Chemosymbiotic bivalves from the late Pliocene Stirone River hydrocarbon seep complex in northern Italy

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Seven species of chemosymbiotic bivalves are described from the late Pliocene Stirone River hydrocarbon seep complex in northern Italy, including one new species and two in open nomenclature. The known species are the solemyid *Acharax doderleini*, the lucinids *Lucinoma persolida* and *Megaxinus ellipticus*, and the vesicomyid *Isorropodon aff. perplexum*; in open nomenclature we report two lucinids, including the largest species of *Lucinoma* known from the Italian Pliocene to date, and a strongly inflated, large *Anodontia* sp. The most abundant species at the Stirone seep complex is the lucinid *Megaxinus stironensis* sp. nov. This Pliocene seep fauna differs from that of the well-known Miocene “Calcari a Lucina” seep deposits by lacking large bathymodiolin mussels and vesicomyid clams; instead, the dominance of the lucinid *Megaxinus stironensis* gives this fauna a unique character. We speculate that at the Stirone seep complex, *Megaxinus* had occupied the ecological niche that *Meganodontia* occupied at the Miocene “Calcari a Lucina” seep sites in the Mediterranean basin, and that the dominance of *Megaxinus* could be a wide-spread feature of Pliocene chemosynthesis-based ecosystems in Mediterranean Pliocene.

**Key words:** Bivalvia, Lucinidae, Vesicomyidae, hydrocarbon seep, chemosymbiosis, Pliocene, Italy, Apennines.

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**Introduction**

Hydrocarbon seeps are places on the seafloor where fluids rich in methane and hydrogen sulfide reach the seabed (Suess 2014). These sites are often inhabited by a highly specialized macrofauna of low diversity but high biomass, which takes advantage of nutrients provided by chemosynthetic bacteria that use methane and sulfide to synthesize organic matter (Paull et al. 1985; Childress et al. 1986; Sibuet and Olu 1998). The fauna inhabiting hydrocarbon-seeps in the Mediterranean Sea has an impoverished character (Olu et al. 2004; Ritt et al. 2010, 2011; Taviani 2011; Taviani et al. 2013) compared to those known from the continental margins of the Atlantic Ocean proper or from the Indian and Pacific oceans (Van Dover et al. 2002). In the Mediterranean basin, seep deposits and associated fauna communities have a long geologic history, with several Mesozoic examples (Lemoine et al. 1982; Gaillard et al. 1992; Kiel and Peckmann 2008), a few Eocene examples (Venturini et al. 1998; Natalicchio et al. 2015), and the well-known “Calcari a Lucina” sites of the Italian Miocene, which host a fauna with “oceanic” character (Moroni 1966; Taviani 1994, 2011, 2014; Kiel and Taviani 2017). However, the timing of the faunal change from an “oceanic” fauna in the Miocene to the present-day impoverished fauna is poorly constrained due to the scarcity of fossil examples from the intervening time interval. A hydrocarbon seep complex of late Pliocene age has been recently described in detail from strata of the Argille Azzurre (Lugagnano Formation) along the Stirone River (Fig. 1) in the Italian Northern Apennines (Cau et al. 2015). The purpose of the present contribution is to provide a detailed report of the associated chemosymbiotic bivalve fauna.

**Institutional abbreviations.—**MGGC, Museo Geologico Giovanni Capellini, University of Bologna, Italy; MSF, Museo Civico di Scienze Naturali, Faenza, Italy; NRM, Swedish Museum of Natural History, Stockholm, Sweden; ZMB, Zoological Museum, University of Bologna, Italy.
Geological setting

This remarkable and to-date unique Pliocene-age situation was first reported on by Taviani et al. (1997) who also provided information on its chemosymbiotic bivalve fauna (Solemya doderleini, Megaxinus cf. incrassatus, and ?Lucinoma sp.). Geological and stratigraphic information on the Stirone River hydrocarbon seep has been added over the years (Monegatti et al. 2001; Barbieri and Cavalazzi 2005; Taviani 2014; Capozzi et al. 2013, 2017) with also some fragmentary information and figures on its chemosymbiotic bivalve fauna (Taviani 2001, 2011; Raineri 2007; Taviani and Ceregato 2009). The detailed work by Cau et al. (2015) summarized and implemented all previous information, further providing detailed lithostratigraphic logs. Discrete hydrocarbon-imprinted carbonate bodies of different thickness and degree of lithification and host muddy sediment punctuate the Stirone section at the Zanclean/Piacenzian boundary, and the benthic foraminifera and mollusk fauna indicate a depositional depth not exceeding 500 m (Cau et al. 2015). Such horizons are the source of the macropaleontological material discussed here (Fig. 1).

Material and methods

The material was collected by the authors and co-workers over many years since 1997. Specimens were coated with ammonium chloride prior to photography.

Systematic palaeontology

Class Bivalvia Linnaeus, 1758
Subclass Protobranchia Pelseneer, 1889
Family Solemyidae Gray, 1840
Genus Acharax Dall, 1908
Type species: Solemya johnsoni Dall, 1891 (by original designation); Recent, Pacific.

Acharax doderleini (Mayer, 1861)

Fig. 2.

Fig. 2. The solemyid bivalve Acharax doderleini (Mayer, 1861) from the late Pliocene of Italy, Stirone River seep deposit; MGGC.22007 view on exterior of right valve.

Subclass Heterodonta Neumayr, 1884
Family Lucinidae Fleming, 1828
Genus Megaxinus Brugnone, 1880
Type species: Lucina transversa Bronn, 1831 (subsequent designation by Pallary 1904, see Glover and Taylor 1997 for a detailed discussion); Pliocene, Italy.

Megaxinus ellipticus (Borson, 1825)

Fig. 3.

Fig. 3. The lucinid bivalve Megaxinus ellipticus (Borson, 1825) from the late Pliocene of Italy, Stirone River seep deposit; MGGC.22008 view on exterior of right valve.
Remarks.—This species is common in the Mediterranean Pliocene (Chirli 2015) and our specimen agrees well with those illustrated by Sacco (1901: pl. 17). It has also been reported from the Pleistocene of Italy (Chirli 2015) but many records may prove in the future to be misidentifications since encompassing too many varieties of habitats, shallow to deep. A similar species is *Megaxinus rostratus* (Pecchioli, 1864), also from the Pliocene of northern Italy, which may or may not be synonym with *M. ellipticus* (Sacco 1901; Glover and Taylor 1997).

Stratigraphic and geographic range.—According to Sacco (1901) this species is found in late Miocene to late Pliocene sediments in northern Italy, and there are unconfirmed reports from the middle Miocene of Poland (Studencka and Studencki 1988), the late Miocene of southern Turkey (Islamoglu and Taner 2003), and the late Pliocene of Crete (Drinia et al. 2005).

*Megaxinus stironensis* sp. nov.

Figs. 4, 5.

1901 *Megaxinus bellardianus* (Mayer); Sacco 1901: 75–76, pl. 17: 30, 36; not 29, 32–35, 37.

Etymology: For the Stirone River.

Type material: Holotype: MGGC.22009, a shell with both valves preserved. Paratypes: articulated and single shells MGGC.22010–22013, MSF 2143, NRM Mo184001–184003; all from the type locality and horizon.

Type locality: The Stirone River seep deposit, Northern Apennines, Italy.

Type horizon: Late Pliocene part of the Lugagnano Formation.

Material.—The type material; hundreds more seen in the field.

Dimensions.—The largest specimens including the holotype reach 45 mm in length and height, individual valves are 13 mm thick.

Diagnosis.—Average-sized, moderately inflated *Megaxinus*, of equal height and length, with small lunule, two incisions on anterodorsal margin, strong and narrow hinge plate, and very long anterior adductor muscle scar.

Description.—Shells of average size for the genus, almost circular, of equal height and length, moderately inflated (T/L = 0.29), with blunt, elevated, prosogyrate umbo. Posterostral area broad, bordered by indistinct, broad, shallow groove resulting in a slight angulation on the posterior margin, another indistinct, broad and shallow groove running from the umbo to the center of the ventral margin, where it results in a shallow indentation. Lunule very small, surrounded by two incisions on the anteroventral margin; ligament external, elongate in a sunken groove. Outer shell surface with irregular commarginal growth increments. Hinge plate strong and narrow, edentulous; nympha plate long and thin. Inner shell surface with few or many pustules that are arranged radially toward the ventral margin; in some specimens the pustules are so numerous that they give the interior a granulate appearance; pallial blood vessel.

Fig. 3. The lucinid bivalve *Megaxinus ellipticus* (Borson, 1825) from the late Pliocene of Italy, Stirone River seep deposit. MGGC.22008, close-up of hinge region of RV (A), view of left valve exterior and right valve hinge (B), view of exterior of right valve (C); dorsal view on right valve (D).
scar usually distinct; anterior adductor muscle scar broad and very elongate, tapering ventrally, reaching the ventral quarter of the shell, close to pallial but detached from it for ¾ of its length; posterior adductor muscle scar indistinct, irregular ovate in outline. Pallial line entire, crenulate; area beyond pallial line crenulate close to pallial line but mostly smooth.

Remarks.—This is the most common species at the Stirone River seep deposit and it also occurs in large numbers in shell beds in the surrounding mudstone. Sacco (1901: 75–76) assigned numerous specimens of middle Miocene to late Pliocene age to *Megaxinus bellardianus* (Mayer, 1864). These specimens show quite a diversity of shapes, some being similar and others being dissimilar to the specimens.
from the Stirone seep deposit. Given the rich and cryptic present-day diversity of this genus and related lucinid genera (Glover and Taylor 1997; Taylor and Glover 2005) we prefer to apply to a more narrow species concept to the Neogene *Megaxinus* than Sacco’s (1901) approach of lumping most specimens into *M. bellardianus*, although a full revision of Neogene *Megaxinus* is beyond the scope of this study. Mayer (1864: 27) wrote that the type specimen of *M. bellardianus* is the specimen illustrated by Michelotti (1847: pl. 4: 10). That specimen is of Miocene age and the provided drawing indicates that it has a more pronounced posterodorsal ridge, and a more pointed and less prosogyrate beak than the specimens from the Stirone seep deposit concerned here. Thus due to the differences in shell shape, we consider the Stirone specimens to be distinct from *M. bellardianus* and introduce the new species *M. stironensis* for them. Among the many specimens of *M. bellardianus* illustrated by Sacco (1901: pl. 17: 30, 36) resemble the Stirone specimens in every aspect, and interestingly are from the late Pliocene (Piacenzian) of Vezza d’Alba, and hence of similar age as the Stirone specimen. We include those specimens from Vezza d’Alba in *Megaxinus stironensis* sp. nov.

The extant *Megaxinus yemenensis* Glover and Taylor, 1997 differs from *M. stironensis* by being elongate-oval rather than round, the extant *M. appendiculatus*, *M. unguiculus*, and the Neogene Italian *M. rostratus* (Pecchioli, 1864) and *M. transversus* (Bronn, 1831) differ from *M. stironensis* in having a truncate posterior margin and a more pointed anterior margin (i.e., Sacco 1901; Glover and Taylor 1997). Among the geographically close, extant species from the Gulf of Oman, *M. omanensis* (Smith, 1906) differs from *M. stironensis* by having a higher and more angular shell and a smaller and less prosogyrate beak, whereas *M. arabicus* Glover and Taylor, 1997 has a lower shell with pointed anterior margin and a narrower anterior adductor muscle scar (Glover and Taylor 1997).

Similar specimens associated with Pliocene whale-fall communities around Italy (Dominici et al. 2009; Danise et al. 2010, 2014) have been assigned to *Megaxinus incrassatus* du Bois de Montpéreux, 1831. However, there are three reasons why we find this questionable. First, the type specimen...
of *M. incrassatus* is from a middle Miocene locality on the Volyn'-Podollya Plateau in the western Ukraine. The specimen was described as “not well preserved” but the illustration shows a very nicely preserved shell (du Bois de Montpéreux 1831: 58, pl. 6: 1–3). The illustration is thus most likely an idealized reconstruction that should not be taken at face value. Second, specimens of *Megaxinus incrassatus* from the type locality or nearby illustrated by Friedberg (1934–1936: pl. 20: 1, 2), which are probably the best available approximation to the type specimens, have a shorter anterior adductor muscle scar and much smaller umbones. Third, the type locality is of Miocene rather than Pliocene age, and Sacco (1901) noted that *M. incrassatus* is rare in the Miocene of Italy, and was unknown to him from Pliocene deposits. The specimen illustrated from the Orciano Pisano whale fall (Danise et al. 2010: fig. 7A) has a shallower lunule and a longer anterodorsal margin than *Megaxinus stironensis*, and may be yet a different (or new) species.

**Stratigraphic and geographic range.**—Late Pliocene of northern Italy.

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Fig. 6. The lucinid bivalve *Lucinoma persolida* (Sacco, 1901) from the late Pliocene of Italy, Stirone River seep deposit. MGGC.22014, left (A) and right (B) valves; in dorsal views (A₁, B₁), views of the interior including hinge and muscle attachment scars (A₂, B₂), and views of exterior (A₃, B₃).
Genus *Lucinoma* Dall, 1901

*Type species:* *Lucina filosa* Stimpson, 1851 (by original designation); Recent, northwest Atlantic.

*Lucinoma persolida* (Sacco, 1901)

Figs. 6, 7.

1901 *Dentilucina persolida* sp. nov.; Sacco 1901: 82–83, pl. 19: 1, 2. 2011 *Lucinoma* sp.; Taviani 2011: fig. 3h.

*Material.*—Four specimens; the largest specimen (MGGC.22014) consists of two disarticulated valves, 50 mm long and 46 mm high; the others (MGGC.22015, NRM Mo184004, NRM Mo184005) are articulated and single valves partly embedded in rock matrix; from the late Pliocene of Italy, Stirone seep complex.

*Remarks.*—*Lucinoma persolida* is distinct from the two extant Mediterranean *Lucinoma* species, *L. kazani* and *L. borealis*, by being less inflated and by having a more distinctive lunula. Among the two, *L. borealis* is more similar to *Lucinoma persolida* because of its similar outline and its straight anterior adductor muscle scar that stays close to the pallial line. The extant *L. gagei* Oliver and Holmes, 2006 from the Oman margin differs from *L. persolida* by its narrower anterior adductor muscle scar and its much more distinct posterodorsal groove.

Among fossil species, *Lucinoma perusina* (Sacco, 1901) from the classical Miocene “Calcare a *Lucina*” deposits throughout Italy (cf. Moroni 1966; Taviani 1994; Kiel and Taviani 2017) is more inflated, has less protruding umbones, and its posterodorsal area is more well-defined, by being broader and flatter, and more angular. Quite similar to *L. persolida* regarding shell outline is the early Miocene Japanese *Lucinoma shinokii* Hirayama, 1954, but it is more strongly inflated than *L. persolida* and its internal features are unknown (Hirayama 1954). *Lucinoma acutilineata* (Conrad 1849), the most common lucinid at Pliocene seep deposits in Japan (Majima et al. 2005a) differs from *L. persolida* by its more angular outline of the whorl. Compared to the recently
described *Lucinoma saetheri* from Miocene seep deposits in New Zealand (Amano et al. 2018), *L. persolida* has sharper ribs and a lower shell with more prominent umbones.

**Stratigraphic and geographic range.**—Late Pliocene of northern Italy.

*Lucinoma* sp.

Fig. 8.

**Material.**—MGGC.22016, a single articulated, thick-shelled specimen, 70 mm long, 67 mm high, and 39 mm thick; from the late Pliocene of Italy, Stirone seep complex.

**Remarks.**—The only known specimen was previously figured as *Lucina* sp. by Taviani and Ceregato (2009). It is very similar to the middle–late Miocene *Lucinoma perusina*, from which it may differ by its broader and shorter lunule and less distinct posterodorsal area. To date, this is the largest deep-water chemosymbiotic lucinid known in the Mediterranean after the Miocene. It could be a new species but we prefer at present to leave in open nomenclature.

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Genus *Anodontia* Link, 1807

*Type species:* *Anodontia alba* Link, 1807 (by monotypy); Recent, Caribbean Sea.

*Anodontia* sp.

Fig. 9.

**Material.**—Two specimens: MGGC.22018 is an articulated shell reaching 25 mm in length, MGGC.22017 is a 23 mm long left valve; from the late Pliocene of Italy, Stirone seep complex.

**Description.**—Small to medium-sized *Anodontia* (sensu lato), strongly inflated (T/L ca. 0.35), umbones elevated, somewhat prosogyrate; posterodorsal area poorly defined; no lunule; external surface with irregular, commarginal growth lines. Hinge plate very narrow and edentulous, ligament long and external, sunken in narrow groove, ligament plate thin.

**Remarks.**—The *Anodontia* group is known to host a remarkable cryptic diversity (Taylor and Glover 2005). Sacco (1901) and Chirli (2015) assigned specimens of *Anodontia* (sensu lato) from the Pliocene of Italy to the extant shallow-water species *Anodontia fragilis* (Philippi, 1836). But most likely the Stirone representative is a new taxon; unfortunately, features of the pallial line and muscle scars are unknown, hence we keep these specimens in open nomenclature. *Anodontia* has been also reported from a Miocene seep deposit in northern Italy (Kiel et al. 2018). However, it is apparently excluded from modern deep-water cold seep situations in the eastern Atlantic Ocean and Mediterranean.

Family Vesicomyidae Dall and Simpson, 1901

Genus *Isorropodon* Sturany, 1896

*Type species:* *Isorropodon perplexum* Sturany, 1896 (by monotypy); Recent, eastern Mediterranean Sea.

*Isorropodon aff. perplexum* Sturany, 1896

Fig. 10.

**Material.**—Two fragmentary specimens, preserving the hinge area of a left and a right valve (MGGC.22019 and MGGC.22020); from the late Pliocene of Italy, Stirone seep complex.

**Description.**—Prodissoconch roundish, about 220 μm wide. Beak somewhat elevated, slightly prosogyrate; surface smooth; lunular incision present. Hinge plate narrow, three teeth in each valve; in right valve, cardinal 1 thin just anterior of beak, elongate, subparallel to dorsal margin, cardinal 3a commencing slightly anterior of beak, elongate and very thin, slightly thickened anteriorly, parallel to dorsal shell margin, bends slightly ventrally at its posterior end; cardinal 3b just posterior of 3a, equally thin, very slightly oblique to dorsal shell margin; in left valve, cardinal 2a short, parallel to anterior shell margin, connected through a thin ridge with the stout, triangular cardinal 2b; 4b elongate, thin, parallel to dorsal shell margin.
Remarks.—The hinge dentition of the two available fragmentary specimens agrees well with that of extant Mediterranean *Isorropodon perplexum* (Sturany, 1896) as figured by Cosel and Salas (2001) and also the prodissococonch shown here on Fig. 9B, is similar in size and shape to that of *Isorropodon perplexum*. A single left valve identified as *Isorropodon aff. perplexum* has been reported from late Miocene (Messinian) sediments near Torino, northern Italy (Janssen and Krylova 2012) and seems to have a slightly shorter hinge plate than the late Pliocene specimens figured here. Two extant *Isorropodon* species are known from mud volcanoes in the Gulf of Cadiz in the NE Atlantic (Oliver et al. 2011). Among them, *Isorropodon megadesmus* Oliver, Rodrigues, and Cunha, 2011 differs from the Pliocene *Isorropodon aff. perplexum* by its higher hinge plate and a curved cardinal 3a; and *Isorropodon* sp. indet. has a hinge similar to that from the Stirone River but its cardinal 3b appears more undulating. Also the hinge dentition of the extant *Isorropodon nyeggaensis* from the North Atlantic off of northern Norway (Krylova et al. 2011) is similar to that of the Stirone specimens, although its nymph appears to be broader. At this Norwegian cold seep site, this vesicomyid (as *Calyptogena* sp.) occurs with *Acharax* and thyasirids in ca. 700 m water depth (Ivanov et al. 2010).

Concluding remarks

The scarcity of *Lucinoma* at the Stirone River seep is remarkable considering that *Lucinoma* is very abundant at many Miocene “Calcari a Lucina” sites (Taviani 1994, 2014) as well as at many seep sites in the present-day Mediterranean Sea (Salas and Woodside 2002; Olu et al. 2004; Taviani et al. 2013). The Stirone River seep fauna is also quite distinct from the middle to late Miocene “Calcari a Lucina” fauna in northern Italy due to the lack of large vesicomyid clams and bathymodiolin mussels. These large “oceanic” species likely disappeared from the Mediterranean basin due to the Messinian Salinity crisis (cf. Taviani 2001, 2003, 2011; Olu et al. 2004; Cau et al. 2015).

The lucinid genus *Megaxinus* that dominates the Stirone seep deposit is still living in the Mediterranean Sea today (Glover and Taylor 1997) but has never been reported from any Mediterranean seep site (Olu et al. 2004; Ritt et al. 2010, 2011; Taviani et al. 2013; Taviani 2014). Although the precise habitat of extant *Megaxinus* is unknown, the lack of extant specimens with preserved soft anatomy might indicate that it is a deep burrower. It could potentially have occupied a similar ecologic niche as *Meganodontia* had in the Miocene “Calcari a Lucina” sites, because also *Meganodontia* is a deep burrower and the extant *M. acetabulum* lives at depths down to 472 m (Bouchet and Cosel 2004). This depth range is similar to both that of extant *Megaxinus* (Glover and Taylor 1997; Cosel and Bouchet 2008) and the estimated paleodepth of the Stirone seep complex (Cau et al. 2015). Whether the dominance of *Megaxinus* is a unique feature

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**Fig. 9.** The lucinid bivalve *Anodontia* sp. from the late Pliocene of Italy, Stirone River seep deposit. **A.** MGGC.22017, smaller specimen, close-up of edentulous hinge area of left valve (A1), dorsal view (A2), and view of outer side of left valve (A3). **B.** MGGC.22018, larger specimen, left valve, exterior view.
of this particular seep complex, or has occurred more widespread in the Mediterranean Pliocene, remains to be tested. However, remarkable in this context is the abundance of a species of Megaxinus associated with a Pliocene whale fall in northern Italy (Dominici et al. 2009; Danise et al. 2010; Danise and Dominici 2014). Lucinids are generally uncommon at whale falls, both fossil (Goedert et al. 1995; Amano and Little 2005; Kiel and Goedert 2006; Amano et al. 2007; Jenkins et al. 2017) and Recent (Smith and Bacca 2003). Thus the abundance of Megaxinus at both the Stirone seep complex and the Orciano Pisano whale fall support the hypothesis that Megaxinus had a prominent role in chemosynthesis-based ecosystems in the Mediterranean Pliocene.

Pliocene seep deposits are not only rare in the Mediterranean Basin but also world-wide. Most records are from the western Pacific and are dominated by “oceanic” taxa such as large vesicomyids, bathymodiolins and lucinids, as well as the thyasirid Conchocele (Nobuhara 2003; Majima et al. 2005a, b, 2007; Wang et al. 2006; Amano and Kiel 2010, 2012). The same applies to three potential Pliocene seep deposits from the eastern Pacific (Olsson 1942; Squires 1991; Campbell 1992). Thus with Megaxinus being unknown not only from Miocene and present-day Mediterranean seeps, but also from extant and fossil seeps world-wide, the Stirone seep fauna has a unique “Mediterranean Pliocene” character.

The uniqueness of the Stirone seep complex and of its valuable accompanying fauna deserves proper measures of protection under the status of a dedicated geosite (Taviani 2014). Fortunately, the site is already located within the Stirone River Regional Park (Raineri 2007) what guarantees its safeguard for the time being.

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