]. Trudy Severo-Vostcvnyi Kompleksnoyi Nauchno-Institut. Akademia Nauk SSSR 65:110 p. In Russian.
- Milova, L. V. 1988. Ranneyurskie Dvustvorchatye Mollyuski Severo-Vostoka SSSR [Lower Jurassic Bivalves of North-Eastern USSR]. *In* Akademiya Nauk SSSR, Dal'nevostochnoe Otdelenie, Severo-Vostochnyj Kompleksnyj Nauchno-Issledovatel' skij Institut. p. 60–128. In Russian.
- Moesch, C. 1874–1875. Monographie der Pholadomyen. Abhandlungen Schweizerische Paläontologisches Gesellschaft 1:1–78, pl. 1–29; 2:79–135, pl. 30–40.

- Mojsisovics, E. v. 1874. Über die Triadischen Pelecypoden-Gattungen Daonella und Halobia. Abhandlungen der Kaiserlich-Königlichen Geologischen Reichsanstalt 7(2):1–38.
- Monari, Stefano. 1994. I bivalvi giurassici dell'Appennino umbro-marchigiano. Studi Geologici Camerti, Volume Speciale "Biostratigrafia dell'Italia centrale." p. 157–187.
- Monari, Stefano. 2003. A new genus and species of fimbriid bivalve from the Kimmeridgian of the Western Pontides, Turkey, and the phylogeny of the Jurassic Fimbriidae. Palaeontology 46(5):857–884.
- Montagu, George. 1808. Supplement to Testacea Britannica. With additional plates. White, London, & Woolmer. Exeter. 183 p., pl. 17–30.
- Moore, Charles. 1861. On the zones of the Lower Lias and the *Avicula contorta* zone. Quarterly Journal of the Geological Society of London 17:283–516, pl. 15–16.
- Mörch, O. A. L. 1853. Catalogus conchyliorum quae reliquit D. Alphonso d'Aguirra et Gadea Comes de Yoldi, (2) Acephala, Annulata, Cirripedia, Echinodermata. Klein. Hafnia. 74 p.
- Morris, John, & John Lycett. 1850. On *Pachyrisma*, a fossil genus of Lamellibranchiate Conchifera. Quarterly Journal of the Geological Society of London 6:399–402.
- Morris, John, & John Lycett. 1851–1855. A monograph of the Mollusca from the Great Oolite, chiefly from Minchinhampton and the coast of Yorkshire. Palaeontographical Society (Monographs), London. Part II, Bivalves: 1–80, pl. 1–8. Part III, Bivalves: 81–147, pl. 9–15.
- Morris, N. J. 1967. Mollusca: Scaphopoda and Bivalvia. *In* H. W. Harland, C. H. Holland, M. R. House, N. F. Hughes, A. B. Reynolds, M. J. S. Rudwick, G. E. Satterthwaite, L. B. H. Tarlo, & E. C. Willey, eds., The Fossil Record. Geological Society of London, Special Publication 2:469–477.
- Morris, N. J. 1978. The infaunal descendants of the Cycloconchidae: an outline of the evolutionary history and taxonomy of the Heteroconchia, superfamilies Cycloconchacea to Chamacea. Philosophical Transactions of the Royal Society of London B 284:259–275.
- Moscoso, Ramón, & Vladimir Covacevich. 1982. Las sedimentitas triásicojurásicas al sur de Canto del Agua, Cordillera de la Costa, Región de Atacama, Chile: descripción de la formación Canto del Agua. Actas del III Congreso Geológico Chileno 3:179–196.
- Muir-Wood, H. M., & G. A. Cooper. 1960. Morphology, classification, and life habits of the Productoidea (Brachiopoda). Geological Society of America Memoir 81:447 p.
- Müller, Johannes. 1842. Beiträge zur Kenntniss der natürlichen Familien der Fische. Archiv für Naturgeschichte 9:292–330.
- Müller, O. F. 1776. Zoologiae Danicae Prodromus, seu Animalium Daniae et Norvegiae Indigenarum Characteres, Nomina, et Synonyma Imprimis Popularium. Impenis Auctoris. Hallageriis. Havniae (Copenhagen). 282 p. In Latin.
- Münster, Georg v. 1841. Beiträge zur Petrefactenkunde. IV. (Beiträge zur Geognosie und Petrefactenkunde des südöstlichen Tirol, vorzüglich der Schichten v. St. Cassian von Dr. Wissmann und Graf Münster unter Mitwirkung des Dr. Braun). Bayreuth. p. 1–152, pl. 1–16.
- Murata, Masafumi, & Yuji Bando. 1975. Discovery of late Permian Araxoceras from the Toyoma Formation in the Kitakami. Transactions and Proceedings of the Palaeontological Society of Japan (new series) 97:22–31.
- Murchison, R. I., J. Buckman, & H. A. Strickland. 1844. Outline of the Geology of the Neighborhood of Cheltenham. J. Murray. London. 109 p., 13 pl.
- Muromtseva, V. A. 1979. Predstaviteli inotseramid v verkhnepermskikh otlozheniyakh Verkhoyan'ya [Members of the Inoceramidae in the upper Permian Deposits of the Verkhoyansk Region]. *In* Verkhnii paleozoi i mezozoi ostrovov i poberezh'ya arkticheskikh morei SSSR [Upper Paleozoic and Mesozoic of the Islands and Coastal Regions of the Arctic Seas of the USSR]. Nauchno-Issled. Institut Geol. Arktiki. Leningrad. p. 34–38. In Russian.

- Muromtseva, V. A. 1984. Rod Vnigripecten Muromtseva, nov. rod, i rod Morrisipecten Muromtseva i Guskov [Genus Vnigripecten Muromtseva, n. gen., and Genus Morrisipecten Muromtseva and Guskov]. In V. A. Muromtseva, ed., Permskiye Morskiye Otlozheniya i Dvustvorchatyie moll'uski Sovietskoi Arktiki [Permian marine deposits and bivalve molluscs of the Soviet Arctic]. Nedra. Leningrad. p. 65–66, 75. In Russian. (*)
- Muster, Heike. 1995. Taxonomie und Paläobiogeographie der Bakevelliidae (Bivalvia). Beringeria 14:161 p.
- Nakazawa, Keiji. 1952. A study on the Pelecypod-Fauna of the Upper Triassic Nabae Group in the Northern Part of Kyoto Prefecture, Japan. Part 1. Pectinids and Limids. Memoirs of the College of Science, University of Kyoto (serie B) 20(2):95–106.
- Nakazawa, Keiji. 1955. A study on the pelecypod-fauna of the Upper Triassic Nabae group in the northern part of Kyoto Prefecture, Japan, part 3. Halobiids and others. Memoirs of the Faculty of Science, University of Kyoto B 22(2):243–260.
- Nakazawa, Keiji. 1956. A study on the Pelecypod-Fauna of the Upper Triassic Nabae Group in the Northern part of Kyoto Prefecture, Japan. Part 4. *Cardinioides, Homomya, Pleuromya* and others, supplement and brief summary. Memoirs of the College of Science, University of Kyoto (Serie B) 23(2):231–253.
- Nakazawa, Keiji. 1959. Permian and Eo-Triassic Bakevellias from the Maizuru zone, southwest Japan. Memoirs, College of Science, University of Kyoto (series B) Geology and Mineralogy 26(2):193–213, pl. 3–4.
- Nakazawa Keiji. 1960. Permian and Eo-Triassic Myophoriidae from the Maizuru Zone, Southwest Japan. Japanese Journal of Geology and Geography 31:49–62.
- Nakazawa, Keiji. 1961. Early and Middle Triassic Pelecypod-fossils from the Maizuru Zone, Southwest Japan. Memoirs of the College of Science, University of Kyoto (series B) Geology and Mineralogy 27(3):249–291.
- Nakazawa, Keiji. 1963. Norian Pelecypod-fossils from Jito, Okayama Prefecture, West Japan. Memoirs of the College of Science, University of Kyoto (series B) 30(2):47–58.
- Nakazawa, Keiji. 1964. On the *Monotis typica* Zone in Japan. Memoirs of the College of Science, University of Kyoto (series B) 30(4):21–39.
- Nakazawa, Keiji. 1967. [Restudy of some described Permian bivalve species from Japan]. With an Appendix: Redescription of *Myophoria japonica* Hayasaka. *In* Committee for the commemoration of Professor Ichiro Hayasaka's 76th birthday, ed., Contributions to celebrate Prof. Ichiro Hayasaka's 76th birthday, Hashimoto Kabundo, Kanazawa. p. 165–174. In Japanese, Appendix in English.
- Nakazawa, Keiji. 1971. The Lower Triassic Kurotaki Fauna in Shikoku and its allied Faunas in Japan. Memoirs of the Faculty of Science, Kyoto University, Series of Geology and Mineralogy 38(1):103–133.
- Nakazawa, Keiji. 1973. On the Permian fossils from Jenga Pass, Pahang, Malay Peninsula. Science Report of Tohoku University, 2nd series (Geology), Special Volume 6:277–296. (*)
- Nakazawa, Keiji. 1991. Mutual Relation of Tethys and Japan during Permian and Triassic Time viewed from Bivalve Fossils. Saito Ho-on Kai Special Publication, Proceedings of Shallow Tethys 3, Sendai. 3:3–20.

Nakazawa, Keiji. 1992. The Permian-Triassic boundary. Albertiana 10:23-30.

- Nakazawa, Keiji. 1994. Occurrence of Triassic bivalves in the Tamba Belt at Izuriha, north of Takatsuki, Osaka Prefecture. Earth Science 48(2):171–173. (*)
- Nakazawa, Keiji. 1996. Lower Triassic bivalves from the Salt Range region, Pakistan. In P. K. S. Guha, K. Ayyasami, S. Sengupta, & R. N. Ghosh, eds., Gondwana Nine, Proceedings of the Ninth International Gondwana Symposium, Hyderabad, 1994. p. 207–229. (*)
- Nakazawa, Keiji. 1999. Permian bivalves from West Spitsbergen. Paleontological Research 3(1):1–17.
- Nakazawa, Keiji. 2002. Permian bivalves from the H. S. Lee Formation, Malaysia. Paleontological Research 6(1):67–72.

- Nakazawa, Keiji, H. M. Kapoor, Ken-ichi Ishii, Yuji Bando, Yuji Okimura, & Takao Tokuoka. 1975. The Upper Permian and the Lower Triassic in Kashmir, India. Memoirs of the Faculty of Science, Kyoto University, Series of Geology and Mineralogy 42(1):1–106.
- Nakazawa, Keiji, & N. D. Newell. 1968. Permian Bivalves of Japan. Memoirs of the Faculty of Science, Kyoto University, Series of Geology and Mineralogy 35(1):1–108.
- Nanjing Institute of Geology and Mineral Resources. 1982. Paleontological atlas of east China. Part 3. Volume of Mesozoic and Cenozoic. Beijing. Geological Publishing House. 405 p. In Chinese.
- Nardo, L. 1840. Sopra un nuovo genere di conchiglie del mare Adriatico. Annali delle Scienze del Regno Lombardo-Venedeto MDCCCXL (Parte I):49–51.
- Nauss, A. L., & P. L. Smith. 1988. *Lithiotis* (Bivalvia) bioherms in the Lower Jurassic of east-central Oregon, United States. Palaeogeography, Palaeoclimatology, Palaeoecology 65:253–268.
- Neave, S.A. 1939. Nomenclator Zoologicus. vol. 1, A–C. The Zoological Society of London. 957 p. Online: http://uio.mbl.edu/Nomenclator-Zoologicus/
- Neri, Claudio., M. Pasini, & Renato Posenato. 1986. The Permian/Triassic boundary and the Early Scythian sequence—Tesero section, Dolomites. *In* Field Conference on Permian and Permian-Triassic Boundary in the South-Alpine segment of the western Tethys. Excursion Guidebook. Società Geologica Italiana. p. 123–145.
- Neri, Claudio, & Renato Posenato. 1985. New biostratigraphical data on uppermost Werfen Formation of western Dolomites (Trento, Italy). Geologie Paläontologie Mitteilungen, Innsbruck 14(3):83–107.
- Neumayr, M. 1891. Beiträge zu einer morphologischen Eintheilung Bivalven. Denkschriften der Kaiserlichen Akademie der Wissenschaften, Wien, Mathematisch-Naturwissenschaftliche Classe 58:701–801.
- Newell, N. D. 1938. Late Paleozoic pelecypods: Pectinacea. State Geological Survey of Kansas, Bulletin 10:1–123, 20 pl.
- Newell, N. D. 1940. Invertebrate fauna of the Late Permian Whitehorse Sandstone. Bulletin of the Geological Society of America 51:261–336, 10 pl.
- Newell, N. D. 1942. Late Paleozoic Pelecypods: Mytilacea. Part II. State Geological Survey of Kansas. University of Kansas Publications 10:1–115.
- Newell, N. D. 1957. Notes on certain primitive Heterodont Pelecypods. American Museum Novitates 1857:1–14.
- Newell, N. D. 1999. A new limoid bivalve from the Texas middle Permian. American Museum Novitates 3264:1–6.
- Newell, N. D., & D. W. Boyd. 1970. Oyster-like Permian Bivalvia. Bulletin of the American Museum of Natural History 143(4):221–278.
- Newell, N. D., & D. W. Boyd. 1975. Parallel evolution in early Trigoniacean Bivalves. Bulletin of the American Museum of Natural History 154:53–162.
- Newell, N. D., & D. W. Boyd. 1985. Notes on Micro-Fabric in Upper Paleozoic Scallops. American Museum Novitates 2816:1–6.
- Newell, N. D., & D. W. Boyd. 1987. Iteration of ligament structures in Pteriomorphian Bivalves. American Museum Novitates 2875:1–111.
- Newell, N. D., & D. W. Boyd. 1989. Phylogenetic Implications of shell Microstructure in the Pseudomonotidae, extinc Bivalvia. American Museum Novitates 2933:1–12.
- Newell, N. D., & D. W. Boyd. 1995. Pectinoid bivalves of the Permian-Triassic crisis. Bulletin of the American Museum of Natural History 227:95 p.
- Newell, N. D., & D. W. Boyd. 1999. A new Lower Triassic *Permophorus* from the central Rocky Mountais. American Museum Novitates 3263:1–5.
- Newell, N. D., & B. Kummel. 1942. Lower Eo-Triassic stratigraphy, Western Wyoming and Southeast Idaho. Bulletin of the Geological Society of America 53:937–996.
- Newton, C. R. 1986. Late Triassic bivalves of the Martin Bridge Limestones, Hell Canyon, Oregon: taphonomy, paleoecology, paleozoogeography.

In T. L. Vallier & H. C. Brooks, eds., Geology of the Blue Mountains Region of Oregon, Idaho, and Washington. Geologic Implications of Paleozoic and Mesozoic Paleontology and Biostratigraphy, Blue Mountains Province, Oregon and Idaho. U. S. Geological Survey Professional Paper 1435:7–22.

- Newton, C. R. 1988 Significance of "Tethyan" fossils in the American Cordillera. Science 242:385–391.
- Newton, C. R. 1989. Biogeography of latest Norian Cordilleran bivalves; implications for extinction analysis and terrane comparisons. Abstracts with Programs - Geological Society of America 21(6):341.
- Newton, C. R., M. T. Whalen, J. B. Thompson, Nienke Prins, & David Delalla. 1987. Systematics and Paleoecology of Norian (Late Triassic) Bivalves from a tropical island arc: Wallowa terrane, Oregon. Journal of Paleontology [Memoir Paleontological Society 22] 61(4):83 p.
- Newton, R. B. 1891. Systematic List of the F.E. Edwards Collection of British Oligocene and Eocene Mollusca in the British Museum (Natural History). British Museum (Natural History). London. xxviii + 365 p.
- Newton, R. B. 1906. Notice on some fossils from Singapore. Geological Magazine 5(3):487–496.
- Nicol, David. 1950. Recent species of the lucinoid pelecypod *Fimbria*. Journal of the Washington Academy of Sciences 40(3):82–87.
- Nicol, David & William T. Allen. 1953. A new pelecypod genus from Upper Triassic strata in Peru? Journal of the Washington State Academy of Sciences 43(11):344-346.
- Nicoll, R. S. 2002. Conodont biostratigraphy and palaeogeography of the Triassic on the western, northwestern and northern margins of the Australian Plate. *In* W. A. Perth, M. Keep, & S. J. Moss, eds., The Sedimentary Basins of Western Australia 3. Proceedings of the Petroleum Exploration Society of Australia Symposium. Perth. p. 167–177.
- Nicora, Alda, Marco Balini, A. Bellanca, Angela Bertinelli, S. A. Bowring, Pietro Di Stefano, Chiara Guaiumi, Maria Gullo, Alexandre Hungerbuehler, Marco Levera, Michele Mazza, C. A. McRoberts, Giovanni Muttoni, Nereo Preto, & Manuel Rigo. 2007. The Carnian/Norian boundary interval at Pizzo Mondello (Sicani Mountains, Sicily) and its bearing for the definition of the GSSP of the Norian stage. Albertiana 36:102–115.
- Nielsen, S. N. 2005. The Triassic Santa Juana Formation at the lower Biobío River, south central Chile. Journal of South American Earth Sciences 19:547–562.
- Niemeyer, Jörg. 2002. Invertebraten und Vertebraten aus dem Muschelkalk von Siles (Jaen), Spanien. Münstersche Forschungen zur Geologie und Paläontologie 94:1–99.
- Niu Zhijun, Xu Guanghong, & Ma Liyan. 2003. Sedimentary characters and ammonoid fauna from the Upper Triassic Bagong Formation in the Geladandong area of the source land of the Yangtze River. Journal of Stratigraphy 27(2):129–137. In Chinese with English abstract.
- Noetling, Fritz. 1880. Die Entwicklung der Trias in Niederschlesien. Zeitschrift der Deutschen Geologischen Gesellschaft 32:300–349.
- Nori, Laetitia, & Bernard Lathuilière. 2003. Form and environment of *Gryphaea arcuata*. Lethaia 36:83–96.
- Ogilvie Gordon, M. M. 1927. Das Grödener-, Fassa- und Enneberggebiet in den Südtiroler Dolomiten. Abhandlungen der Geologischen Bundesanstalt Band XXIV, I und II Teil, 376 p., III Teil, 89 p.
- Okuneva, T. M. 1985. Norijskie dvustvorchatye mollyuski i ammonoidei Zabajkaljya [Norian bivalves and ammonoids of Zabajkal]. In E. A. Modzalevskaya & T. M. Okuneva, eds., Ezhegodnik Vsesoyuznogo Paleontologicheskogo Obshchestva, Akademiya Nauk SSSR [Annual All-Sovietic Union Palaeontological Association, USSR Academy of Sciences] XXVIII:148–168. In Russian.
- Okuneva, T. M. 1986. Rod *Otapiria* (Monotidae, Bivalvia) [The *Otapiria* genus (Monotidae, Bivalvia)]. *In* Y. D. Zakharov & Y. I. Onoprienko, eds., Permo-Triasovye sobytiya v razvitii organicheskogo mira sevrovostochnoj Azii [Permian-Triassic events during the evolution of the

North-East Asia Biota]. Dalnevostochnyi Nauchnyi Centr Akademii Nauk SSSR [Far-Eastern Scientific Centre, USSR Academy of Sciences]. Vladivostok. p. 57–73. In Russian.

- Okuneva, T. M. 1987. Norijskie galobii Yuzhnykh rajonov Vostoka SSSR [Norian halobiids from the south regions of East USSR]. *In* Y. D. Zakharov & Y. I. Onoprienkko, eds., Problemy biostratigrafii permi i triasa vostoka SSSR [Problems of the Permian and Triassic biostratigraphy of East USSR]. Dalnevostochnyi Nauchnyi Centr Akademii Nauk SSSR [Far Eastern Scientific Centre, USSR Academy of Sciences]. Vladivostok. p. 88–104. In Russian.
- Oliver, P. G., & A. M. Holmes. 2006. The Arcoidea (Mollusca: Bivalvia): a review of the current phenetic-based systematic. *In* Rüdiger Bieler FLS, ed., Bivalvia–a look at the Branches. Zoological Journal of the Linnean Society 148:237–251.
- Onoue, Tetsuji, & Hitoshi Tanaka. 2002. Discovery of Upper Triassic bivalves from Sambosan Subterrane, Itsuki-mura area, Kumamoto Prefecture, and its geologic implication. Journal of Geological Society of Japan 108(9):610–613. In Japanese with English abstract.
- Onoue, Tetsuji, & Hitoshi Tanaka. 2005. Late Triassic bivalves from Sambosan accretionary complex, southwest Japan, and their biogeographic implications. Paleontological Research 9(1):15–25.
- Oppel, A. 1856–1858. Die Juraformation Englands, Frankreichs und des südwestlichen Deutschlands. Jahrbuch Verein für Vaterländische Naturkunde Württemberg, Stuttgart. 12:121–556 (= 1-438) [1856]; 13:141–396 (= 439-694) [1857]; 14:129–291 (= 695-857) [1858].
- d'Orbigny, Alcide D. 1844–1848. Tome 3, Terrains Crétacés, Lamellibranches. *In* A. D. d'Orbigny, G. H. Cotteau, Mr. Piette, E. Eudes-Deslongchamps, P. de Loriol-Le Fort, G. Saporta, E. de Fromentel, & Mr. Ferry, 1840–1894, Paléontologie Française. Description Zoologique et Géologique de Tous le Animaux Mollusques et Rayonnés Fossiles de France: Comprenant leur Application à la Reconnaissance des Couches. Published by the author. Paris. 807 p., pl. 237–489, 242 bis, 247 bis, 186 bis.
- d'Orbigny, Alcide D. 1850. Prodrome de Paléontologie Stratigraphique Universelle des Animaux and Mollusques et Rayonnés: Faisant Suite au Cours Élémentaire de Paléontologie et de Geologie Stratigraphiques, vol. 1. Victor Masson. Paris. 394 p.
- d'Orbigny, Alcide D. 1852. Prodrome de Paléontologie Stratigraphique Universelle des Animaux and Mollusques et Rayonnés: Faisant Suite au Cours Élémentaire de Paléontologie et de Geologie Stratigraphiques, vol. 3. Victor Masson. Paris. 189 p.
- d'Orbigny, Alcide D. 1853. Mollusques. *In* Arthus Bertrand, ed., Histoire Physique, Politique et Naturelle de l'ile de Cuba, tome Premier. Paris. 264 p.
- Özdikmen, Hüseyin. 2008. *Neodietrichia* nom. nov., a replacement name for the genus *Dietrichia* Crosby & Bishop, 1933 (Arenea: Linyphiidae) non Reck, 1921. Munis Entomology and Zoology 3(1):537–538.
- Paleobiology Database (PBDB). 2005. Fürsich, F. T. Primary collection data of Cassian Formation used for publication of Fürsich and Wendt (1977). http://paleodb.org/cgi-bin/bridge.pl?a=displayReference&reference_ no=9188 Checked January 2013.
- Palmer, Charles P. 1975. The British Lower Jurassic species of the bivalve genus *Cardinia*. Bulletin of the British Museum (Natural History), Geology 26(1):1–44.
- Palmer, T. J. 1979. The Hampen Marly & White Limestone formations: Florida-type carbonate in the Jurassic of central England. Palaeontology 22:189–228.
- Palmer, T. J. 1984. Revision of the bivalve family Pulvinitidae Stephenson, 1941. Palaeontology 27(4):815–824.
- Parkinson, J. 1811. Organic Remains of a Former World, vol. 3. London. 479 p., 22 pl.
- Parona, C. F. 1888. Contributo allo studio dei Megalodonti. Atti della Società Italiana di scienze naturali 30:1–10.

- Parona, C. F. 1889. Studio monográfico della fauna Raibliana di Lombardia. Memoria dal Regio Istituto Lombardo di Science e Lettere. 156 p.
- Parona, C. F. 1890. I fossili del Lias inferiore di Saltrio in Lombardia. Atti della Società Italiana di Szience Naturali 33:1–37.
- Patte, Étienne. 1926. Etudes paléontologiques relatives a la géologie de l'est du Tonkin (Paléozoïque et Trias). Bulletin du Service Géologique de l'Indochine 15(1):1-231. (*)
- Patte, Etienne. 1935. Fossiles Paléozoîques et Mésozoîques du sud-ouest de la China. Palaeontologia Sinica (series B) 15(2):1–50.
- Payevskaya, E. B. 1985. Pozdnetriasovye Mollyuski Roda Monotis Bronn (Opyt logiko-matematicheskogo issledovaniya) [The late Triassic Molluscan genus Monotis Bronn (an experiment in logical-mathematical research)]. In A. N. Oleynikov, ed., Nedra. Leningrad. Tom 324. 144 p. In Russian.
- Pérez [-d'Angelo], Ernesto. 1982. Bioestratigrafía del Jurásico de Quebrada Asientos, Norte de Potrerillos, Región de Atacama. Boletín del Servicio Nacional Geológico y Minero de Chile 30:1–149.
- Pérez[-d'Angelo], Ernesto, Martin Aberhan, Renato Reyes[-Bianchi], & Axel von Hillebrandt. 2008. Lower Jurassic Bivalvia of northern Chile. Part III. Order Trigonioida. Beringeria 39:51–102.
- Pérez[-d'Angelo], Ernesto, & Renato Reyes[-Bianchi]. 1991. El orden Trigonioida (Mollusca Bivalvia) en el Mesozoico de Sudamérica. 6º Congreso Geológico Chileno 1991. Actas. Resúmenes expandidos, vol. 1. Servicio Nacional de Geología y Minería. Santiago. p. 72–76.
- Pérez[-d'Angelo], Ernesto, & Renato Reyes[-Bianchi]. 1994. Presencia de *Maoritrigonia* Fleming, 1962 (Bivalvia, Minetrigoniidae) en el Triásico superior de Sudamérica: descripción de dos nuevas especies. Revista Geológica de Chile 21(1):105–117.
- Pérez[-d'Angelo], Ernesto, Renato Reyes[-Bianchi], & S. E. Damborenea. 1995. El género *Groeberella* Leanza, 1993 y Groeberellidae nov. (Bivalvia; Trigonioida) del Jurásico de Chile y Argentina. Revista Geológica de Chile 22:143–157.
- Pérez-Barría, L. P. 2004. Estudio taxonómico de la fauna de Invertebrados del Triásico del Biobío (VII Región, Chile): una aproximación preliminar. MSc Thesis. Universidad de Concepción. 87 p. Unpublished.
- Pérez-Barría, L. P. 2006. Revisión de la fauna de moluscos del Triásico del Bío-Bío (Triásico Superior de la VIII Región, Chile): implicancias paleobiogeográficas, paleooceanográficas y sobre las afinidades presentes y pasadas entre Chile y Nueva Zelanda. *In S. Palma, P. Arana & A.* Guerrero, eds., Resúmenes del XXV Congreso de Ciencias del mar y XI Congreso Latinoamericano de Ciencias del mar. p. 177–188.
- Pérez-López, A. D. 1991. El Trias de Fácies Germánica del Sector Central de la Cordillera Bética. PhD thesis. Universidad de Granada. 400 p.
- Pérez-López, A. D., J. Fernández, Nuria Solé de Porta, & Ana Márquez-Aliaga. 1991. Bioestratigrafía del Triásico de la zona Subbética (Cordillera Bética). Revista Española de Paleontología No. Extra. p. 139–151.
- Pérez-Varela, J. A., Sonia Ros, Fernando Pérez-Varela, L. A. Pérez-Varela, M. R. Alcalde-Fuentes, & Ana Márquez-Aliaga. 2007. Hallazgo de Fimbriidae y Pachycardiidae (Bivalvia) en areniscas y lutitas de Facies Keuper del Triásico Superior de Calasparra (Murcia). *In* J. C. Braga, A. Checa, & M. Company, eds., Libro de Resúmenes. XXIII Jornadas de la Sociedad Española de Paleontología. Instituto Geológico y Minero de España y Universidad de Granada. p. 179–180.
- Petrova, G. T. 1947. Class Lamellibrachiata. In G. Krimholtz, ed., Atlas rukovodyashchikh form iskopaemÿkh faun SSSR [Atlas of the guide forms of the fossils faunas of the USSR]. vol. VIII. The Lower and Middle Jurassic. USSR Ministry of Geology. Moscow. 102–141. In Russian.
- Philippi, Emil. 1898. Die fauna des unteren Trigonodus-Dolomits vom Hühberfeld bei Schwieberdingen und des sogenannten "Cannstatter Kreidemergels." Jahreshefte des Vereins für Vaterländische Naturkunde in Württemberg 54:145–227.
- Philippi, Emil. 1899. Beiträge zur Morphologie und Phylogenie der Lamellibranchier. Zeitschrift der Deutschen geologischen Gesellschaft 50(4):597–622, table 19.

- Philippi, E. 1900. Beiträge zur Morphologie und Phylogenie der Lamellibranchier. III. *Lima* und ihre Untergattungen. Zeitschrift der Deutschen Geologischen Gesellschaft 52(4):619–639.
- Phillips, John. 1829. Illustrations of the Geology of Yorkshire; or a Description of the Strata and Organic Remains of the Yorkshire Coast: Accompanied by a Geological Map, Sections, and Plates of the Fossil Plants and Animals. Thomas Wilson & Sons. York. 192 p., 14 pl.
- Phillips, John. 1836. Illustrations of the Geology of Yorkshire; or a Description of the Strata and Organic Remains: Accompanied by a Geological Map, Sections, and Diagrams, and Figures of the Fossils. Part II. The Mountain Limestone District. John Murray. London. 253 p., 24 pl.
- Pictet, F. J., & G. Campiche. 1864–1867. Descriptions des fossils du terrain Crétacé des environs de Sainte-Croix, troisième partie. H. Georg Libraire. Geneve. 558 p.
- Pinna, Giovanni, & Giorgio Teruzzi. 1991. Il giacimento paleontologico di Besano. Natura (Milan) 82(1):1–55.
- Pisera, Andrzej. 1987. Boring and nestling organisms from Upper Jurassic coral colonies from Northern Poland. Acta Palaeontologica Polonica 32(1–2):83–104.
- Poel, Luc van de. 1959. Faune malacologique du Hervien. Troisième note (première partie). Bulletin de l'Institut Royal des Sciences Naturelles de Belgique, Sciences de la Terre 35:1–26, 1 pl.
- Pojeta, Jr., John. 1971. Review of Ordovician pelecypods. U. S. Geological Survey Professional Paper 695:46 p.
- Pojeta, Jr., John. 1988. The origin and Paleozoic diversification of solemyoid pelecypods. New Mexico Bureau of Mines and Mineral Resources Memoir 44:201–271.
- Pojeta, Jr., John, & T. J. Palmer. 1976. The origin of rock boring in mytilacean pelecypods. Alcheringa 1:167–179.
- Pojeta, Jr., John, Ian Speden, Alan Beu, & Bruce Runnegar. 1971. Review of Treatise on Invertebrate Paleontology: Part N, Mollusca 6, Bivalvia. Journal of Paleontology 45(1):144–149.
- Polubotko, I. V. 1966. Rod Ochotomya (Bivalvia) iz Verkhnego Triasa Severo-Vostoka SSSR [Genus Ochotomya (Bivalvia) from Upper Triassic of North-eastern Russia]. Paleontologicheskij Zhurnal [Paleontological Journal] 3:13–21. In Russian.
- Polubotko, I. V. 1968a. Nekotorye pozdnetriasovye *Lima* i *Plicatula* Severo-Vostoka SSSR [Some Upper Triassic *Lima* and *Plicatula* from North-Eastern USSR]. *In* V. P. Markovskii, ed., Novye vidy drevnikh rasteniï i bespozvonochnykh SSSR, Vypusk II [New species of prehistoric plants and invertebrates of the USSR, part 2]. Chast'I. VSEGEI, Izd-vo 'Nedra'. Moskva. p. 224–227. In Russian.
- Polubotko, I. V. 1968b. Dvustvorchatye Mollyuski nizhnej i srednej yury [Bivalve Mollusks from the Lower and Middle Jurassic]. *In* A. F. Efimova, V. R. Kinasov, K. V. Paraketzov, I. V. Polubotko, Yu. S. Repin, & A. S. Dagis, eds., Polevoj Atlas Yurskoj Fauny I Flory Severo-Vostoka SSSR [Field Atlas of Jurassic fauna and flora of North-Eastern USSR]. Magadans Knizh- Izd-vo. Magadan. p. 29–50, 59–99. In Russian.
- Polubotko, I. V. 1984. Zonal and correlation Significance of Late Triassic halobiids. Sovetskaya Geologiya [Sovietic Geology] 6:40–51. In Russian. (*)
- Polubotko, I. V. 1988. On the morphology and systematics of the Late Triassic Halobiidae (bivalve mollusks). Annual of the All-Union Paleontological Society 31:90–103. In Russian.
- Polubotko, I. V. 1990. In I. V. Polubotko, A. I. Alabushev, & Y-M- Bychkov, Late Triassic halobiids (Bivalve Mollusks) from the Kenkern Range (Northeast USSR). Annual of the All-Union Paleontological Society 33:122–139. In Russian.
- Polubotko, I. V. 1992. Inotseramovye dvustvorki nizhnej i srednej yury severo-vostoka SSSR i severa Sibiri [Lower and Middle Jurassic inoceramid bivalves of the northeastern USSR and northern Siberia]. *In* T. D. Zonova & K. O. Rostovtsev, eds., Atlas Rukovodyashchikh grupp fauny Mezozoya yuga i vostoka SSSR [Atlas of the main groups of the Mesozoic fauna from South and East USSR]. Trudy Vsesoyuznyy Ordena Lenina

Nauchno-Issledovatel'-skiy Geologichsesky Institut im A. P. Karpinskogo (VSEGEI), Novaya Seriya 350:56–79. In Russian.

- Polubotko, I. V. 2010. K Voprosu o Norijskikh i Rehtskikh otlozheniyakh na Severe Sibirskoj Platformy [On the occurrence of Norian and Rhaetian deposits in the North of Siberian Platform]. Novosti paleontologii i stratigrafii (Prilogenie k zhurnalu Geologiya I Geofizika t. 51) [News on Paleontology and Stratigraphy (Geology and Geophysics, Supplement to vol. 51)] 14:51–60. In Russian with English abstract.
- Polubotko, I. V., & L. V. Milova. 1986. Sinemyurskie i Plinsbakhskie Pektinidy severo-vostoka SSSR i ikh stratigraficheskoe znachenie [Sinemurian and Pliensbachian pectinids of the Northeastern USSR and their stratigraphic importance]. *In* A. L. Yanshin & A. S. Dagys, eds., Biostratigrafiya mezozoya Sibiri i Dal'nego Vostoka [Mesozoic biostratigraphy of Siberia and Far East]. Trudy Instituta Geologii i Geofiziki [Proceedings of the Institute of Geology and Geophysics] (Novosibirsk) 648:118–126. In Russian.
- Polubotko, I. V., E. B. Payevskaya, & Y. S. Repin. 2001. Class Bivalvia. In A. Y. Rozanov & A. A. Shevyrev, eds., Atlas triasovykh bespozvonochnykh Pamira [Atlas of the Triassic invertebrates of Pamirs]. Hayka. 190 p. In Russian.
- please provide publisher
- Polubotko, I. V., & Y. S. Repin. 1990. The Triassic/Jurassic boundary in Northeast Asia (Sedimentological and biotic features). Les Cahiers de l'Université Catholique de Lyon (série Sciences) 3:191–205.
- Pompeckj, J. P. 1901. Über Aucellen und Aucellenähnliche Formen. Neues Jahrbuch für Mineralogie, Geologie und Paläontologie 14:319–368.
- Portlock, J. E. 1843. Report on the Geology of the county Londonderry, and Parts of Tyrone and Fermanagh. A. Milliken. Dublin & London. 784 p., 38 pl.
- Posenato, Renato. 1988. The Permian/Triassic boundary in the Western Dolomites, Italy. Review and Proposal. Scienze della Terra 1(3):31–45.
- Posenato, Renato. 1989. Un' associazione oligotipica a *Neoschizodus ovatus* (Goldfuss) della formazione di Werfen (Triassico inf. - Dolomiti). 3º Simposio di Ecologia e Paleoecologia delle Comunità Bentoniche. p. 141–153.
- Posenato, Renato. 2002. Bivalves and other macrobenthic fauna from the Ladinian "Muschelkalk" of Punta del Lavatoio (Alghero, SW Sardinia). Rendiconti della Società Paleontologica Italiana 1:185–196.
- Posenato, Renato. 2008a. Patterns of bivalve biodiversity from Early to Middle Triassic in the Southern Alps (Italy): Regional vs. global events. Palaeogeography, Palaeoclimatology, Palaeoecology 261:145–159.
- Posenato, Renato. 2008b. Anisian (Middle Triassic) bivalves from the Dolomites (Italy). Neues Jahrbuch für Geologie und Paläontologie Abhandlungen 247(1):93–115.
- Posenato, Renato, & Marco Avanzini. 2004. "*Lithiotis*" beds of the Rotzo Formation (Calcari Grigi Group, Lower Jurassic) from Albaredo (Rovereto, Trento). Studi Trentini di Scienze Naturali, Acta Geologica 81:23–28.
- Posenato, Renato, & Antonio Ietto. 1995. Late Triassic Megalodontidae from northern Calabria (Italy). Rivista Italiana di Paleontologia e Stratigrafia 101(3):325–340.
- Posenato, Renato, Pál Pelikán, & Kinga Hips. 2005. Bivalves and brachiopods near the Permian-Triassic boundary from the Bükk Mountains (Bálvány-North section, Northern Hungary). Rivista Italiana di Paleontologia e Stratigrafia 111(2):215–232.
- Posenato, Renato, Dario Sciunnach, & Eduardo Garzanti. 1996. First report of *Claraia* (Bivalvia) in the Servino Formation (Lower Triassic) of the Western Orobic Alps, Italy. Rivista Italiana di Paleontologia e Stratigrafia 102(2):201–210.
- Posenato, Renato, L. Simone, Max Ürlichs, & Angelo Ibba. 2002. The Ladinian Muschelkalk of Punta del Lavatoio (Alghero, NW Sardinia). Rendiconti della Società Paleontologica Italiana 1:283–291.
- Poulton, T. P. 1976. Some lower Jurassic trigoniid bivalves from southwestern British Columbia. Geological Survey of Canada Bulletin 256:41–53.

- Poulton, T. P. 1979. Jurassic trigonid bivalves from Canada and western United States of America. Geological Survey of Canada Bulletin 282:81 p.
- Poulton, T. P. 1991. Hettangian through Aalenian (Jurassic) guide fossils and biostratigraphy, Northern Yukon and adjancent Northwest territories. Geological Survey of Canada Bulletin 410:95 p.
- Pozzi, Renato, Romano Gelati, & Andrea Allasinaz. 1962. Osservazioni stratigrafiche e paleontologiche sulla bassa valle dello Spöl (Livigno-Alpi Retiche). Rivista Italiana di Paleontologia e Stratigrafia 68(1):39–66.
- Prado-Velazco, Isabel. 1991. Los trigoniidae depositados en el Museo de Historia Natural "Javier Prado". 7º Congreso Peruano de Geología (Lima), Volumen de Resúmenes Extendidos. INGEMMET, Lima. p. 443–447.
- Prinz, Peter. 1985. Stratigraphie und Ammonitenfauna der Pucara-Gruppe (Obertrias-Unterjura) von Nord-Peru. Palaeontographica (Abt. A) 188:153–197.
- Ptchelincev, V. F. [P elincev]. 1960 Semejstwo Prospondylidae fam. nov. [Family Prospondylidae fam. nov.]. In A. G. Eberzin, ed., Osnovy Paleontologii. Mollyuski-Pantsirnye, Dvustvorchatye, Lopatonogie [Fundamentals of Paleontology. Mollusks-Loricata, Bivalvia, Scaphopoda]. Isdastelstvo Akademii Nauk SSSR. Moscow. p. 78–87.
- Pugaczewska, Halina. 1986. Bivalvia of the Polish Middle Jurassic and remarks on their paleoecology. Acta Palaeontologica Polonica 31(1–2):27–83.
- Pulteney, Richard. 1799. Catalogues of the Birds, Shells, and Some of the More Rare Plants, of Dorsetshire. From the New and Enlarged Edition of Mr. Hutchins's History of that Country. Nichols. London. 92 p.
- Quenstedt, F. A. 1851–1852. Handbuch der Petrefaktenkunde. Laupp and Siebeck. Tübingen. Volume 1, p. 1–528 (1851), Volume 2, p. 529–792 (1852).
- Quenstedt, F. A. 1856–1858. Der Jura. Laupp and Siebeck. Tübingen. 842 p.
- Quenstedt, Werner. 1930. Die Anpassung an die grabende Lebensweise in der Geschichte der Solenomyiden, und Nuculaceen. Geologische und Paleontologische Abhandlungen (neue Folge) 18(1):120 p., 3 pl.
- Quintero, Indalecio, Antonio Almela, Evaristo Gómez, C. Martínez, & Hermenegildo Mansilla. 1977. El Trías de facies germánica de Alpera. Cuadernos de Geología Ibérica 4:447–454.
- Quiroz-Barroso, S. A., & M. C. Perrilliat. 1998. Pennsylvanian bivalves from the Ixtaltepec Formation, Mexico. Journal of Paleontology 72(6):1011–1024.
- Rafinesque, C. S. 1815. Analyse de la Nature, ou Tableau de l'Univers et des Corps Organisés, etc. C. S. Rafinesque. Jean Barravecchia. Palermo. 223 p.
- Ramovs, Anton, & Bogdan Jurkovsek. 1983a. Ausbildung der Ladin-Schichten oberhalb des Supca-Aufsichtspunktes, Südlich des Vrsic-Pases (Julische Alpen). Geoloski Zbornik 4:81–91.
- Ramovs, Anton, & Bogdan Jurkovsek. 1983b. Ausbildung der Anis-schichten bei Trzic (Sudkarawanken) mit besonderer rücksicht auf die neu entdeckte "illyr"- unterstufe. Geoloski Zbornik 4:37–45.
- Rangel, César. 1978. Fósiles de Lircay-Uruto. Boletín, Serie D: Estudios Especiales, Instituto Geológico Minero y Metalúrgico de Perú 6:35 p.
- Rau, Antonio, & Marco Tongiorgi. 1966. I. Lamellibranchi Triassici del Verrucano dei Monti Pisani. Nuova Revisione. Palaeontographia Italica 61(31):185–234.
- Reck, H. 1921. Über eine neue Faunula im Juragebiet der deutsch-ostafrikanischen Mittellandbahn. Zentralblatt für Mineralogie, Geologie und Palentologie. Stuttgart. 1921:431–436.
- Récluz, C. A. 1848–1849. Description d'un noveau genre de coquilles bivalves nommé septifère (Septifer). Société Cuviérienne, Revue Zoologique 11(9):275–279; (2)1(3):117–137.
- Rehder, Harald A. 1967. Valid zoological names of the Portland Catalogue. Proceedings of the United States National Museum 121(3579):1–51.
- Reid, R. G. B. 1998. Subclass Protobranchia. In P. L. Beesley, G. J. Ross, & A. Wells, eds., Mollusca: The southern synthesis. Fauna of Australia. Volume 5, Part A. CSIRO Publishing. Melbourne. p. 235–247.

- Reis, O. M. 1926. Die Fauna des Wettersteinkalks. III: Gastropoden, Bivalven, Brachiopoden etc. Geognostische Jahreshefte (München), Jahrgang 39:87–188. (*)
- Repin, Y. S. 1996. New Late Triassic bivalves from Iran and a taxonomy of the Superfamily Spondylacea. Paleontologicheskij Zhurnal 1996(3)3–8. Translation to English in Paleontological Journal 30(4):363–369.
- Repin, Y. S. 2001. New taxa of Late Triassic Bivalve Mollusks from Central Iran. Paleontological Journal 35(3):233–242.
- Retallack, G. J., & R. J. Ryburn. 1982. Middle Triassic deltaic deposits in Long Gully, near Otematata, north Otago, New Zealand. New Zealand Journal of Geology and Geophysics 12(3):207–227.
- Retzius, Anders J. 1788. Dissertatio historico-naturalis sistens nova testaceorum genera. Berlingianis. Lundae. 23 p.
- Reyes[-Bianchi], Renato, & Ernesto Pérez[-d'Angelo]. 1980. Quadratojaworskiella nov., a Liassic subgenus of Trigoniidae from Chile. Pacific Geology 14:87–93.
- Riccardi, A. C., S. E. Damborenea, M. O. Mancefiido, & S. C. Ballent. 1991. Hettangian and Sinemurian (Lower Jurassic) biostratigraphy of Argentina. Journal of South American Earth Sciences 4(3):159–170.
- Riccardi, A. C., S. E. Damborenea, M. O. Manceñido, & M. P. Iglesia Llanos. 2004. The Triassic/Jurassic boundary in the Andes of Argentina. Rivista Italiana di Paleontologia e Stratigrafia 110(1):69–76.
- Riccardi, A. C., S. E. Damborenea, M. O. Manceñido, R. Scasso, Silvia Lanés, & M. P. Iglesia Llanos. 1997. Primer registro de Triásico marino fosilífero de la Argentina. Revista de la Asociación Geológica Argentina 52(2):228–234.
- Rieber, Hans. 1968. Die Artengruppe der *Daonella elongata* Mojs. aus der Grenzbitumenzone der mittleren Trias des Monte San Giorgio (Kt. Tessin, Schweiz). Paläontogische Zeitschrift 42(1/2):33–41.
- Rieber, Hans. 1969. Daonellen aus der Grenz-bitumenzone der mittleren Trias des Monte S. Giorgio (Kt. Tessin, Schweiz). Eclogae Geologicae Helvetiae 62(2):657–683.
- Riegraf, Wolfgang. 1977. Goniomya rhombifera (Goldfuss) in the Posidonia Shales (Lias epsilon). Neues Jahrbuch für Geologie und Paläontologie, Monatshefte 7:446–448.
- Röder, H. A. 1882. Beiträge zur Kenntniss des Terrain à Chailles und seiner Zweischaler in der Umgegend von Pfirt im Ober-Elsass. Schultz. Strassburg. 110 p., 4 pl.
- Rodland, D. L., & D. J. Bottjer. 2001. Biotic Recovery from the End-Permian Mass Extinction: Behavior of the Inarticulate Braquiopod *Lingula* as a Disaster Taxon. PALAIOS 16:95–101.
- Roemer, Friedrich Adolph. 1839. Die Versteinerungen des Norddeutschen Oolithen-Gebirges. Ein Nachtrag. Hahn'schen Hofbuchhandlung. Hannover. 59 p., pl. 17–20.
- Roemer, Ferdinand. 1842. De Astartarum genere et speciebus, quae e saxis jurassicis atque cretaceis proveniunt. Unpublished Dissertatio Palaeontologica. Sittenfeld, Berlin. 24 p.
- Roemer, Ferdinand. 1851. Ueber einige neue Versteinerungen aus dem Muschelkalke von Willebadessen. Palaeontographica 1(6):311–315.
- Rollier, L. 1911–1918. Fossiles nouveaux ou peu connus des terrains secundaires (Mésozoiques) du Jura et des contrées environnantes. Abhandlungen Schweizerisches Paläontologisches Gesellschaft 37–44:1–696, 49 pl.
- Romanov, L. F. 1985. Yurskie Pektinoidy Yuga SSSR [Jurassic Pectinoids of southern USSR]. Shtiintsa. Kishinev. 231 p. In Russian.
- Ros, Sonia. 2009. Dinámica de la paleodiversidad de los Bivalvos del Triásico y Jurásico Inferior. PhD Thesis. University of Valencia. Valencia. 564 p., http://www.tesisenred.net/handle/10803/9952. Checked August 2012.
- Ros, Sonia, S. E. Damborenea, & Ana Márquez-Aliaga. 2009. Parainoceramus Cox, 1954 [ex Voronetz, 1936] partim (Bivalvia, Jurásico): un grupo de especies en busca de género. In P. Palmqvist & J. A. Pérez-Clarós, coordinators, Libro de Resúmenes XXV Jornadas de la Sociedad Española de Paleontología. Universidad de Málaga. p. 324–325.

- Rossi Ronchetti, Carla. 1959. Il Trias in Lombardia. I. Lamellibranchi ladinici del grupo delle Grigne. Rivista Italiana di Paleontologia e Stratigrafia 65(4):269–357.
- Rossi Ronchetti, Carla, & Andrea Allasinaz. 1965. Il Trias in Lombardia. (Studi geologici e paleontologici) XI. *Curionia*, nuovo genere di Lamellibranco Eterodonte triassico. Rivista Italiana di Paleontologia e Stratigrafia 71(2):351–412.
- Rossi Ronchetii, Carla, & Andrea Allasinaz. 1966. Il Trias in Lombardia. (Studi geologici e paleontologici) XX. *Pseudomyoconcha*, nuovo genere Triasico di Lamellibranchi Eterodonti. Rivista Italiana di Paleontologia e Stratigrafia 72(4):1083–1132.
- Ruban, D. A. 2006a. Diversity dynamics of the Triassic marine biota in the Western Caucasus (Russia): A quantitative estimation and a comparison with the global patterns. Revue de Paléobiologie 25(2):699–708.
- Ruban, D. A. 2006b. Taxonomic diversity dynamics of the Jurassic bivalves in the Caucasus: Regional trends and recognition of global patterns. Palaeogeography, Palaeoclimatology, Palaeoecology 239:63–74.
- Rubilar, Alfonso. 1998. La superfamilia Ostreacea en Chile y su importancia cronoestratigráfica, paleogeográfica y paleoecológica (Triásico Superior-Jurásico). Unpublished PhD Thesis. Universidad Nacional de La Plata. 363 p.
- Runnegar, Bruce. 1965. The bivalves *Megadesmus* Sowerby & Astartila Dana from the Permian of Eastern Australia. Australian Journal of Earth Sciences 12(2):227–252.
- Runnegar, Bruce. 1974. Evolutionary history of the bivalve subclass Anomalodesmata. Journal of Paleontology 48:904–939.
- Runnegar, Bruce, & N. D. Newell. 1974. *Edmondia* and the Edmondiacea shallow-burrowing Paleozoic Pelecypods. American Museum Novitates 2533:1–19.
- Saadi, Zouhra, Bouaza Fedan, Mohamed Laadila, & Abdelhadi Kaoukaya. 2003. Les tidalites liasiques de la Haute Moulouya et du Moyen Atlas meridional (Maroc): Dynamique sédimentaire et contexte paléogéographique. Bulletin de l'Institut Scientifique, Rabat, section Sciences de la Terre 25:55–71.
- Sacco, Federico. 1897. I molluschi dei terreni Terziarii del Piemonte e della Liguria. Parte XXIV (Pectinidae). Carlo Clausen. Torino. 116 p.
- Saeki, Shiro. 1925. On some new species of Jurassic Trigonia from Province of Tamba, Japan. Journal of the Geological Society, Tokyo 32(373):35–36.
- Saint-Seine, R. 1951. Mimetisme ou "pseudomorphose" chez des lamellibranches fixes sur echinides. Bulletin de la Société Géologique de France 6(8):653–656.
- Salomon, Wilhelm. 1895. Geologische und palaeontologische Sudien uber die Marmolata. Palaeontographica 42:1–210.
- Sánchez, T. M. 2002. Ordovician Bivalvia and Rostroconchia of Argentina: An updated synthesis. *In* F. G. Aceñolaza, ed., Aspects on the Ordovician System. Serie Correlación Geológica 16:195–208.
- Sandberger, F. 1864. Beobachtungen im mittleren Jura des badischen Oberlandes. Würzburger naturwissenschaftliche Zeitschrift 5:1–22.
- Sanin, V. Ya. 1973. Taimyrodon–Novyj Rod paleotaksodont iz nizhnemelovykh otlozhenij na severe Sibiri [Taimyrodon–New genus of paleotaxondont from Lower Cretaceous deposits of northern Siberia]. Geologiya Geofizika [Soviet Geology and Geophysics] 9:92–98. In Russian.
- Sano, Shin-ichi, Yojiro Taketani, Muneo Taira, Yasuo Yamaki, Yoshimi Ara, Yoshihiro Morino, & Yasuo Kondo. 2010. Discovery of a pinnid bivalve, *Trichites*, from the Kimmeridgian-Tithonian Nakanosawa Formation of the Somanakamura Group, Northeast Japan. Bulletin of Fukushima Museum 24:31–40. In Japanese with English abstract.
- Sato, Tadashi, & G. E. G. Westermann. 1991. Jurassic taxa ranges and correlation charts for the Circum Pacific. 4. Japan and south-east Asia. Newsletters on Stratigraphy 24(1–2):81–108.
- Savazzi, Enrico. 1984. Functional morphology and autecology of *Pseudoptera* (Bakevelliid bivalves, Upper Cretaceous of Portugal). Palaeogeogaphy, Palaeoclimatology, Palaeoecology 46:313–324.

- Savazzi, Enrico. 2001. A review of simbiosis in the Bivalvia, with special attention to macrosymbiosis. Paleontological Research 5(1):55–73.
- Scandone, Paolo, & Paola Capoa de Bonardi. 1966. Sulla posizione stratigrafica e l'età dei livelli a *Daonella* e ad *Halobia* in Lucania. Bulletino della Società dei Naturalisti in Napoli 75:30–39.
- Scarlato, Orest A., & Yaroslav I. Starobogatov. 1979. Osnovnye cherty evoliutsii i sistema klassa Bivalvia [General evolutionary patterns and the systematics of the Class Bivalvia]. *In* Yaroslav I. Starobogatov, ed., Morfologiia, sistematika i filogeniia molliuskov [Morphology, Systematics and Phylogeny of Mollusks]. Akademiia Nauk SSSR, Zoologicheskii Institut, Trudy 80:5–38.
- Schafhäutl, K. E. 1852. Geognostische Bemerkungen über den Krammeberg usf. Neues Jahrbuch für Mineralogie, Geologie und Paläontologie 1852:129–286.
- von Schafhäutl, K. F. E. 1863. Süd-Bayern's Lethea Geognostica. Der Kressenberg und die Südlich von ihm Gelegenen Hochalpen Geognostich Betrachtet in ihren Petrefacten. Leopold Voss, Leipzig, Williams and Norgate, London, and F. Savy, Paris. Vo. 1, Text: xvi p., 487 p. Vol. 2: Atlas zu Süd-Bayern's Lethea Geognostica. Einhundert Tafeln Enthaltend 1758 Original Abbildungen und 2 Karten, viii p., 100 pl., 2 maps.
- Schatz, Wolfgang. 2001a. Taxonomic significance of biometric characters and the consequences for classification and biostratigraphy, exemplified through moussoneliform daonellas (*Daonella*, Bivalvia; Triassic). Paläontologische Zeitschrift 75(1):51–70.
- Schatz, Wolfgang. 2001b. Revision der Untergattung Daonella (Pichlerella) (Bivalvia, Ladin). Eclogae Geologicae Helvetiae 94:389–398.
- Schatz, Wolfgang. 2004. Revision of the subgenus Daonella (Arzelella) (Halobiidae; Middle Triassic). Journal of Paleontology 78(2):300–316.
- Schatz, Wolfgang. 2005. Palaeoecology of the Triassic black shale bivalve Daonella —New insights into an old controversy. Palaeogeography, Palaeoeclimatology, Palaeoecology 216:189–201.
- Schinz, H. R. 1822. Das Thierreich eingetheilt nach dem Bau der Thiere als Grundlage ihrer Naturgeschichte und der vergleichenden Anatomie. Mit vielen Zusätzenversehen von H. R. Schinz. Das Thierreich vol. 2. Cotta. Stuttgart & Tübingen. p. i–xvi + 1–835. A translation, with emendations, of Cuvier's *Règne animal.*
- Schlotheim, E. F. v. 1820. Die Petrefaktendunde auf ihrem jetzigen Standpunkte durch die Beschreibung seiner Sammlung versteinerter und fossiler Überreste des Thier- und Pflanzenreichs der Vorwelt erfäutert. Becker. Gotha, Germany. 437 p.
- Schlotheim, E. F. v. 1822–1823. Nachträge zur Petrefactenkunde, zweite Abteilung. Becker'sche Buchhandlung. Gotha. 114 p., 16 pl.
- Schmarda, L. K. 1859. Neue wirbellose Thiere. 1(1). Neue Turbellarien, Rotatorien und Anneliden. Verlag von Wilhelm Engelmann, Leipzig. 66 p, 15 pl.
- Schmidt, Martin. 1935. Fossilien der spanischen Trias. Abhandlungen der Heidelberger Akademie der Wissenschaften, Mathematisch-Naturwissenschaftliche Klasse 22:1–140.
- Schneider, J. A. 1995. Phylogeny of the Cardiidae (Mollusca, Bivalvia): Protocardiinae, Laevicardiinae, Lahilliinae, Tulongocardiinae subfam. n. and Pleuriocardiinae subfam. n. Zoologica Scripta 24(4):321–346.
- Schneider, J. A., & J. G. Carter. 2001. Evolution and phylogenetic significance of Cardioidean shell microstructure (Mollusca, Bivalvia). Journal of Paleontology 75(3):607–643.
- Schneider, J.A., & A. Kaim. 2012. Early ontogeny of Middle Jurassic hiatellids from a wood-fall association: implications for phylogeny and palaeoecology of hiatellidae. Journal of Molluscan Studies 78:119-127.
- Schnetzer, R. 1934. Die Muschelkalkfauna des Öfenbachgrabens bei Saalfelden. Palaeontographica 81:1–160.
- Scholz, Annemarie, Martin Aberhan, & C. M. González-León. 2008. Early Jurassic bivalves of the Antimonio Terrane (Sonora, NW Mexico): Taxonomy, biogeography, and paleogeographic implications. *In* R. B. Blodgett & G. D. Stanley, eds., The Terrane Puzzle: New Perspectives

on Paleontology and Stratigraphy from the North American Cordillera. Geological Society of America Special Paper 442:267–310.

- Schubert, J. K. 1993. Rebound from the Permian–Triassic mass extinction event: paleoecology of Lower Triassic carbonates in the Western U.S.A. PhD Thesis. University of Southern California. Los Angeles. 396 p.
- Schubert, J. K., & D. J. Bottjer. 1995. Aftermath of the Permian-Triassic mass extinction event: Paleoecology of Lower Triassic carbonates in western United States. Palaeogeography, Palaeoclimatology, Palaeoecology 116:1–39.
- Schumacher, Crétien Frédéric. 1817. Essai d'un nouveau système des habitations de vers testacés. Schultz. Copenhagen. 278 p., 22 pl.
- Scopoli, J. A. 1777. Introductio ad historiam naturalem sistens genera Lapidum, Plantarum, et Animalium hactenus detecta, caracteribus essentialibus donata, in Tribus divisa, subinde ad leges naturae. Wolfgangum Gerle. Prague. 506 + 36 p.
- Scott, P. J. B. 1988. Initial settlement behaviour and survivorship of *Lithophaga bisulcata*. Journal of Molluscan Studies 54:97–108.
- Seeling, Jens, & Peter Bengtson. 1999. Cenomanian oysters from the Sergipe Basin, Brazil. Cretaceous Research 20:747–765.
- Seguí, Julio. 1999. Una nueva *Pinna* (Bivalvia) del Triásico de Alcover-Montral. Batalleria 9:21–22.
- Seilacher, Adolf. 1954. Ökologie der Triassischen Muschel *Lima lineata* (Schloth.) und ihrer Epöken. Neues Jahrbuch für Geologie und Paläontologie Monatshefte 4:163–183.
- Seilacher, Adolf. 1972. Divaricate patterns in pelecypod shells. Lethaia 5:325–343.
- Seilacher, Adolf. 1984. Constructional morphology of bivalves: Evolutionary pathways in primary versus secondary soft-bottom dwellers. Palaeontology 27(2):207–237.
- Seilacher, Adolf. 1990. Aberrations in bivalve evolution related to photo- and chemosymbiosis. Historical Biology 3:289–311.
- Seilacher, Adolf, B. A. Matyja, & A. Wierzbowski. 1985. Oyster beds: morphologic response to changing substrate conditions. *In* U. Bayer & A. Seilacher, eds., Sedimentary and evolutionary cycles. Lecture Notes in Earth Sciences 1:421–435.
- Senkowiczowa, Hanna. 1985. Fauna z osadów retu i wapienia muszlowego na Ni u Polskim [The Raetian and Muschelkalk fauna in the Polish Lowlands]. Prace Instytutu Geologicznego CXVII:5–41. In Polish.
- Sepkoski, Jr., J. J. 2002. A compendium of fossil marine animal genera. In D. Jablonski & M. Foote, eds., Bulletins of American Paleontology 363:560 p.
- Septfontaine, Michel. 1995. Catalogue des types paléontologiques déposés au Musée cantonal de Géologie, Lausanne. Mémoires de Géologie (Lausanne) 26:69 p.
- Sey, I. I., Y. S. Repin, E. D. Kalacheva, T. M. Okuneva, K. V. Paraketsov, & I. V. Polubotko. 1981. Eastern Russia. *In* G. E. G. Westermann, ed., The Jurassic of the Circum-Pacific. World and Regional Geology 3. Cambridge University Press. Cambridge. p. 225–245.
- Sha Jingeng. 1991. Larval shell of *Aucellina* and its ecological, biogeographical and classificatory significance. *In* Jin Y., Wang J., & Xu S., eds., Palaeoecology of China. Nanjing University Press. Nanjing. 1:249–265.
- Sha Jingeng, ed. 1995. Palaeontology of the Hoh Xil Region, Qinghai. The series of comprehensive scientific expedition to the Hoh Xil Region. Science Press. Beijing. p. 1–177.
- Sha Jingeng. 1996. Antitropicality of the Mesozoic Bivalves. In Pang Z. H., Zhang J., & Sun J., eds., Advances in Solid Earth Sciences. Science Press. Beijing. p. 90–98.
- Sha Jingeng. 1998. Characteristics of stratigraphy and palaeontology of Hohxil, Qinghai: Geographic significance. Acta Palaeontologica Sinica 37(1):85–96. In Chinese with English abstract.
- Sha Jingeng. 2002. Distribution patterns of the Jurassic ostreids (Bivalvia) from Tanggula of China. Science in China (series D) 44 Supp:112–120.

- Sha Jingeng. 2003. Plankton and pseudoplankton of the marine mesozoic bivalves. Acta Palaeontologica Sinica 42(3):408–416. In Chinese with English abstract.
- Sha Jingeng, Chen Chuzhen, & Qi Liangzhi. 1990. Bivalves of the Middle and Upper Triassic in Yushu Region, Qinghai. *In* Quinghai Institute of Gological Sciences, Nanjing Institute of Geology and palaeontology, Academia Sinica, ed., Devonian-Triassic stratigraphy and palaeontology from Yushu Region of Qinghai, China. Vol. I. Najing University Press. Nanjing. p. 133–234. In Chinese with English abstract.
- Sha Jingeng, & F. T. Fürsich, 1993. Biostratigraphy of the Upper Jurassic– Lower Cretaceous bivalves *Buchia* and *Aucellina* of eastern Heilongjiang, northeast China. Geological Magazine 130(4):533–542.
- Sha Jingeng, & F. T. Fürsich. 1994. Bivalve faunas of eastern Heilongjiang, Norheastern China. II. The Late Jurassic and Early Cretaceous buchiid fauna. Beringeria 12:3–93.
- Sha Jingeng, F. T. Fürsich, P. L. Smith, & Lijun Wang. 1998. Palaeotaxodonta, Pteriomorphia, and Isofilibranchia (Bivalvia) from the Jurassic of the main ridge of the Tanggula Mountains, Qinghai–Xizang Plateau, China. Beringeria 21:3–55.
- Sha Jingeng, & J. A. Grant-Mackie. 1996. Late Permian to Miocene bivalve assemblages from Hohxil, Qinghai-Xizang Plateau, China. Journal of the Royal Society of New Zealand 26(4):429–455.
- Sha Jingeng, & Jiang Baoyu. 2004. Potential nonmarine Triassic/Jurassic boundary in Xinjiang, NW China. In Abstracts of 32nd IGC Florence 2004, Part 2:1138.
- Sha Jingeng, A. L. A. Johnson, & F. T. Fürsich. 2004. From deep-sea to high mountain ranges: palaeogeographic and biotic changes in Hohxil, the source area of the Yangtze River (Tibet Plateau) since the Upper Palaeozoic. Neues Jahrbuch für Geologie und Paläeontologie Abhandlungen 233:169–195.
- Sha Jingeng, Lin Li, Zhu Lidong, Pang Yanchun, Fu Xiugen, & Wang Xinli. 2005. Distribution pattern of Upper Triassic *Pergamidia* (Bivalvia) and its significance. Geo-Temas 8:189–191.
- Sha Jingeng, P. L. Smith, & F. T. Fürsich. 2002. Jurassic Ostreoida (Bivalvia) from China (Tanggula Mountains, Qinghai-Xizang Plateau) and their paleobiogeographic context. Journal of Paleontology 76(3):431–446.
- Sha Jingeng, Wang JianPo, Galina Kirillova, Pan YanHong, Cai HuaWei, Wang YaQiong, Yao XiaoGang, & Peng Bo. 2009. Upper Jurassic and Lower Cretaceous of Sanjiang-Middle Amur basin: Non-marine and marine correlation. Science in China (series D: Earth Sciences) 52(12):1873–1889.
- Sharpe, Daniel. 1850. On the secondary District of Portugal which lies to the North of the Tagus. Quarterly Journal of the Geological Society of London 6:135–195.
- Shen Shuzhong, He Xilin, & Shi Guangrong. 1995. Biostratigraphy and correlation of several Permian-Triassic boundary sections in southwestern China. Journal of Southeast Asian Earth Sciencies 12(1-2):19-30.
- Shimer, H. W. 1926. A Triassic coral reef fauna in British Columbia. Museum Bulletin Geological Survey of Canada, 42 Geology (series 45):85–89.
- Shrank, F. von P. 1798. Durchgedachte Geschichte der in Baieren einheimischen und zahmen Thiere. Nürnberg. Fauna Boica 1:1–720.
- Shurygin, B. N., & O. A. Lutikov. 1991. Nizhneyurskie pektinidy severa aziatskoi chasti SSSR. Trudy Instituta Geologii i Geofiziki, Sibirskoe Otdelenie 769:47–78, pl. 13–18. In Russian.
- Siblík, Miloš, István Szente, Radek Mikulaš, & Harald Lobitzer. 2010. An Invertebrate Faunula in the Kössen Beds of Starnkogel (Bad Ischl, Upper Austria). Abhandlungen der Geologischen Bundesanstalt 65:57–64.
- Silberling, N. J. 1961. Upper Triassic marine mollusks from the Natchez Pass Formation in Northwestern Nevada. Journal of Paleontology 35(3):535–542.

- Silberling, N. J., J. A. Grant-Mackie, & K. M. Nichols. 1997. The Late Triassic bivalve *Monotis* in accreted terranes of Alaska. U. S. Geological Survey Bulletin 2151:1–21.
- Silberling, N. J., & K. M. Nichols. 1982. Middle Triassic molluscan fossils of biostratigraphic significance from the Humboldt Range, northwestern Nevada. U. S. Geological Survey Professional Paper 1207:1–77.
- Simões, M. G., & F. C. Fittipaldi. 1987. Bivalves do Grupo Passa Dois, Permiano da Bacia do Paraná: Sinopse das pesquisas. Atas do 6º Simposio Regional de Geologia, SBG/SP. Rio Claro. p. 281–295.
- Simões, M. G., A. C. Marques, L. H. Mello, & L. E. Anelli. 1997. Phylogenetic analysis of the genera of extinct family Megadesmidae (Bivalvia, Anomalodesmata), with remarks on its paleoecology and taxonomy. Journal of Comparative Biology 2(2):75–90.
- Simonelli, V. 1884. Faunula del calcare ceroide di Campiglia Marittima (Lias Inferiore). Atti Società Toscana Scienze Naturale, Pisa, Memorie 6(1):111–128.
- Simpson, Martin. 1855. The fossils of the Yorkshire Lias: Described from nature, with a carefully measured section of the strata and the fossils peculiar to each. Whittacker. London. 149 p. (*)
- Singh, C. S. P., & S. Kanjilal, 1977. *Habonucula*, a new nuculid (Bivalvia) genus from Jurassic rocks of Kutch (Gujarat), W. India. Journal of the Geological Society of India 18(4):189–193.
- Sirna, Giuseppe. 1968. Fossili retici dei Monti di Amelia (Umbria). Rivista Italiana di Paleontologia e Stratigrafia 74(3):747–802.
- Skeat E. G., & V. Madsen. 1898. On Jurassic, Neocomian and Gault boulders found in Denmark. Danmarks geologiske Undersøgelse 2(8):1–213. Copenhagen.
- Skelton, P. W. 1978. The evolution of functional design in rudists (Hippuritacea) and its taxonomic implications. Philosophical Transactions of the Royal Society of London (Series B) 284:305–318.
- Skelton, P. W., & M. J. Benton. 1993. Mollusca: Rostroconchia, Scaphopoda and Bivalvia. In M. J. Benton, ed., The Fossil Record 2. Chapman & Hall. London. p. 237–263.
- Skwarko, S. K. 1967. First Upper Triassic and ?Lower Jurassic marine Mollusca from New Guinea. *In* S. K. Skwarko, ed., Mesozoic Mollusca from Australia and New Guinea. Bureau of Mineral Resources, Geology and Geophysics 75:41–82.
- Skwarko, S. K. 1981. Spia, a new Triassic bakevelliid bivalve from Papua New Guinea. Bureau of Mineral Resources, Geology and Geophysics, Palaeontological Papers 209:63–64.
- Skwarko, S. K. 1983. Somareoides hastatus (Skwarko), a new Late Triassic bivalve from Papua New Guinea. Bureau of Mineral Resources, Australia, Bulletin 217:67–71
- Smith, J. P. 1927. Upper Triassic Marine Invertebrate faunas of North America. U. S. Geological Survey Professional Paper 141:1–262.
- Smith, P. L., & H. W. Tipper. 1986. Plate tectonics and paleobiogeography: Early Jurassic (Pliensbachian) endemism and diversity. PALAIOS 1(4):399–412.
- Sowerby, G. B. I. 1821–1825. The Genera of Recent and Fossil Shells, For the Use of Students in Conchology and Geology, vol. 1. G. B. Sowerby. London. 126 pl. + text unpaginated.
- Sowerby, G. B. II. 1872. Conchologia Iconica or, Illustrations of the Shells of Molluscous Animals. 18. Monograph of the genus Lima. L. Reeve & Co. London. 11 p., 5 pl.
- Sowerby, James. 1813. The Mineral Conchology of Great Britain; or Coloured Figures and Description of those Remains of Testaceous Animals or Shells Which Have Been Preserved at Various Times and Depths in the Earth, vol. 1, no. 3–8. B. Meredith. London. p. 33–96, pl. 10–44.
- Sowerby, James. 1814. The Mineral Conchology of Great Britain; or Coloured Figures and Description of those Remains of Testaceous Animals or Shells Which Have Been Preserved at Various Times and Depths in the Earth. B. Meredith. London. 1(9–14):97–178, pl. 45–78.

- Sowerby, James. 1816. The Mineral Conchology of Great Britain; or Coloured Figures and Description of those Remains of Testaceous Animals or Shells Which Have Been Preserved at Various Times and Depths in the Earth. B. Meredith. London. 2(21–26):29–116, pl. 115–150.
- Sowerby, James. 1817. The Mineral Conchology of Great Britain; or Coloured Figures and Description of those Remains of Testaceous Animals or Shells Which Have Been Preserved at Various Times and Depths in the Earth. B. Meredith. London. 2(27–32):117–194, pl. 151–186.
- Sowerby, James. 1818. The Mineral Conchology of Great Britain; or Coloured Figures and Description of those Remains of Testaceous Animals or Shells Which Have Been Preserved at Various Times and Depths in the Earth. B. Meredith. London. Vol. 3(36–38):1–40, pl. 204–221.
- Sowerby, James. 1819. The Mineral Conchology of Great Britain; or Coloured Figures and Description of those Remains of Testaceous Animals or Shells Which Have Been Preserved at Various Times and Depths in the Earth. B. Meredith. London. 3(39–44):41–98, pl. 222–253.
- Sowerby, James. 1842–1844. Mineral-Conchologie Grossbritanniens, oder ausgemalte Abbildungen u. Beschreibungen der Schalthier-Ueberreste, welche zu verschied. Zeiten und in verschied. Tiefen der Erde erhalten worden sind. Deutsch bearbeitet von Ed. Desor. Durchgesehen und mit Anmerkungen und Berichtigungen versehen von L. Agassiz. Solothurn. 689 p., 389 pl.
- Sowerby, James de C. 1823. The Mineral Conchology of Great Britain; or Coloured Figures and Description of those Remains of Testaceous Animals or Shells Which Have Been Preserved at Various Times and Depths in the Earth. B. Meredith. London. Vol. 5(71–76):1–64, pl. 408–443.
- Sowerby, James de C. 1824. The Mineral Conchology of Great Britain; or Coloured Figures and Description of those Remains of Testaceous Animals or Shells Which Have Been Preserved at Various Times and Depths in the Earth. B. Meredith. London. 5(77–83):65–138, pl. 444–485.
- Sowerby, James de C. 1825. The Mineral Conchology of Great Britain; or Coloured Figures and Description of those Remains of Testaceous Animals or Shells Which Have Been Preserved at Various Times and Depths in the Earth. B. Meredith. London. 5(84–86):139–168, pl. 486–503.
- Sowerby, James de C. 1826. The Mineral Conchology of Great Britain; or Coloured Figures and Description of those Remains of Testaceous Animals or Shells Which Have Been Preserved at Various Times and Depths in the Earth. B. Meredith. London. Vol. 6(87–93):1–86, pl. 504–545.
- Sowerby, James de C. 1827. The Mineral Conchology of Great Britain; or Coloured Figures and Description of those Remains of Testaceous Animals or Shells Which Have Been Preserved at Various Times and Depths in the Earth. B. Meredith. London. 6(94–99):87–156, pl. 546–580.
- Spath, L. F. 1930. The Eo-Triassic Invertebrate Fauna of East Greenland. Meddelelser om Grønland 83(1):1–87.
- Speden, I. G. 1970. The Type Fox Hills Formation, Cretaceous (Maestrichtian), South Dakota. Part 2. Systematics of the Bivalvia. Peabody Museum of Natural History Bulletin 33:1–222.
- Staesche, Karl. 1926. Die Pectiniden der schwäbischen Jura. Geologische und Palaeontologische Abhandlungen, N. F. 15(1):1–136.
- Stampfli, G. M., & G. D. Borel. 2002. A plate tectonic model for the Paleozoic and Mesozoic constrained by dynamic plate boundaries and restored synthetic oceanic isochrones. Earth and Planetary Science Letters 196:17–33.
- Stampfli, G. M., & G. D. Borel. 2004. The TRANSMED transects in space and time: constraints on the paleotectonic evolution of the Mediterranean domain. *In* W. Cavazza, F. Roure, W. Spakman, G. M. Stampfli, P. A. Ziegler, eds., The TRANSMED Atlas. The Mediterranean Region from Crust to Mantle. Mediterranean Consortium for the 32nd International Geological Congress. p. 53–80.
- Stampfli, G. M., Jon Mosar, Philippe Favre, Alain Pillevuit, & J. C. Vannay. 2001. Permo-Mesozoic evolution of the western Tethys realm: the Neo-Tethys East Mediterranean Basin connection. *In* P. A. Ziegler, W. Cavazza, A. H. F. Robertson & S. Crasquin-Soleau, eds., Peri-Tethys Memoir 6:

Peri-Tethyan Rift/Wrench Basins and Passive Margins. Mémoires du Muséum national d'Histoire naturelle 186:51-108.

- Stanley, G. D. 1997. Upper Triassic fossils from the Antimonio formation, Sonora and their implications for paleoecology and paleogeography. *In* C. M. González-León, & G. D. Stanley, Jr., eds., US - Mexico Cooperative Research: International Workshop on the Geology of Sonora Memoir. Publicaciones Ocasionales 1:62–65.
- Stanley, G. D., C. Gonzalez-León, M. R. Sandy, Baba Senowbari-Daryan, Peter Doyle, Minoru Tamura, & D. H. Erwin. 1994. Upper Triassic Invertebrates from the Antimonio Formation, Sonora, Mexico. Journal of Paleontology [Memoir 36] 68(4):33 p.
- Stanley, S. M. 1968. Post-Paleozoic adaptive radiation of infaunal Bivalve molluscs- A consequence of mantle fusion and siphon formation. Journal of Paleontology 42(1):214–229.
- Stanley, S. M. 1969. Bivalve mollusk burrowing aided by discordant shell ornamentation. Science 166:634–635.
- Stanley, S. M. 1970. Relation of shell form to life habits of the Bivalvia (Mollusca). Geological Society of America Memoir 125:296 p.
- Stanley, S. M. 1972. Functional morphology and evolution of byssally attached bivalve Mollusks. Journal of Paleontology 46(2):165–212.
- Stanley, S. M. 1975. Why clams have the shape they have: an experimental analysis of burrowing. Paleobiology 1:48–58.
- Stanley, S. M. 1977. Coadaptation in the Trigoniidae, a remarkable family of burrowing bivalves. Palaeontology 20(4):869–899.
- Stanley, S. M. 1978. Aspects of the adaptive morphology and evolution of the Trigoniidae. Philosophical Transactions of Real Society of London B 284:247–258.
- Stefanov, Stefan. 1942. Die fauna aus der Trias von Golo-Bardo in SW Bulgarien. 3 Lamelibranchiata. Review of the Bulgarian Geological Society XIV(1):1–11.
- Stefanov, Stefan. 1963. Vertreter der familie Halobiidae aus dem Ladin von Golo-Bardo (SW Bulgarien). Travaux sur la Géologie de Bulgarie (serie Paleontologie) 5:89–107.
- Steinmann, G. 1881. Zur Kenntnis der Jura- und Kreideformation von Caracoles (Bolivia). Neues Jahrbuch für Mineralogie, Geologie und Paläontologie, B. 1(2):239–301, pl. 9–14.
- Steinmann, G. 1929. Geologie von Peru. Carl Winters Universitätsbuchhandlung. Heidelberg. 448 p.
- Stephenson, L. W. 1941. The larger invertebrate fossils of the Navarro Group of Texas (Exclusive of corals and crustaceans and exclusive of the fauna of the Escondido Formation). The University of Texas Publication 4101:641 p., 95 pl., 7 tables.
- Stenzel, H. B. 1971. Oysters. In R. C. Moore, ed., Treatise on Invertebrate Paleontology, Part N, Mollusca, 6, Bivalvia, vol. 3. Geological Society of America & University of Kansas. Boulder, Colorado & Lawrence, Kansas. p. 953–1224.
- Sterren, A. F. 2000. Moluscos bivalvos en la Formación Río del Peñón (Carbonifero Tardío - Pérmico Temprano), provincia de La Rioja, Argentina. Ameghiniana 37(4):421–438.
- Sterren, A. F. 2004. Bivalvos pérmicos de la Formación Tupe en la Quebrada de la Herradura, provincia de San Juan. Ameghiniana 41(4):57–74.
- Stewart, F. J., & C. M. Cavanaugh. 2006. Bacterial endosymbioses in Solemya (Mollusca: Bivalvia)–Model systems for studies of symbiont-host adaptation. Antonie van Leeuwenhoek 90:343–360.
- Stewart, R. B. 1930. Gabb's California Cretaceous and Tertiary type lamellibranchs. The Academy of Natural Sciences of Philadelphia, Special Publication 3:1–314, 17 pl.
- Stiller, Frank. 2000. Two lower Millericrinids and an unusual Crinoid of uncertain systematic position from the lower Upper Anisian (Middle Triassic) of Qingyan, Southwestern China. Journal of Paleontology 74(1):32–51.
- Stiller, Frank. 2001. Fossilvergesellschaftungen, Paläoökologie und paläosynökologische Entwicklung im Oberen Anisium (Mittlere Trias) von

Qingyan, insbesondere Bangtoupo, Provinz Guizhou, Südwestchina. Münstersche Forschungen zur Geologie und Paläontologie 92:1–523. (*)

- Stiller, Frank. 2006. Early Jurassic shallow marine bivalves from Xiaping, southern Hunan, China. Palaeontographica Abteilung A 274:1–70.
- Stiller, Frank, & Chen Jinhua. 2004. Eophilobryoidella sinoanisica new genus and species, an early philobryid bivalve from the Upper Anisian (Middle Triassic) of Qingyan, Southwertern China. Journal of Paleontology 78(2):414–419.
- Stiller, Frank, & Chen Jinhua. 2006. New Mysidiellidae (Bivalvia) from the Anisian (Middle Triassic) of Qingyan, South-West China. Palaeontology 49(1):213–227.
- Stoliczka, Ferdinand. 1866. Geological Sections across the Himalayan Mountains, from Wangto-bridge on the river Sutlej to Sungdo on the Indus: with an account of the formations in Spiti, accompanied by a revision of all known fossils from that district. Memoirs of the Geological Survey of India V:1-154, pl. 1-10.
- Stoliczka, Ferdinand. 1870–1871. The Pelecypoda, with a review of all known genera of this class, fossil and Recent. *In* Thomas Oldham, ed., Cretaceous Fauna of Southern India. Vol. 3, Palaeontologia Indica, Being Figures and Descriptions of the Organic Remains Procured During the Progress of the Geological Survey of India. Trübner and Co. Memoirs of the Geological Survey of India, Calcutta, (serie 1) 3:xxii + 535 p., 50 pl. Parts dates of issue: pts. 1–4:1–222, pl. 1–12 [1870]; parts. 5–13:223–535, pl. 13–50 [1871].
- Stoppani, Antonio. 1858–1860. Les pétrifications d'Ésino ou description des fossiles appartenant au dépôt triasique supérieur des environs d'Ésino en Lombardie. Paléontologie Lombarde 1. :1–151, 32 pl. Impr. J. Bernardoni. Milano.
- Stoppani, Antonio. 1860–1865. Géologie et paléontologie des couches à Avicula contorta en Lombardie. Paleontologie Lombarde 3:1–267. Impr. J. Bernardoni. Milano.
- Stur, D. 1868. Eine Exkursion in die Umgegend von St. Cassian. Jahrbuch der k. k. Geologischen Reichsanstalt in Wien 18(3):529–568.
- Suarez-Riglos, Mario, & Alejandra Dalenz-Farjat. 1993. Pteriomorphia (Bivalvia) Noriano de la Formación Vitiacua, del área de Villamontes (Tarija). *In* R. Suarez-Soruco, ed., Fósiles y Facies de Bolivia. Vol. II Invertebrados y Paleobotánica. Revista Técnica de YPFB 13–14(1–4):155–160.
- Sugawara, Kazuhiro, & Yasuo Kondo. 2004. Brackish and shallow-marine benthic associations of the Lower Jurassic Niranohama Formation in the Shizugawa area, South Kitakami Belt, Northeast Japan. Research Reports of the Kochi University (Natural Science) 53:21–40.
- Swainson, W. 1840. A Treatise on Malacology or Shells and Shell-fish. Longman. London. viii + 419 p.
- Szajnocha, Władysław. 1889. *Pholadomyocardia Jelskii* novum genus, nova species z pokład w jurajskich północnej Peruwii. Pami tnik Akademii Umiej tno ci w Krakowie, Wydziału Matematyczno-Przyrodniczy 16:88.
- Szente, István. 1990. Középsö Liàsz bivalviàk paleobiogeogràfiai értékelése az Alp-Kàrpati Régiòban [Palaeogeographic evaluation of Middle Liassic bivalves in the Alpine-Carpathian region]. Altalanos Foldtani Szemle [General Geological Review] 25:223–229. In Hungarian. (*)
- Szente, István. 1996. Bivalve assemblages from the Austrian and Hungarian Hierlatzkalk (Lower Jurassic): A comparison. *In* E. Dudich, & H. Lobitzer eds., Advances in Austrian-Hungarian Joint Geological Research. Budapest. p. 137–145.
- Szente, István. 1997. Bivalve assemblages from the Middle Triassic Muschelkalk of the Mecsek Mts, South Hungary: An overview. Acta Geologica Hungarica 40(4):411–424.
- Tamura, Minoru. 1959. On *Kumatrigonia*, a new subgenus of *Frenguelliella* and a *Tosapecten* from the Carnic Tanoura Formation in Kyushu, Japan. Memoir, Faculty of Education, Kumamoto University 7:212–218.
- Tamura, Minoru. 1960. Upper Jurassic Pteriacea from the Soma group in north Japan. Palaeontological Society of Japan, Transactions and Proceedings (new series) 37(379): 223–229, pl. 26.

- Tamura, Minoru. 1973. Pectinids from Malayan Triassic. Contributions to the geology and palaeontology of Suotheast Asia, CXX. Geology and Palaeontology of Southeast Asia 12:115–131.
- Tamura, Minoru. 1981. Preliminary report on the Upper Triassic Megalodonts discovered in South Kyushu, Japan. Proceedings of the Japan Academy (series B: Physical and Biological Sciences) 57:290–295.
- Tamura, Minoru. 1990. The distribution of Japanese Triassic bivalve faunas with special reference to parallel distribution of inner Arcto-Pacific fauna and outer Tethyan fauna in Upper Triassic. *In* K. Ichikawa, S. Mizutani, I. Hara, S. Hada & A. Yao, eds., Pre-Cretaceous Terranes of Japan. Publication IGCP Project 224:347–359.
- Tamura, Minoru. 1996. Upper Triassic Tethyan bivalves from Japan and trigonian bioprovinces. *In* G. Braga, F. Finotti, & G. Piccoli, eds., Reports of shallow Tethys 4; International Symposium. Proceedings of the International Symposium on Shallow Tethys 4:185–198.
- Tamura, Minoru, Wataru Hashimoto, Hisayoshi Igo, Takeshi Ishibashi, Junichi Iwai, Teiichi Kobayashi, Toshio Koike, Kaset Pitakpaivan, Tadashi Sato, & E. H. Yin. 1975. The Triassic System of Malaysia, Thailand and some adjacent areas. *In* T. Kobayashi & R. Toriyama, eds., Contributions to the Geology and Palaeontology of Southeast Asia, CLI. Geology and Palaeontology of Southeast Asia XV:103–149.
- Tamura, Minoru, Shigema Kawada, Atsushi Takeda, Hisakazu Oguri, Tei Saito, Kozo Tsuchida, & Makoto Inoue. 1978. A find of Triassic molluscs from the Buko Limestone Formation, Chichibu, Saitama Prefecture. Proceedings of the Japan Academy (series B: Physical and Biological Sciences) 54(2):41–44.
- Tamura, Minoru, & C. A. McRoberts. 1993. A new species of *Myophorigonia* from the Upper Triassic of Oregon, with a reference to the Minetrigoniidae of the circum-Pacific. Memoirs of the Faculty of Education, Kumamoto University, Natural Science 42:29–34.
- Tamura, Minoru, & Eiichi Nishimura. 1994. Description Upper Triassic Kyushutrigonia hachibarensis, new genus and new species with revision of Trigonian species from Sambosan terrane, Japan. Memoirs of the Faculty of Education Kumamoto University Natural Science 43:15–23.
- Tanabe, Kazushige. 1983. Mode of life of an inoceramid bivalve from the Lower Jurassic of West Japan. Neues Jahrbuch für Geologie und Paläontologie Monatshefte 7:419–428.
- Tanaka, Hitoshi. 1989. Mesozoic formations and their moluscan faunas in the Haidateyama area, Oita Prefecture, southwest Japan. Journal of Science of the Hiroshima University (series C: Geology and Mineralogy) 9(1):1–45.
- Tang Zhaoyang, Yao Huazhou, Niu Zhijun, Duan Qifa, Zhao Xiaoming, & Wang Jian. 2007. Preliminary discussion on bivalves assemblagers and their environments of the Bagong Formation of Upper Triassic in Geladandong area, Yangtze Source Region. Journal of Palaeogeography 9(1):59–68
- Tanner, L. H., S. G. Lucas, & M. G. Chapman. 2004. Assessing the record and causes of Late Triassic extinctions. Earth-Science Reviews 65:103–139.
- Tashiro, Masayuki. 1986. Lower Cretaceous bivalves from the Sakawa area, Shikoku. Transactions and Proceedings of Palaeontological Society of Japan (new series) 142:366–392.
- Tate, Ralph, & J. F. Blake. 1876. The Yorkshire Lias. John Van Voorst, Paternoster Row. London. 475 p.
- Taylor, D. G., Karen Boelling, & Jean Guex. 2000. The Triassic/Jurassic System in the Gabbs Formation, Nevada. In R. L. Hall & P. L. Smith, eds., Advances in Jurassic Research 2000. GeoResearch Forum 6:225–236.
- Taylor, D. G., & Jean Guex. 2002. The Triassic/Jurassic System boundary in the Hohn Day Inlier, east-central Oregon. Oregon Geology 64(1):3–28.
- Taylor, J. D., R. J. Cleevely, & N. J. Morris. 1983. Predatory gastropods and their activities in the Blackdown Geensand (Albian) of England. Palaeontology 26(3):521–553.

- Taylor, J. D., & E. A. Glover. 2000. Functional anatomy, chemosymbiosis and evolution of the Lucinidae. *In* E. M. Harper, J. D. Taylor, & J. A. Crame, eds., Evolutionary Biology of the Bivalvia. Geological Society, London, Special Publications 177:207–225.
- Taylor, J. D., & E. A. Glover. 2006. Lucinidae (Bivalvia) the most diverse group of chemosymbiotic molluscs. *In* Rüdiger Bieler FLS, ed., Bivalvia - a look at the Branches. Zoological Journal of the Linnean Society 148:421–438.
- Taylor, J. D., W. J. Kennedy, & Anthony Hall. 1969. The shell structure and mineralogy of the Bivalvia. Introduction. Nuculacea-Trigonacea. Bulletin of the British Museum (Natural History) Zoology Series Supplement 3:1–125.
- Taylor, J. D., W. J. Kennedy, & Anthony Hall. 1973. The shell structure and mineralogy of the Bivalvia. II. Lucinacea-Clavagellacea, conclusions. Bulletin of the British Museum (Natural History) Zoology Series 22:225–294.
- Taylor, J. D., S. T. Williams, & E. A. Glover. 2007. Evolutionary relationships of the bivalve family Thyasiridae (Mollusca: Bivalvia), monophyly and superfamily status. Journal of the Marine Biological Association of the UK 87:565–574.
- Taylor, J. D., S. T., Williams, E. A. Glover, & Patricia Dyal. 2007. A molecular phylogeny of heterodont bivalves (Mollusca: Bivalvia: Heterodonta): New analyses of 18S and 28S rRNA genes. Zoologica Scripta 36(6):587–606.
- Taylor, P. D., & M. A. Wilson. 2003. Palaeoecology and evolution of marine hard substrate communities. Earth-Science Reviews 62:1–103.
- Teichert, Curt. 1990. The Permian-Triassic boundary revisited. In E. G. Kauffman & O. H. Walliser, eds., Extinction Events in Earth History. Lecture Notes in Earth Sciences 30:199–238.
- Teichert, Curt, Bernhard Kummel, & W. C. Sweet. 1973. Permian–Triassic strata, Kuh-e-Ali Bashi, northwestern Iran. Bulletin of Museum of Comparative Zoology 145(8):359–472.
- Teller, Friedrich. 1886. Die Pelecypod–Fauna von Werchojansk in Ostsiberien. In E. Mojsisovics, ed., Arktische Triasfaunen. Mémoires de l'Académie Impériale des Sciences de St.-Pétersbourg (series 7) 33(6):103–137.
- Tëmkin, Ilya. 2006. Morphological perspective on the classification and evolution of Recent Pterioidea (Mollusca: Bivalvia). *In* Rüdiger Bieler FLS, ed., Bivalvia - a look at the Branches. Zoological Journal of the Linnean Society 148:253–312.
- von Teppner, Wilfried. 1922. Lamellibranchia Tertiaria, Anisomyaria II. In C. Diener, ed., Fossilium Catalogus, I. Animalia, Pars. W. Junk. Berlin. (15):67–296.
- Termier, Geneviève, & J. J. Verriez. 1974. Les Bivalves carniens d'Atalanti (Locride, Grece). Annales de la Société Géologique du Nord 93(2):157–162.
- Termier, Henri & Geneviève Termier. 1977. Paléontologie des Invertébrés, in Monographie paléontologique des affleurements permiens du Djebel Tebaga (Sud Tunisien). Palaeontographica A 156(1-3):25-99.
- Terquem, Olry. 1855. Paléontologie de l'étage inférieur de la formation Liasique de la province de Luxembourg, Grande-Duché (Hollande) et de Hettange, du département de la Moselle. Mémoires de la Société Géologique de France (série 2) 3:219–343.
- Terquem, Olry, & E. Piette. 1865. Le Lias inférieur de l'est de la France comprenant la Meurthe, la Moselle, le Grand Duché de Luxembourg, la Belgique et la Meuse. Mémoires de la Société géologique de France (série 2) 8:1–175.
- Terranini, D. 1958. Studio paleontologico sul Norico di Songazzo (Bergamo). Rivista Italiana di Paleontologia e Stratigrafia 64(2):143–182.
- Tevesz, M. J. S. 1975. Structure and habits of the 'living fossil' pelecypod *Neotrigonia*. Lethaia 8:321–327.
- Thévenin, Armand. 1909. Types du Prodrome de Paléontologie Stratigraphique Universelle d'Alcide d'Orbigny. Tome 1: Silurien–Bathonien. Annales de Paléontologie 4:65–92.

- Thiele, Johannes, & Siegfried Jaeckel. 1931. Muscheln der Deutschen Tiefsee-Expedition. Wissenschaftliche Ergebnisse der Deutschen Tiefsee-Expedition auf dem Dampfer "Valdivia" 21(1):159–268.
- Thiele, Ricardo. 1967. El Triásico y Jurásico del departamento de Curepto en la provincia de Talca. Publicaciones de la Universidad de Chile, Facultad de Ciencias Físicas y Matemáticas, Departamento de Geología 28:27–46.
- Thomas, R. D. K. 1978. Shell form and the ecological range of living and extinct Arcoida. Paleobioloby 4(2):181–194.
- Thurmann, Jules. 1833. Essai sur les soulèvements jurassiques du Porrentruy. Mémoire Société d'Histoire Naturelle, Strassbourg 1(2):1–85, pl. 1–5.
- Tichy, Gottfried. 1970. Typen-Katalog. Verzeichnis der in der Geologisch-Paläontologischen Abteilung des Naturhistorischen Museums in Wien aufbewahrten Typen sowie der Abbildungs-originale. Annalen des Naturhistorischen Museums in Wien 74:607–655.
- Tichy, Gottfried. 1974. Ueber Vorkommen und Altersstellung von *Pachyrisma* (*Pachyrisma*) leonardii (Ruggieri, 1959) von Sciacca (Sizilien). Rivista Italiana di Paleontologia e Stratigrafia 80(4):603–164.
- Tichy, Gottfried. 1975. Fossilfunde aus dem Hauptdolomit (Nor, Trias) der östlichen Gailtaler Alpen (Kärnten, Österreich). Annalen des Naturhistorischen Museums in Wien 79:57–100.
- Tichy, Gottfried. 1980a. Gastropoden und Bivalven aus dem Karnischen Ramsaudolomit súdlich von Bad Reichenhall (Oberbayern, Bundesrepublik Deutschland). Geologie Paläontologie Mitteilungen, Innsbruck 9:221–238.
- Tichy, Gottfried. 1980b. Zur Stratigraphie und Ontogenese von Neomegalodon (N.) triqueter triqueter (WULFEN, 1793) (Bivalvia) aus der Trias der Gailataler Alpen (Kärnten, Österreich). Annalen des Naturhistorischen Museums in Wien 83:303–328.
- Tichy, Gottfried, & J. M. Schramm. 1983. Fossilfunde aus dem Hauptdolomit (Trias: Nor) der nördlichen Osterhorn-gruppe (Salzburg, Österreich). Jahrbuch der geologischen Bundesanstalt 126(2):289–293.
- Todd, J. A., & T. J. Palmer. 2002. The Jurassic bivalve genus *Placunopsis*: new evidence on anatomy and affinities. Palaeontology 45(3):487–510.
- Tokuyama, A. 1958. Late Triassic *Palaeopharus* in Japan. Palaeontological Society of Japan, Transactions and Proceedings (new series) 32:291–298.
- Tokuyama, Akira. 1959a. "*Bakevellia*" and "*Edentula*" from the upper Triassic Mine series in west Japan. Transactions and Proceedings of the Palaeontological Society of Japan (new series) 35:147–155.
- Tokuyama, Akira. 1959b. Late Triassic pelecypod fauna of the Aso Formation in West Japan. Japanese Journal of Geology and Geography 31(1):23–38.
- Tokuyama, A. 1959c. Late Triassic Pteriacea from the Atsu and Mine Series, West Japan. Japanese Journal of Geology and Geography 30:1–19, pl. 1.
- Tomašových, Adam. 2004. Microfacies and depositional environment of an Upper Triassic intra-platform carbonate basin: the Fatric Unit of the West Carpathians (Slovakia). Facies 50:77–105.
- Tomašových, Adam. 2006a. Brachiopod and bivalve ecology in the Late Triassic (Alps, Austria): onshore-offshore replacements caused by variations in sediment and nutrient supply. PALAIOS 21:344–368.
- Tomašových, Adam. 2006b. Linking taphonomy to community-level abundance: Insights into compositional fidelity of the Upper Triassic shell concentrations (Eastern Alps). Palaeogeography, Palaeoclimatology, Palaeoecology 235:355–381.
- Tomašových, Adam. 2006c. Differential effects of environmental factors on ecology of brachiopods and bivalves during the Late Triassic and Jurassic. PhD Thesis. Universität Würzburg. 430 p.
- Tomlin, J. R. le B. 1930. Some preoccupied generic names. II. Malacological Society of London, Proceedings 19(1):22–24.
- Tommasi, Annibale. 1890. Revista della Fauna Raibliana del Friuli. Annali del Reale Istituto technico Antonio Zanon in Udina (Series 2) anno 8:1–77.
- Tommasi, Annibale. 1896. La fauna del Trias inferiore nel versante meridionale delle Alpi. Palaeontographia Italica, Memorie di Paleontologia, Pisa 1:43–76, pl. 3–4.

- Tommasi, Annibale. 1911. I fossili della lumachella triassica di Ghegna in Valsecca presso Roncobello, Part prima. Palaeontographia Italica 17:1–36.
- Tong Jinnan. 2005. Studies on the Triassic in Chaohu, Anhui Province, China. Albertiana 32:57-63.
- Tong Jinnan, & Liu Zhili. 2000. The Middle Triassic stratigraphy and sedimentary paleogeography of South China. Albertiana 24:37–47.
- Tong Jinnan, Wu Shunbao, Li Zhiming, Guo Gang, & Zhang Jianjun. 2006. Lower Triassic bivalves from Chaohu, Anhui Province, China. Albertiana 34:42–51.
- Tong Jinnan, & Yin Hongfu. 2002. The Lower Triassic of South China. Journal of Asian Earth Sciences 20:803–815.
- Tong Jinnan, Y. D. Zakharov, M. J. Orchard, Yin Hongfu, & H. J. Hansen. 2004. Proposal of Chaohu Section as the GSSP Candidate of the Induan-Olenekian Boundary. Albertiana 29:13–25.
- Toula, F. 1913. Die Kalke vom Jägerhaus unweit Baden (Rauchstallbrunnengraben) mit nordalpiner St. Cassianer Fauna. Jahrbuch der k. k. Geologischen Reichsanstalt 63(1):78–126.
- Tozer, E. T. 1961. Triassic stratigraphy and faunas Queen Elizabeth Islands, Arctic Archipelago. Geological Survey of Canada, Memoir 316:116 p.
- Tozer, E. T. 1962. Illustrations of Canadian fossils. Triassic of western and Arctic Canada. Geological Survey of Canada Paper 62, 19:1–26.
- Tozer, E. T. 1970. Marine Triassic Faunas. In R. J. W. Douglas, ed., Biochronology: standard of Phanerozoic time. Geological Survey of Canada Economic Geology Report 1. Chapter 9. p. 633–640.
- Tozer, E. T. 1980. Latest Triassic (Upper norian) ammonoid and *Monotis* faunas and correlations. Rivista Italiana di Paleontologia e Stratigrafia 85(3–4):843–876.
- Tozer, E. T., & J. R. Parker. 1968. Notes on the Triassic biostratigraphy of Svalbard. Geological Magazine 105(6):526–541. (*)
- Trechmann, C. T. 1918. The Trias of New Zealand. Quarterly Journal of the Geological Society of London 73(3):165–246.
- Trechmann, C. T. 1923. The Jurassic rocks of New Zealand. Quarterly Journal of the Geological Society of London 79(3):246–286.
- Troedsson, Gustaf. 1951. On the Högänas Series of Sweden (Rhaeto–Lias). Lunds Universitets Årsskrift Ny Följd Avd. 2, 47(1) [Kungl. Fysiografiska Sällskapets Handlingar. N.F., 62(1)]:1–268.
- Tronkov, D.A., & Z. K. Damyanov. 1993. Triassic fossil remains in the siderite ore of the Kremikovci iron-ore deposit. Geologica Balcanica 23(5):34.
- Trushchelev, A. M. 1984. Novye taksony triasovykh dvustvorok Iakutii. [The new Triassic bivalve taxons of Jakutia]. Rossiiskaya Akademiya Nauk Sibirskoe Otdelenie, Trudy. Instituta Geologii i Geofiziki, Akademiia Nauk SSSR 600:64–72, pl. 10–11. In Russian.
- Tuchkov, I. I. 1956. Fauna morskogo reta Severo-Vostoka Azii [Fauna of the marine Rhaetian of north-east Asia]. Ezhegodnik Vsesoyuznogo Paleontologicheskogo Obshchestva 15:177–212, 5 pl. (*)
- Tullberg, S. A. 1881. Ueber Versteinerungen aus den Aucellen–Schichten Novaja–Semljas. Bihang Till Kunglige Svenska Vetenskaps-Akademiens Handlingar 6(3):1–25.
- Turculet, I. 1972. Contributions on the study of *Daonella* genus, with a special view on the Ladinian halobiid fauna of mthe Raráu Region. Analele tiin ifice Universit ii Al. I. Cuza Ia i (Geologie) 18:115–123. In Romanian.
- Turculet, Ilie. 1988. Précisions bionomiques concernant l'espèce Chlamys (Praechlamys) subalternicostata (Bittner, 1901; Bivalvia, Trias). Analele stiintifice ale Universitatii, Al. I. Cuza" din Iasi 34(2):34–36.
- Twitchett, R. J. 2006. The palaeoclimatology, palaeoecology and palaeoenvironmental analysis of mass extinction events. Palaeogeography, Palaeoclimatology, Palaeoecology 232:190–213.
- Twitchett, R. J., Leo Krystyn, Aymond Baud, J. R. Wheeley, & Sylvain Richoz. 2004. Rapid marine recovery after the end-Permian mass-extinction event in the absence of marine anoxia. Geology 32(9):805–808.

- Ürlichs, Max. 1978. Über zwei alpine Ammoniten aus dem Oberer Muschelkalk SW-Deutchlands. Stuttgarter Beiträge zur Naturkunde (Geologie und Paläontologie) (Serie B) 39:13 p.
- Ürlichs, Max. 1992. Vermeintlicher Zwergwuchs bei Muscheln aus dem untersten Trochitenkalk (Oberer Muschelkalk, Mitteltrias) Württembergs. Carolinea 50:9–26.
- Ürlichs, Max, & Gottfried Tichy. 1998. Correlation of the Bleiglanzbank (Gipskeuper, Grabfeld Formation) of Germany with Upper Ladinian beds of the Dolomites (Italy). *In* G. H. Bachmann & I. Lerche, eds., Epicontinental Triassic. E. Schweizerbart'sche Verlagsbuchhandlung (Nägele u. Obermiller). Stuttgart. Zentralblatt für Geologie und Paläontologie 2(9–10):997–1007.
- Valls, Jorge, M. J. Comas-Rengifo, & Antonio Goy. 2004. Bivalvos del Pliensbachiense en la Sección de Almonacid de la Cuba (Cordillera Ibérica, España). Coloquios de Paleontología 54:145–178.
- Végh-Neubrandt, E. 1960. Petrologische Untersuchung der Obertrias-Bildungen des Gerecsegebirges in Ungarn. Geologica Hungarica (series Geologica) 12:132 p. (*)
- Végh-Neubrandt, E. 1969. Bemerkungen zur Gattung Paramegalodus und Aufstellung des neuen Genus Rhaetomegalodon. Anzeiger der Oesterreichischen Akademie der Wissenschaften, Mathematisch-Naturwissenschaftliche Klasse 106(1–14):120–125.
- Végh-Neubrandt, E. 1974. Uj Megalodontacea fajok a magyarorszagi fels triaszbol [Megalodontacea new species from the Upper Triassic of Hungary]. Földtani Közlöny [Bulletin of the Hungarian Geological Society] 104:10–39. In Hungarian.
- Végh-Neubrandt, E. 1982. Triassische Megalodontaceae. Entwicklung, Stratigraphie und Paläontologie. Akadémiai Kiadó. Budapest. 526 p.
- Végh-Neubrandt, E., J. F. Dumont, Marcel Gutnic, Jean Marcoux, Olivier Monod, & André Poisson. 1976. Megalodontidae du Trias supérieur dans la Chaine Taurique (Turquie meridionale). Geobios 9(2):199–222.
- Venzo, Sergio. 1934. Il Ladinico superiore dell'Isola di Rodi (Egeo), II: La fauna. Palaeontographia Italica 34:142–170.
- Venzo, Sergio. 1942. *Trigonia (Laevitrigonia) stefaninii* n.sp. del batoniano dell'Oltregiuba; revisione del sottogenere e sua distribuzione. Atti della Societa Italiana di Scienze Naturali e del Museo Civico di Storia Naturale de Milano 81:210–229.
- Vera, J. A. 1994. Estratigrafía. Principios y Métodos. Ed. Rueda. Madrid. 806 p.
- Vía, Luis, & J. F. Villalta. 1975. Restos de crustáceos decápodos en el Triásico de Montral-Alcover (Tarragona). Boletín Geológico y Minero LXXXVI–V:485–497.
- Vía, Luis, J. F. Villalta, & Mateu Esteban. 1977. Paleontología y Paleoecología de los yacimientos fosilíferos del Muschelkalk Superior entre Alcover y Montral (Montañas de Prades, Provincia de Tarragona). Cuadernos de Geología Ibérica 4:247–256.
- Vialov, O. S. 1936. Sur la classification des huitres. Doklady Akademii Nauk SSSR (series 2) 4(1):17–20.
- Vigh, G. 1914. Adatok az esztergomvidéki triász ismeretéhez [Beiträge zur Kenntnis der Trias in Komitate Esztergom]. Földtani Közlony (Geologische Mitteilungen) 44(10–12):572–577 + 599–604.
- Villamil, Tomas, E. G. Kauffman, & H. A. Leanza. 1998. Epibiont habitation patterns and their implications for life habits and orientation among trigoniid bivalves. Lethaia 31:43–56.
- Virgilli, Carmina. 1958. El triásico de los Catalánides. Boletín del Instituto Geológico y Minero de España 69:1–831.
- Vokes, H. E. 1945. Supraspecific groups of the Pelecypod Family Corbulidae. Bulletin of the American Museum of Natural History 86(1):1–32.
- Vokes, H. E. 1956. Notes on the Nucinellidae (Pelecypoda) with description of a new species from the Eocene of Oregon. Journal of Paleontology 30(3):652–671.

- Vokes, H. E. 1967. Genera of the Bivalvia: A systematic and bibliographic catalogue. Bulletins of American Paleontology 51(232):[103–111] + 112–394.
- Vokes, H. E. 1980. Genera of the Bivalvia: a Systematic and Bibliographic Catalogue (Revised and Updated). Paleontological Research Institution. Ithaca. 307 p.
- Voronetz, N. S. 1936. Mezozoiskaya fauna Kharaulakhskogo khrebta [The Mesozoic fauna of the Kharaulakh mountain range]. Trudy arkt. nauchno-issled. Institut [Transactions of the Arctic Institution] 37:7–36. In Russian.
- Voronetz, N. S. 1938. Fauna morskogo mezozoia Bureinskogo Basseina. Mater. po Geol. Bureinskogo Kamennoug. Basseina. 2 (1937). Trudy VIMS 123:47–85.
- Vörös, Attila. 1971. The lower and middle Jurassic bivalves of the Villany Mountains. Annales Universitatis Scientiarum Budapestinensis de Rolando Eoetvoes Nominatae, Sectio Geologica 14:167–208.
- Vörös, Attila. 1981. A survey of the Rhaetian (Upper Triassic) Bivalvia from Borzavár (Bakony Mts., Hungary). Annales Historico-Naturales Musei Nationalis Hungarici 73:33–54.
- Vörös, Attila, & József Pálfy. 2002. New data to the stratigraphy of the Pelsonian Substage at Köveskál (Middle Triassic, Balaton Highland, Hungary). Fragmenta Palaeontologica Hungarica 20:53–60.
- Vozin, V. F., & V. V. Tikhomirova. 1964. Polevoi atlas dvystorchatkh i goloxonogikh molliuskov Triasovykh otlozhenii severo-vostoka SSSR [Field atlas of Bivalvia and Cephalopoda in the Triassic deposits of northeastern USSR]. Institute of Geology, Siberian Branch, USSR Academy of Sciences. Moscow. 94 p. In Russian.
- Vu Khuc, Dang. 1977a. A revision of certain bivalve genera from the Upper Triassic of North Vietnam. Doklady Akademia Nauk USSR 232(3):676–679. In Russian.
- Vu Khuc, Dang. 1977b. Novyy rod *Songdaella* (Bivalvia) iz verkhnetriasovykh otlozheniy Severnogo V'yetnama [The new genus *Songdaella* (Bivalvia) from the Upper Triassic of North Vietnam]. Paleontologicheskii Zhurnal (2): 49-56 [= Paleontological Journal 11(2):179–187].
- Vu-Khuc, Dang, A. S. Dagys, L. D. Kiparisova, N. B. Nguyen, T. C. Bao, & I. N. Srebrodolskaia. 1965. Les Fossiles caractéristiques du Trias du Nord Viêt-Nam. Direction Générale de Géologie de la RDV. Hanoi. 117 p.
- Vu Khuc, Dang, & D. T. Huyen. 1998. Triassic correlation of the Southeast Asian mainland. Palaeogeography, Palaeoclimatology, Palaeoecology 143:285–291.
- Waagen, Lukas. 1907. Die Lamellibranchiaten der Pachycardientuffe der Seiser Alm nebst vergleichend paläontologischen und phylogenetischen Studien. Abhandlungen der Kaiserlich-Königlichen Geologischen Reichsanstalt 18(2):1–180.
- Waller, T. R. 1978. Morphology, morphoclines and a new classification of the Pteriomorphia (Mollusca: Bivalvia). Philosophical Transactions of the Royal Society of London (series B) 284:345–365.
- Waller, T. R. 2006. Phylogeny of families in the Pectinoidea (Mollusca: Bivalvia): importance of the fossil record. *In* Rüdiger Bieler FLS, ed., Bivalvia - a look at the Branches. Zoological Journal of the Linnean Society 148:313–342.
- Waller, T. R., & Louie Marincovich, Jr. 1992. New species of *Camptochlamys* and *Chlamys* (Mollusca: Bivalvia: Pectinidae) from near the Cretaceous/ Tertiary boundary at Ocean Point, North Slope, Alaska. Journal of Paleontology 66(2):215–227.
- Waller, T. R., & G. D. Stanley, Jr. 1998. New marine Middle Triassic bivalves from the New Pass Range, west-central Nevada. Abstracts with Programs - Geological Society of America 30(7):287.
- Waller, T. R., & G. D. Stanley, Jr. 2005. Middle Triassic Pteriomorphian Bivalvia (Mollusca) from the New Pass Range, West-Central Nevada: Systematics, Biostratigraphy, Paleoecology and Paleobiogeography. Journal of Paleontology [Memoir Paleontological Society 61] 79(1):64 p.

- Walter, J. C. 1953. Paleontology of Rustler Formation, Culberson County, Texas. Journal of Paleontology 27(5):679–702.
- Walther, Karl. 1927. Zwölf Tafeln der verbreitetsten Fossilien aus dem Buntsandstein und Muschelkalk der Umgebung von Jena, 2nd ed. Verlag von Gustav Fischer. Jena. 47 p.
- Wang Mingqian. 1993. Bivalve fauna from Uppermost Permian and Lowermost Triassic of Fenghai, Yong'an, Fujian. Acta Palaeontologica Sinica 32(4):458–476. In Chinese with English abstract.
- Wang Xiaofeng, Gerard H. Bachmann, Hans Hagdorn, P. M. Sanders, Gilles Cuny, Chen Xiaohong, Wang Chuanshang, Chen Lide, Cheng Long, Meng Fansong, & Xu Guanghong. 2008. The Upper Triassic black shales of the Guanling Area, Guizhou Province, South-West China: a unique marine reptile and pelagic crinoid fossil lagerstätte. Palaeontology 51(1):27–61.
- Wang Yigang, & P. L. Smith. 1986. Sinemurian (Early Jurassic) ammonite fauna from the Guangdong Region of Southern China. Journal of Paleontology 60(5):1075–1085.
- Wang Yue, Shen Shuzhong, Cao Changqun, Wang Wei, Charles Henderson, & Jin Yugan. 2006. The Wuchiapingian–Changhsingian boundary (upper Permian) at Meishan of Changxing County, South China. Journal of Asian Earth Sciences 26(6):575–583.
- Ward, P. D., G. H. Garrison, J. W. Haggart, D. A. Kring, & M. J. Beattie. 2004. Isotopic evidence bearing on Late Triassic extinction events, Queen Charlotte Islands, British Columbia, and implications for the duration and cause of the Triassic/Jurassic mass extinction. Earth and Planetary Science Letters 224:589–600.
- Warrington, Geoffrey, & H. C. Ivimey-Cook. 1990. Biostratigraphy of the Late Triassic and Early Jurassic: A review of type sections in Southern Britain. Les Cahiers de l'Université Catholique de Lyon (série Sciences) 3:207–213.
- Warth, Manfred. 1990. The bivalves of the Schilfsandstein (Upper Triassic, Carnian, Middle Keuper) of Southwest Germany and their stratigraphical range. Neues Jarbuch für Geologie und Paläontologie Monatshefte 181(1–3):107–115.
- Warth, Manfred. 1994. Vorkommen von "Perna" keuperina Blanckenhorn (Lamellibranchiata) in der Rottweiler Bank (Stubensandstein, Nor, Trias) von Baden-Wurttemberg (SW Germany). Stuttgarter Beitraege zur Naturkunde Serie B (Geologie und Palaeontologie) 212:1–13.
- Waterhouse, J. B. 1958. The occurrence of *Atomodesma* Beyrich in New Zealand. New Zealand of Geology and Geophysics 1:166–177.
- Waterhouse, J. B. 1959. Notes on New Zealand species of Atomodesma Beyrich. New Zealand of Geology and Geophysics 2:262–264.
- Waterhouse, J. B. 1960. Some Carnian pelecypods from New Zealand. Transactions of the Royal Society of New Zealand 88(3):425–442.
- Waterhouse, J. B. 1963. *Etheripecten*, a new aviculopectinid genus from New Zealand. New Zealand Journal of Geology and Geophysics 6:193–196.
- Waterhouse, J. B. 1964. Palaeotaxodont bivalves from the Permian of New Zealand. Palaeontology 7(4):630–655.
- Waterhouse, J. B. 1966. On the validity of the Permian bivalve family Pachydomidae Fischer (1887). Journal of the Geological Society of Australia 13(2):543–559, pl. 15–17.
- Waterhouse, J. B. 1969. The Permian bivalve genera *Myonia, Megadesmus, Vacunella* and thier allies and their occurrences in New Zealand. Paleontological Bulletin, New Zealand Geological Survey 41:141 p.
- Waterhouse, J. B. 1978. Permian brachiopoda and Mollusca from North-West Nepal. Palaeontographica, Abteilung A, Paläozoologie - Stratigraphie 160(1–6):1–175.
- Waterhouse, J. B. 1979a. The Upper Triassic bivalve Oretia Marwick 1953 (Note). New Zealand Journal of Geology and Geophysics 22(5):621–625.
- Waterhouse, J. B. 1979b. A new species of *Permophorus* Chavan (Bivalvia) from the early Triassic of New Zealand. New Zealand Journal of Geology and Geophysics 22(6):743–749.

- Waterhouse, J. B. 1980a. Permian bivalves of New Zealand. Journal of the Royal Society of New Zealand 10(1):97–133.
- Waterhouse, J. B. 1980b. A new bivalve species (Buchiidae) from the Lower Triassic of New Zealand. Alcheringa 4:1–10.
- Waterhouse, J. B. 1983. Systematic description of Permian brachiopods, bivalves and gastropods below Wall Sandstone Member, northern Bowen Basin. Papers - Department of Geology, University of Queensland 10(3):155–179.
- Waterhouse, J. B. 1987. Late Palaeozoic Mollusca and correlations from the Southeast Bowen Basin, East Australia. Palaeontographica (Abt. A) 198:129–233.
- Waterhouse, J. B. 2000. Early Triassic Pectinidina (Mollusca: Bivalvia) from Dolpo and Manang, Nepal Himalaya. Records of the Canterbury Museum 14:155–186.
- Waterhouse, J. B. 2002. The stratigraphic succession and structure of Wairaki Downs, New Zealand and its implications for Permian biostratigraphy and marine Permian of eastern Australia and New Zealand. Earthwise 4:1–262.
- Waterhouse, J. B. 2008. Aspects of the evolutionary record for fossils of the Bivalve subclass Pteriomorphia Beurlen. Earthwise 8:220 p.
- Waterhouse, J. B., & Chen Zhong-qiang. 2006. Stratigraphy and Mollusca of the Late Permian Senja Formation, Manang area, Nepal Himalaya. Palaeontographica (Abt. A) 275(1–3):55–96.
- Watson, Hugh. 1930. On the anatomy and affinities of *Plicatula*. Malacological Society of London, Proceedings 19:25–30.
- Watson, J. S. 1982. The occurrence of *Discinisca* on *Dacryomya ovum*: An example of commensalism from the Upper Lias of Yorkshire. Proceedings of the Yorkshire Geological Society 44:45–51.
- Wen Shixuan. 1982. Jurassic Bivalvia of Xizang. Institute of Geology and Paleontology, Chinese Academy of Sciences, ed., Paleontology of Xizang. In the collection: The series of the scientific expedition to the Qinghai-Xizang Plateau, Book IV. Science Press. Beijing. p. 225–254. In Chinese with English abstract.
- Wen Shi-xuan. 1999. Cretaceous bivalve biogeography in Qinghai-Xizang Plateau. Acta Paleontologica Sinica 38(1):1–30. In Chinese with English abstract.
- Wen Shi-xuan, Lan Xiu, Chen Jin-hua, Zhang Zuo-ming, Chen Chu-chen, & Gu Chi-wei. 1976. Fossil Lamellibranchia from the Mount Jolmo Lungma Region. In Lan X. & Chen C., eds., A report of scientific expedition in the Mount Jolmo Lungma region (1966–1968), Paleontology, Fasc. 3. Science Press. Beijing, p. 1–210. In Chinese.
- West, I. M. 2007. Burton Bradstock–Bridport Sands and Inferior Oolite. Geology of the Wessex Coast, http://www.soton.ac.uk/~imw/burton. htm. Checked April 2012.
- Westermann, G. E. G. 1970. Occurrence of *Monotis subcircularis* Gabb in Central Chile and the dispersal of *Monotis* (Triassic Bivalvia). Pacific Geology 2:35–40.
- Westermann, G. E. G. 2000. Marine faunal realms of the Mesozoic: review and revision under the new guidelines for biogeographic classification and nomenclature. Palaeogeography, Palaeoclimatology, Palaeoecology 163:49–68.
- Westermann, G. E. G., & Kazem Seyed-Emami. 1981. Occurrence of the Upper Triassic bivalve *Monotis* in Iran. Paläontologische Zeitschrift 55(2):173–174.
- Westermann, G. E. G., & Harish Verman. 1967. The Norian Pine River Bridge section, British Columbia and the succession of *Monotis*. Journal of Paleontology 41(3):798–803.
- Westwood, J. O. 1840. Observations on the genus *Derbe* of Fabricius. Proceedings of the Linnean Society of London 1:82–85.
- White, C. A. 1887. On new generic forms of Cretaceous Mollusca and their relation to other forms. Academy of Natural Sciences of Philadelphia, Proceedings 39:32–37, pl. 2.

- Whitfield, Robert Parr. 1885. Brachiopoda and Lamellibranchia of the Raritan Clays and Greensand Marls of New Jersey. U. S. Geological Survey Monograph 9: 338 p., 34 pl., map.
- Whitfield, Robert Parr. 1902. Description of a new form of Myalina from the Coal Measures of Texas. Bulletin of the American Museum of Natural History 16(4):63–66.
- Wignall, P. B. 1990. Benthic palaeoecology of the upper Jurassic Kimmeridge Clay of England. Special Papers in Palaeontology 43:74 p.
- Wignall, P. B., & D. P. G. Bond. 2008. The end-Triassic and Early Jurassic mass extinction records in the British Isles. Proceedings of the Geologists' Association 119:73–84.
- Wignall, P. B., R. Morante, & Robert Newton. 1998. The Permo-Triassic transition in Spitsbergen: ¹³C_{org} chemostratigraphy, Fe and S geochemistry, facies, fauna and trace fossils. Geological Magazine 135(1):47–62.
- Wignall, P. B., & M. J. Simms. 1990. Pseudoplankton. Palaeontology 33(2):359–378.
- Wignall, P. B., & R. J. Twitchett. 2002. Permian–Triassic sedimentology of Jameson Land, East Greenland: incised submarine channels in an anoxic basin. Journal of the Geological Society, London 159:691–703.
- Wignall, P. B., J. P. Zonneveld, R. J. Newton, Ken Amor, M. A. Sephton, & S. Hartley. 2007. The end Triassic mass extinction record of Williston Lake, British Columbia. Palaeogeography, Palaeoclimatology, Palaeoecology 253:385–40.
- Wilckens, Otto. 1927. Contributions to the Palaeontology of the New Zealand Trias. Palaeontological Bulletin 12:65 p.
- Wilckens, Rudolf E. 1909. Paläontologische Untersuchung triadischer Faunen aus der Umgebung von Predazzo in Südtirol. Verhandlungen des Naturhistorisch-medizinischen Vereins zu Heidelberg (neue folge) 10:81–230.
- Wilkes, J. 1810. Conchology. In Encylopaedia Londinensis; or, Universal Dictionary of Arts, Sciences, and Literature 5. J. Adlard. London. p. 14–41.
- Wilton, C. P. N. 1830. Memoir on the geology of the shore of the Severn, in the parish of Awre, Glocestershire. Quarterly Journal of Science, Literature and Art (Royal Institute, London) 72:64–73.
- Winchell, Alexander. 1862. Description of Fossils from the Marshall and Huron Groups of Michigan. Proceedings of the Academia of Natural Sciences. Philadelphia 33:405–430.
- Winkler, G. G. 1859. Die Schichten der Avicula contorta inner- und ausserhalb der Alpen. Habilitationschrift Univers. München. 51 p., 2 pl.
- Winkler, G. G. 1861. Der Oberkeuper, nach Studien in den bayrischen Alpen. Zeitschrift der Deutschen geologischen Gessellschaft XIII:459–521.
- Winkler, G. G. 1886. Neue Nachweise über den unteren Lias in den bairischen Alpen. Neues Jahrbuch f
 ür Mineralogie, Geologie und Pal
 äontologie II:1–34.
- Wirth, E. 1936. Beiträge zur Kenntnis der Trias in der Provinz Szechuan, West-China. Neues Jahrbuch für Mineralogie, Geologie und Paläontologie, Abhandlungen B 75:412–445.
- Wissmann, H. L., & G. V. Münster. 1841. Beiträge zur Geognosie und Petrefakten-Kundes der Südöstlichen Tirol's, vozuglich der Schichten von St. Cassian, IV. Bayreuth. 152 p., 16 pl.
- von Wittenburg, P. W. 1908. Beiträge zur Kenntnis der Werfener Schichten Südtirols. Abhandlungen der Geologie und Paläontologie (new series) 8(5):44 p.
- von Wittenburg, P. W. 1909. Einige Lamellibranchiata der Salt Range, mit Berücksichtigung der Lamellibranchiata der Süd-Ussuri-Gebietes. Neues Jahrbuch für Mineralogie 1:6–13.
- von Wöhrmann, S. F. 1889. Die Fauna der sogenannten *Cardita*-und Raibler-Schichten in den Nordtiroler und bayerischen Alpen. Jahrbuch der Kaiserlich-Königlichen Geologischen Reichsanstalt 39:181–258.

- von Wöhrmann, S. F. 1894. Die Raibler Schichten nebst kritischen Zusammenstellung ihrer Fauna. Jahrbuch der k.k. Geologischen Reichsanstalt, Wien 43(3–4):617–768.
- von Wöhrmann, S. F., & E. Koken. 1892. Die Fauna der Raibler Schichten von Schlernplateau. Zeitschrift der Deutschen Geologischen Gesellschaft (Hannover) 44:167–223, Taf. 6–10.
- Wood, Searles Valentine. 1839. Descriptions of the species of the genus *Lima*, from the Coralline Crag, in the cabinet of Searles Valentine Wood, Esq., late Curator to the Geological Society of London. Magazine of Natural History (new series) 3(29):233–236, pl. 3.
- Wood, Searles Valentine. 1840. On the fossil shells of the crag. Magazine of Natural History 4(5):230–234, pl. 13.
- Wood, Searles Valentine. 1851–1882. A Monograph of the Crag Mollusca, or descriptions of shells from the middle and upper Tertiaries of the east of England. The Palaeontographical Society Monographs, London. Vol. II, Bivalves, p. 1–150, pl. 1–12 [1851]; p. 151–216, pl. 13–20 [1853]; p. 217–342, pl. 21–31 [1857]; [note], p. 1–2 [1861]. Vol. III, Supplement, Bivalves, p. 99–231, pl. 7–11 and Addendum Plate [1874]. Vol. IV, Second Supplement, title page, preface, p. 1–58, pl. 1–6 [May, 1879]. Third Supplement, p. 1–24, pl. 1 [1882].
- Woodring, W. P. 1925. Contributions to the Geology and Palaeontology of the West Indies. Miocene Mollusks from Bowden, Jamaica. Pelecypods and Scaphopods. Carnegie Institution Publication. Carnegie Institution of Washington, D.C. 366:1–122, 28 pl.
- Woods, M. A., & T. E. Yancey. 2004. Wallowaconcha in the United Arab Emirates–a Late Triassic bivalve a long way from home. In Abstracts, 48th Annual Meeting The Paleontological Association. Newsletters 57:193–194.
- Woodward, S. P. 1868. A Manual of the Mollusca: a treatise on recent and fossil shells, with an Appendix of recent and fossil conchological discoveries to the present time by Ralph Tate, 2nd edit. Virtue & Co. London. Appendix, 86 p.
- Wright, A. D., & M. J. Benton. 1987. Trace fossils from Rhaetic shore-face deposits of Staffordshire. Palaeontology 30(2):407–428.
- Wu Fa-ming. 1985. New material of bivalves from the Early Triassic in Fujian. Acta Palaeontologica Sinica 24(4):395–401. In Chinese with English abstract.
- Wu, Shunbao. 1981. Notes on the Jurassic System and bivalves of Fengjiachong-Lingling, Hunan. Geological Review 27(5):375–383. In Chinese with English abstract.
- Wu Shunbao, Li Zhiming, Guo Gang, & Tong Jinnan. 2005. Lower Triassic bivalve sequence of Chaohu, Anhui Province. Albertiana 33(1):91.
- Wulfen, F. X. 1793. Abhandlungen von k\u00e4rnthnerischen pfauenschweifigen Helmintholith oder opalisierenden Muschelmarmor. J. Palm. Erlangen. 125 p.
- Wurm, Adolf. 1911. Untersuchungen über den Geologischen Bau und die Trias von Aragonien. Zeitschrift der Deutsche Geolgische Gesellschaft 63:37–175.
- Wurm, Adolf. 1913. Beiträge zur Kenntnis der iberisch balearischen Triasprovinz. Verhandlungen des Naturhistorisch-Medizinischen vereins zu Heidelberg, N. F. 12(4):477–599.
- Yamani, S. A. 1983. Die Bivalvenfauna der Schwammkalke von Biburg (Oberoxford, Südliche Frankenalb) Pteriomorphia II. Mitteilungen der Bayerischen Staatssammlung für Paläontologie und Historische Geologie 23:3–33.
- Yancey, T. E. 1978. Brachiopods and molluscs of the lower Permian Arcturus Group, Nevada and Utah. Part 1: Brachiopods, scaphopods, rostroconchs, and bivalves. Bulletins of American Paleontology 74(303):253–367.
- Yancey, T. E. 1985. Bivalvia of the H. S. Lee formation (Permian) of Malaysia. Journal of Paleontology 59(5):1286–1297.
- Yancey, T. E., & G. D. Stanley, Jr. 1987. Alatoform megalodontid bivalves from the Late Triassic of eastern Oregon. Abstracts with Programs of the Geological Society of America 19(7):901.

- Yancey, T. E., & G. D. Stanley, Jr. 1996. Giant alatoform bivalves of the Late Triassic in western North America. Abstracts with Programs of the Geological Society of America 28(7):430.
- Yancey, T. E., & G. D. Stanley, Jr. 1999. Giant alatoform bivalves in the Upper Triassic of Western North America. Palaeontology 42(1):1–23.
- Yancey, T. E., G. D. Stanley Jr., W. E. Piller, & M. A. Woods. 2005. Biogeography of the Late Triassic wallowaconchid megalodontoid bivalves. Lethaia 38:351–365.
- Yang Fengqing, Peng Yuanqia, & Gao Yongqun. 2001. Study on the Late Permian *Claraia* in South China. Science in China (series D) 44(9):797–807.
- Yang Shouren, Wang Xinping, & Hao Weicheng. 1986. Early Triassic bivalve assemblage of western Guangxi. Journal of Stratigraphy 10(2):88–97. In Chinese with English abstract.
- Yang Zunyi, & Yin Hongfu. 1979. Marine Triassic faunas from Shihchienfeng Group in the northern Weihe River basin, Shaanxi Province. Acta Palaeontologica Sinica 18(5):465–474. In Chinese with English abstract.
- Yang Zunyi, Yin Hongfu, Wu Shunbao, Yang Fengqing, Ding Meihua, & Xu Guirong. 1987. Permian-Triassic boundary stratigraphy and fauna of South China. People's Republic of China Ministry of Geology and Mineral Resources, Geological Memoirs 2(6):295–379. In Chinese with English abstract.
- Yao Huazhou, Sha Jingeng, Duan Qifa, Niu Zhijun., Zeng Bofu, & Zhang Renjie. 2003. A new genus *Quemocuomegalodon* of Megalodontidae from the Upper Triassic in the source area of the Yangtze river, Western China. Acta Palaeontologica Sinica 42(3):393–407. In Chinese with English abstract.
- Yao Huazhou, Zhang Renjie, John Pojeta, Sha Jingeng, & Wang Jianxiong. 2007. Late Triassic megalodontids (Bivalvia) from the Headwaters of the Yangtze River, Qinghai Province, West China. Journal of Paleontology 81(6):1327–1347.
- Yao Zhaoqi, Xu Jun-tao, Zheng Zhuoguan, Zhao Xiugu, & Mou Zhuangguan. 1980. Biostratigraphy of Late Permian and the boundary of Permian-Triassic in western Guizhou and eastern Yunnan. *In* Nanjing Institute of Geology and Palaeontonlogy, Academia Sinica, eds., The Palaeontology and the Coal-Bearing Strata of Late Permian in Western Guizhou and Eastern Yunnan. Science Press. Beijing. p. 1–69. In Chinese.
- Yarnell, J. M., George Stanley, & C. J. R. Hart. 1999. New paleontological investigations of Upper Triassic shallow-water reef carbonates (Lewes River Group) in the Whitehorse area, Yukon. *In C. F. Roots & D.* S. Emond, eds., Yukon Exploration and Geology 1998. Exploration and Geological Services Division, Yukon, Indian and Northern Affairs Canada. p. 179–184.
- Yehara, S. 1921. On some new species of *Trigonia* from the Lias of Province Nagato, and the Cretaceous of Province Awa. Japanese Journal of Geological Society 28:7–11. (*)
- Yin Hongfu. 1974. Middle-Upper Triassic lamellibranchs from Qingyan and Zengfeng of Guizhou Province. Geological Science and Technology Information 5:19–60. In Chinese. (*)
- Yin Hongfu. 1983. Uppermost Permian (Changxingian) Pectinacea from South China. Rivista Italiana di Paleontologia e Stratigrafia 88(3):337– 386.
- Yin Hongfu. 1985. Bivalves near the Permian-Triassic boundary in south China. Journal of Paleontology 59(3):572–600.
- Yin Hongfu. 1990. Paleogeographical distribution and stratigraphical range of the Lower Triassic *Claraia, Pseudoclaraia* and *Eumorphotis* (Bivalvia). Journal of China University of Geoscience 1(1):98–110.
- Yin Hongfu, & Nie Zhetong. 1990. Triassic bivalves of the Ngari Area. In Yang Z. & Nie Z., eds., Paleontology of Ngari, Tibet (Xizang). The China University of Geosciences Press. Beijing. p. 100–113. In Chinese with English summary, p. 254–257. (*)
- Yin Hongfu, & Yin Jia-ren. 1983. Bivalves. *In* Yang Zun-yi, Yin Hong-fu, Xu Gui-rong, Wu Shun-bao, He Yuan-liang, Liu Guang-cai, & Yin Jia-

ren, eds., Triassic of the South Qilian Mountains. Geological Publishing House. Beijing, p. 128–175. In Chinese. (*)

- Yin Jiarun, & Raimond Enay. 2000. The Earliest Jurassic psiloceratids in the Eastern Tethyan Himalaya, South Tibet. Earth and Planetary Sciences 331:601–608.
- Yin Jiarun, Raimond Enay, & Wan Xiaoqiao. 1999. The first report of the Late Triassic-Early Jurassic passage beds in the Eastern Tethyan Himalaya dans l'Himalaya tethysien oriental. Comptes Rendus de l'Academie des Sciences, ser. IIA, Earth and Planetary Science 329(2):125–133.
- Yin Jiarun, & J. A. Grant-Mackie. 2005. Late Triassic–Jurassic bivalves from volcanic sediments of the Lhasa block, Tibet. New Zealand Journal of Geology and Geophysics 48:555–576.
- Yin Jiarun, & C. A. McRoberts. 2006. Latest Triassic-Earliest Jurassic bivalves of the Germig Formation from Lanongla (Tibet, China). Journal of Paleontology 80(1):104–120.
- Yin Jiarun, Paul L. Smith, József Pálfy, & Raymond Enay. 2007. Ammonoids and the Triassic/Jurassic boundary in the Himalayas of Southern Tibet. Palaeontology 50(3):711–737.
- Yin Jiarun, Yao Huazhou, & Sha Jingeng. 2004. First record of the Early Jurassic *Lupherella* fauna (Bivalvia) in eastern Guangdong, southeast China. New Zealand Journal of Geology and Geophysics 47:321–326.
- Yin Yugan, Wang Yue, Charles Henderson, B. R. Wardlaw, Shen Shuzhong, & Cao Changqun. 2006. The Global Boundary Stratotype Section and Point (GSSP) for the base of Changhsingian Stage (upper Permian). Episodes 29(3):175–182.
- Yokoyama, Matajiro. 1904. On some Jurassic fossils from Rikuzen. Journal of the College of Sciences, Imperial University, Tokyo 18(6):1–13, pl. 1–2.
- Yonge, C. M. 1953. Form and habit in *Pinna carnea* Gmelin. Philosophical Transactions of the Royal Society of London (series B) 237:335–374.
- Young, G., & J. Bird. 1822. A Geological Survey of the Yorkshire coast: Describing the Strata and Fossils Occurring between the Humber and the Tees, from the German Ocean to the Plain of York. [first edition] R. Kirby and Longman & Co. London. p. 1–235, 17 pl.
- Zakharov, V. A. 1962. Novye Monotidae nizhnego Leiasa s poberezhya Okhotskogo Morya i ikh stratigraficheskoe znachenie [New Monotidae from the Lower Lias of the Okhotsk Sea coast and their stratigraphic significance]. Geologiya i Geofizika 3:23–31. In Russian with English abstract.
- Zakharov, V. A. 1965. New Upper Jurassic and Lower Cretaceous Camptonectid (Pectinidae, Bivalvia) from Arctic Siberia. *In V. N. Saks*, ed., Stratigraphy and Paleontology of the Mesozoic deposits of Northern Siberia. Nauka. Moscow. p. 72–80. In Russian.
- Zakharov, V. A., Yu. I. Bogomolov, V. I. Il'ina, A. G. Konstantinov, N. I. Kurushin, N. K. Lebedeva, S. V. Meledina, B. L. Nikitenko, E. S. Sobolev, & B. N. Shurygin. 1997. Boreal zonal standard and biostratigraphy of the Siberian Mesozoic. Russian Geology and Geophysics 38(5):965–993.
- Zakharov, V. A., N. I. Kurushin, & V. P. Pokhialainen. 1996. Paleobiogeographic criteria of terrane geodynamics of Northeastern Asia in Mesozoic. Russian Geology and Geophysics 37(11):1–22.
- Zakharov, V. A., B. N. Shurygin, V. I. Il'ina, & B. L. Nikitenko. 2006. Pliensbachian-Toarcian biotic turnover in North Siberia and the Arctic Region. Stratigraphy and Geological Correlation 14(4):399–417.
- Zapfe, Helmuth. 1963. Beiträge zur Paläontologie der nordalpinen Riffe. Zur Kenntnis der Fauna des oberrhätischen Riffkalkes von Adnet, Salzburg (exkl. Riffbildner). Annalen des Naturhistorischen Museums in Wien 66:207–259.
- Zapfe, Helmuth. 1967. Beiträge zur Paläontologie nordalpiner Riffe. Die Fauna der Zlambach-Mergel der Fischwiese bei Aussee, Steiermark. Annalen des Naturhistorischen Museums Wien 71:413–480.
- Zapfe, Helmuth. 1969. Beitraege zur Palaeontologie der nordalpinen Riffe; rhaetomegalodonten aus dem Dachsteinkalk der niederooesterreichischen Voralpen. Annalen des Naturhistorischen Museums in Wien 73:141–159.

- Zapfe, Helmuth. 1972. Cornucardia hornigii (Bittner) in einer "Dachsteinkalk-Fazies" der Nordalpen. Annalen des Naturhistorischen Museums in Wien 76:587–603.
- Zapfe, Helmuth. 1973. *Otapiria* (Monotidae, Bivalvia) aus der alpinen Trias. Annalen des Naturhistorischen Museums in Wien 77:149–158.
- Zapfe, Helmuth. 1981. Ein eigenartiges *Dicerocardium* (Bivalvia) aus der obersten Trias der Tethys. Anzeiger der Akademie der Wissenschaften in Wien, Mathematisch-Naturwissenschaftliche Klasse 118(5):112–119.
- Zardini, Rinaldo. 1981. Fossili Cassiani (Trias Medio-Superiore). Atlante dei Bivalvi della Formazione di S. Cassiano raccolti nella regione Dolomitica attorno a Cortino d'Ampezzo. Edizione Ghedina. Cortina d'Ampezzo. 16 p. (*)
- Zenker, J. C. 1833. Beiträge zur Naturgeschichte der Urwelt. Organische Reste (Petrefacten) aus der Altenburger Braunkohlen-formation, dem Blankenburger Quadersand-stein, jenaischen bunten Sandstein und böhmischen Übergangsgebirge. Friedrich Mauke. Jena. 67 p.
- Zhang Jinghua. 1983. New data of Late Triassic (Noric) bivalves from Lanping and Weixi, Yunnan. Geological Review 30(4):377–381.
- Zhang Ren-jie. 1980. On the ligament area, systematics, position and evolutionary relationship of *Claraia*. Acta Palaeontologica Sinica 19(6):433–443.
- Zhang Renjie, & J. Pojeta, Jr. 1986. New bivalves from the Datang stage, Lower Carboniferous, Guangdong Province, China. Journal of Paleontology 60(3):669–679.
- Zhang Renjie, Wang Deyou, & Zhou Zuren. 1977. Mesozoic and Cenozoic bivalves. *In* Hubei Institute of Geological Sciences, ed., Palaeontological Atlas of Central South China, vol. 3. Geological Publishing House. Nanjing. p. 4–65. In Chinese.
- Zhang, Renjie, & Yan Daoping. 1993. Stratigraphic and Paleobiogeographic summary of Carboniferous marine bivalves of China. Journal of Paleontology 67(5):850–856.
- Zhang Wan-ping, & J. A. Grant-Mackie. 2001. Late Triassic-Early Jurassic palynofloral assemblages from Murihiku strata of New Zealand, and comparisons with China. Journal of the Royal Society of New Zealand 31(3):575–683.
- Zhang Yu-xiu. 1981. Late Permian bivalves from Yuanjia of Jiahe, Hunan Province. Acta Palaeontologica Sinica 20(3):260–265. In Chinese with English abstract.

- Zhang Zuo-ming. 1980. Discussion on the ligament structure of *Claraia* and its evolution. Acta Palaeontologica Sinica 19(6):433–443. In Chinese with English abstract.
- Zhao Jinke, Sheng Jinzhang, Yao Zhaoqi, Liang Xiluo, Chen Chuzhen, Rui Lin, & Liao Zhuoting. 1981. The Changhsingian and Permian-Triassic boundary of south China. Bulletin of Nanjing Institute of Geology and Palaeontology, Academia Sinica 2:1–112. In Chinese with English abstract.
- Zheng Y. J. 1993. Bivalves of Early Permian Yujiabeigou Formation in Chifeng area of Nei Mongol. Memoirs of Shenyang Institute of Geology Mineral Resources Chinese Academy of Geological Sciences 2:89–102. In Chinese.
- Zhu Minda, J. P. Cuif, Alain Denis, & Lucien Jaillard. 1990. Aragonitic structure in a *Palaeoneilo* (Mollusca, Pelecypoda, Nuculid) from the upper Trias of Yunnan (south China). Acta Palaeontologica Sinica 29(1):35–42. In Chinese with English abstract.
- Ziegler, A. M., M. L. Hulver, & D. B. Rowley. 1997. Permian world topography and climate. *In* I. P. Martini, ed., Upper Glacial and Postglacial Environmental Changes-Quaternary, Carboniferous-Permian and Proterozoic. Oxford University Press. New York. p. 111–146.
- von Zieten, C. H. 1830–1833. Die Versteinerungen Württembergs. Stuttgart. I–VIII, 1–16, pl. 1–12 (1830); p. 17–32, pl. 13–24 (1831); p. 33–64, pl. 25–48 (1832); p. 65–102, pl. 49–72 (1833).
- Zimmerman, E. H. 1886. Ein neuer Monomyarier aus dem ostthüringischen Zechstein (*Prospondylus liebeanus*). Jahrbuch der Preussischen Geologischen Landesanstalt zu Berlin 1885:105–119.
- Zittel, K. A. 1864. Fossile Mollusken und Echinodermen aus Neu-Seeland. Reise der Osterreichischen Fregatte Novara um die Erde. Novara Expedition Geologischer Theil 1(2):17–28.
- Zittel, K. A. 1881. Handbuch der Palaeontologie. Schimper, Schenk & Scudder, eds., Abt. 1, Palaeozoologie, Band 2 [Mollusca and Arthropoda]. Oldenburg, München and Leipzig. 893 p.
- Zittel, K. A., & E. Goubert. 1861. Description des fossiles du coral-rag de Glos. Journal of Conchyliologie 9:192–208.
- Zorn, Helmunt. 1971. Die Triasfauna der Tessiner Kalkalpen; XXI. Paläontologische, stratigraphische und sedimentologische Untersuchungen des Salvatoredolomits. (Mitteltrias) der Tessiner Kalkalpen. Schweizerische Paläontologische Abhandluungen 91:1–90.

Bupecten 158

INDEX

Acanomyphoria 116, 170 Acharax 19, 156 Actinostreon 63, 64, 65, 166 Adula 21 Agerchlamys 96, 100 Agonisca 120 Aguilerella 42, 43, 44, 45 Aguileria 157 Alectryonia 62 Amerinumopecten 157 Amonotis 52, 53, 55 Ampezzania 126, 128 Amphijanira 69, 73 Anadara 30 Anatina 163 Angustella 157 Anningella 77, 79 Anningia 79 Annulinectes 98, 99, 160 Anomalopleura 149 Anomalopleuroides 147, 149, 153 Anomia¹⁵⁷ Anradulonectites 157 Anshunopecten 157 Antijanira 69, 72, 73 Antiquicorbula 124, 125 Antiquilima 60, 61, 66 Aparimella 91, 92, 94 Aphanaia 157 Arcavicula 38, 39 Arcomya 149, 151 Arcomytilus 157 Arctomytiloides 35, 37 Arctotis 157 Arzelella 91 Asoella 84, 85, 86 Astarte 136, 137, 138, 139, 140, 142, 161, 162 Astartella 157 Astartellopsis 173 Astartopis 157 Astartopsis 136, 139 Atalantia 119, 121, 122, 130 Atomodesma 157, 173 Atreta 65, 66, 160 Atrina 53, 54, 59 Aucellina 78, 84 Avichlamys 96, 99 Avicula 38, 40, 46, 78, 81, 83 Avicularca 82, 83 Aviculolima 56, 61 Aviculomyalina 33, 34, 35, 162 Aviculopecten 72 Aviculopinna 54 Badiotella 56, 61 Bakevellia 40, 41, 42, 157 Bakevelloides 40, 157 Balantioselena 138, 140, 141 Bapristodia 27, 30 Barbatia 158 Baryvellia 41, 44 Bittneria 77, 78 Bittnericlaraia 2, 76 Bleta 173 Boreionectes 98, 164 Bositra 50, 52, 53, 55, 91, 167, 168 Bosniopecten 158 Botula 25 Botulopsis 23, 24, 25 Brachidontes 158 Bucardiomya 158

Burckhardtia 46, 47 Bureiomya 158 Burmesia 152, 153 Caenodiotis 52, 53, 55, 160 Caledogonia 119, 121 Calvaentolium 99, 103, 105, 158 Camptochlamys 98, 100 Camptonectes 96, 98, 99, 103, 160, 162, 164 Canadonectites 96, 100 Cardinia 66, 138, 140, 141, 142, 161 Cardiomorpha 158 Cardita 120, 121, 131, 135, 136, 137, 158 Cardium 128, 135, 143, 144, 146, 172 Carinocardia 129, 130, 131 Carnidia 173 Cassianella 38, 46, 47, 48 Catella 27, 28, 29 Ceratomya 153, 154, 159 Cercomya 155, 156, 163 Cerkesia 134 Chaenocardia 79, 159 Chiron 159 Chironopecten 159 Chlamys 72, 96, 97, 99, 100, 101, 103, 158, 160, 162, 168 Chuluaria 159 Cingentolium 103, 159, 165 Claraia 2, 68, 70, 71, 72, 76, 159, 161, 162, 167, 168, 169, 170 Claraioides 70, 159, 161, 169 Cochlearites 46 Coelastarte 136, 138, 139 Coelopis 136, 137, 140 Comatahalobia 91, 159 Conchodon 125, 126, 127, 128, 129, 130, 131 Corbula 125, 159 Corculum 130 Cornucardia 129, 130, 131 Cornutoentolium 103 Cortinia 147, 149 Cosmetodon 28, 29, 30 Costatoria 117, 118 Costentolium 103, 159, 165 Costibakevellia 40 Costicamptonectes 98 Costigervillia 45, 160 Cowperesia 173 Crenamussium 96, 99, 105, 158 Crenella 36 Crittendenia 2, 74, 76 Cruciella 173 Cryptocoelopis 137 Ctenostreon 60, 61, 62 Cucullaea 29, 30, 31, 32 Cultriopsis 41, 42, 157, 160 Cuneigervillia 42, 43, 50 Cuneolus 22 Curionia 107, 108, 109, 110, 111 Cuspidaria 156, 160, 170 Cyclopellatia 160 Cyrtopinna 173 Cyrtorostra 173 Dacryomya 11, 15, 18 Daonella 91, 92, 93, 94, 164, 169 Datta 48, 51 Dentiterquemia 88

Desiderinectes 160

Dianocuspidaria 160

Dianomya 147, 151 Dianucula 13, 15 Dicerocardium 126, 127, 129, 130 Dietrichia 160 Dimyodon 65, 66, 160 Diotis 52, 53, 160 Dipleurites 91 Durga 126, 127 Echinopecten 99 Edentula 42, 50 Elegantarca 32, 33 Elegantinia 119, 121, 163 Eleganuculana 10, 12 Ellesmerella 53, 55 Enantiostreon 62, 63, 70, 86, 160 Ennucula 9 Enormihalobia 73 Enosolen 156, 161 Ensia 173 Enteropleura 91, 92, 93, 172 Entolioides 104, 105, 106 Entolium 103, 104, 105, 106, 107, 157, 158, 159, 165, 166 Entomonotis 79 Eolimea 57 Eomonotis 79, 161 Eopecten 96, 100, 101, 103, 171, 172 Eophilobryoidella 30, 31 Eoplicatula 67, 68, 70, 168 Eoschizodus 161, 173 Eotrapezium 147 Epiclaraia 70, 161 Erugonia 119, 122 *Etalia* 84, 85, 86 Etheripecten 173 Eumorphotis 35, 69, 71, 72, 73, 74, 83, 84 Falcimytilus 20, 21, 22 Fengjiachonia 161 Filamussium 95, 161 *Filopecten* 104, 105, 106 *Fimbria* 134, 161 Frenguelliella 113, 114, 116, 118 Gemmellarodus 128, 161, 170 Geratrigonia 114, 115 Gerlus 173 Gervillancea 41, 43, 44 Gervillaria 41, 42, 43, 44 Gervilleiognoma 46 Gervilleioperna 43, 45, 46, 49 Gervillella 40, 41, 43, 45 Gervillia 40, 41, 42, 43, 44, 45, 48, 50, 157, 160 Gibboconcha 161 Gingillum 173 Glyptoleda 11, 12, 161, 166 Gonilia 162 Goniomya 147, 149 Grabella 91 Grammatodon 27, 28, 29, 30 Granulochlamys 97, 168 Gresslya 154, 155 Groeberella 123, 124 Gruenewaldia 119, 120, 121, 163 Gryphaea 46, 62, 63, 65, 66, 169 Gryphellina 162 Guichiella 70, 162 Guineana 114, 116 Guizhoumyophoria 119, 122, 123, 170 Gythemon 168

Habonucula 162 Halobia 73, 91, 92, 93, 94, 159, 162, 164, 166, 167, 173 Harpax 67, 68, 168 Healeya 23, 26, 28, 111, 112 Hemimenion 162 Heslingtonia 113 Heteropecten 71 Hiatella 125 Hijitrigonia 115 Hippopodium 112, 113 Hoernesia 41, 42, 43, 44, 47 Hoernesiella 47, 48 Hoferia 31, 32, 33 Hokonuia 77, 78 Homomya 147, 149, 151, 152 Hunanonectes 103, 162 Imposidonia 162 Indigirohalobia 91, 162, 168 Indigiropecten 101 Indogrammatodon 29 Indepecten 66, 96, 98, 162 Inflatomonotis 79 Inoceramus 36, 162, 165 Inoperna 20, 21 Integribakevellia 40 Integricardium 144, 172 Iranopecten 162 Isocardioides 133, 163 Isocyprina 146, 147, 148 Isognomon 46, 49, 51 Isolimea 57 Isopristes 138, 141 Janeia 19 Janopecten 96, 101, 102 Jaworskiella 114, 116 Jianchuania 82, 84 Joannina 23, 24, 26 Jurassicardium 163 Kalentera 108, 109, 110 *Kija* 161 Kolymonectes 94, 95 Krumbeckia 127, 133, 163 Krumbeckiella 23, 89, 90 Kumatrigonia 113, 116 Kyushutrigonia 114, 116 Labayophorus 110 Laevitrigonia 163 Langsonella 42, 43 Langvophorus 146, 148 Lapteviella 13, 15 Laternula 156, 163 Laubeia 129, 131 Lecompteus 163 Leda 11, 18 Ledoides 12 Leidapoconcha 23, 24, 26, 28 Leproconcha 49, 50, 51 Leptochondria 72, 75, 76, 77, 85, 100 Leviconcha 163 Lilangina 47, 48 *Lima* 56, 60 *Limatula* 57, 61 *Limea* 57, 61 Liostrea 64, 65, 66 Liotrigonia 119, 122 Lithioperna 46 Lithiotis 46, 166, 167 Lithophaga 20, 21, 22

Lockeia 12 *Loemmelella* 91 *Longidaonella* 91 *Lopha* 62, 63, 64, 65, 66, 173 Loxochlamys 96, 102 *Lucinia* 31, 133 *Luciniola* 131, 133 *Lupherella* 81, 163 Lycettia 20, 22 Lyriomyophoria 121, 163 Lywea 98 Macrodontella 27, 28 Mactromya 173 Mactromyella 164 Mactromyopsis 163 Magnolobia 91, 164 Malletia 164 Manticula 89, 90 Maorimonotis 79 Maoritrigonia 122, 123, 124 Martesia 164 Marwickiella 77, 78, 79 Mclearnia 98, 164 Meekopinna 54 Megalodon 128, 161, 164 Meleagrinella 82, 83, 84 Mesomiltha 164 Mesoneilo 11, 15, 17, 18 Mesosaccella 14 Minepharus 138, 142 Minetrigonia 122, 123 Modestella 113 Modiolopsis 110, 112 Modiolus 20, 21, 64, 107, 109, 111, 165 Molukkana 164 Monotis 79, 80, 81, 83, 100, 109, 161 Moussonella 91 Musculus 165 Multisidonia 53 Mya 149, 151, 152, 153, 155 Myacites 152 Myalina 33, 34, 35, 44, 90 Myalinella 33, 34 Myoconcha 22, 66, 107, 110, 111 Myonia 166 Myophoria 47, 118, 119, 120, 121, 123, 124, 142, 158, 163 Myophoricardium 141, 142, 143 Myophorigonia 122, 123 Myophoriopis 141, 142, 143, 158 Mysidia 23, 25 Mysidiella 23, 24, 25, 90, 165 Mysidioptera 35, 57, 58, 61, 98 Mytiliconcha 23, 25, 26, 171 Mytiloconcha 23, 171 Mytiloides 36, 165 *Mytiloperna* 46, 49 *Mytilus* 20, 21, 22, 23, 25, 37, 44, 90, 159, 163, 165, 170 Nacrosolemya 156 Neobakevellia 40 Neocrassina 136, 138, 139 Neoentolium 103, 159, 165 Neomegalodon 126, 128, 129, 161, 164, 170 Neomorphotis 69, 73, 74 Neopecten 165 Neoschizodus 119, 120, 121, 163 Neotrigonia 113, 114, 118 Neptunella 165 Neptunopecten 165

Nevadapecten 96, 102

Newaagia 86, 87, 88, 171 Nicaniella 136, 139 Ningliconcha 13, 16 Nodomytilus 163 Nucinella 17, 19 Nucula 8, 9, 14, 16, 18, 143, 159, 165 Nuculana 9, 10, 11, 12, 14, 16, 18 Nuculoma 8, 9, 162 Nuculopsis 8, 165 Nucundata 12, 162, 166 Oceanopieris 28 Ochotochlamys 96, 101 Ochotomya 154 Okunominetania 120 *Opis* 25, 26, 136, 140 *Opisoma* 137, 166 *Oretia* 89, 90, 91 Ornithopecten 69, 72 Orthomya 155, 156 Orthotrigonia 115 Osteomya 150, 151, 152 Otapiria 80, 81, 163, 167, 169, 172 Ouamouia 110 Oxypteria 69, 72 Oxytoma 38, 73, 81, 82, 83, 84 Pachymegalodon 126, 127 Pachymya 147, 149, 151, 164 Pachymyonia 166 Pachyopis 140 Pachyrisma 126, 127 Pacifihalobia 91, 166 Pacimonotis 79 Palaeocardita 128, 134, 135, 137 Palaeoentolium 166 Palaeolima 56, 61 Palaeolopha 63, 166 Palaeoneilo 12, 13, 14, 15, 16, 18, 164, 171 Palaeontolium 103 Palaeonucula 8, 9, 18, 165 Palaeopharus 142 Palmoxytoma 82, 84 Panopea 173 Parahalobia 91, 166 Parainoceramus 35, 36, 162 Parallelodon 28, 29 Paramegalodus 126, 127, 129 Parapergamidia 89, 90 Parvamussium 94, 95, 97, 161, 168 Paullia 166 Pecten 56, 57, 72, 73, 76, 78, 84, 95, 97, 98, 99, 101, 102, 103, 105, 106, 158, 159, 160, 161, 165, 172 Pectinula 106, 167 Pegmavalvula 87, 88, 89 Perampliata 167 Pergamidia 89, 90 Peribositra 167 Peribositria 50, 167 Periclaraia 96, 102, 103 Perihalobia 91, 167 Permoperna 50 Permophorus 107, 108, 109, 110 Perna 40, 44, 49 Pernopecten 105, 166 Persia 58, 87, 88 Perugonia 122, 123, 124 Phaenodesmia 11, 12, 13, 14, 18 Phestia 10, 11, 15, 18, 168 Philippiella 88 Pholadomya 147, 148, 149, 151, 152, 153, 154, 155, 158 Pholadomyocardia 167

Ros-Franch, Márquez-Aliaga, & Damborenea—Triassic and Early Jurassic Bivalvia

Phymodonucula 167 Physocardia 129, 130 Pichlerella 91, 93 Pichleria 32 Pinna 50, 53, 54, 59, 66, 93 Placunopsis 70, 157, 167 Plagiostoma 58, 61, 62, 66 Platymyoidea 167 Plesiocyprina 167 Pleuromya 149, 150, 152, 153 Pleuromysidia 81, 167 Pleuronectites 96, 99, 100, 102, 103, 166 Plicatostylus 167 Plicatula 67, 68, 70, 168 Polidevcia 11, 168 Pomarangina 126, 127 Posidonia 50, 53, 162, 168 Posidonotis 53, 104, 106, 167 Praechlamys 97, 168 Praeconia 136, 139, 140 Praegonia 114, 115 Praeotapiria 81, 172 Praesaccella 14 Primahalobia 91, 168 Primahinnites 69, 73 Prolaria 152, 153 Promyalina 33, 34, 35, 72 Promysidiella 23, 24, 25, 165, 170 Promytilus 20, 21, 33 Pronoella 168 Propeamussium 95, 161, 168 Prorotrigonia 114, 115 Prosogyrotrigonia 114, 115 Prosoleptus 13, 14 Prospondylus 168 Protocardia 143, 144 Protodiceras 126, 168 Protopis 23, 24, 25, 26, 171 Protostrea 65, 66, 67 Psammoconcha 173 Pseudacesta 57 Pseudastarte 138, 142 Pseudoburmesia 152, 153 *Pseudoclaraia* 70, 159, 168 *Pseudocorbula* 141, 143 Pseudolimea 57 Pseudomonotis 2, 38, 62, 71, 74, 76, 81, 83, 168, 169 Pseudomyoconcha 110, 111, 112 Pseudomytiloides 35, 36, 37 Pseudopachymytilus 169 Pseudopecten 96, 99, 100 Pseudopis 169 Pseudoplacunopsis 67, 70, 157, 167, 168 Pseudosaxicava 124, 125 Pseudotrapezium 146, 148 Pseudovola 98 Pteria 37, 38, 39, 56, 83, 84, 169, 170

Pterinea 46 Pteroclaraia 70, 169 Pterohalobia 169 Pteromya 153, 154 Pteronites 54 Pteroperna 37, 169 Pteropiria 169 Ptychostolis 169 Pulvinites 170 Qingyaniola 23, 24, 26, 28 Quadratia 122, 170 Quadratojaworskiella 116 Quemocuomegalodon 126, 128 Radiastarte 136, 139 Radulonectites 96, 100, 103, 162 Rebusum 173 Retroceramus 170 Reubenia 47, 48 Rhaetavicula 38, 39 Rhaetidia 144, 145, 146 Rhaetomegalodon 126, 127, 129, 130 Rhynchopterus 37, 170 Rollieria 15, 16 Rossiodus 128, 161, 170 Rugiclaraia 70, 170 Ryderia 15, 16, 18, 171 Sakawanella 145, 146 Saturnella 170 Saturnopecten 170 Schafhaeutlia 127, 132, 133, 134, 135, 163 Schizocardita 134, 137 Scythentolium 104, 105, 106 Semuridia 89, 90 Septifer 170 Septocardia 134, 135, 136 Serania 58, 60, 61 Sichuania 77, 78, 79 Sichuantrigonia 116, 170 Sinbadiella 131, 132, 133 Solemya 17, 19, 156 Solemyatuba 19 Solenomorpha 160, 170 Somapteria 171 Somareoides 107, 108, 109 Songdaella 43, 44, 45 Sowerbya 145 Sphaeriola 132, 134, 135 Ŝpia 40 Śpondylopecten 171 Spondylus 88, 171 Stefaninia 38, 39, 40 Steinmanella 22 Steinmannia 53, 55 Storthodon 132, 133 Streblochondria 2, 171, 172

Streblopteria 74, 76, 171 Streblopterinella 171 Taeniodon 173 Taimyrodon 171 Tancredia 145, 146, 148 Teinonuculana 16, 171 Terquemia 63, 86, 87, 88, 160 Terraia 173 Thracia 154, 155, 156 Timoria 89 Tirolidia 60, 61 Tommasina 23, 25, 26, 171 Torastarte 138, 142 Tosapecten 96, 98, 101, 102 Towapteria 43, 45, 83 Triadomegalodon 126, 128, 164 Triaphorus 107, 108, 109 Triasoperna 21 Trichites 172 Trigonia 113, 114, 115, 116, 118, 120, 121, 122, 123, 124, 140, 163 Trigonopis 140 Trigonucula 8, 9 Tulongella 155, 156, 161 Tulongocardium 144, 172 Tutcheria 134, 135 Umbrostrea 62, 63, 64, 86, 161 Unicardium 173 Unionites 35, 72 Variamussium 95 Vaugonia 114, 115, 116 Veldidenella 52, 55 Ventalium 172 Veteranella 10, 11, 12, 161, 166 Vietnamicardium 144, 172 Virgellia 43, 45 Vokesella 172 Waagenoperna 42, 50, 51 Waijiaoella 23, 24, 26, 28 Wallerobia 172 Wallowaconcha 129, 130, 131, 132 Weixiella 108, 110 Weyla 96, 98, 101 Xiaoschuiculana 10 Xinanopecten 172 Yokoyamaina 144, 172 Yoldia 15 Yongshengia 13, 16 Yunnanomya 151 Zandaia 172 Zittelihalobia 91, 173