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# Indian Ocean echinoderms collected during the *Sindbad Voyage* (1980–81): 3. Ophiuroidea and Echinoidea

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**SYNOPSIS.** At least 44 ophiuroid and 11 echinoid species are recorded from echinoderm collections made during an international expedition, the *Sindbad Voyage*, from Oman to China. Sampling localities include the little known Lakshadweep (Laccadive), Islands and Pula Wé (Sumatra) from which 71% of the species were recorded. Following the zoogeographic subdivisions of Clark & Rowe (1971), range extensions are recorded for ten of the ophiuroids: W. India (*Amphioplus* (*Lymanella*) sp.); Sri Lanka (*Ophiactis modesta*, *Ophiarachna robillardi*, *Ophiodyscrita instratus*); Maldives area (*Cryptopelta granulifera*, *Ophiochaeta hirsuta*); and Indonesia / East Indies (*Amphiura* (*Amphiura*) *dejectoides*, *Amphiura* (*Amphiura*) *micra*, *Amphioplus* (*Amphioplus*) *stenaspis*, *Ophiogymna pellicula*). In addition to the taxonomic treatment, ecological information for each echinoderm species (habitat types, depth range) is provided and broadly analysed.

## INTRODUCTION

The systematics and distribution of Indian Ocean ophiuroids and echinoids are treated in detail by Clark & Rowe (1971). Regions for which limited information is available include the Lakshadweep (Laccadive) Islands, Sumatra and other parts of SE Asia. Recent studies including the systematics and zoogeography of ophiuroids and echinoids for the Lakshadweeps include Nagabhusanam & Rao (1972) and James (1989), the latter yielding many new species records for both echinoderm classes, which fill in gaps in the distribution records of Clark & Rowe (1971). Recent work has also been undertaken in SE Asia, including Indonesia (Aziz, 1981) and the west coast of Thailand (Bussarawit & Rowe, 1985; Bussarawit, in prep.).

This paper reports on collections of ophiuroids and echinoids from these areas and other localities during an international, transdisciplinary voyage across the Indian Ocean from Oman to China. The expedition, *Sindbad Voyage*, was undertaken in 1980–81 aboard a replica of an ancient Arab sailing vessel, 'Sohar'. In addition to a systematic account, the zoogeographic significance of the results and the ecology of each species are broadly assessed. Details of the holothurian collections (Price & Reid, 1985) and asteroid collections (Marsh & Price, 1991) resulting from the expedition have already been published. Details of the crinoids collected are also being prepared (Marshall & Price, in prep.), and a detailed analysis of the ecology and biogeography of all five echinoderm classes will follow.

## MATERIALS AND METHODS

Specimens were collected by one of us (A.R.G.P.) and other expedition members from localities at Muscat, Oman; Chetlat, Lakshadweeps (Laccadives); SW India; SW Sri Lanka; and Pula Wé, Sumatra. Details of the sampling localities are shown in Figure 1. Sampling was undertaken principally on coral reefs using scuba. At each locality details of habitat type and depth range were recorded, along with the number of individuals of each species. The number of specimens collected is placed in parenthesis after each station number in the Material lists for each species.

Material was fixed and preserved using standard methods (Lincoln & Shields, 1979). Although several specimens (inadvertently included with the asteroid collections) had been identified earlier by L.M. Marsh and a few preliminary identifications were made in the field by A.R.G.P., specimens were mostly identified by and all species confirmed by F.W.E.R. who is also responsible for taxonomic comments. The ophiuroid and echinoid collections are deposited at the Natural History Museum, London, where the holothurian collection (Price & Reid, 1985) and a representative collection of the asteroids (Marsh & Price, 1991) have also been lodged.

Following recent practice (e.g. Sloan, Clark & Taylor, 1979), systematic references are kept to a minimum by citing major works when possible (e.g. Clark & Rowe, 1971; Clark & Courtman Stock, 1976; Cherbonnier & Guille, 1978; Rowe & Gates, 1995) from which the original species descriptions, recent authoritative diagnoses and taxonomic decisions can be traced. In some instances, further references are given (e.g. Sloan *et al.*, 1979) to provide additional systematic or biological information.

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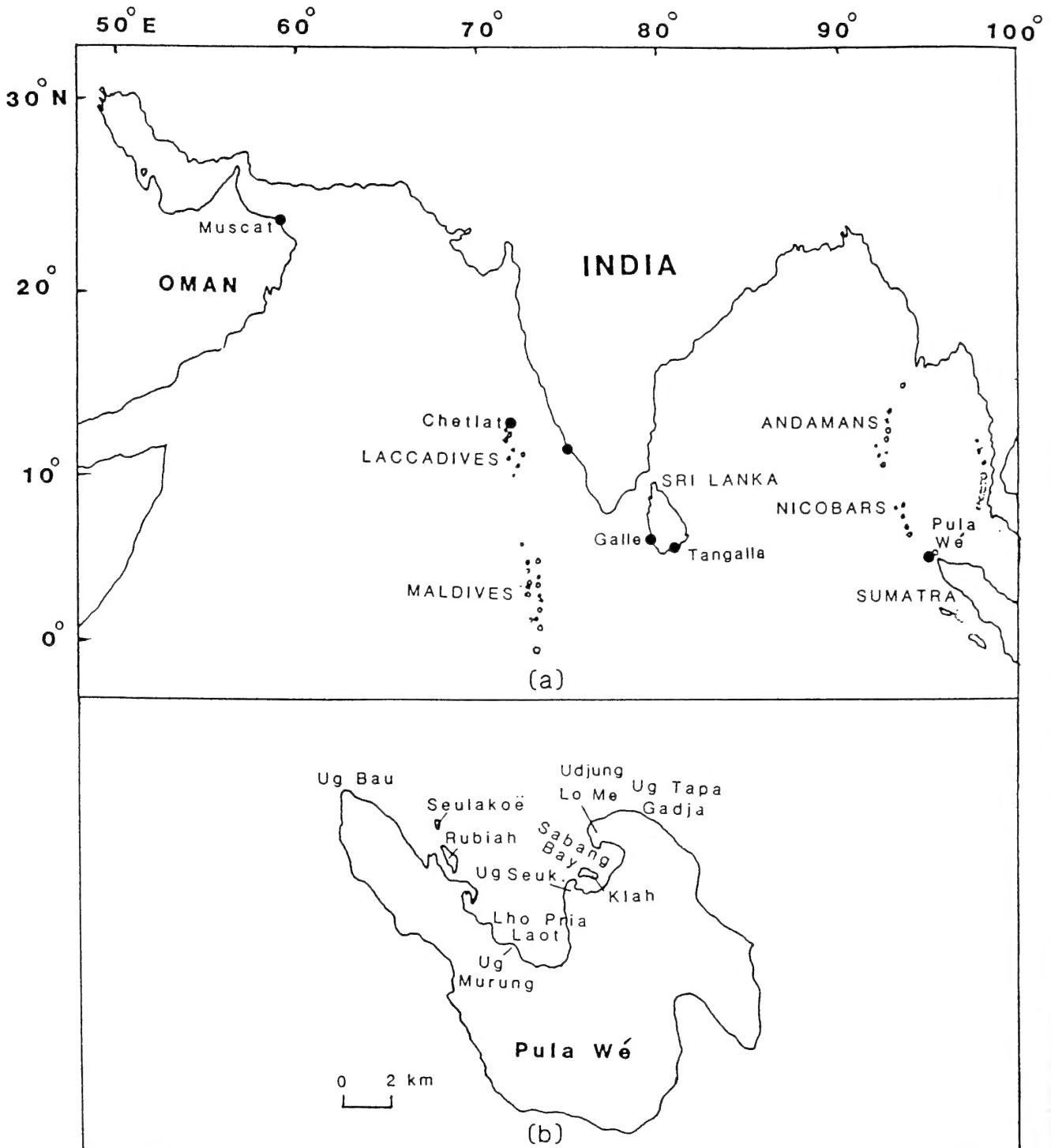


Fig. 1. (a) Map of northern Indian Ocean showing sampling areas (●) during Sindbad Voyage, with insert (b) for Pula Wé Sumatra.

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**RESULTS**


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**Class Ophiuroidea**
**Family GORGONOCEPHALIDAE**
**1. *Astroboa nuda* (Lyman, 1874)**

SEE. Clark & Rowe, 1971: 78; 92; Clark & Courtman Stock, 1976: 108; 130; Baker, 1980: 60; Rowe & Gates, 1995: 364.

MATERIAL. 810504D/3 (1), 810504D/4 (1).

COLLECTION SITES. NW Klah, Pula Wé, Sumatra.

HABITAT AND DEPTH. Subtidal rock/coral; 13 m.

**Family AMPHIURIDAE**
**2. *Amphiura (Amphiura) dejectoides* H.L. Clark, 1939**

SEE. Clark & Rowe, 1971: 80; 97; Cherbonnier & Guille, 1978: 33.

MATERIAL. 810501E/8 (4); ?810428D/5 (2).

COLLECTION SITES. Ug Tapa Gadjá & Ug Seukundo, Pula Wé, Sumatra.

HABITAT AND DEPTH. Subtidal rock / coral, coral reef; 10 & 15 m.

REMARKS. The specimens from stn 810501E/8 appear to accord well with the original description (Clark, 1939) of the species and descriptions in Clark & Rowe (1971) and Cherbonnier & Guille (1978). The 2 specimens from stn 810428D/5 differ in their firmer disc, with coarser ventral scaling and in having 6 arm spines proximally instead of fine ventral scaling and 5 proximal arm spines. They are identified as *A. dejectoides* with reservation. Clark (in Clark & Rowe, 1971) suspected *Amphiura inhaensis* Balinsky may be conspecific with *A. dejectoides* H.L. Clark, a conclusion confirmed by Cherbonnier & Guille (1978). The records included herein extend the distribution of this species eastwards across the Indian Ocean from the Red Sea, East Africa and Madagascar to the Indo-Malayan region. In the latter region it may prove to be widespread.

**3. *Amphiura (Amphiura) micra* H.L. Clark, 1938**

SEE. Clark & Rowe, 1971: 80; 97; Cherbonnier & Guille, 1978: 46; Rowe & Gates, 1995: 350.

MATERIAL. 810423B/2 (1).

COLLECTION SITES. Ug Bau, Pula Wé, Sumatra.

HABITAT AND DEPTH. Coral, coral reef; 10–30 m.

REMARKS. This species is recorded across the tropical coast of Australia and from Madagascar. Its discovery at Pula Wé suggests a wider distribution in the Indo-Malayan region for this very small species.

**4. *Amphiura (Amphichilus) ochroleuca* (Brock, 1888)**

SEE. Clark & Rowe, 1971: 78; 100; Rowe & Gates, 1995: 344.

MATERIAL. 810502C/1 (1), 810427B/3 (1).

COLLECTION SITES. SE Klah, SE Lho Pria Laot, Pula Wé, Sumatra.

HABITAT AND DEPTH. In sponge and on subtidal rock; 0–10 m.

REMARKS. This species is known from the Indo-Malayan region and more or less circumscribes the Australian continental coastline (Rowe & Gates, 1995). Pula Wé appears to be the most westerly point of its distribution known to date.

**5. *Amphioplus (Amphioplus) stenapsis* H.L. Clark, 1938**

SEE. Clark & Rowe, 1971: 78; 101; Rowe & Gates, 1995: 344.

MATERIAL. 810422B/4 (1).

COLLECTION SITES. Nr. Klah / Seukundo, Pula Wé, Sumatra (disc only).

HABITAT AND DEPTH. Coral reef, 2–8 m.

REMARKS. Although the single specimen comprises only a complete disc with the bases of 2 arms (6 & 9 segments respectively), there is little doubt of its identity. This record extends the range of the species to Pula Wé from its type locality, Darwin, N Australia. The record of *A. stenapsis* from Madagascar by Cherbonnier & Guille (1978) is almost certainly based on a misidentification, judging by the very small size of the radial shields of their specimens. The confirmation of this species in the western Indian Ocean therefore requires confirmation.

**6. *Amphioplus (Lymanella)* sp.**

MATERIAL. 810109A/1b (2).

COLLECTION SITES. Bepore, India (west coast).

HABITAT AND DEPTH. Subtidal mud, 9 m.

REMARKS. Only the mouthparts and bases of the arms are available to identify this taxon, which clearly represents a species of *Amphioplus (Lymanella)*. The dorsal arm plates are trilobed, suggesting either species *A. (L.) andreae* (Lütken, 1872) or *A. (L.) laevis* (Lyman, 1874) in the key provided by Clark & Rowe (1971: 102). Cherbonnier & Guille (1978) indicate that *laevis* has a wide range in the Indian Ocean and Indo-Malay region, whereas Clark & Rowe (1971) record *andreae* only from the Malay region. Without complete specimens it is not possible to determine the species nor indeed whether *andreae* and *laevis* are taxonomically separable.

**Family OPHIACTIDAE**
**7. *Ophiactis modesta* Brock, 1888**

SEE. Clark & Rowe, 1971: 105; Rowe & Gates, 1995: 379.

MATERIAL. 810206A/8 (3).

COLLECTION SITES. Negombo, Sri Lanka.

HABITAT AND DEPTH. Coral/rock; 5 m.

REMARKS. The 3 specimens were collected in a batch of 6 from stn 810206A/8 with 3 specimens of *O. savignyi*. They run unequivocally to the species *O. modesta* in the key provided by Clark & Rowe (1971: 105) where the relationship with other congeners is discussed by A.M. Clark (notes 25–27, pp. 103–105). Rowe (in Rowe & Gates, 1995) preferred to follow the views of Clark & Rowe (1971) regarding recognising the validity of the species *O. modesta*, a course also followed herein, rather than accept the sweeping synonymy of *O. savignyi*, which includes *O. modesta*, given by Cherbonnier & Guille (1978).

**8. *Ophiactis savignyi* (Müller & Troschel, 1842)**

SEE. Clark & Rowe, 1971: 82; 103; Clark & Courtman Stock, 1976: 164; Cherbonnier & Guille, 1978: 125; Sloan *et al.*, 1979: 102; Price, 1983: 61; Rowe & Gates, 1995: 380.

MATERIAL. 810206A/3 (5), 810206A/8 (3), 810426A/2 (1), 810501E/1 (1), 810501G/3 (1), 810501I/2 (1), 810502F/4 (2).

COLLECTION SITES. Negombo, Sri Lanka; W Rubiah, Ug Seukundo, E Klah, Pula Wé, Sumatra.

HABITAT AND DEPTH. Sponge, coral/rock, subtidal rock, coral reef; 5–14 m.

REMARKS. The specimens identified here are fissiparous and 6-armed with trilobed dorsal arm plates and proximally 6 arm spines, according well with the key characters given in Clark & Rowe (1971).

Family **OPHIOTRICHIDAE**

**9. *Gymnolophus obscura* (Ljungman, 1867)**

SEE. Clark & Rowe, 1971: 82; 117; Rowe & Gates, 1995: 411.

MATERIAL. 810124A/11 (1), 810125A/2 (1), 810430A/20d (2).

COLLECTION SITES. Ala Gala & Deumba Gala, Galle, Sri Lanka; Ug Seukundo, Pula Wé, Sumatra.

HABITAT AND DEPTH. Coral reef and epizoic on crinoids on subtidal rock; 8–15 m.

REMARKS. This species is commonly epizoic on comasterid crinoids, the host species for which have not been identified for the specimens recorded herein.

**10. *Ophiothela danae* Verrill, 1869**

SEE. Clark & Rowe, 1971: 84; 116; Clark & Courtman Stock, 1976: 141; Price, 1983: 63; Rowe & Gates, 1995: 419.

MATERIAL. 810125B/1 (3), 810206A/3/78 (4), 810425F/8 (10+), 810428A/2 (7), 810428A/14 (3), 810501A/3 (30+), 810501A/4 (1), 810428B/1 (2), 810428D/3c (3), 810501E/9 (15+).

COLLECTION SITES. Ala Galla, Galle & Negombo, Sri Lanka; N. Udjung Lo Me (NE Sabang Bay), Ug Bau, Ug Seukundo, Ug Tapa Gadja, Pula Wé, Sumatra

HABITAT AND DEPTH. Epizoic on macroalgae, gorgonian, fire coral (*Millepora* sp.) sponge and on holothurians (*Thelenota ananas*), all on rock /coral; 2–30 m.

**11. *Ophiothrix exigua* Lyman, 1874**

SEE. Clark & Rowe, 1971: 84; 110; Cherbonnier & Guille, 1978: 140; Rowe & Gates, 1995: 422.

MATERIAL. 810206A/8 (5), 810502C/1 (2), 810502E/2 (3).

COLLECTION SITES. Negombo, Sri Lanka; E Klah, Pula Wé, Sumatra.

HABITAT AND DEPTH. Coral/rock, sponge on coral reef and subtidal rock; 5–10 m.

REMARKS. One specimen from stn 810206A/8, both from 810502C/1 and two from 810502E/3 are very juvenile specimens.

**12. *Ophiothrix savignyi* (Müller & Troschel, 1842)**

SEE. Clark & Rowe, 1971: 84; 109; Cherbonnier & Guille, 1978: 142; Price, 1983: 65.

MATERIAL. 801114B/1 (1).

COLLECTION SITES. Muscat, Oman (1 specimen).

HABITAT AND DEPTH. Coral reef; 2 m.

**13. *Ophiothrix trilineata* Lütken, 1869**

SEE. Clark & Rowe, 1971: 84; 111; Clark & Courtman Stock, 1976: 145; Sloan *et al.*, 1978: 103; Rowe & Gates, 1995: 423.

MATERIAL. 810420A/5 (2), 810422B/3 (3), 810426A/2 (1), 810428A/9 (2), 810428C/7 (1), 810428D/3a&b (6&2), 810430A/8 (1), 810430A/18 (1), 810430A/21c (1), 810501D/2 (1), 810501E/2 (3), 810501E/5 (1), 810501E/8 (1), 810501E/9 (2), 810501E/11 (1), 810502F/2 (1), 810501F/3 (1), 810501G/1 (2), 810501G/3 (1 & 1 juvenile), 810501G/5 (3), 810502C/2 (1 very juvenile), 810502D/4 (4), 810502E/3 (4 juvenile), 810502F/2 (2 juvenile), 810502G/3 (2 juvenile).

COLLECTION SITES. Klah / E Klah, Nr. Seukundo, Ug Seukundo, Pula Wé, Sumatra, W Rubiah, Ug Bau, Ug Tapa Gadja, Pula Wé, Sumatra.

HABITAT AND DEPTH. Coral reef, soft coral / gorgonian, subtidal rock, sponge/subtidal rock, sponge; 2–30 m.

REMARKS. The majority of specimens exhibit the characteristic arm colour pattern of 5 longitudinal lines alternating white and dark blue. However, a few specimens bear a wide median pale longitudinal line along the arms, and at least one specimen (810501E/2) is distinctively patterned with cream blotches, the linear pattern being discernable only near the ends of the arms (see Clark & Rowe, 1971: 111). Next to *O. (A.) purpurea*, this is the most common ophiuroid species collected.

**14. *Ophiothrix (Acanthophiothrix) armata* Koehler, 1905**

SEE. Clark & Rowe, 1971: 84; 111; Rowe & Gates, 1995: 423.

MATERIAL. 810422B/4 (15+), 810502F/6 (2).

COLLECTION SITES. Klah / Nr. Seukundo, Pula Wé, Sumatra.

HABITAT AND DEPTH. Coral reef, 2–8 m.

REMARKS. This species is recorded from the Indo-Malay region, tropical Australian coasts and the South Pacific (Clark & Rowe,

1971). The present record is the most westerly so far known for the species.

**15. *Ophiothrix (Acanthophiothrix) purpurea* von Martens, 1867**

SEE. Clark & Rowe, 1971: 86; 112; Cherbonnier & Guille, 1978: 148; Sloan *et al.*, 1978: 103; Rowe & Gates, 1995: 423.

MATERIAL. 810204A/4 (3), 810421A/2 (1), 810421A/9 (1 juvenile), 810421B/1 (1), 810422D/3 (2), 810423A/4 (1), 810423B/2 (1), 810425D/2 (1), 80425D/4b (1), 810425F/7 (7), 810425F/8 (1), 810426A/2 (3), 810427A/2 (2), 810428A/2 (2), 810428A/7 (3), 810428A/8 (4), 810428A/9 (2), 810428D/3f (23), 810430A/3 (1), 810430A/21c (6), 810430A/22b (1), 810430A/26 (5), 810501A/1 (1), 810501A/2 (1 juvenile), 810501A/4 (1 juvenile), 810501A/6 (9), 810501E/7 (2), 810501E/9 (1), 810501E/14 (1), 810501E/15 (1), 810504B/2 (1).

COLLECTION SITES. SW Kalpitiya, Sri Lanka; Ug Bau, Rubiah, Ug Seukundo, Ug Tapa Gadja, N Klah, ?N. Udjung Lo Me, NE Sabang Bay, Pula Wé, Sumatra.

HABITAT AND DEPTH. Coral reef, soft coral, fire coral (*Millepora* sp.), subtidal rock/coral (epizoic on soft coral / gorgonian & crinoid, sponge); 2–30 m.

REMARKS. The most common ophiuroid species collected.

**16. *Ophiothrix (Acanthophiothrix) spinosissima* Koehler, 1905**

SEE. Clark & Rowe, 1971: 86; 112.

MATERIAL. 810422B/4 (4), 810501G/5 (2), 810502F/6 (1).

COLLECTION SITES. Ug Seukundo, Klah / Ug Seukundo, Pula Wé, Sumatra.

HABITAT AND DEPTH. Coral reef, 2–8 m.

REMARKS. The specimens run down well to *O. (A.) spinosissima* in the key provided by Clark & Rowe (1971). However, 2 specimens from stn 810422B/4 have a single dark line running the length of the dorsal side of the arms, rather than a series of dark spots.

**17. *Macrophiothrix aspidota* (Müller & Troschel, 1842)**

SEE. Clark, 1968: 285; Clark & Rowe, 1971: 114; Clark & Courtman Stock, 1976: 137; Hoggett, 1992: 91.

MATERIAL. 810123B/6 (1), 810124A/8 (1), 810206A/6 (1), 810206A/7 (1).

COLLECTION SITES. Negombo, Closenburg Point, Galle, Ala Gala, Galle, Sri Lanka.

HABITAT AND DEPTH. Subtidal rock, coral / rock; 5–15 m.

**18. *Macrophiothrix demessa* (Lyman, 1861)**

SEE. Clark, 1968: 289; Clark & Rowe, 1971: 82; 114; Hoggett, 1991: 1089; 1992: 117; Rowe & Gates, 1995: 412.

MATERIAL. 801212B/1 (1), 810424B/4 (2).

COLLECTION SITES. Chetlat, Lakshadweep (Laccadive) Islands; Seulakoe, Pula Wé, Sumatra.

HABITAT AND DEPTH. Coral rubble, coral reef; 8 m & 20–30 m.

REMARKS. A.M. Clark (1968) transferred this species to the genus *Macrophiothrix* referring *Ophiothrix (Amphiothrix)* H.L. Clark, 1946, of which *demessa* is type (and only) species to the synonymy of *Macrophiothrix* H.L. Clark, 1938.

**19. *Macrophiothrix elongata* (H.L. Clark, 1938)**

SEE. Clark, 1968: 291; Clark & Rowe, 1971: 82; 114; Price, 1983: 61; Hoggett, 1992: 125.

MATERIAL. 801111A/5 (1), 801114A/5 (2), 801114B/2 (1).

COLLECTION SITES. Muscat harbour, Oman.

HABITAT AND DEPTH. Coral reef, subtidal rock/coral/sand; 0.5–3 m.

**20. *Macrophiothrix longipeda* (Lamarck, 1816)**

SEE. Clark, 1968: 300; Clark & Rowe, 1971: 82; 114; Clark & Courtman Stock, 1976: 139; Hoggett, 1991: 1103; Hoggett, 1992: 151; Rowe & Gates, 1995: 413.

MATERIAL. 801212B/3 (1), 810206A/5 (1), 810430A/24a (1).

COLLECTION SITES. Chetlat, Lakshadweep (Laccadive) Islands; Negombo, Sri Lanka; Ug Seukundo, Pula Wé, Sumatra.

HABITAT AND DEPTH. Coral/rock, coral rubble; 5–10 m.

**21. *Macrophiothrix lorioli* A.M. Clark, 1968**

SEE. Clark, 1968: 302; Clark & Rowe, 1971: 82; 115; Hoggett, 1991: 1108; Hoggett, 1992: 161; Rowe & Gates, 1995: 414.

MATERIAL. 810502D/2 (1), 810502E/1 (1).

COLLECTION SITES. E Klah, Pula Wé, Sumatra.

HABITAT AND DEPTH. Coral reef, coral reef/subtidal rock; 5 & 10 m.

**22. *Macrophiothrix nereidina* (Lamarck, 1816)**

SEE. Clark & Rowe, 1971: 86; 107 (as *Ophiothrix (Keystonea) nereidina*); Hoggett, 1992: 228 (as *Macrophiothrix*); Rowe & Gates, 1995: 426 (as *O. (Keystonea) nereidina*).

MATERIAL. 810421A/4 (1), 810422B/3 (2), 810430A/20d (1), 810430A/21c (2), 810430A/22b (1), 810501E/6 (1).

COLLECTION SITES. Ug Seukundo, Pula Wé, Sumatra.

HABITAT AND DEPTH. Coral reef, coral / rock; 2–10 m.

REMARKS. The species *nereidina* (which was placed in the subgenus *Ophiothrix (Keystonea)* by A.M. Clark, 1967) is included herein in the genus *Macrophiothrix*. Hoggett (1991) stated that 'it is particularly difficult to determine the respective boundaries between *Macrophiothrix* H.L. Clark, 1938 and two subgenera of *Ophiothrix*, *O. (Placophiothrix)* H.L. Clark, 1938 and *O. (Keystonea)* A.M. Clark, 1967.' The differences between these taxa have traditionally relied principally on arm length, shape of dorsal arm plates, relative spinular armament of the disc plates including cover of the radial plates (see A.M. Clark, 1967; Clark & Rowe, 1971). The difficulty in recognising the supraspecific limits of these taxa is made all the more obvious by the treatment of the species *Macrophiothrix*

*propinqua*, placed in the subgenus *Keystonea* by A.M. Clark (1967), whilst Devaney (1974) described *O. (Placophiothrix) westwardi* which has been considered conspecific with *propinqua* by Hoggett (1991). Later, in a far-reaching and critical treatment of *Macrophiothrix*, Hoggett (1992: PhD thesis) commits both the taxa *Placophiothrix* and *Keystonea* to the synonymy of *Macrophiothrix*, transferring the included species of the former two taxa to the latter taxon and to whom this move is herein credited.

### 23. *Macrophiothrix propinqua* (Lyman, 1861)

SEE. Clark & Rowe, 1971: 86; 107 (as *Ophiothrix (Keystonea) propinqua*); Clark, 1980: 537; Hoggett, 1991: 1130; Hoggett, 1992: 204; Rowe & Gates, 1995: 415.

MATERIAL. 810428D/5 (5), 810501C/3 (1 juvenile), 810501D/1 (1 juvenile), 810501E/13 (1), 810501DF/1 (1 juvenile), 810501G/1 (1), 810501G/5 (3), 810501I/2 (2), 810502F/3 (1), 810502G/3 (1).

COLLECTION SITES. Ug Tapa Gadjaja, Ug Seukundo, E Klah, Pula Wé, Sumatra.

HABITAT AND DEPTH. Subtidal rock/coral, coral rubble, coral reef, soft coral; 2–20 m.

REMARKS. This species was first transferred to the genus *Macrophiothrix* H.L. Clark, 1938 by A.M. Clark (1980) from *Ophiothrix (Keystonea)* A.M. Clark, 1967.

### 24. *Macrophiothrix variabilis* (Duncan, 1887)

SEE. Clark, 1968: 308; Clark & Rowe, 1971: 115; Hoggett, 1991: 1138; Hoggett, 1992: 218; Rowe & Gates, 1995: 416.

MATERIAL. 810206A/4 (1).

COLLECTION SITES. Negombo, Sri Lanka.

HABITAT AND DEPTH. Coral / rock, 5 m.

### 25. *Macrophiothrix virgata* (Lyman, 1861)

SEE. Clark & Rowe, 1971: 86; 113 (as *Ophiothrix (Placophiothrix) virgata*); Hoggett, 1992: 236.

MATERIAL. 810423D/2 (1).

COLLECTION SITES. Ug Bau, Pula Wé, Sumatra.

HABITAT AND DEPTH. Coral reef; 2–8 m.

REMARKS. See remarks under *Macrophiothrix nereidina*.

### 26. *Ophiogymna pellicula* (Duncan, 1876)

SEE. Clark & Rowe, 1971: 84; 117; Clark & Courtman Stock, 1976: 140 (as *O. fulgens*); Rowe & Gates, 1995: 417.

MATERIAL. 810504B/2 (1).

COLLECTION SITES. Rubiah, Pula Wé, Sumatra.

HABITAT AND DEPTH. Soft coral; 10 m.

REMARKS. The single specimen has a d.d. = 2.7 mm, a.l. = c. 20 mm. The disc is mottled cream and pink, and the arms are banded with wide pink and narrow cream bands. An irregular longitudinal line of cream spots is evident along the dorsal midline of the arms. The disc is covered (except for the radial shields) with minute,

pointed granules with larger, conical spines interradially towards the edge of the disc. Clark & Courtman Stock (1976) include *Placophiothrix phriza* H.L. Clark as a synonym of *O. fulgens* (Koehler) which in turn is included in the synonymy of *O. pellicula* by Rowe (in Rowe & Gates, 1995). The species therefore appears to be distributed from the Gulf of Aden to the Indo-Malay region and the NW coast of Australia in depths of 10–116 m.

### 27. *Ophiopteron elegans* Ludwig, 1888

SEE. Clark & Rowe, 1971: 84; 115; Rowe & Gates, 1995: 419.

MATERIAL. 810428D/3e (2), 810428D/5 (8).

COLLECTION SITES. Ug Tapa Gadjaja, Pula Wé, Sumatra.

HABITAT AND DEPTH. Coral reef, subtidal rock/coral; 15 m.

## Family OPHIOCOMIDAE

### 28. *Ophiarthrum pictum* Müller & Troschel, 1842

SEE. Clark & Rowe, 1971: 86; 121; Rowe & Gates, 1995: 385.

MATERIAL. 810502H/1 (1).

COLLECTION SITES. E Klah, Pula Wé, Sumatra.

HABITAT AND DEPTH. Subtidal rock, 2m.

### 29. *Ophiocomella sexradia* (Duncan, 1887)

SEE. Clark & Rowe, 1971: 86; 118; Devaney, 1974: 162; Cherbonnier & Guille, 1978: 179; Rowe & Gates, 1995: 389.

MATERIAL. 81042D8/3d (1).

COLLECTION SITES. Ug Tapa Gadjaja, Pula Wé, Sumatra.

HABITAT AND DEPTH. Coral reef; 15 m.

REMARKS. The single specimen from Pula Wé measures d.d. = 3 mm; a.l. = 10+ mm (broken near tip). With the exception that the uppermost of the 4 arm spines is distinctly longer than the lower ones, the character separating *Ophiomastix sexradiata* A.H. Clark 1952 (known only from its type locality: Bikini Atoll, Marshall Is, SW Pacific) from *Ophiocomella sexradia* (Duncan) (identified throughout the Indo-West Pacific region and possibly tropicopolitan) in Clark & Rowe's (1971) key, all other skeletal characters of the present specimen accord with those described as fitting *O. sexradia* (note 65, p. 118) by A.M. Clark. Cherbonnier & Guille (1978), following their study of Malagasy material, concur with the comments expressed by A.M. Clark (in Clark & Rowe, 1971) and tentatively consider *Ophiomastix sexradiata* to be conspecific with *Ophiocomella sexradia* simultaneously agreeing with A.M. Clark that the status of the genus *Ophiocomella* as distinct from *Ophiocoma* remains doubtful. Although the present specimen does nothing to clarify the generic status of *Ophiocomella*, it does support strongly the view that the two nominal species are conspecific and confirm the synonymy proposed by Cherbonnier & Guille (1978).

### 30. *Ophiocoma dentata* Müller & Troschel, 1842

SEE. Devaney, 1970: 13; Clark & Rowe, 1971: 86; 119; Rowe & Gates, 1995: 386.

MATERIAL. 810502H/2b (1).



COLLECTION SITES. E Klah, Pula Wé, Sumatra.

HABITAT AND DEPTH. Subtidal rock, 2m.

REMARKS. This single specimen (d.d. c. 11.3 mm, distorted) exhibits two of the described colour forms for the species. At the centre of the dorsal surface of the disc is a cream spot (c. 1.1 mm diameter). This is surrounded by an irregular ring (c. 2.2 mm wide) which is uniformly dusky/grey with darker spots. The remainder of the disc dorsally and ventrally is reticulated dusky/grey on a cream background.

### 31. *Ophiocoma erinaceus* Müller & Troschel, 1842

SEE. Clark & Rowe, 1971: 114;119; Clark & Courtman Stock, 1976: 173; Sloan *et al.*, 1979: 106; Bussarawit & Rowe, 1985: 1 (as *O. similanensis* n. sp.); Rowe & Gates, 1995: 387.

MATERIAL. 801212A/3 (1), 810422B/3 (10), 810427B/1 (1), 810427D/5 (1), 810428C/6 (2 juveniles), 810428E/5 (1 juvenile), 810430A/3 (2), 810501F/1 (1), 810501G/5 (3), 810501G/6 (2), 810501K/1 (2).

COLLECTION SITES. Chetlat, Lakshadweep (Laccadive) Islands; Ug Seukundo, Ug Bau, Lho Pria Laot, Ug Murung, Ug Tapa Gadj, Pula Wé, Sumatra.

HABITAT AND DEPTH. Coral reef, coral conglomerate, coral rubble, subtidal rock; 2–25 m.

REMARKS. This is the commonest species of *Ophiocoma* collected. The collection comprises some 26 specimens, ranging in size from d.d. = 3.6–22.2 mm, which show clearly both colour changes and development of disc granulation with growth. Juveniles up to d.d. = c. 5 mm bear no granules and are usually marked radially across each radial shield with a cream line, as described by Bussarawit & Rowe (1985) for their new species *O. similanensis*. By d.d. = 5.7 mm granules are developed at the centre of the disc and along 10 radiating lines to the edge of the disc where an irregular line of granules is developed around the periphery joining these radiating lines and thus leaving bare the dorsal interradial and radial portions of the disc. By d.d. = c. 11 mm granules are developed over the interradial but not radial regions of the disc, but granules are still not developed ventrally. By d.d. = 12.5 mm granules cover the whole surface of the disc except for the radial shields which remain bare, while granules begin to extend in a wedge shape, on the ventral side of the disc. This arrangement may remain in specimens up to d.d. = 14 mm but generally from about d.d. = > 13 mm the radial shields become covered by granules. In specimens up to d.d. = 12.5 mm some central and peripheral granules may be more prominent by their slightly more elongate shape, but in larger specimens granules are more evenly rounded and more or less evenly sized. The cream colour pattern disappears with increased d.d. and is absent in specimens with d.d. > 11 mm.

It is very clear that *Ophiocoma similanensis* Bussarawit & Rowe, 1985 is based on juvenile specimens of *O. erinaceus*, to the synonymy of which *O. similanensis* is herein committed.

### 32. *Ophiocoma pica* Müller & Troschel

SEE. Devaney, 1970: 25; Clark & Rowe, 1971: 86; 118; Clark & Courtman Stock, 1976: 173; Sloan *et al.*, 1979: 106; Rowe & Gates, 1995: 387.

MATERIAL. 801212A/3 (1).

COLLECTION SITES. Chetlat, Lakshadweep (Laccadive) islands.

HABITAT AND DEPTH. Coral reef; 20 m.

### 33. *Ophiocoma pusilla* (Brock, 1888)

SEE. Devaney, 1970: 25; Clark & Rowe, 1971: 86; 118; Clark & Courtman Stock, 1976: 174; Sloan *et al.*, 1979: 106; Rowe & Gates, 1995: 388.

MATERIAL. 810422E/4 (1), 810425C/2 (1), 810430A/21c (1), 810501E/3 (1), 810501K/4 (1).

COLLECTION SITES. N Klah island, Sabang Bay, Ug Seukundo, Pula Wé, Sumatra.

HABITAT AND DEPTH. Coral/sand, coral conglomerate, coral reef; 2–10 m.

REMARKS. The 5 specimens range in size from d.d. = 3–7.5 mm. The characteristic, enlarged, tissue-covered arm spines (see Clark & Rowe, 1971) appear on specimens from d.d. > 5 mm.

### 34. *Ophiomastix annulosa* (Lamarck, 1816)

SEE. Clark & Rowe, 1971: 86; Rowe & Gates, 1995: 390.

MATERIAL. 810123A/3 (1), 810212A/3 (2), 810213A/4 (1).

COLLECTION SITES. Kakoni rocks, Pigeon Island & Unawatuna, Galle, Tangalla, Sri Lanka.

HABITAT AND DEPTH. Coral reef, subtidal rock/coral, subtidal rock; 3–10 m.

### 35. *Ophiomastix caryophyllata* Lütken, 1869

SEE. Clark & Rowe, 1971: 86; 120; Cherbonnier & Guille, 1978: 185; Rowe & Gates, 1995: 390.

MATERIAL. 810425C/1 (2), 810428C/7 (1/2), 810430A/21b (2).

COLLECTION SITES. E Sabang Bay, Ug Bau, Ug Seukundo, Pula Wé, Sumatra

HABITAT AND DEPTH. Coral reef, coral/rock; 3–10 m.

## Family OPHIONEREIDAE

### 36. *Ophionereis dubia* (Müller & Troschel, 1842)

SEE. Clark & Rowe, 1971: 122; Clark & Courtman, 1976: 179; Price, 1983: 67; Rowe & Gates, 1995: 408.

MATERIAL. 810502D/5 (1).

COLLECTION SITES. E Klah, Pula Wé, Sumatra.

HABITAT AND DEPTH. Subtidal rock/sand; 10 m.

### 37. *Ophionereis fusca* Brock, 1888

SEE. Clark, A.M. 1953: 69; 78; Clark & Rowe, 1971: 88; 122; Rowe & Gates, 1995: 408.

MATERIAL. 810421B/2 (1).

COLLECTION SITES. Nr. Seukundo, Pula Wé, Sumatra.

HABITAT AND DEPTH. Subtidal sand; 10–20 m.

## Family OPHIODERMATIDAE

**38. *Cryptopelta granulifera* H.L. Clark, 1909**

SEE. Clark & Rowe, 1971: 88; 128; Rowe & Gates, 1995: 394.

MATERIAL. 801212B/1 (1).

COLLECTION SITES. Chetlat, Lakshadweep (Laccadive) Islands.

HABITAT AND DEPTH. Coral rubble; 8 m.

REMARKS. Originally described from Mauritius. Rowe & Gates (1995) describe the distribution as including tropical Australia, the Indo-Malayan region and Philippine Islands. The specimen is identified here from the Laccadive Islands for the first time.

**39. *Ophiarachna affinis* Lütken, 1869**

SEE. Clark & Rowe, 1971: 88; 123; Sloan *et al.*, 1979: 111; Rowe & Gates, 1995: 395.

MATERIAL. 810425C/1a,b (1).

COLLECTION SITES. E. Sabang Bay, Pula Wé, Sumatra.

HABITAT AND DEPTH. Coral conglomerate; 3–6 m.

REMARKS. A.M. Clark (in Clark & Rowe, 1971: Note 83, p. 123) describes in detail colour variation in *Ophiarachna affinis* and *O. mauritiensis* de Loriol, concluding that specific distinction between the two is difficult to make. In the present specimens with d.d. = 22 mm, the disc is uniformly brownish-grey and the dorsal side of the arms has a broad longitudinal central dusky band either side of which is a narrower pale band, the 3 bands being demarcated by 4 irregular, very narrow longitudinal dark bands. This corresponds to A.M. Clark's form C colour pattern. If *O. affinis* and *O. mauritiensis* are conspecific the species is clearly widely distributed in the Indo-West Pacific region.

**40. *Ophiarachna robillardi* de Loriol, 1893**

SEE. Clark & Rowe, 1971: 88; 123.

MATERIAL. 810126B/4 (1), 810213A/3 (2).

COLLECTION SITES. Galle, Tangalla, Sri Lanka.

HABITAT AND DEPTH. Coral reef, 3–5 m.

REMARKS. This is a significant extension of range for this species described from Mauritius. The species is recorded as having 5 arm spines (H.L. Clark, 1909; size not recorded) but the present 3 specimens have 7–9 arm spines at d.d. = 21.5 mm; 9–10 arm spines at d.d. 31.5 mm and 10–11 arm spines at d.d. = 36.5 mm.

**41. *Ophiochaeta hirsuta* Lütken, 1869**

SEE. Clark & Rowe, 1971: 88; 127; Sloan *et al.*, 1979: 115; Rowe & Gates, 1995: 398.

MATERIAL. 801212B/1 (1), 810425C/2 (1).

COLLECTION SITES. Chetlat, Lakshadweep (Laccadive) Islands; E. Sabang Bay, Pula Wé, Sumatra.

HABITAT AND DEPTH. Coral rubble, 3–8 m.

REMARKS. Sloan *et al.* (1979) concluded that variation in the

occurrence of spinelets on the discs of 4 specimens from Aldabra, western Indian Ocean and 2 specimens from Palau in the western Pacific Ocean which they examined suggested that *Ophiochaeta boschmai* A.H. Clark, 1964 is a synonym of *Ophiochaeta hirsuta* Lütken, 1869. They pointed out that Cherbonnier & Guille, 1978, had described a new species, *O. crinita*, based on a single specimen, from Madagascar, but did not comment further. In the present collection the specimen from the Lakshadweep Islands accords with the description of *boschmai* in that the disc is granule-covered dorsally but bears spinelets on its ventral surface. The specimen from Pula Wé, on the other hand, accords with the description of *crinita* in that the disc is covered dorsally and ventrally by elongate spines. Considering the comments by Sloan *et al.* (1979), the two specimens reported herein are referred to *O. hirsuta*, with the implication that *O. crinita* Cherbonnier & Guille should also be referred to the synonymy on the basis that it exhibits the extreme spiny form of *O. hirsuta*.

**42. *Ophiodyscrita instratus* (Murakami, 1944) n. comb.**

SEE. Murakami, 1944: 272 (as *Ophiostegastus instratus*); A.M. Clark, 1968: 320 (as *Ophiostegastus instratus*; discussion)

MATERIAL. 810124A/4 (1).

COLLECTION SITES. Ala Gala, Galle, Sri Lanka.

HABITAT AND DEPTH. Subtidal rock, 10–15 m.

REMARKS. This specimen, apart from its smaller size and fewer naked disc plates, accords so well with Murakami's (1944) description of *Ophiostegastus instratus* that its identity is in no doubt. The species is, however, transferred to the genus *Ophiodyscrita* H.L. Clark, 1938 (type-species *O. acosmeta* H.L. Clark), with *Ophiostegastus* Murakami, 1944 (of which *instratus* is the type-species) reduced to a junior synonym of *Ophiodyscrita*. The distinctness of two genera has been questioned by A.M. Clark (1968) and Guille & Vadon (1985) on the grounds of variation of granulation with increased specimen size. Tabulation of measurements (Table 1) taken from original species descriptions and similar details of the specimens from Sri Lanka, shows an interesting picture. From this table it becomes apparent that only two species can be recognised: a) *Ophiodyscrita instratus* (Murakami, 1943) (d.d. = 7 mm) of which the larger *Ophiostegastus novaecaledoniae* Guille & Vadon (d.d. = 9–11 mm) is a synonym, being an extreme form of *instratus* in which many disc plates, including the radial shields, have become prominent (convex) and naked of granules; this species possesses supplementary oral shields (granule covered in small specimens < 7 mm d.d.) and b) *O. acosmeta* H.L. Clark (d.d. = 5 mm) with which *O. pacifica* (Murakami, 1943) (d.d. 4 mm) and *Ophiostegastus compsus* A.M. Clark (1968) type locality Bahrain (d.d. = 8–10.5 mm) appear to be conspecific. *O. acosmeta* has an even covering of granules over the disc which are gradually lost only from the oral shields (as in the type of *acosmeta*; d.d. = 5 mm) and adoral plates (as in the type series of *compsus*; d.d. = 8–10.5 mm; though A.M. Clark (1968) does note that the largest paratype of *compsus* (d.d. = 10.5 mm) has a small bare patch dorsally at the base of 4 of the arms). Supplementary oral shields are absent from *compsus*, according to A.M. Clark, and are not recorded for either *acosmeta* or *pacifica*. In both *instratus* and *acosmeta*, as recognised herein, it is clear that arm spine number increases with size.

There is clearly insufficient justification for recognising *Ophiodyscrita* and *Ophiostegastus* as separate genera on the basis of the extent of disc granulation, for it is clear (see Clark & Rowe, 1971) that such differences occur between species included within

**Table 1.** Details of species *Ophiodyscrita acosmeta* H.L. Clark\* and *O. instratus* (Murakami)\*

Taxon	d.d. (mm)	a.sp.	l.a.sp.	s.o.s.	Granulation (disc)
<i>Ophiocryptus</i> * <i>pacificus</i> Murakami, 1943	4.00	6-5	1/3 seg.	- ?	Complete cover dorsal and ventral
<i>Ophiodyscrita</i> * <i>acosmeta</i> H.L. Clark, 1938	5.0	8(7)	1/2 seg.	- ?	Complete cover dorsal and ventral except 2 oral shields (Clark & Rowe (1971: 135)
Sindbad spec.*	5.3	6-5	<1/2 seg.	+ (granule covered)	Complete cover dorsal and ventral except radial plate at base of deach arm, each mid marginal plate and ventrally, each of the oral shields
<i>Ophiostegastus</i> * <i>instratus</i> Murakami, 1944	7.00	7	1/3 seg.	+ (naked)	Complete except 3 plates at base of each arm; each mid marginal; each oral shield and supplementary oral shield
<i>Ophiostegastus</i> * <i>compus</i> A.M. Clark, 1968	8-10.5	9	<1/2 seg.	-	Complete except each oral shield and adoral shield (A.M. Clark notes a small bare patch as base of 4 arms of one paratype d.d. = 10.5 mm)
<i>Ophiostegastus</i> * <i>novaecaledoniae</i> Guille & Vadon, 1985	9-11	9-10	<1/2 seg.	+ (naked)	Many dorsal plates, including radial shields, marginal plates and ventral plates including oral, supplementary oral and adoral shields bare of granules

d.d. = disc diameter; a.sp. = number of arm spines; l.a.sp. = length of arm spines; s.o.s. (+/-) = presence/absence of supplementary oral shields.

the recognised limits of other ophiidermatid genera (e.g. *Ophiopoeza* & *Ophiarachnella*). Similarly, the occurrence of supplementary oral shields is also a variable character. The recognition of *Ophiodyscrita* within the family appears, therefore, to rely, more or less solely, on the extension of granulation along the arms. The genus is closely related to *Ophiopoeza* with which it shares the possession of a triangle of 3 plates between the radial shields (see Vail & Rowe, 1989). The record of *O. instratus* from Sri Lanka greatly extends the known distribution of the species from Japan and New Caledonia (S.W. Pacific). The distribution of the genus, *Ophiodyscrita* (syn: *Ophiostegastus*) is clearly widespread in the Indo-West Pacific region.

#### 43. *Ophiopsammus yoldii* (Lütken, 1856)

SEE. A.M. Clark, 1968: 317; Clark & Rowe, 1971: 90; 127; Vail & Rowe, 1989: 277; Rowe & Gates, 1995: 402.

MATERIAL. 810430A/22b (1).

COLLECTION SITES. Ug Seukundo, Pula Wé, Sumatra.

HABITAT AND DEPTH. Coral rubble; 9 m.

REMARKS. Nearly half of the disc of this small specimen has been lost leaving the remaining disc (d.d. = c. 6 mm) and three of the original arms, which are also damaged (a.l. = c. 17 mm; d.d./a.l. = c. 3+ : 1). Arising from the damaged edge of the disc are three new, minute arms at slightly different stages of growth judging by the relative development of the ventral arm plates on each arm. These arms are also damaged but the longest is judged to have been not more than c. 3-4 mm in length. Following Vail & Rowe's (1989) revision of the genus *Ophiopsammus*, there is no reason for not identifying the specimen from Pula Wé as *O. yoldii*, for it appears to match their criteria for the species even though it is of small size. The species is not known to be fissiparous. Although this may be the first observation of fissiparity in *O. yoldii*, the development of six arms, in this case, may be an unusual response to severe damage, rather than being related to an asexual reproductive strategy. This matter requires further investigation.

#### Family OPHIURIDAE

#### 44. *Ophiolepis cincta* Müller & Troschel, 1842

SEE. Clark & Rowe, 1971: 90; 129; Clark & Courtman Stock,

1976: 189; Sloan *et al.*, 1979: 115-117; Rowe & Gates, 1995: 399.

MATERIAL. 801212B/1 (1).

COLLECTION SITES. Chetlat, Lakshadweep (Laccadive) Islands.

HABITAT AND DEPTH. Coral rubble; 8 m.

### Class Echinoidea

#### Family CIDARIDAE

#### 1. *Eucidaris metularia* (Lamarck, 1816)

SEE. Clark & Rowe, 1971: 140; 150; Clark & Courtman Stock, 1976: 215; Sloan *et al.*, 1979: 117; Rowe & Gates, 1995: 195.

MATERIAL. 810426A/4 (1), 810501E/12 (1).

COLLECTION SITES. Rubiah, Ug Seukundo, Pula Wé, Sumatra.

HABITAT AND DEPTH. Coral reef, coral aggregate; 10 & 14 m.

#### Family DIADEMATIDAE

#### 2. *Diadema setosum* (Leske, 1778)

SEE. Clark & Rowe, 1971: 140; 153; Clark & Courtman Stock, 1976: 226; Sloan *et al.*, 1979: 118; Price, 1983: 73; Rowe & Gates, 1995: 207.

MATERIAL. 801027A/1 (1), 801030A/1a,b (4), 810212A/1 (1), 810420A/1 (1), 810420A/4 (1), 810426A/3 (1), 810426B/5 (2), 810426B/12 (1), 810427D/5 (1 juvenile), 810428D/4 (2), 810501G/4 (1 juvenile), 810501I/1 (1), 810502D/1 (1, broken).

COLLECTION SITES. Muscat, Oman; Unawatuna, nr Galle, Sri Lanka; Rubiah, Klah, Ug Murung, Ug Tapa Gadjaja, Ug Seukundo, Pula Wé, Sumatra.

HABITAT AND DEPTH. Subtidal rock, subtidal rock / sand, subtidal rock / coral, coral reef; 0-20 m.

REMARKS. A number of these specimens are juveniles, as small as 10 mm h.d., and with banded spines. The characteristic elongate, tridentate pedicellariae of *D. setosum* (see Clark & Rowe, 1971) are

absent from all specimens examined. The remaining character of the red (or in some cases a faded, cream) ring on the anus is the only means of distinguishing this species from *D. savignyi*. In the field the two species are easily distinguished by the colour pattern, *D. savignyi* lacking the red ring and having characteristic, iridescent blue lines along the upper interamulacra.

### 3. *Echinothrix calamaris* (Pallas, 1774)

SEE. Clark & Rowe, 1971: 140; 153; Clark & Courtman Stock, 1976: 226; Rowe & Gates, 1995: 208.

MATERIAL. 801111A/2 (1), 810422B/1 (1), 810425C/4 (1), 810426B/4 (1), 810426B/13 (1), 810501K/2 (1), 810501K/3 (1).

COLLECTION SITES. Muscat, Oman; S. Sabang Bay (to S. of Klah), E. Sabang Bay, Rubiah, Ug Seukundo, Pula Wé, Sumatra.

HABITAT AND DEPTH. Subtidal rock, subtidal rock / coral / sand, coral reef; 0–20 m.

### Family STOMECHINIDAE

### 4. *Stomopneustes variolaris* (Lamarck, 1816)

SEE. Clark & Rowe, 1971: 140; 153; Clark & Courtman Stock, 1976: 228; Sloan *et al.*, 1976: 118; Rowe & Gates, 1995: 246.

MATERIAL. 810123B/4 (1).

COLLECTION SITES. Galle, Sri Lanka.

HABITAT AND DEPTH. Subtidal rock; 5 m.

### Family TEMNOPLEURIDAE

### 5. *Mespilia globulus* (Linnaeus, 1758)

SEE. Clark & Rowe, 1971: 140; 155; Rowe & Gates, 1995: 250.

MATERIAL. 810502F/5 (1).

COLLECTION SITES. Klah, Pula Wé, Sumatra.

HABITAT AND DEPTH. Subtidal rock; 5 m.

### 6. *Microcyphus ceylanicus* Mortensen, 1942

SEE. Clark & Rowe, 1971: 140; 156.

MATERIAL. 820204A/11 (1).

COLLECTION SITES. SW Kalpitiya, Sri Lanka.

HABITAT AND DEPTH. Coral reef; 3–5 m.

### 7. *Salamacis bicolor* L. Agassiz, 1846

SEE. Clark & Rowe, 1971: 140; 156; Clark & Courtman Stock, 1976: 232.

MATERIAL. 810126B/1 (1).

COLLECTION SITES. Galle, Sri Lanka.

HABITAT AND DEPTH. Coral reef; 4–5 m.

### Family TOXOPNEUSTIDAE

### 8. *Toxopneustes piloleus* (Lamarck, 1816)

SEE. Clark & Rowe, 1971: 142; 156; Clark & Courtman Stock, 1976: 234; Rowe & Gates, 1995: 258.

MATERIAL. 801114C/1 (1); 810424A/1 (1).

COLLECTION SITES. Muscat, Oman; W. Klah, Pula Wé, Sumatra.

HABITAT AND DEPTH. Coral reef; 10–12 m.

### Family PARASALENIIDAE

### 9. *Parasalenia gratiosa* A. Agassiz, 1863

SEE. Clark & Rowe, 1971: 142; 157; Rowe & Gates, 1995: 233.

MATERIAL. 810422B/5 (2).

COLLECTION SITES. S. Sabang Bay (to S. of Klah), Sumatra.

HABITAT AND DEPTH. Coral conglomerate; 2–8 m.

### Family ECHINOMETRIDAE

### 10. *Echinometra mathaei* (de Blainville, 1825)

SEE. Clark & Rowe, 1971: 142; 157; Clark & Courtman Stock, 1976: 239; Sloan *et al.*, 1979: 119; Price, 1983: 76; Rowe & Gates, 1995: 211.

MATERIAL. 801111A/1 (5), 810204A/10 (1).

COLLECTION SITES. Muscat, Oman; SW Kalpitiya, Sri Lanka.

HABITAT AND DEPTH. Coral reef, rock / coral / sand; 0–5 m.

REMARKS. Two forms of this species are represented. The specimen from Sri Lanka is relatively large, with h.d. = 48.7 mm. It has 4 pore-pairs per arc, the spines are uniformly pale blue/green and the test, when cleaned, is whitish in colour. The 5 specimens from Muscat, Oman are smaller, ranging in size from h.d. = 10–21 mm. They have 5 pore-pairs per arc, the spines are dark olive green tipped with lilac/brown and the test, when cleaned, is greenish in colour. *Echinometra mathaei* is clearly a complex species which is in need of critical investigation to determine whether, as it is currently identified, it comprises a single species or more than one closely related species (see Mortensen, 1943; Tsuchiya & Nishira, 1984).

### 11. *Echinostrephus molaris* (de Blainville, 1825)

SEE. Clark & Rowe, 1971: 142; 157; Clark & Courtman Stock, 1976: 239–240; Sloan *et al.*, 1979: 119; Rowe & Gates, 1995: 212.

MATERIAL. 801210B/6 (1), 810423A/5 (1), 810427B/3 (1), 810427D/5 (1), 810502G/4 (1)

COLLECTION SITES. Chetlat, LakshawEEP (Laccadive) Islands; Ug Bau, Lho Pria Laot, Ug Murung, Klah, Pula Wé, Sumatra.

HABITAT AND DEPTH. Coral reef, subtidal rock, subtidal rock / coral; 0–40 m.

## DISCUSSION

Echinoderm collections from the *Sindbad Voyage* have yielded at least 44 species of ophiuroids and 11 species of echinoids. Species totals for each area sampled, with the corresponding zoogeographic subdivision used by Clark & Rowe (1971), are given in Table 2. Despite the small size of Pula Wé (c. 20 km x 12 km), 32 ophiuroids and 7 echinoids (71% of all species recorded) were encountered at, although not necessarily restricted to, this island. This high species richness is partly a reflection of the sampling intensity in Pula Wé, but is equally or more an indication of high biodiversity known for coral reefs in the SE Asia region (Sheppard, 1987; Wells & Price, 1992).

Of the ophiuroids collected, *O. (A.) purpurea* and *O. trilineata* were the most common, occurring in more than 20% of the ophiuroid

**Table 2.** Ophiuroid and echinoid species numbers for each area of the Indian Ocean sampled during the *Sindbad Voyage*. (total ophiuroid species recorded for all regions = 44; total echinoid species recorded for all regions = 11)

Sampling area and equivalent zoogeographic subdivision	No. of species recorded	
	OPHIUROIDS	ECHINOIDS
Oman (SE Arabia)	2	4
S India (W. India & Pakistan)	1	0
Laccadive (Maldivian area)	7	1
Sri Lanka (Sri Lanka area)	7	1
Sumatra (Indonesia / East Indies)	31	7

Of the 44 ophiuroid species collected, ten result in new area records (Table 3), as follows: W India (*Amphioplus (Lymanella)* sp.); Sri Lanka (*Ophiactis modesta*, *Ophiarachna robillardi*, *Ophiodyscrita instratus*); Maldives area (*Cryptopelta granulifera*, *Ophiochaeta hirsuta*); and Indonesia / East Indies (*Amphiura (Amphiura) dejectoides*, *Amphiura (Amphiura) micra*, *Amphioplus (A.) stenaspis*, *Ophiogymna pellicula*).

**Table 3.** New area records and previously known distribution of ophiuroid species recorded in the Indian Ocean during the *Sindbad Voyage*

Species	New area record	Previously known distribution
<i>Amphiura (Amphiura) dejectoides</i>	Indonesia/East Indies	Red Sea; E. Africa (Madagascar)
<i>Amphiura (Amphiura) micra</i>	Indonesia/East Indies	N. Australia and possibly E. Africa/Madagascar
<i>Amphioplus (A.) stenaspis</i>	Indonesia/East Indies	N. Australia and possibly E. Africa/Madagascar
<i>Amphioplus (Lymanella)</i> sp.	West India (Beyport)	<i>A. (L.) andreae</i> from Indonesia/East Indies, and <i>A. (L.) laevis</i> from Indo-West Pacific
<i>Ophiactis modesta</i>	Sri Lanka	W. India & Pakistan, and eastwards from Bay of Bengal to Hawaiian Is. but not Philippines, and possibly also E. Africa/Madagascar Bay of Bengal and N. Australia
<i>Ophiogymna pellicula</i>	Indonesia/East Indies	
<i>Cryptopelta granulifera</i>	Maldives	Mascarene Is. (Mauritius, Réunion, Rodrigues group) and N. Australia
<i>Ophiarachna robillardi</i>	Sri Lanka	Mascarene Is.
<i>Ophiochaeta hirsuta</i>	Maldives	Is. of W. Indian Ocean, Indonesia/East Indies, N. Australia and S. Pacific Is.
<i>Ophiodyscrita instratus</i>	Sri Lanka	Japan and New Caledonia (SW Pacific)

samples. These species also occupied a wide range of substrata and depths (2–30m). Other species occurred in less than 10% of the ophiuroid samples, and generally occupied fewer habitats and a narrower depth range. The echinoid fauna was less diverse, although some species were very common, in particular *D. setosum*, *E. calamaris* and *E. molaris* (a coral rock borer), which occurred in 36%, 19% and 14% of the echinoid samples respectively. These echinoids were found in a wide range of habitats and depths, up to 40 m in the case of *E. molaris*. A more comprehensive ecological analysis of echinoderms of Pula Wé, Sumatra is to be undertaken following completion of the taxonomic appraisal of the crinoids.

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# Rare cyclopoid copepods (Crustacea) from Mediterranean littoral caves

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**SYNOPSIS.** Three cyclopoid copepods are reported from anchihaline cave habitats on Mallorca. Both sexes of *Cyclopina esilis* Brian are redescribed. Sexual dimorphism in the mandibular exopod, as discovered in *C. esilis*, has not previously been reported for any cyclopoid. The male of the primitive marine cyclopoid *Neocyclops (Protoneocyclops) mediterraneus* (Kiefer) is described in detail for the first time and the characters distinguishing this species are discussed. Both sexes of *Euryte longicauda* Philippi are redescribed. The genus *Euryte* Philippi is briefly reviewed and the characters used to differentiate species are critically reevaluated. It is concluded that all three genera, the cyclopoid genus *Cyclopina* Claus and the primitive cyclopoids *Neocyclops* Gurney and *Euryte*, are in urgent need of revision, but that this process will be hampered by the inadequacy of published descriptions and the lack of available type material.

## INTRODUCTION

Anchihaline cave habitats are rich sources of interesting and unusual crustaceans. Copepods of immense phylogenetic significance have been discovered in such caves in the last decade, including the platycopoid *Antriscopia* Fosshagen, the misophrioid *Speleophriopsis* Jaume & Boxshall, and the calanoid *Erebionectes* Fosshagen (Fosshagen & Iliffe, 1985; Jaume & Boxshall, 1996). There have been few reports of cyclopoid copepods in anchihaline caves. Recently, however, Rocha & Iliffe (1991, 1994) described a new family of cyclopoids, the Speleoithonidae, and the primitive cyclopoid *Troglocyclops*, from caves on the Bahamas Islands. During our studies of the copepod fauna of Mallorcan caves we discovered numerous cyclopoids, including the three rare species described in this account. All three species were originally described from the Mediterranean Basin. All have been the subject of considerable taxonomic confusion because their original descriptions were inadequate and we have attempted to clear up some of this confusion at the same time as presenting full redescrptions.

## THE CAVES

The copepods were collected from four caves located on the East coast of Mallorca, less than 20 m inland. These caves occur in two different types of substratum. Cova 'C' de Cala Varques and Es Secret des Moix are in Tortonian (10 Myr BP), coral reef-derived, porous calcarenites and mixing-zone corrosion processes seem to have played an important role in their development. Cova de na Barxa and Cova de na Mitjana are in Triassic, fissured limestones. All these caves have subaerial entrances; the difficulty of access can be deduced from their topographies, published elsewhere (see below). The water conditions varied from cave to cave: In Es Secret des Moix, the sampled lake (that located closest to the entrance) was completely marine (i.e., in water salinity, a detectable swell, the nature of the accompanying fauna), as was the lake in Cova de na Mitjana. Cova 'C' de Cala Varques and Cova de na Barxa are typical anchihaline caves (in the sense of Stock *et al.*, 1986), with a thin layer of fresh water on the top of the deeper saline waters of the lakes.

Sampling was undertaken using meat-baited traps placed at different depths in the cave lakes and left for several days, and by using a hand-held plankton net with an extensible handle. The terminology used in the descriptions follows Huys & Boxshall (1991).

## SYSTEMATICS

Order **CYCLOPOIDA** Burmeister, 1834  
 Family **CYCLOPINIDAE** Sars, 1913  
 Subfamily **CYCLOPININAE** Kiefer, 1927  
 Genus *Cyclopina* Claus, 1863

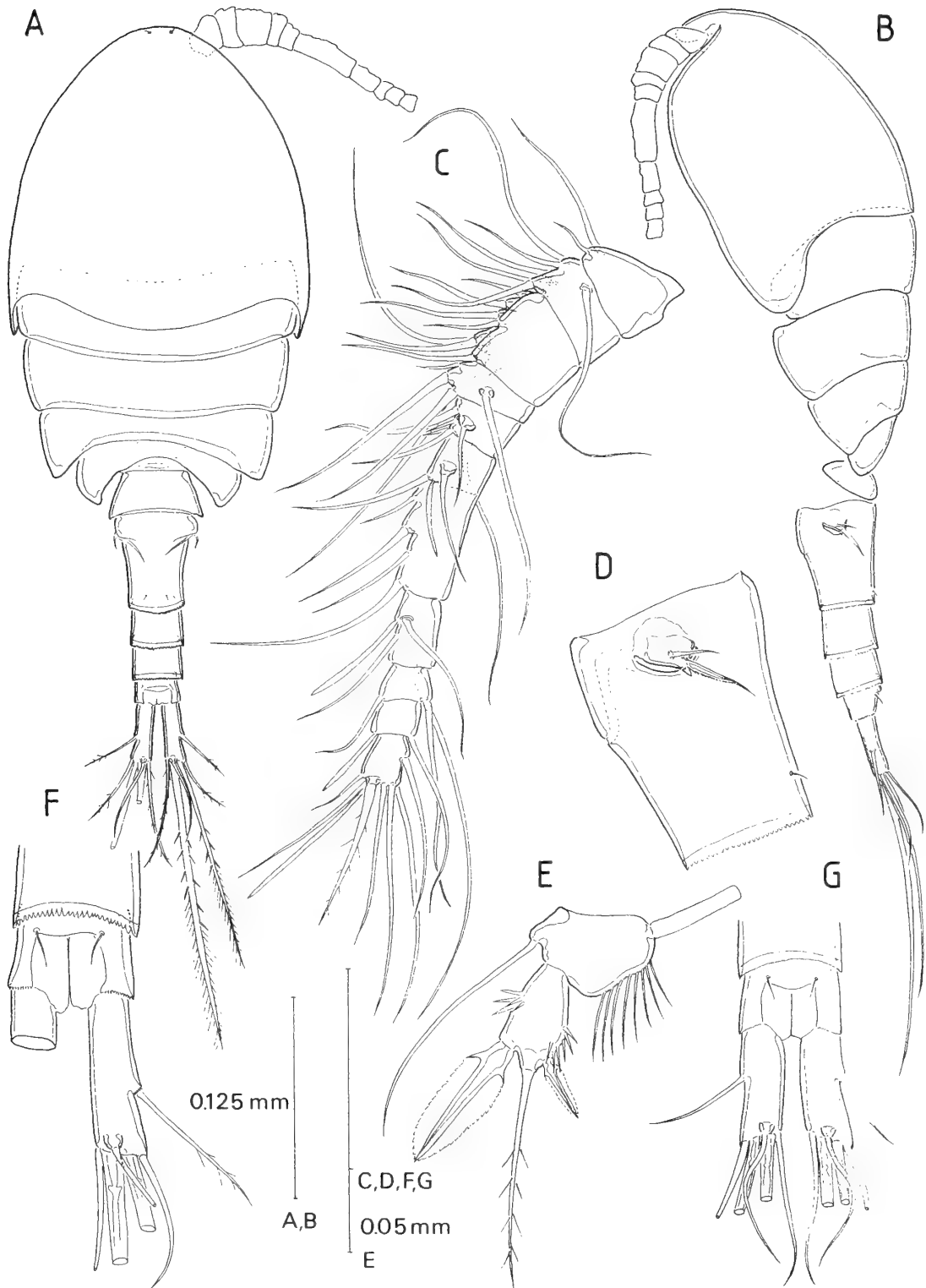
*Cyclopina esilis* Brian, 1938

(Figs 1–4)

*Cyclopina cf. kieferi*: Herbst (1953; 1962)

**MATERIAL EXAMINED.** Cova de na Mitjana (Capdepera). UTM coordinates: 539.10; 4390.95. Topography published by Ginés *et al.* (1975): 96 individuals, both sexes (BMNH 1995. 1331–1340). Collected by D. Jaume, 17 July 1994.

**ADULT FEMALE.** Body (Figs 1A, B) cyclopiform, up to 0.37 mm long. Prosome 5-segmented, about 1.7 times longer than urosome. Rostrum developed, oval. Posterolateral margins of cephalosome vaulted. First pedigerous somite free, partially concealed by dorsal and posterolateral extensions of cephalosome. Second to fourth pedigerous somites with evenly rounded posterolateral angles. Urosome 5-segmented, with genital and first abdominal somites completely fused to form genital double-somite. Serrate hyaline frill adorning posterodorsal margin of fifth pedigerous somite, posterior margins of genital double-somite and abdominal somites 2 and 3, and posterolateral margins of anal somite; degree of serration varying directly with body size. Genital double-somite (Fig. 1D) symmetrical, 1.6 times longer than wide, expanded anteriorly. Single copulatory pore opening mid-ventrally at about two-fifths of distance along double-somite. Paired gonopores located laterally, each covered by operculum armed with short spinous process, 1



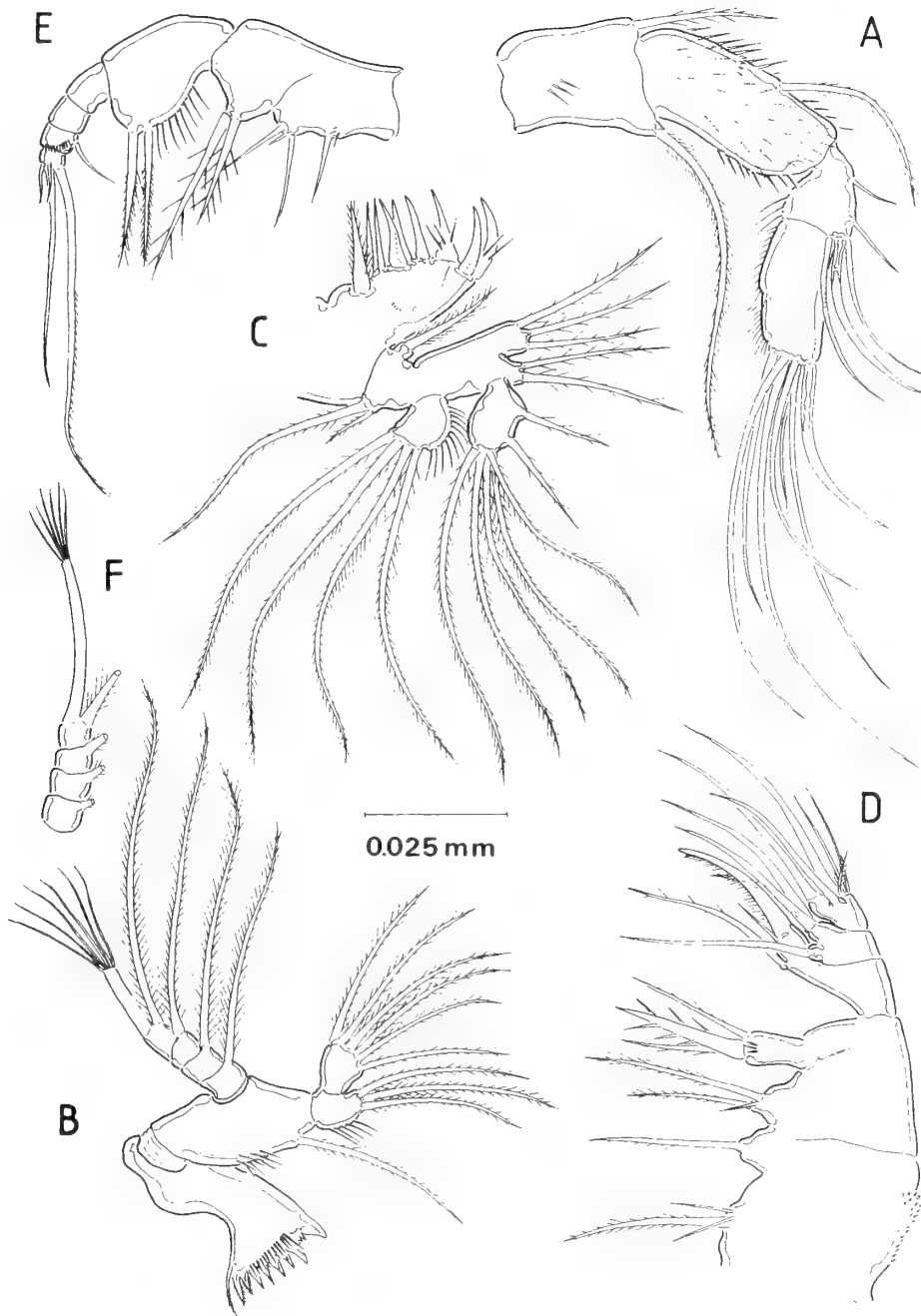
**Fig. 1.** *Cyclopina esilis* Brian, 1938, adult female. A, body, dorsal view; B, lateral; C, antennule; D, genital double-somite, lateral; E, fifth leg; F-G, dorsal view of anal somite and caudal rami, showing variation with body size in the position of the lateral seta of the caudal ramus.

seta, and 1 long flanged spine (Fig. 1D). Anal somite (Figs 1F, G) bearing smooth operculum. Caudal rami longer than anal somite, about 2.6 to 3.3 times longer than wide; proportional length related to body size. Armature consisting of 6 setae; position of seta II

variable, ranging from two-fifths (Fig. 1G) to three-fifths of distance along distal margin (Fig. 1F).

Antennules (Fig. 1C) symmetrical, 10-segmented, shorter than prosome (Fig. 1B). Segmental fusion pattern and armature as fol-





**Fig. 2.** *Cyclopina esilis* Brian, 1938, A and C–F, adult female, B, adult male. A, antenna; B, mandible; C, maxillule; D, maxilla; E, maxilliped; F, mandibular exopod.

lows: segment 1 (corresponding to fused ancestral segments I and II), 3 setae; segment 2 (corresponding to fused ancestral segments III to V), 5 setae; segment 3 (fused segments VI to IX), 8 setae; segment 4 (fused segments X and XI), 4 setae; segment 5 (fused segments XII to XIV), 6 setae; segment 6 (fused segments XV to XX, partial suture present between segments XVI and XVII), 6 setae + aesthetasc; segment 7 (fused segments XXI to XXIII), 3 + aesthetasc; segment 8 (XXIV), 2 setae; segment 9 (XXV), 1 + aesthetasc; segment 10 (fused segments XXVI to XXVIII), 7 + aesthetasc.

Antenna (Fig. 2A) 4-segmented. Fused coxa and basis short, about 1.5 times longer than wide, armed with 1 inner basal seta distally and 1 long seta (representing exopod) on outer margin.

Endopod 3-segmented. Proximal segment with 1 seta at about two-thirds of distance along inner margin; segment covered by long spinules on anterior surface. Second segment with 2 lateral and 3 distal setae (one of them claw-like) along inner margin. Distal segment with 7 distal setae, one of them claw-like. Secondary ornamentation on segments as figured.

Mandible (Figs 2B, F) with gnathobase armed with 10 unequal, sharp teeth plus 1 dorsal seta; row of 13 spinules located subdistally. Palp well developed; basis elongate, with patch of setules and 1 subdistal seta along inner margin. Exopod (Fig. 2F) inserted at about midway of distance along outer margin of basis; 4-segmented; setal formula 1,1,1,2; distal, brush-like seta somewhat shorter and thicker

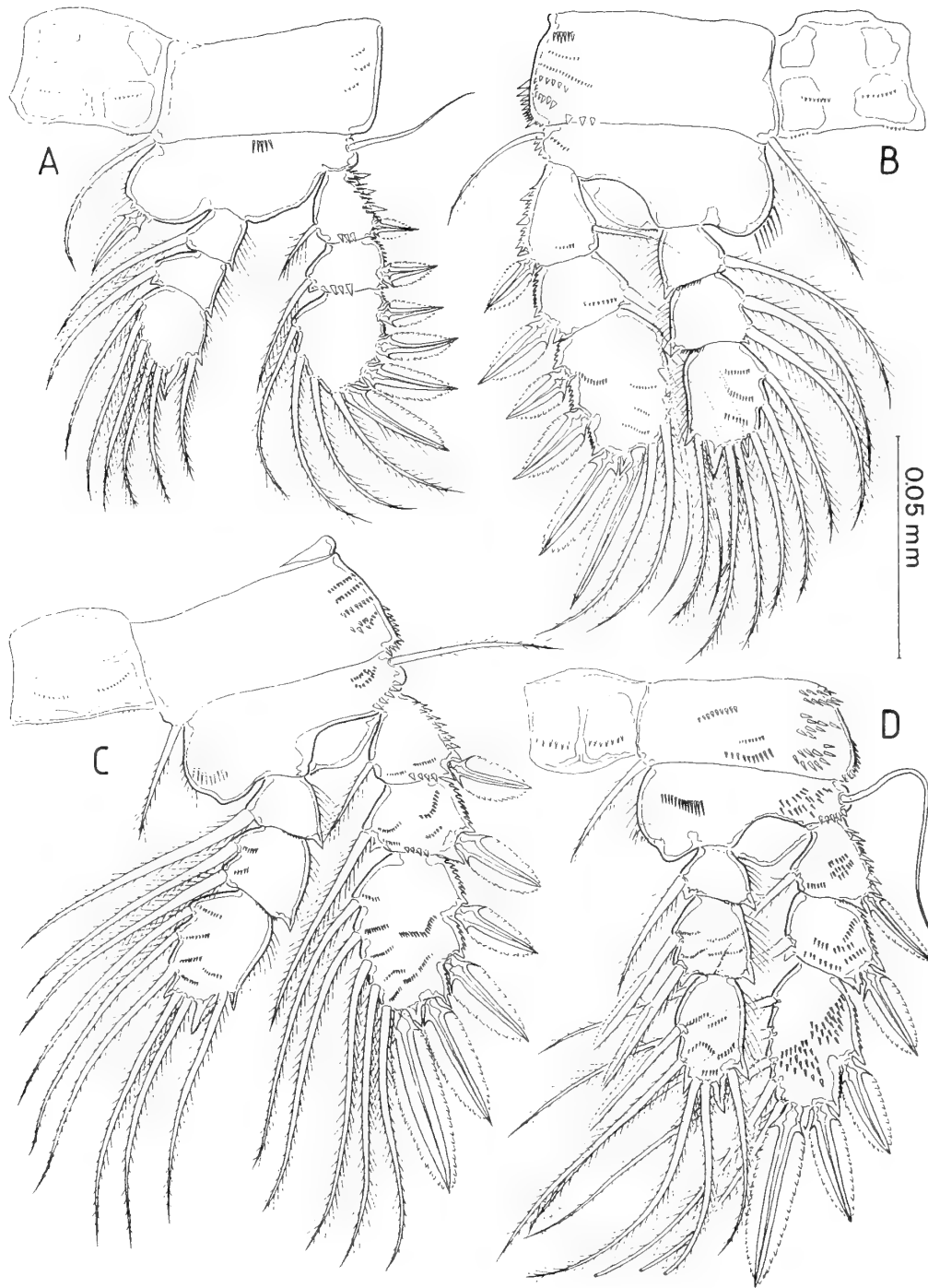


Fig. 3. *Cyclopina esilis* Brian, 1938, adult female swimming legs, posterior view. A, leg 1; B, leg 2; C, leg 3; D, leg 4.

than others, with brush of 6 setules on tip. Endopod 2-segmented, shorter than exopod, setal formula 3,6.

Maxillule (Fig. 2C) with well developed praecoxal arthrite, armed with 9 thick, unequal spines around distal margin plus isolated seta on posterior surface. Coxa and basis fused; coxal epipodite represented by 2 unequal setae; coxal endite represented by small cylindrical knob armed with 1 seta. Proximal and distal endites of basis discrete, bearing 3 and 2 setae respectively. Endopod 1-segmented, bearing 7 setae. Exopod 1-segmented, armed with 4 distal setae and marginal row of long setules.

Maxilla (Fig. 2D) well developed, 4-segmented. Praecoxa and coxa partially fused, endites with setal formula 3,1,3,3. Basis with large endite bearing claw-like spine plus 2 unequal setae. Endopod 2-segmented; proximal segment representing fused first and second ancestral endopodal segments; distal representing fused third and fourth ancestral segments; endopod setation formula (2+2),(2+4).

Maxilliped (Fig. 2E) slender, 6-segmented. Praecoxa and coxa fused forming syncoxa, bearing 3 (coxal) endites with setal formula 1,3,2. Basis with medial margin swollen, ornamented with marginal row of long setules; 2 setae implanted subdistally on medial margin.

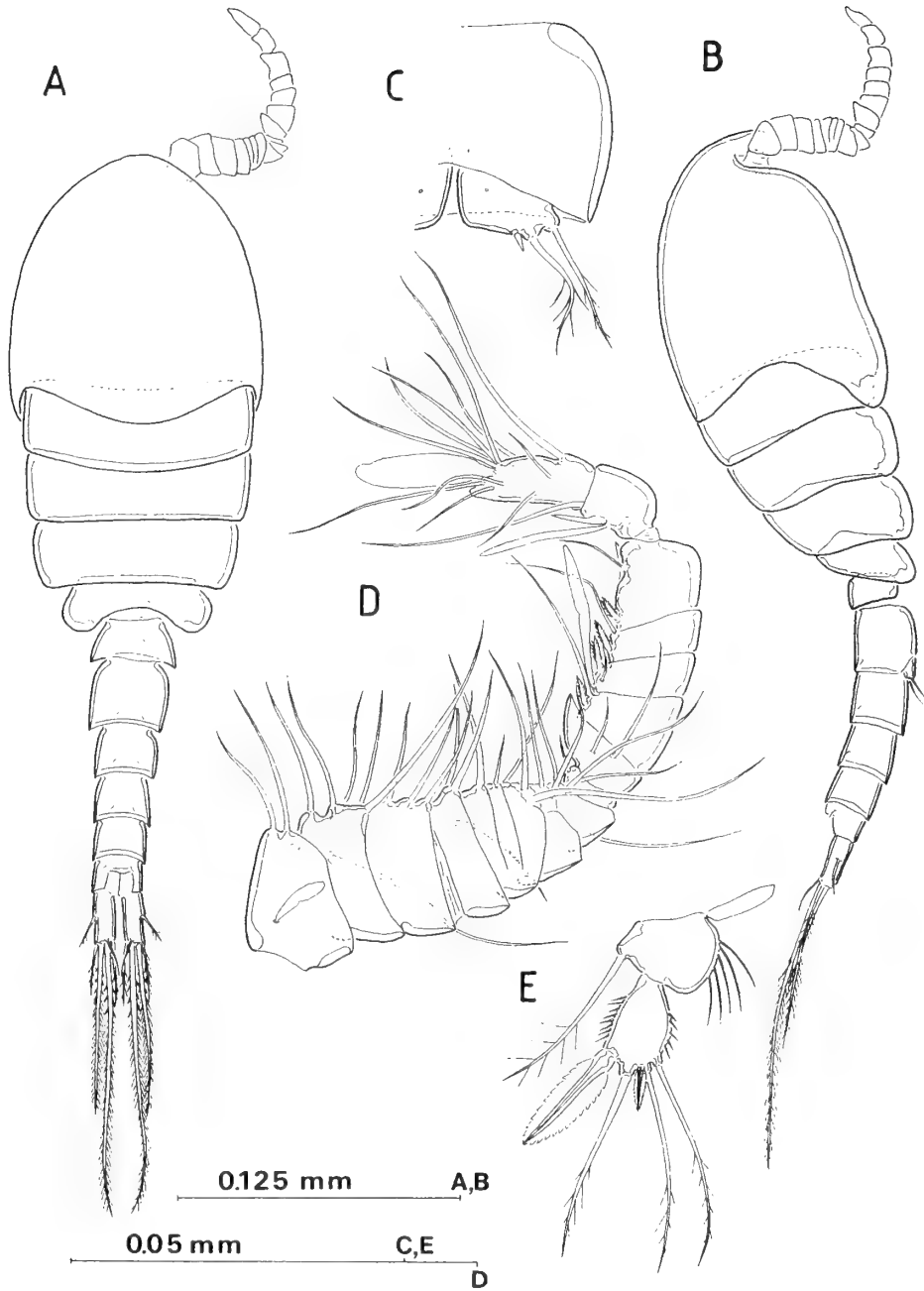


Fig. 4. *Cyclopina esilis* Brian, 1938, adult male. A, body, dorsal; B, lateral; C, genital somite, ventral; D, antennule; E, fifth leg.

Endopod 4-segmented, setal formula 0,0,1,4; transverse row of setules on segment 3.

Swimming legs 1 to 4 (Fig. 3) biramous, both rami 3-segmented. Legs subequal in size except first, somewhat reduced. All legs richly ornamented with denticles, as figured; anterior surface of coxae covered by small denticles, omitted from figures. Spines on exopodal segments flanged with serrate hyaline fringe; distal spine on third exopodal segment of legs 1 and 2, and spines on endopod of leg 4 flanged only on one side. Armature as follows:

	Coxa	Basis	Exopod	Endopod
Leg 1	0-1	1-I	I-1;I-1;III,I,4	0-1;0-1;1,2,3
Leg 2	0-1	1-0	I-1;I-1;III,I,5	0-1;0-2;1,2,3
Leg 3	0-1	1-0	I-1;I-1;III,I,5	0-1;0-2;1,2,3
Leg 4	0-1	1-0	I-1;I-1;II,I,5	0-1;0-II;1,2,1+I

Fifth legs (Fig. 1E) uniramous, 2-segmented, joined by naked intercoxal sclerite. Coxa and basis fused forming trapezoidal protopodal segment; inner margin with row of long setules; outer margin with long, smooth seta subdistally. Distal segment (exopod) about 1.6 times as long as wide, produced distally into median

process bearing 1 long, plumose seta; 1 flanged spine present subdistally on each side of process, outer stouter, longer than segment; inner spine less than half length of outer spine; spinule ornamentation on segment as figured.

**ADULT MALE.** Body (Figs 4A, B) up to 0.38 mm long, more slender than female. Urosome 6-segmented, with genital somite (Fig. 4C) symmetrical, slightly expanded laterally; paired gonopores opening ventrally at posterior border of somite; genital opercular flaps each armed with tiny inner spine plus 2 long, outer setae.

Antennules (Fig. 4D) 15-segmented, symmetrical, digeniculate. Geniculations between segments homologous with ancestral segments XV and XVI (9 and 10), and between XX and XXI (13 and 14). Segment 9 (XV) cup-shaped, forming sheath around proximal half of segment 10 (XVI). Segmental fusion pattern and armature as follows: segment 1 (corresponding to fused ancestral segments I and II), 3 setae + aesthetasc; segment 2 (fused ancestral segments III to V), 5 setae; segment 3 (fused ancestral segments VI and VII), 4 setae; segment 4 (VIII), 2 setae; segment 5 (partially fused ancestral segments IX to XI), 6 setae; segments 6 to 9 (XII to XV), 2 setae each; segment 10 (XVI), 1 pectinate spine, 1 seta + aesthetasc; segments 11 and 12 (XVII and XVIII), 1 pectinate spine and 1 seta each; segment 13 (fused ancestral segments XIX and XX), 1 pectinate spine, 1 modified flattened spine, and 1 seta; segment 14 (fused ancestral segments XXI and XXII), 1 modified flattened, spine plate, 1 seta + aesthetasc; segment 15 (fused ancestral segments XXIII to XXVIII), 11 + aesthetasc.

Segmentation and setation of other cephalosomic appendages and swimming legs 1 to 4 as in female, except mandibular palp (Fig. 2B); distal brush-like seta on exopod much shorter and thicker, setules on tip longer than in female.

Fifth legs (Fig. 4E) resembling female condition, but with 2 additional setae implanted subdistally along inner margin of exopod.

**REMARKS.** The *Cyclopina* from the cave on Mallorca belongs to the group of species in the genus that displays a female leg 5 with the inner spine of distal segment less than half the length of the outer spine, the latter being longer than the segment itself. This group comprises *Cyclopina esilis* Brian, 1938, *C. americana* Herbst, 1982, and *C. cuipora* Lotufo, 1994. The taxon from Mallorca differs clearly from *C. cuipora*. The female antennule is 10-segmented (not 12-segmented as in *C. cuipora*), and the intercoxal sclerite of leg 4 is almost completely smooth (not powerfully ornamented with several rows of thick spinules as in *C. cuipora*) (Lotufo, 1994). Differences from *C. americana* include the short, subquadrate caudal rami (Herbst, 1982) which contrast with the elongate (2.5 to 3.2 times as long as wide) caudal rami of the Mallorcan taxon.

The *Cyclopina* from Mallorca is identified as *C. esilis*, based on the segmentation of the female antennule, the setation of leg 5 and the proportions of the caudal rami. We noted variation in the proportional length of the caudal rami within the Mallorcan population. A similar degree of variability in length of the caudal rami has been reported in *C. esilis* (Brian, 1938; Monchenko, 1979).

Apparent differences in the armature of mouthparts have not been evaluated since we suspect that the armature of mouthparts in *C. esilis* (as in most species of *Cyclopina*) were inadequately described in the original descriptions. This is evident in the presence of a coxal endite, armed with 1 seta, on the maxillule of the *Cyclopina* from Mallorca. This character state (presence of coxal endite) had never previously been noted in *Cyclopina* and was known only from *Cyclopinodes elegans* (T. Scott, 1894) in the family Cyclopinidae (Huys & Boxshall, 1991). The coxal endite may well be present on the maxillule of all *Cyclopina* species. It is present in *C. gracilis* Claus, 1863 from the coast of Scotland (BMNH 1986.377) (pers.

obs.), and perhaps also in *C. oblivia* Monchenko, 1981, according to Monchenko (1989: Fig. 9).

The *Cyclopina* reported by Herbst (1953; 1962) from Banyuls (South France) and from Brittany (NW France) as *C. cf. kieferi* Schäfer, 1936 was recorded living as a commensal with polychaetes (Bretagne) and free-living in the marine interstitial (Banyuls). The illustrations provided by Herbst differ from the original description by Schäfer (1936) in an important character for the taxonomy of the genus, the relative length of the female leg 5 exopodal spines. This discrepancy was already noted by Lotufo (1994) when he presented the differential diagnosis of *C. yutimaete* Lotufo in comparison to *C. kieferi*. As the female leg 5 of *C. cf. kieferi* figured by Herbst (1953; 1962) is identical to that of *C. esilis*, we here consider Herbst's material as belonging to *C. esilis*.

The distribution of *Cyclopina esilis* thus encompasses the littoral zone from the Black Sea to the western approaches of the English Channel; a distribution equivalent to the Mediterranean and Lusitanian provinces of classical marine biogeography (Ekman, 1953).

Family **CYCLOPIDAE** Dana, 1846

Subfamily **HALICYCLOPINAE** Kiefer, 1927

Genus *Neocyclops* Gurney, 1927

*Neocyclops (Protoneocyclops) mediterraneus* (Kiefer, 1960)

(Figs 5–7)

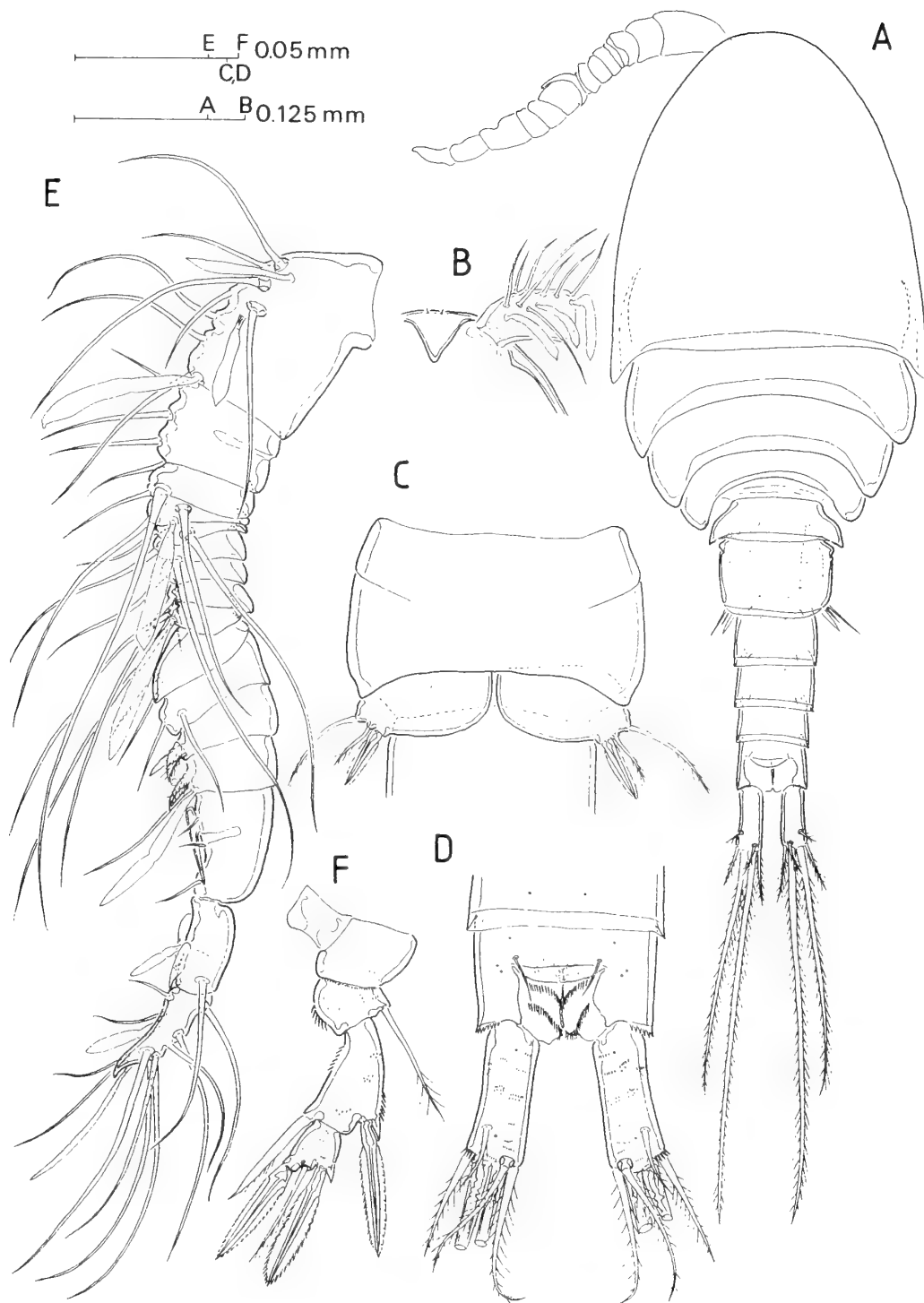
*Pareuryte mediterranea*: Kiefer (1960).

*Neocyclops remanei mediterraneus*: Pesce & Galassi (1987)

**MATERIAL EXAMINED.** Cova 'C' de Cala Varques (Manacor). UTM coordinates: 525.27; 4372.19. Topography in Trias & Mir (1977): Two adult males (one not preserved) and 2 copepodids (one not preserved) (BMNH 1995. 1329–1330). Collected by authors, 29 March 1995.

**ADULT MALE.** Body (Fig. 5A) cyclopidiform, up to 0.58 mm long, colourless. Nauplius eye absent. Prosome about 1.4 times as long as urosome, 5-segmented, first pedigerous somite completely concealed by carapace-like, posterior extension of cephalosome. Rostrum (Fig. 5B) triangular in frontal view. Urosome 6-segmented, robust. Fifth pedigerous somite with pointed posterolateral angles; entire hyaline frill adorning posterodorsal margin of somite. Genital somite (Fig. 5C) slightly expanded laterally, with ventrolateral fold each side of somite at about one-third distance from posterior margin; folds slightly projecting dorsally. Paired gonopores opening ventrally, each covered by opercular flap derived from sixth leg; flaps each armed with 1 inner flanged spine and 2 outer plumose setae. Abdominal somites 1 to 3 subequal, with posterior margins adorned with entire hyaline frill. Anal somite bearing operculum dorsally at about midlength; operculum ornamented with serrate hyaline frill; 4 rows of transverse setules adorning sides of anal cleft; posterolateral margins of somite bearing serrate hyaline frill. Caudal rami 2.6 times as long as wide, inserted widely separate from each other; secondary ornamentation of pores and tiny spinules distributed as figured; armature consisting of 7 setae; seta I reduced, tiny, implanted ventrolaterally about one-third of distance along ramus; seta II implanted dorsolaterally at about three-quarters of distance along ramus.

Antennules (Fig. 5E) 16-segmented, not extending beyond posterior margin of prosome, symmetrical, digeniculate (Fig. 5A).



**Fig. 5.** *Neocyclops (Protoneocyclops) mediterraneus* (Kiefer, 1960), adult male. A, body, dorsal; B, detail of rostral plate; C, genital somite, ventral; D, anal somite and caudal rami, dorsal; E, antennule; F, fifth leg.

Geniculations between segments homologous with ancestral segments XV and XVI (10 and 11), and between XX and XXI (14 and 15). Segment 10 cup-shaped, forming sheath around proximal half of segment 11. Segmental fusion pattern and armature as follows: segment 1 (corresponding to fused ancestral segments I to V), 8 setae + 3 aesthetascs; segment 2 (partially fused ancestral segments VI and VII), 4 setae; segment 3 (VIII), 2 setae; segment 4 (IX), 2 setae +

aesthetasc; segments 5 to 8 (X to XIII), 2 setae each; segment 9 (XIV), 2 setae + aesthetasc; segments 10 and 11 (XV and XVI), 2 setae each; segment 12 (XVII), 2 setose spines; segment 13 (XVIII), 1 setose spine, 1 seta + aesthetasc; segment 14 (fused segments XIX and XX), 1 modified flattened spine and 3 setae; segment 15 (partially fused segments XXI and XXII), 2 modified flattened spines, 1 seta + aesthetasc; segment 16 (fused segments XXIII to

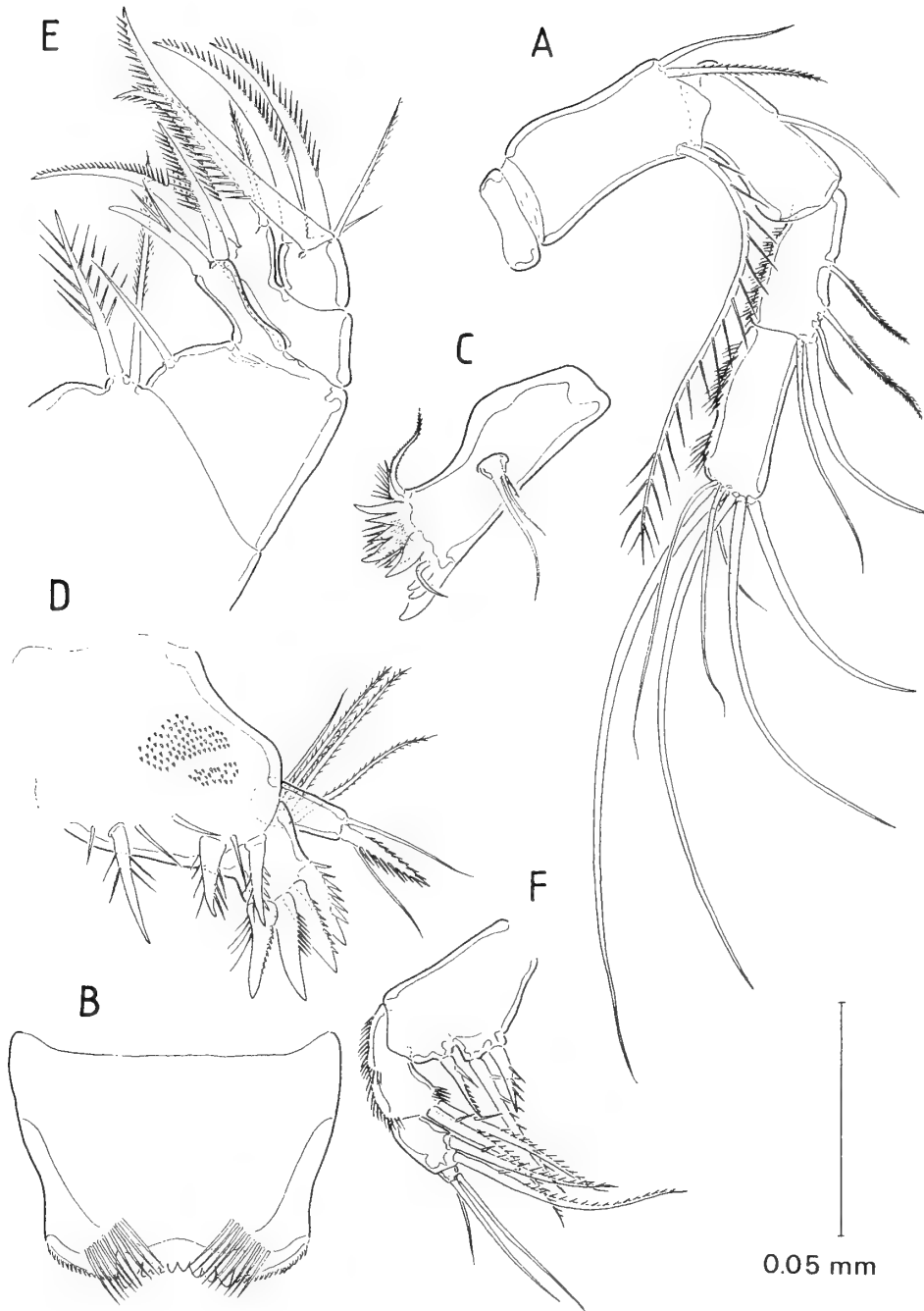


Fig. 6. *Neocyclops (Protoneocyclops) mediterraneus* (Kiefer, 1960), adult male. A, antenna; B, labrum, ventral; C, mandible; D, maxillule; E, maxilla; F, maxilliped.

XXVIII), 11 + 2 aesthetascs.

Antenna (Fig. 6A) well developed, 5-segmented. Coxa and basis separate, coxa small, unarmed. Basis with 2 distal setae on inner margin and long distal seta, representing exopod, on outer margin. First endopod segment with seta inserted midway along inner margin; second endopod segment with 2 lateral and 3 distal setae along inner margin; outer margin covered by patch of setules. Third endopod segment with 7 unequal setae on tip; outer margin adorned with 2 patches of setules, as figured.

Labrum (Fig. 6B) with laterally serrate distal margin and row of 9 rounded teeth midway along margin; paired transverse rows of long

spinules located subdistally on anterior surface of labrum.

Mandible (Fig. 6C) with coxal gnathobase armed with 9 sharp, unequal teeth, 2 dorsal setae, and 1 naked setiform element located between third and fourth ventralmost teeth; inner dorsal seta with long spinules along one side; outer dorsal seta longer, with short spinules on both sides; transverse row of 7 long spinules disposed subdistally on cutting blade. Mandibular palp reduced to knob bearing 3 unequal setae.

Maxillule (Fig. 6D) with well developed praecoxal arthrite; armature consisting of 3 stout spines, 1 tiny spine and 3 slender setae proximally, and distal lobe armed with 4 thick, denticulate spines; 2

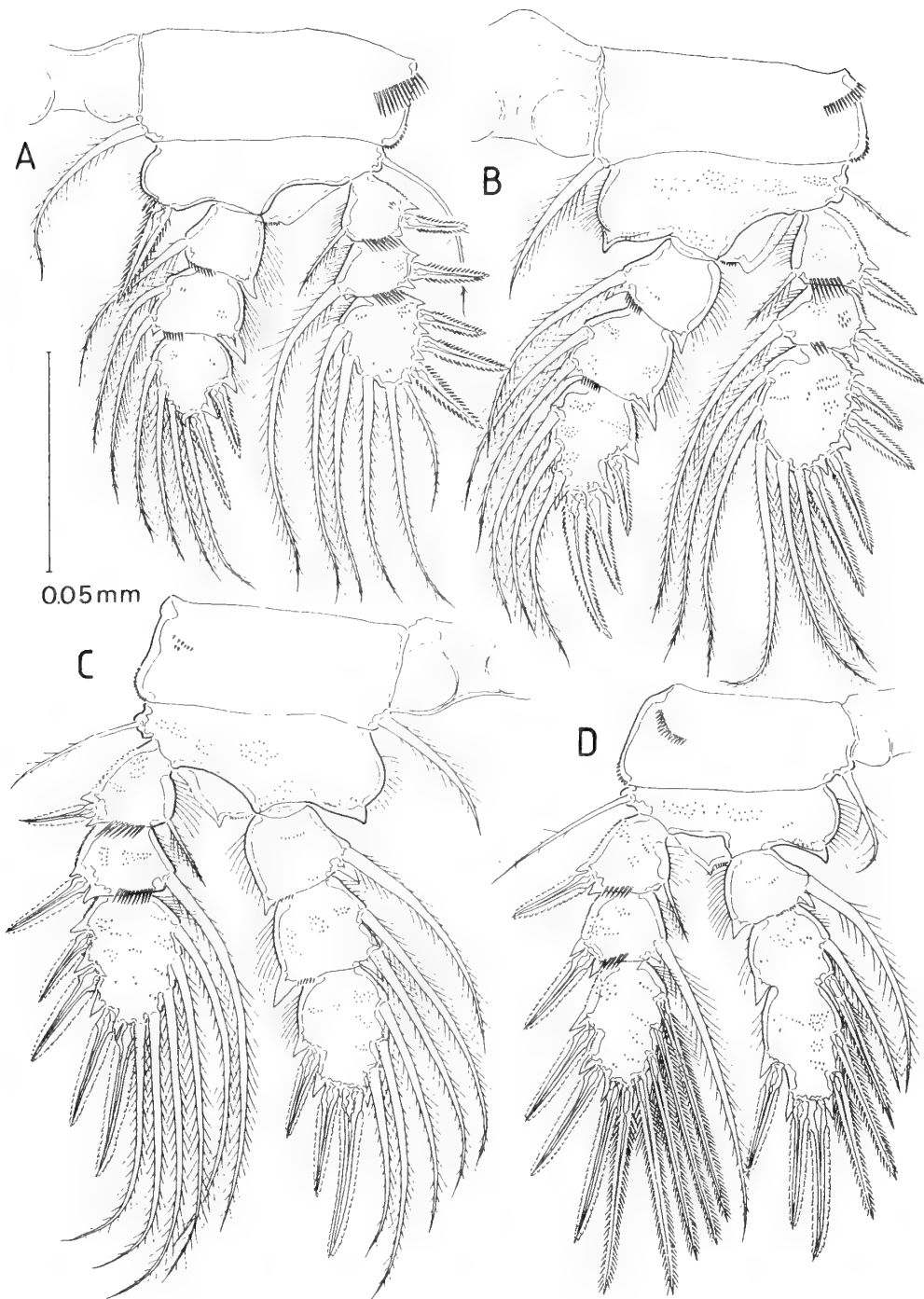


Fig. 7. *Neocyclops (Protoneocyclops) mediterraneus* (Kiefer, 1960), adult male swimming legs, posterior view. A. leg 1; B. leg 2; C. leg 3; D. leg 4.

patches of tiny denticles on segment, as figured. Palp comprising coxobasis with 1 spinulate spine and 2 setae distally and 1 seta (representing exopod) implanted on outer margin, and 1-segmented endopod bearing 3 setae (obscured in Fig. 6D).

Maxilla (Fig. 6E) 4-segmented, powerfully developed. Praecoxa and coxa separate. Praecoxa with single, distal endite armed with 2 elements. Proximal coxal endite represented by single seta; distal coxal endite discrete, armed with 1 spine fused to endite and ornamented with 2 strong spinules, and 1 pectinate spine. Basis with endite bearing 2 stout pectinate elements, one fused to segment, plus

reduced pinnate seta. Endopod 1-segmented, bearing 3 stout pectinate elements plus 2 reduced, unequal setae.

Maxilliped (Fig. 6F) somewhat reduced in size, 4-segmented. Syncoxa bearing 2 weakly developed endites with spine formula 2,1. Basis with 2 distal setae on inner margin; secondary ornamentation of spinules and setules on segment as figured. Endopod 2-segmented, setal formula 2,3. Ornamentation on spines and setae of maxilliped as figured; setae on distal endopodal segment naked.

Swimming legs 1 to 4 (Fig. 7) biramous, both rami 3-segmented, except leg 4 with 2-segmented endopod; distal endopodal segment

subdivided by posterior surface suture marking plane of fusion between second and third segments. Legs subequal in size except first somewhat reduced. Intercostal sclerites lacking ornamentation. Legs richly ornamented with spinules, setules and denticles, as figured. Armature as follows:

	Coxa	Basis	Exopod	Endopod
Leg 1	0-1	1-1	1-1;I-1;III,I,4	0-1;0-2;I,I+1,3
Leg 2	0-1	1-0	1-1;I-1;III,I,5	0-1;0-2;I,II,3
Leg 3	0-1	1-0	1-1;I-1;III,I,5	0-1;0-2;I,II,3
Leg 4	0-1	1-0	1-1;I-1;III,I,IV	0-1;I,II,III+1

Spines on legs 1 and 2 adorned with spinules, those on leg 3 flanged with serrate hyaline frill. Outer spines on exopod and outer and distal spines on endopod of leg 4 ornamented with serrate hyaline frill; inner margin setae on both rami modified, spine-like, ornamented with short pinnules proximally and serrate membrane distally.

Fifth legs (Fig. 5F) 4-segmented, joined by smooth intercoxal sclerite. Coxa and basis separate, coxa unarmed, basis bearing plumose seta on outer margin. First exopodal segment elongate, about as long as coxa and basis combined, outer margin with subdistal spine flanged with serrated hyaline frill; spinous process just anterior to insertion point of spine; flanged spine longer than segment; inner margin of segment bearing 1 distal spine adorned with sparse setules and about as long as segment. Distal margin of second exopodal segment bearing 1 seta flanked by 2 spines flanged with serrate hyaline frill; spinous process just proximal to insertion point of outer spine; outer spine slightly longer than inner, and longer than first exopod segment; inner spine about as long as first endopod segment; seta shorter than spines. Secondary ornamentation on fifth leg segments as figured.

**REMARKS.** The genus *Neocyclops* Gurney, 1927 contains 15 species distributed in coastal waters of the Northeast and Tropical Atlantic (including the Caribbean), the Mediterranean, the Black and Red Seas, the Indian Ocean, as well as the Pacific (Papua New Guinea) (Petkovski, 1986; Fiers, 1986; Pesce & Galassi, 1993; Lotufo & Rocha, 1993; Rocha, 1995). Petkovski (1986) has split the genus into two subgenera according to the number of exopodal segments of the male fifth legs. The subgenus *Neocyclops*, characterized by a 3-segmented male leg 5, embraces the following species: *N. medius* Herbst, 1955, *N. vicinus* (Herbst, 1955), *N. affinis* (Plesa, 1961), *N. salinarum* (Gurney, 1927) and *N. remanei* (Herbst, 1952).

The subgenus *Protoneocyclops*, with 4-segmented male fifth legs, comprises *P. stocki* Pesce, 1985, *P. geltrudeae* Pesce & Galassi, 1993, *P. papuensis* Fiers, 1986, *P. mediterraneus* (Kiefer, 1960), *P. herbsti* Petkovski, 1986, *P. wellsi* Petkovski, 1986 and *P. ferrarii* Rocha, 1995. This subgenus displays the so-called full Tethyan pattern of distribution (Stock, 1993), i.e., circum-tropical in the entire region of the former Tethys Sea.

Three other species, viz. *N. improvisus* Plesa, 1973 from Cuba, and *N. magnus* (Sewell, 1949) and *N. parvus* (Sewell, 1949) from islands in the Indian Ocean, cannot be assigned to either subgenus as their males are unknown.

Three representatives of the genus are known so far from the Mediterranean region. *Neocyclops (N.) salinarum*, originally described from the Suez Canal, was reported also from the Camargue (South France) and the Sirbonian lagoon (Mediterranean coast of Sinai) (Gurney, 1927a; 1927b; Aguesse & Dussart, 1956; Por, 1973). As Petkovski (1986) pointed out, the identity of the French population needs to be confirmed. Similarly the single copepodid from the

Andaman Islands (Indian Ocean), assigned by Sewell (1949) to this species, should be reexamined. *Neocyclops (N.) vicinus*, a species distributed along the coasts of Brazil and the Lesser Antilles (Pesce & Galassi, 1993; Lotufo & Rocha, 1993), has been also reported from the Black Sea (as *Eurycyclops remanei vicinus*) by Plesa (1963) and Monchenko (1975). As pointed out by Lotufo & Rocha (1993), this record is dubious since their material seems more closely related to *N. (N.) remanei* than to *N. (N.) vicinus*.

The single representative of the subgenus *Protoneocyclops* in Mediterranean waters is *Neocyclops (P.) mediterraneus*, originally described by Kiefer (1960) as *Pareuryte mediterranea* from an anchihaline cave on Menorca (Balearic Islands). Later, Pesce & Galassi (1987) reported it from an anchihaline cave in Southern Italy. Plesa (1981) cited the same species from Cuba, but this record has been reassigned by Petkovski (1986) to *N. (N.) stocki*, a taxon widespread in the Caribbean region (Pesce & Galassi, 1993).

The *Neocyclops* from Mallorca has been identified on the basis of the 4-segmented condition of the male fifth legs and the relative lengths of the armature elements on this leg. Mallorca is also close to the type-locality of the species (Menorca). Other characters could not be checked against Kiefer's (1960) original description since this contained only 5 drawings (viz. female anal somite and caudal rami, distal segment of endopod of female leg 4; fifth leg of both sexes, and genital operculum of male). In addition, Kiefer did not designate types for the species. Pesce & Galassi (1987) had only 2 females at their disposal for their supplementary description.

A differential diagnosis of *Neocyclops (Protoneocyclops) mediterraneus* (Kiefer, 1960) can be constructed based on characters of the male fifth leg. It differs from *N. (P.) geltrudeae* Pesce & Galassi (1993) from Curaçao (Antilles) in the number of armature elements on the distal segment (3, compared to 4 in *N. (P.) geltrudeae*). Differences from *N. (P.) papuensis* Fiers, 1986 from New Guinea and *N. (P.) ferrarii* Rocha, 1995 from Brazil involve the relative lengths of the spines on the distal segment (the inner spine is clearly longer than the outer in both these species whereas in *N. (P.) mediterraneus* the outer spine is subsimilar, slightly longer than the inner). Differences from *N. (P.) herbsti* Petkovski (1986) from the Red Sea, and *N. (P.) stocki* Pesce, 1985 from the Caribbean, are based on the relative lengths of the flanged spines on the 2 distal segments of leg 5 (these are clearly shorter than the first exopodal segment whereas in *N. (P.) mediterraneus* they are longer than the segment). In addition, in *N. (P.) herbsti* the armature element on the inner margin of the first exopodal segment is a plumose seta, whereas in *N. (P.) mediterraneus* it is a thick spine. Differences from *N. (P.) wellsi* Petkovski (1986) from Mozambique lie only in the nature of the armature element on the inner margin of the first exopodal segment, which is also a seta in this species instead of a thick spine.

Subfamily **EURYTEINAE** Monchenko, 1975

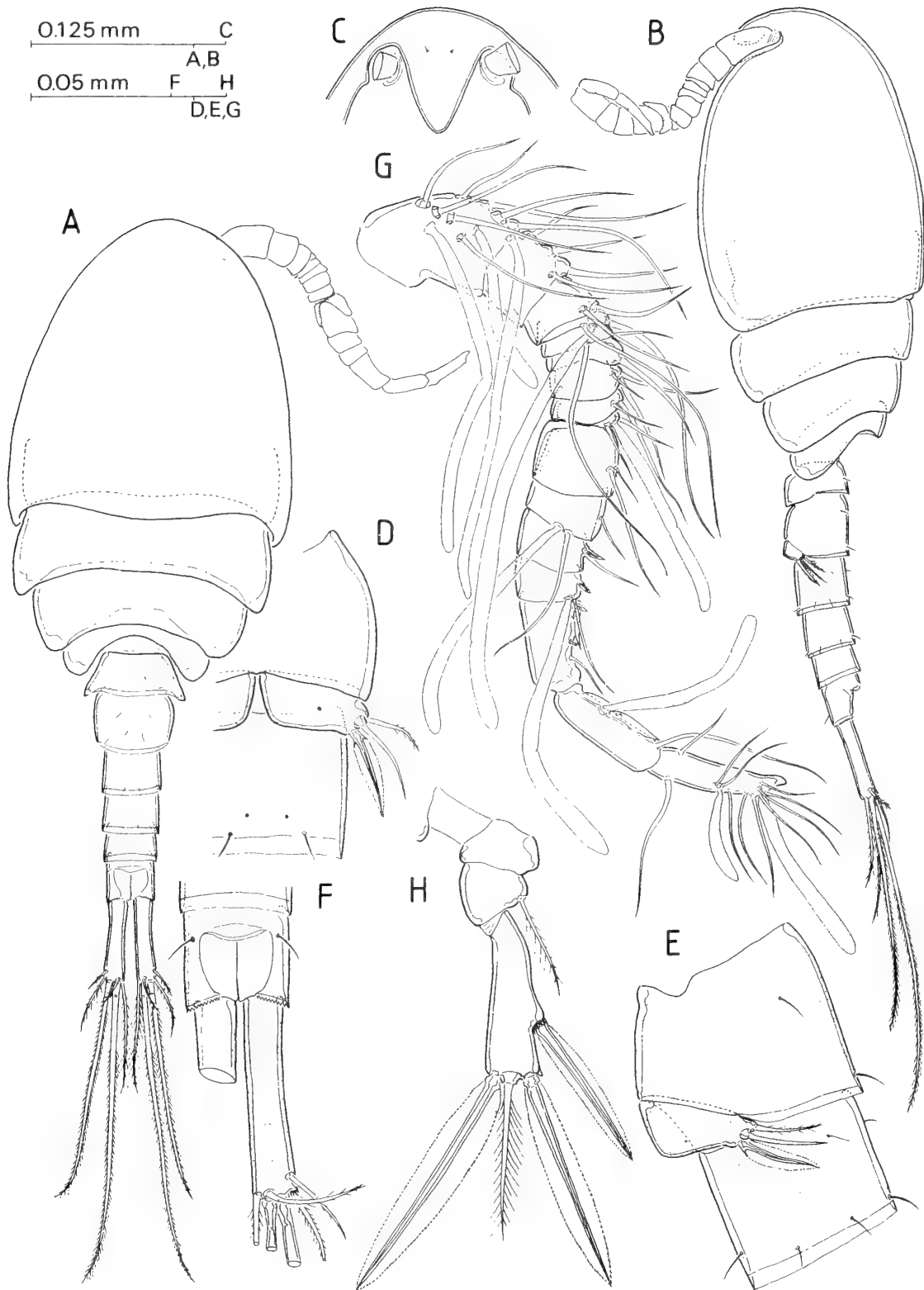
Genus *Euryte* Philippi, 1843

***Euryte longicauda* Philippi, 1843 emend. Giesbrecht, 1900**

(Figs 8-11)

**MATERIAL EXAMINED.** Cova de na Barxa (Capdepera). UTM coordinates: 539.30; 4393.10. Topography in Andrews *et al.* (1989): Two adult females, 1 adult male, and 1 copepodid (BMNH 1995. 1323-1326). Collected by authors, 3 April 1995. - Cova de na Mitjana (Capdepera): 19 adult males, 7 adult females, and 5

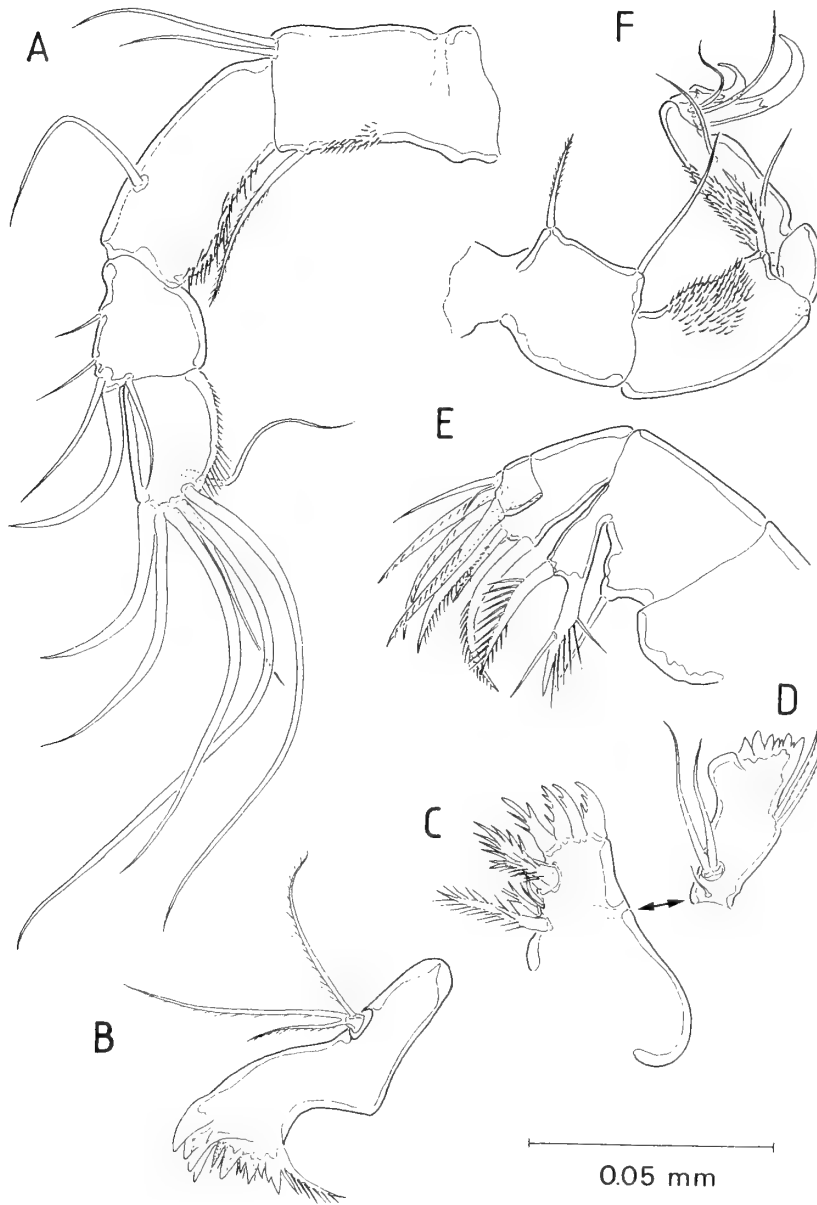




**Fig. 8.** *Euryte longicauda* Philippi, 1843, adult male. A, body, dorsal; B, lateral; C, detail of rostral plate; D, genital and first abdominal somites, ventral; E, same, lateral; F, anal somite and caudal rami, dorsal; G, antennule; H, fifth leg.

copepodids (BMNH 1995. 1313–1322). Collected by authors, 1 April 1995. – Es Secret des Moix (Manacor). Coordinates: 523.69; 4365.53. Topography in Ginés *et al.* (1975): One adult female

(BMNH 1995. 1327). Collected by D. Jaume, 25 May 1994. ADULT MALE. Body (Figs 8A, B) cyclopiform, up to 0.61 mm long, body surface completely covered by tiny cuticular granulations.

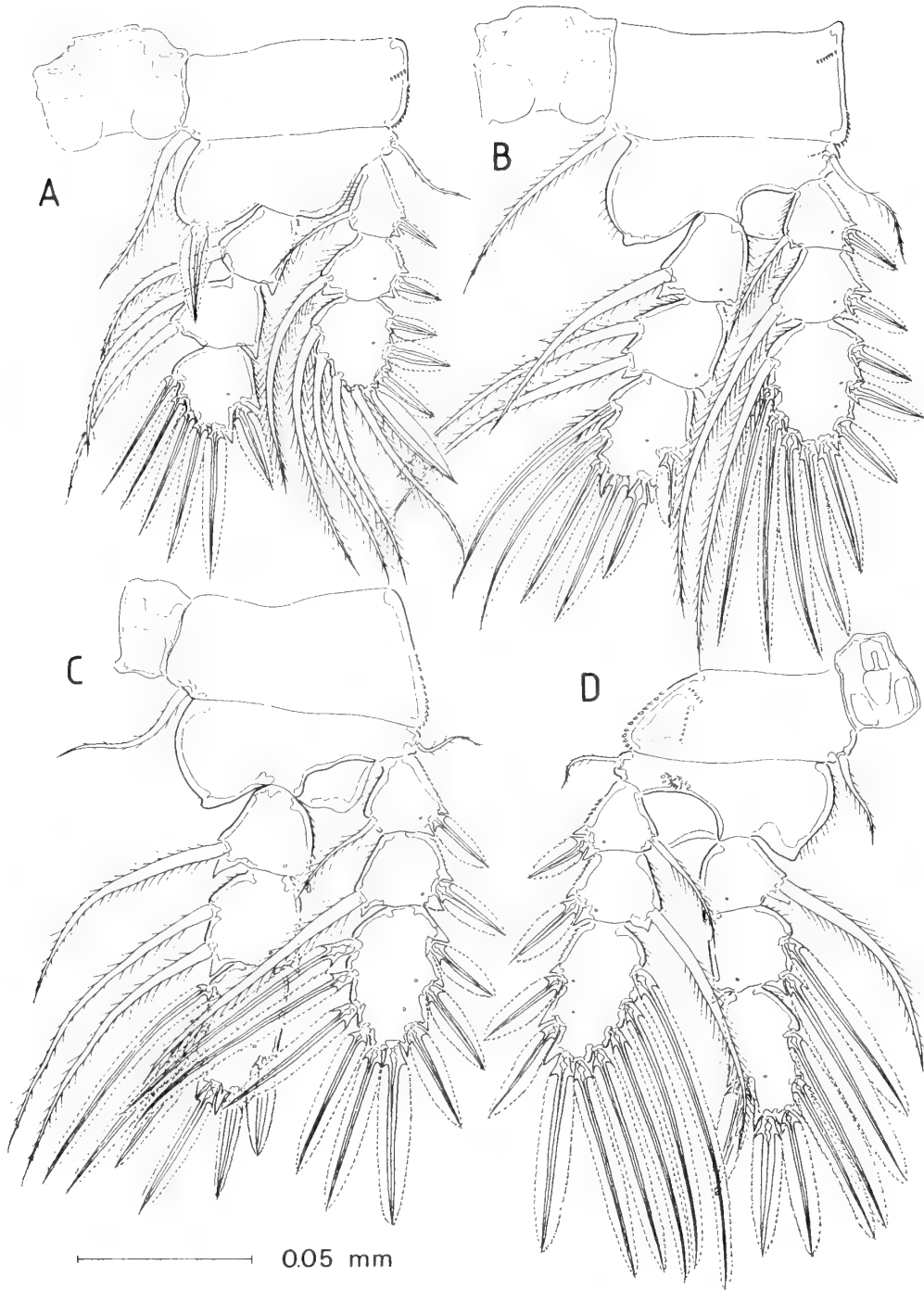


**Fig. 9.** *Euryte longicauda* Philippi, 1843, adult male. A, antenna; B, mandible; C, maxillule praecoxal arthrite; D, maxillary palp; E, maxilla; F, maxilliped.

Prosoma about 1.4 times longer than urosome, comprising cephalosome plus 4 free pedigerous somites; first pedigerous somite completely concealed beneath posterior extension of cephalosome, with lateral margins weakly sclerotized; second to fourth somites with evenly rounded posterolateral corners. Rostrum (Fig. 8C) fused at base, well developed, subtriangular in frontal view. Urosome 6-segmented; first abdominal somite with pointed posterolateral angles; entire hyaline frill adorning posterodorsal margin. Genital somite (Figs 8D,E) symmetrical, laterally expanded, with entire hyaline frill around posterodorsal margin; pair of gonopores opening ventrally at posterior border of somite; opercular flaps each armed with 1 inner flanged spine plus 2 outer setae. Third to fifth urosome somites subequal, narrower than genital somite, with entire hyaline frill adorning posterior margin. Anal somite (Fig. 8F) about same size as preceding somites; smooth anal operculum present at one third of distance along somite; serrate hyaline frill around posterolateral margins of

somite. Caudal rami (Fig. 8F) of variable length, from 4.4 to 6.4 times longer than wide, slightly divergent; distal part slightly wider; armature consisting of 6 setae; seta II located subdistally.

Antennules (Fig. 8G) 16-segmented, symmetrical, digeniculate with geniculations between segments homologous with ancestral segments XV and XVI (10 and 11), and segments XX and XXI (14 and 15). Segment 10 cup-shaped, forming sheath around proximal half of segment 11. Segmental fusion pattern and armature as follows: Segment 1 (corresponding to fused ancestral segments I to V), 8 setae + 3 aesthetascs; segment 2 (corresponding to fused ancestral segments VI and VII), 4 setae; segment 3 (VIII), 2 setae; segment 4 (IX), 2 + aesthetasc; segments 5 to 8 (X to XIII), 2 setae each; segment 9 (XIV), 2 + aesthetasc; segment 10 (XV), 2 setae; segment 11 (XVI), 2 + aesthetasc; segment 12 (XVII), 1 short denticulate spine and 1 seta; segment 13 (XVIII), 1 short denticulate spine, 1 seta + aesthetasc; segment 14 (fused ancestral segments XIX and XX), 1 short denticulate spine, 1 modified flattened spine,



**Fig. 10.** *Euryte longicauda* Philippi, 1843, adult male swimming legs, posterior view. A, leg 1; B, leg 2; C, leg 3; D, leg 4.

1 seta + minute aesthetasc; segment 15 (fused ancestral segments XXI to XXIII), 2 modified flattened spines, 1 seta + aesthetasc; segment 16 (fused segments XXIV to XXVIII), 11 setae + 2 aesthetascs.

Antenna (Fig. 9A) 4-segmented. First segment representing partially fused coxa and basis, armed with 2 inner basal setae distally, and 1 outer seta representing exopod; patch of setules present midway along outer margin of segment. Endopod 3-segmented; first segment about as long as coxa and basis combined, armed with 1 seta at three-quarters of distance along inner margin; outer margin

ornamented with setules. Segments 2 and 3 subsimilar in size, each about half length of segment 1. Segment 2 armed with 2 lateral and 3 distal setae, one of latter (seta VIII in scheme of Boxshall & Evstigneeva, 1994) claw-like, along inner margin. Segment 3 armed with 7 distal setae, one claw-like; outer margin with 2 rows of setules, as figured.

Mandible (Fig. 9B) with large coxal gnathobase bearing 11 unequal, sharp blades, plus 2 dorsal spines; outer dorsal spine spinulate; transverse row of 6 thin spinules located adjacent to cutting edge. Palp reduced to knob bearing 3 setae.

Maxillule with praecoxal arthrite (Fig. 9C) well developed, armed distally with 4 stout, denticulate spines, plus 6 more proximal elements, ranging from a tiny seta to a thick denticulate spine. Palp (Fig. 9D) comprising coxobasis with medial gnathobase-like structure and minute endopodal segment bearing 2 setae; distal margin of coxobasal gnathobase provided with 9 irregular blades; coxobasis with 2 setae located subapically on dorsal margin and single seta, representing exopod, located proximally on distal surface.

Maxilla (Fig. 9E) well developed, 4-segmented. Praecoxa and coxa incompletely separate. Praecoxa naked, lacking endites. Coxa with proximal endite represented by single seta; distal endite powerfully developed, bearing 2 stout, spinulate spines, proximal spine bearing single, conspicuous strong spinule on outer margin and row of thinner spinules on inner margin. Basis with endite bearing 3 unequal, claw-like setae. Endopod 1-segmented, armed with total of 3 stout spine-like setae, 1 naked seta and 1 very reduced seta.

Maxilliped (Fig. 9F) 5-segmented, prehensile. Syncoxa bearing 2 weakly developed endites provided with single seta each. Basis with inner margin covered by patch of setules and single seta positioned distally. Endopod 3-segmented, first segment short, unarmed; second segment elongate, inner margin covered by patch of setules, armed with 2 setae laterally; small distal segment with 2 stout, curved claws plus 2 accessory setae.

Swimming legs 1 to 4 (Fig. 10) biramous, both rami 3-segmented. Legs subequal in size except first somewhat smaller. Intercoxal sclerites lacking ornamentation and getting progressively narrower from legs 1 to 4. All spines on segments flanged bilaterally with serrate hyaline frill except distalmost spine on exopod of leg 1, which is flanged with frill on outer side only, inner side adorned with row of setules. Secondary ornamentation and pore pattern on segments as figured; pores on coxa and basis possibly overlooked. Armature as follows:

	Coxa	Basis	Exopod	Endopod
Leg 1	0-1	1-I	I-1;I-1;III,I,4	0-1;0-2;I-II-III
Leg 2	0-1	1-0	I-1;I-1;III,I,IV+1	0-1;0-2;I-II-III
Leg 3	0-1	1-0	I-1;I-1;III,I,V	0-1;0-2;I-II-III
Leg 4	0-1	1-0	I-1;I-1;II,I,V	0-1;0-II;I-II-III

Fifth legs (Fig. 8H) uniramous, 3-segmented, joined by smooth intercoxal sclerite. Coxa and basis separate, former naked, latter with single seta on outer margin. Distal segment (exopod) elongate, about 2.5 times as long as wide; armature consisting of flanged spine as long as segment located two-thirds of distance along outer margin, plus 2 flanged spines and single seta on distal margin; distal seta shorter than both spines and segment itself; distal spines located either side of seta, flanged, both clearly longer than segment, inner longer, about 1.5 times as long as segment. Secondary ornamentation of spinules present on outer margin of exopod of some individuals, similar to that figured on female leg 5 (Fig. 11E), not present in figured specimen (Fig. 8H).

**ADULT FEMALE.** Body (Figs 11A, B) up to 0.74 mm long, resembling male. Urosome 5-segmented; genital and first abdominal somites partially fused to form genital double-somite. Genital double-somite (Fig. 11C) symmetrical, subdivided dorsally by partial suture line; single copulatory pore opening mid-ventrally about one-third of distance along somite, connected via copulatory duct to fused seminal receptacles. Paired gonopores located laterally; gonopores covered by opercula, each consisting of lobe projecting dorsally bearing 2 setae and 1 tiny spine. Tapering soft lobe pointing posteriorly located at both sides of somite just behind

operculum. Entire hyaline frill present along posterior margin of somite.

Antennules (Fig. 11D) 21-segmented, not extending beyond posterior margin of cephalosome (Figs 11A, B), symmetrical. Segmental fusion pattern and armature as follows: Segment 1 (corresponding to fused ancestral segments I to V), 8 setae; segment 2 (fused ancestral segments VI and VII), 4 setae; segments 3 to 9 (VIII to XIV), 2 setae each; segments 10 to 13 (XV to XVIII), 1 seta each; segment 14 (XIX), naked; segment 15 (XX), 1 seta; segment 16 (XXI), 1 seta + aesthetasc; segment 17 (XXII), naked; segment 18 (XXIII), 1 seta; segment 19 (XXIV), 2 setae; segment 20 (XXV), 2 + aesthetasc; segment 21 (fused XXVI to XXVIII), 7 + aesthetasc.

Segmentation and setation of other cephalosomic appendages and swimming legs 1 to 4 as in male.

Fifth legs (Fig. 11E) resembling those of male, but with shorter exopod, about twice as long as wide; inner distal spine almost twice as long as segment; spine on outer margin clearly longer than segment. Secondary ornamentation of spinules on outer margin of exopod not discernible in some individuals.

**REMARKS.** The genus *Euryte* typically contains shallow water hyperbenthic species, although Brady (1910) reported the genus from depths of 320 m in the Antarctic and some species have been found living in the interstices of coarse sand, or in association with seaweed or corals. Ten species are currently recognized, distributed worldwide (Gurney, 1927b; Sewell, 1949; Herbst, 1989; Humes, 1991; 1992), with the possible exception of the Pacific coast of South America. Apart from the original contributions by Giesbrecht (1900) and Sars (1913-1918), new species have been described mainly on the basis of a biometric analysis of characters that have otherwise proved to exhibit high intra-populational variability (such as the relative length of caudal rami), or that may vary significantly in their measurements simply according to the precise angle of observation (such as the relative length of the armature elements on the distal segment of the endopod of leg 4). Such characters are widely used in the two identification keys available for the genus (Vervoort, 1964; Herbst, 1989), and their use has resulted in the false impression of cosmopolitanism of some taxa (viz. *E. longicauda* and *E. robusta* Giesbrecht, 1900; see Kiefer (1929) and Sewell (1949)).

The type material for most species of *Euryte* is no longer extant. This hampers the necessary revision of the genus, that could permit the critical reevaluation of all those taxa established on the basis of variable characters.

Using material from the type locality of both species (the Gulf of Naples), Giesbrecht (1900) differentiated *E. longicauda* Philippi, 1843 from *E. robusta* Giesbrecht, 1900 mainly by the proportions of the caudal rami and by details of the armature of the male antennule. The proportions of the caudal rami of the Mallorcan population overlap the characteristic values for both species given by Giesbrecht (1900). The armature of the male antennule, however, corresponds to that of *E. longicauda*: the cup-shaped segment 10 carries 2 slender setae, whereas in *E. robusta* it carries a characteristic robust, S-shaped spine plus a seta. On this basis we have assigned the *Euryte* from the Mallorcan caves to *E. longicauda*.

The differential diagnosis separating *E. longicauda* from Mallorca from *E. robusta* can be completed as follows (see the detailed illustrations of the latter species in Huys & Boxshall, 1991): in *E. longicauda* the proximal spine on the distal coxal endite of the maxilla of both sexes is armed with a single, strong spinule on one side and a row of thinner spinules on the other side; in *E. robusta* both sides are armed with thin spinules. Additionally, in *E. longicauda* there is a transverse dorsal suture midway along the female genital double-somite that seems to be absent in *E. robusta*.

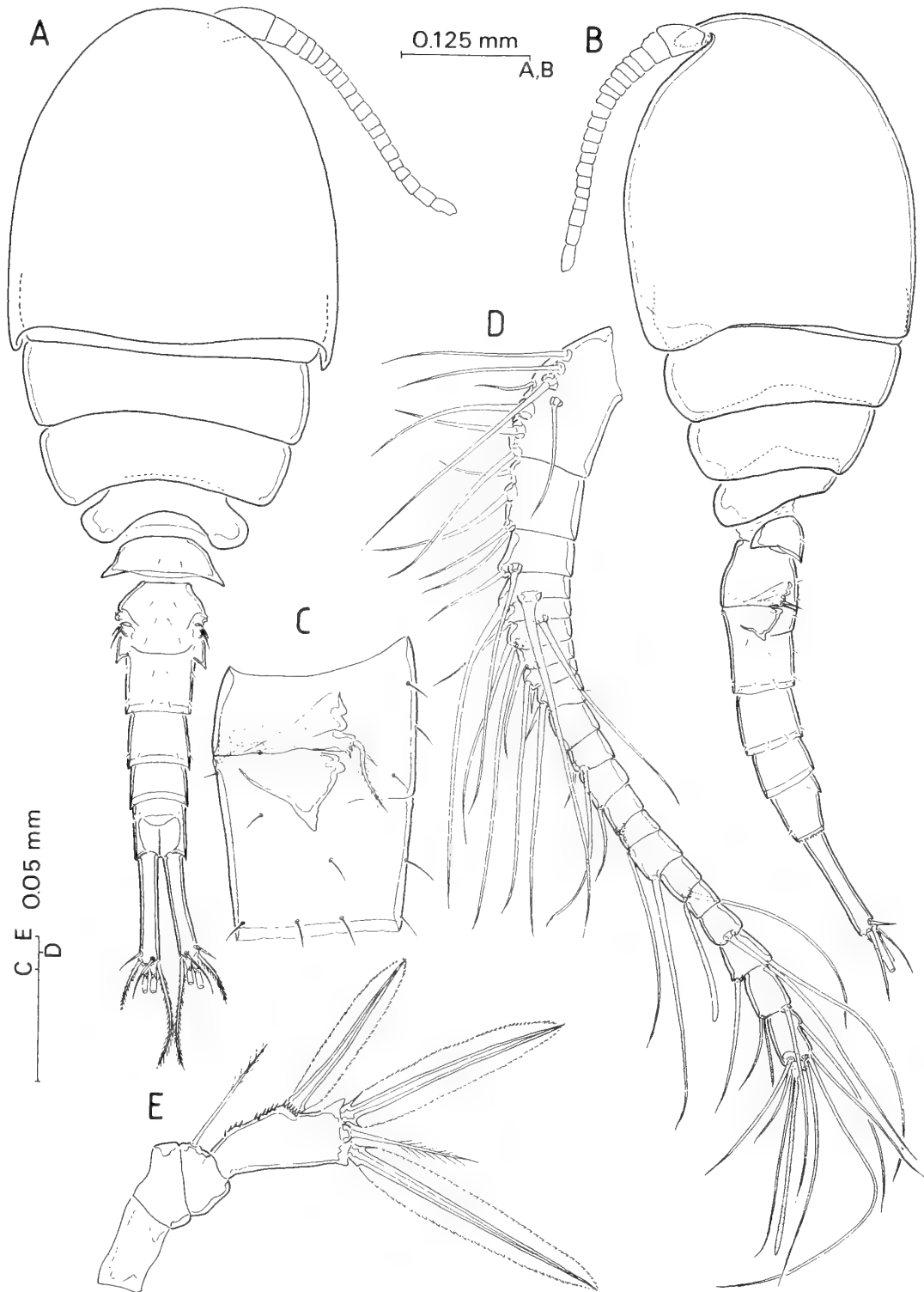


Fig. 11. *Euryte longicauda* Philippi, 1843, adult female. A, body, dorsal; B, lateral; C, genital double-somite, lateral; D, antennule; E, fifth leg.

The distribution of *E. longicauda* apparently encompasses all European coasts from the Black Sea to the Arctic Ocean (Franz-Joseph Land) and the East coast of Greenland (Giesbrecht, 1900; Herbst, 1989) but many of the records of this species are accompanied by inadequate descriptions, if any. Geographically remote records from outside the European region, such as those of Gurney

(1927b) from the Suez Canal and Samoa, or those of Thomson (1882) (as *Thorellia brunnea* var. *antarctica*) from New Zealand, require verification.

The body size of the *Euryte longicauda* from Mallorca falls in the range characteristic of the variety *E. longicauda* var. *minor* Scott, 1905, which was elevated, rather inconsistently, to full specific

status by Sars (1919–1921). This variety is supposed to inhabit deeper waters and never to occur in the littoral zone (Sars, 1913–1918; 1919–1921).

According to Sars (1913–1918) *E. longicauda* carries only modified flanged spines on the inner margin of the third exopodal segment of leg 2. In our material the proximalmost element on the inner margin of this segment is a plumose seta (Fig. 10B). However, this may be an observational error by Sars since material of *E. longicauda* from Raunefjorden in Norway (BMNH 1986.387) and from Scotland (BMNH 1951.8.10.587) in the collections of the Natural History Museum possesses a plumose seta in this position, as in the Mallorcan material.

## REVIEW OF EURYTE SPECIES

The eight remaining species of *Euryte* are briefly reviewed here, in order to facilitate the identification of representatives of this problematic genus. The review is essentially comparative and emphasises the most robust and reliable characters available in published descriptions.

*E. curticornis* Sars, 1913 is characterised by short, 21-segmented female antennules and the shortened third segment of the maxilliped; the curved distal claws on this appendage are also reduced in size and subsimilar in length. These features contrast with the maxilliped of *E. longicauda*, which is provided with an elongate third segment and with long, unequal distal claws. The distal spines on the third segment of endopod of leg 1 are clearly unequal in length in *E. curticornis*, whereas in *E. longicauda* they are about equal.

*E. longicauda* can be distinguished from *E. grata* Herbst, 1889 and *E. verecunda* Humes, 1992 by some features of the maxilla and maxilliped. In *E. verecunda*, the proximal spine on the distal coxal endite of maxilla is adorned on both sides with slender spinules. On the maxilliped, the armature element on the proximal syncoxal endite is a seta in *E. longicauda*, whereas in *E. grata* and *E. verecunda* this endite is represented by a stout spine. *E. verecunda* differs additionally in the setose condition of the armature elements on the inner margin of the second endopodal segment of leg 4; these elements are flanged spines in *E. longicauda*. The generic placement of *E. verecunda* needs verification since, according to Humes (1992), this species displays a 2-segmented condition of leg 5. This is a characteristic of the genus *Ancheuryte* Herbst, 1989, whereas in *Euryte* leg 5 is 3-segmented in both sexes.

In *E. pseudorobusta* Vervoort, 1964 two distal setae are present in the outer margin of the antennary coxobasis, whereas there is only 1 seta in *E. longicauda*. The proximal spine on the distal coxal endite of maxilla has a different armature in the two species, with a row of setules along each side in *E. pseudorobusta*. Finally, the caudal rami of *E. pseudorobusta* are short, about as long as the anal segment, and differ significantly from the elongate caudal rami of *E. longicauda*.

Two other species, each described from a single female from the Addu Atoll (Maldives), viz. *E. brevicauda* Sewell, 1949 and *E. sewelli* Vervoort, 1964 (= '*Euryte* sp.' of Sewell, 1949) also differ from *E. longicauda* in their very short caudal rami. The status of *E. sewelli* Vervoort, 1964 as a distinct species from *E. brevicauda* is equivocal (Vervoort, 1964; Sewell 1949). The main difference between them is the apparently 18-segmented female antennule in the former species. Unfortunately, Sewell's (1949) original material is not preserved, thus precluding verification. However, if Sewell's illustrations are accurate, the 18-segmented antennule, combined with the absence of the inner seta on the proximal segment of exopod of leg 1, can be used as diagnostic characters of this taxon.

*E. longicauda* differs from *E. bellatula* Humes, 1991 in the nature of the two armature elements on the inner margin of the second endopodal segment of leg 4; these are flanged spines in the former species, whereas in the latter they are setae. *E. bellatula* also has the proximal spine on the distal coxal endite of maxilla armed with a row of thin spinules on both sides. As commented above for *E. verecunda*, the generic placement of *E. bellatula* must be confirmed due to the apparently 2-segmented condition of the leg 5. The association with corals of the two taxa described by Humes is similar to the life-style of *Ancheuryte*, a closely related genus characterized by its 2-segmented leg 5.

The status of *E. similis* Scott, 1912, originally described from the South Orkneys and never found since, is debatable. Scott pointed out its similarity to *E. robusta*, and that it appeared '... to differ in one or two minor points, such as in the armature of the first and fourth pairs of thoracic legs and in the proportional lengths of the abdominal segments' (Scott, 1912). The differences in the armature of the swimming legs mentioned by Scott in the text do not correspond with his figures. Also, as Sewell (1949) already pointed out, it seems certain that Scott had confused the legs so that his second leg is in reality the fourth, and his fourth leg is either the second or third. In fact, the original description is very superficial and does not permit any conclusion other than that the taxon belongs to *Euryte*. The only apparent diagnostic features displayed by this taxon could be the lack of an inner seta on both the first endopodal and first exopodal segments of leg 4 (Scott's leg 2). This is unreliable, however, since the number of armature elements on the swimming legs is a very conservative character at the generic level in the Cyclopidae. In our opinion, given the lack of type material, this taxon should be considered *species inquirendum*.

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# Studies on the deep-sea Protobranchia (Bivalvia): the family Neilonellidae and the family Nuculanidae.

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**SYNOPSIS.** The morphology and distribution of four deep-water protobranch species of the family Neilonellidae and ten deep-water protobranch species of the family Nuculanidae are described. These include four new species. The evolution of the nuculanoid siphonate form is discussed.

## INTRODUCTION

This is the ninth paper in a series on the deep-sea species of Protobranchia of the Atlantic. While a few undescribed species of families reported upon in earlier papers have been found in later samples and remain to be described, this paper is the last of our major descriptive accounts of the morphologies of this exceptionally important group of deep sea bivalves.

In this paper we give an account of a number of nuculanacean species belonging to the families Neilonellidae and Nuculanidae present in our collections from the deep Atlantic. Related species already described are referred to only when essential for descriptive and comparative purposes. Unlike previous papers in this series most of the species described below are known, but only from their shell characters. This is in part due to the fact that many species come from upper-slope depths, and thus more likely to have been sampled

in the past, and in part due to the fact that they are relatively large and thus less likely to have been lost due to the coarseness of nets used in earlier expeditions. Many earlier descriptions of the shell are far from adequate and new descriptions are given.

We have followed the methods and approach of earlier papers in this series (Allen and Hannah, 1989; Allen and Sanders, 1973, 1982; Allen, Sanders and Hannah, 1995; Rhind and Allen, 1992; Sanders and Allen, 1973, 1977, 1985). From stereoscopic microscope examination of shell features and whole mounts of the body stained in haemotoxylin, from dissected specimens and serial sections stained with irichrome techniques, we give detailed descriptions of the shell and internal morphology of key species. Descriptions of related species are limited to points of difference and importance.

Much of the material was taken by ourselves on numerous expeditions by research vessels of the Woods Hole Oceanographic Institution. These include the 'Atlantis', 'Atlantis II', 'Chain' and 'Knorr'. Other samples were taken by JAA on the British research

vessels 'Sarsia' and 'Discovery'. In addition to these, deep-sea bivalves from a series of French expeditions using the research vessels 'La Perle', 'Cryos' and 'Jean Charcot' were kindly donated by our French colleagues. At present this material, other than the types of new species and cited specimens, is housed at the Woods Hole Oceanographic Institution and the University Marine Biological Station Millport. On completion of our studies, it is our intention to return the French collections to the Museum National d'Histoire Naturelle, Paris and our own collections to the U.S. National Museum and the Natural History Museum, London.

Museums are abbreviated as follows:- ANSP – Academy of Natural Sciences, Philadelphia; BMNH – Natural History Museum, London; MCZ – Museum of Comparative Zoology, Harvard; MNHN – Museum National d'Histoire Naturelle, Paris; SAM – South Africa Museum; USNM – United States National Museum, Washington, D.C.; ZMHU – Zoologisches Museum, Humboldt-Universität, Berlin; ZMUC – Zoological Museum, University of Copenhagen.

Sampling gear is abbreviated as follows:- AD – anchor dredge; CP – beam trawl; CV – Blake trawl; DS – epibenthic sledge Sanders (COB); ES – epibenthic sledge (WHOI); KG – USNEL boxcorer; OS – epibenthic sledge (SMBA); WS – epibenthic sledge (IOS).

## ABBREVIATIONS TO THE TEXT-FIGURES

AA	anterior adductor muscle
AN	anus
AR	anterior pedal retractor muscle
AS	anterior sense organ
BG	'byssal' gland
CG	cerebral ganglion
CM	caecum
CV	cerebro-visceral commissure
DD	digestive diverticula
DH	dorsal hood
DT	digestive ducts
FA	feeding aperture
FG	fused gill tissue
FR	faecal rod
FT	foot
GA	gill axis
GI	gill
GP	gill plate
GS	gastric shield
HG	hind gut
HP	hinge plate
HT	hinge teeth
LI	internal ligament or resilium
LO	opisthodontic external ligament
MG	combined mid gut
MM	mantle margin
OE	oesophagus
PA	posterior adductor muscle
PE	sensory papilla
PG	pedal ganglion
PL	palp
PP	palp proboscis
PR	posterior pedal retractor muscle
PS	posterior sorting area
SE	siphonal embayment
SP	combined siphon

SR	siphonal ridge
SS	style sac
ST	sensory tentacle
SM	stomach
VG	visceral ganglion

## Family Neilonellidae Schileyko 1989

Shell usually robust, inflated, moderately large, not particularly elongate, more or less ovate, shell posteriorly extended to some degree and may be acutely rounded or carinate, usually marked concentric sculpture; umbo prominent, anterior to midline; hinge teeth taxodont, continuous or interrupted below the umbo; ligament external, sometimes with an internal component; siphons present; hind gut either a single loop or a series of loops and coils to the left and right of the body.

The family comprises four genera, *Neilonella*, *Pseudotindaria*, *Neilo* and *Protonucula* (Allen and Hannah, 1986). By reason of their external ligament, the neilonellids in the past have been included in the family Mallettiidae, however this family was shown to be an assemblage of unrelated taxa and is now much restricted (Sanders and Allen, 1985). The robust, ovate, concentrically ridged shells of the neilonellids are in marked contrast to the thin, translucent, elongate shells, lacking any marked concentric sculpture, of the mallettiids.

### Genus *NEILONELLA* Dall 1881

TYPE SPECIES. *Leda (Neilonella) corpulenta* Dall 1881; OD.

Shell moderately robust, relatively wide, strong concentric sculpture, anterior margin rounded, posterior margin slightly attenuate and may be acutely rounded, not carinate; umbo prominent, anterior to the midline; hinge plate moderately strong, many chevron teeth, small gap between anterior and posterior series; external ligament opisthodontic, inserting in an external gutter in each valve, with minute internal resilifer, siphonal embayment shallow; hind gut single loop to the right of the body.

### Genus *PSEUDOTINDARIA* Sanders and Allen 1977

TYPE SPECIES. *Pseudotindaria erebus* (Clarke 1959); OD.

Shell robust, relatively wide, anteriorly and posteriorly rounded, not carinate, strong concentric sculpture; hinge plate strong with numerous chevron teeth continuous below umbo; ligament external, anterior part short, posterior part elongate; hind gut with complex series of loops and coils to the right and left of the body.

### Genus *NEILO* Adams 1854

TYPE SPECIES. *Neilo cumingii* Adams 1854; by monotypy.

Shell robust, relatively wide, moderately elongate and posteriorly extended, postero-dorsal margin straight or slightly concave, carinate, posteriorly truncate or slightly rostrate, postero-ventral margin may be somewhat sinuous, two rounded radial ridges from umbo to posterior margin, concentric sculpture; hinge plate well-developed; numerous chevron teeth, anterior and posterior series separate and may differ in size (anterior larger); ligament external.

Genus *PROTONUCULA* Cotton 1930

TYPE SPECIES. *Protonucula verconis* Cotton 1930; OD.

Shell relatively fragile, oval, compressed, glossy, concentric sculpture; umbo prominent; hinge plate narrow, chevron teeth squat, broad, continuous; ligament relatively small, external, amphidetic.

***Neilonella salicensis*** (Seguenza 1877)

TYPE SPECIMEN. Whereabouts unknown.

TYPE LOCALITY. Upper Pliocene fossil, southern Italy.

CITED SPECIMENS. BMNH 1995051.

*Leda pusio* var. *laticostata* Jeffreys 1876a, p.190, nom. nud.; 1876b, p.430.

*Leda pusio* Jeffreys 1879, p.578, in part.

*Leda pusio* var. *salicensis* Seguenza 1877, p.1178, pl.4, fig.20.

*Yoldia sericea* var. *striolata* Verrill 1884, p.226; 1885, p.576.

*Leda salicensis* Locard 1898, p.348, pl.14, figs 22–25.

*Neilonella subovata* Verrill and Bush 1897, p.57, figs 7, 8, 22; type locality: Georges Bank to Cape Hatteras, 125–1731 fms; syntype: USNM 34826 (examined by HLS).

*Tindaria* (*Neilonella*) *guineensis* Thiele 1931, in Theile and Jaekel 1931, p.50, pl.3, fig.70; type locality: Gulf of Guinea, 710–2492m; type: ZMHU (not seen).

*Neilonella guineensis* Knudsen 1970, p.59, text-fig.39, pl.5, fig.3, pl.6, figs 8–10.

*Neilonella schepmani* Prashad 1932, p.27, pl.1, figs 50, 51; type locality: Banda Sea, Siboga sta.221, 2798m; ZMHU (not seen).

*Pseudoneilonella salicensis* Laghi 1986, p.191, text fig.1, pl.5, figs 1–7.

*Pseudoneilonella laticostata* Laghi 1986, p.192, text fig.1, pl.7, figs 7a–c.

*Pseudoneilonella salicensis atlantica* Laghi 1986 p.192, text fig.1, pl.6, figs 1–7.

*Pseudoneilonella montanaroe* Laghi 1986, p.193, text fig.1, pl.9, figs 4–8; type locality: off Portugal, Porcupine Expedition, 400m, USNM No. 199739 (not seen).

*Neilonella striolata* (Brugnone) Warén 1989, p.252, figs 3a, 16a–d.

Warén (1989) gives a lengthy and detailed discussion of much of the nomenclatorial history of this species although, the history is even more complex than he reports. Warén (1989) synonymized *N. salicensis* with an earlier described species *N. striolata* (Brugnone, 1876). The type of *N. striolata* is lost but from the original description and figure by Brugnone (1876) and those of Seguenza (1877) (who synonymized it with *Yoldia abyssicola* Torrell) and later authorities (e.g. Jeffreys, 1879; Locard, 1898) it is clear that the posterior part of the shell of *N. striolata* is significantly different in form to that of *N. salicensis*. Whereas the posterior shell margin of *N. salicensis* is acutely rounded that of *N. striolata* is sub-rostrate with the posterior limit of the shell more ventral in position and the distal postero-dorsal margin more steeply angled (Figs 1.2 & 4).

The brief description of *N. guineensis* by Theile in Theile and Jaekel (1931) was enlarged upon by Knudsen (1970) who compared specimens from the Galathea Expedition with those of the Valdivia and Siboga Expeditions. From our examination of the Galathea material from 02°17'S 08°10'E (2770m) and our own specimens from 02°32'S 08°18'E (2514m) we conclude that *N. guineensis* and *N. salicensis* are the same species. Knudsen (1970) also synonymized *N. schepmani* with *N. guineensis*. Although *N. schepmani* is from the Banda Sea (Prashad, 1932) which might cast doubt on this synonymy, it must be remembered that *N. salicensis* is a very widespread species and we believe that it is comparable to

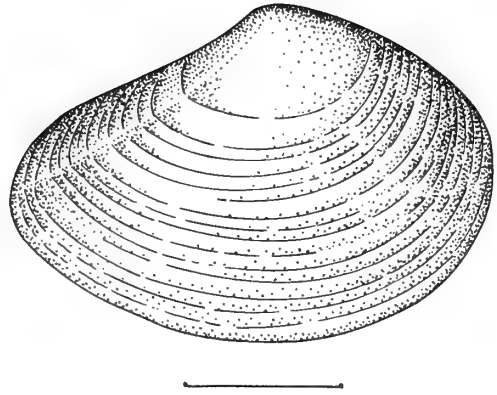


Fig. 1. '*Yoldia abyssicola*' Torell. Right valve, redrawn from Seguenza (1877), synonymized with *Leda producta* Monterosato and *Yoldia striolata* Brugnone. Scale = 1mm.

*Ledella sublevis*, which also has a widespread distribution in the Atlantic and extends into the southwest Pacific (Allen and Hannah, 1989).

Further complexity in the synonymy of *N. salicensis* concerns so-called varieties of '*Leda pusio* Philippi' a species that has recently been investigated by Laghi (1986).

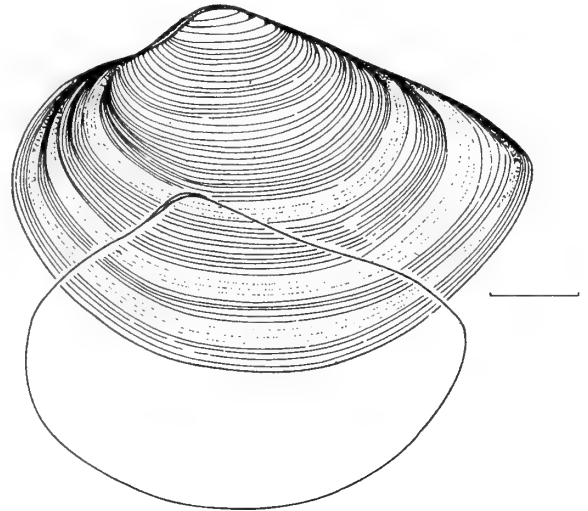


Fig. 2. *Neilonella salicensis*. a, view of shell, from Atlantis II station 73 in the North America Basin, drawn from the left side; b, outline of shell from Biogas VI station DS 86 in the West European Basin to show difference in shape. Scale = 1mm.

*L. pusio* is rostrate and not a neilonellid, but a ledellid, possibly synonymous with *L. acuminata* (Jeffreys 1870) (Laghi, 1986). Jeffreys (1879) reporting on *Leda pusio* taken by the 'Lightning' and 'Porcupine' Expeditions, refers to what he thought was a variety of this species which he had previously named *L. pusio* var. *laticostata* (Jeffreys, 1876) and which he changed to *L. pusio* var. *salicensis* of Seguenza (1877). Laghi (1986) proposed a new genus *Pseudoneilonella* to accommodate this latter and raised to species level a number of the records of Jeffreys (1876, 1879). Since then, Warén (1989) has synonymized these with *Neilonella salicensis* and this we confirm.

Jeffreys (1879) also listed a further variety which he called *semistriata* and which is now regarded as a separate species (Warén, 1989). In his detailed analysis Warén (1989, figs 17E & F) also

figures what he describes as '*Neilonella latior* (Jeffreys)?', young syntype of *Leda sericea* Jeffreys, *Valorous* Expedition, Stn 12, USNM 199595? These valves are equilateral in shape and differ in outline from *N. salicensis*. We have specimens of this species in our collections and these we intend to describe in a later paper.

Locard (1898), not mentioned by Warén (1989), came to the same conclusion as ourselves and recognized three entities, *Leda pusio*, *L. striolata* and *L. salicensis*, accurately separating them on shell characters, the most significant of these being the more elongate, triangular form and greater post-umbonal length of *L. salicensis* as compared with *L. pusio*, both of which he figures.

There is a relatively wide degree of variation in the height, length and post-umbonal length of the shell in *N. salicensis* and this variation is similar in populations from different basins (Figs 4 & 5). This may, in part, explain why this species has been described anew so many times.

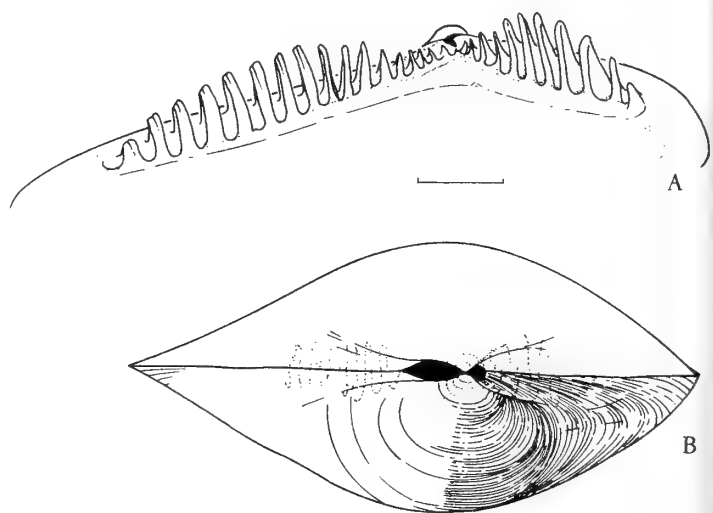
#### MATERIAL:

Cruise	Sta	Depth (m)	No	Lat	Long	Date	Gear
<b>NORTH AMERICA BASIN</b>							
AtlantisII 12	62	2496	13	39°26.0'N	70°33.0'W	21.8.64	ES
	64	2886	2	38°46.0'N	70°06.0'W	21.8.64	ES
	73	1330-1470	495	39°46.5'N	70°43.3'W	25.8.64	ES
Chain 50	87	1102	6	39°48.7'N	70°40.8'W	6.7.65	ES
Chain 58	103	2022	217	39°43.6'N	70°37.4'W	4.5.66	ES
AtlantisII 30	131	2178	119	39°38.5'N	70°36.5'W	18.12.66	ES
Chain 88	210	2024-2064	48	39°43.0'N	70°55.5'W	23.2.69	ES
<b>GUYANA BASIN</b>							
Knorr 25	293	1456-1518	2	08°58.0'N	54°04.3'W	27.2.72	ES
	295	1000-1022	1575	08°04.2'N	54°21.3'W	28.2.72	ES
	297	508-523	194	07°45.3'N	54°24.0'W	28.2.72	ES
	299	1942-2076	1	07°55.1'N	55°42.0'W	29.2.72	ES
	307	3835-3862	2	12°34.4'N	58°59.3'W	3.3.72	ES
<b>WEST EUROPEAN BASIN</b>							
Sarsia	S33	1537-1830	6	43°41.0'N	03°36.0'W	13.7.67	ES
	S37	1739	2	43°39.2'N	03°30.2'W	15.7.67	ES
	S40	860	4	43°35.6'N	03°24.8'W	15.7.67	ES
	S44	1739	60	43°40.8'N	03°35.2'W	16.7.67	ES
	S50	2379	16	43°46.7'N	03°38.0'W	18.7.67	ES
	S65	1922	2	46°15.0'N	04°50.0'W	25.7.67	ES
Chain 106	313	1491-1500	432	51°32.2'N	12°35.9'W	17.8.72	ES
	316	2173-2209	583	50°58.7'N	13°01.6'W	18.8.72	ES
La Perle	DS11	2205	1	47°35.5'N	08°33.7'W	8.8.72	DS
Biogas I	DS25	2096	9	44°08.2'N	04°15.0'W	1.11.72	DS
Polygas	DS32	2138	2	47°32.2'N	08°05.3'W	19.4.73	DS
Biogas II	DS38	2138	2	47°32.2'N	08°35.8'W	25.8.73	DS
Biogas III	DS49	1845	70	44°05.9'N	04°15.6'W	31.8.73	DS
	DS52	2006	16	44°06.3'N	04°22.4'W	18.2.74	DS
Biogas IV	DS63	2126	5	47°32.8'N	08°35.0'W	26.2.74	DS
Cryos	CP07	2170	5	44°09.8'N	04°16.4'W	21.6.74	CP
Biogas V	DS77	4240	1	47°31.8'N	09°34.6'W	24.10.74	DS
J.Charcot	DS80	4720	4	46°29.5'N	10°29.5'W	26.10.74	DS
Biogas VI	DS82	4462	1	44°25.4'N	04°50.8'W	29.10.74	DS
	DS86	1950	198	44°04.8'N	04°18.7'W	31.10.74	DS

Incal	DS87	1913	173	44°05.2'N	04°19.4'W	1.11.74	DS
	DS88	1894	31	44°05.2'N	04°15.7'W	1.11.74	DS
	DS01	2091	98	57°59.2'N	10°41.3'W	15.7.76	DS
	DS02	2081	91	57°58.2'N	10°48.5'W	16.7.76	DS
	CP01	2068	19	57°57.7'N	10°55.0'W	16.7.76	CP
	DS05	2503	3	56°28.1'N	11°11.7'W	18.7.76	DS
	OS01	2634	4	50°14.4'N	13°10.9'W	30.7.76	OS
	WS02	2498	1	50°19.3'N	12°55.8'W	30.7.76	WS
	CP10	4823	2	48°25.5'N	15°10.7'W	31.7.76	CP
	WS03	4829	1	48°19.2'N	15°23.3'W	1.8.76	WS
Challenger 7/83	232	2195	3	57°17.0'N	10°16.0'W	19.5.83	ES
<b>CANARY BASIN</b>							
Discovery	6701	1934	8	27°45.2'N	14°13.0'W	16.3.68	ES
<b>SIERRE LEONE BASIN</b>							
AtlantisII 31	142	1624-1796	20	10°30.0'N	17°51.5'W	5.2.67	ES
	147	2934	16	10°38.0'N	17°52.0'W	23.2.67	ES
<b>ANGOLA BASIN</b>							
AtlantisII 42	191	1546-1559	52	23°05.0'S	12°31.5'E	17.5.68	ES
	200	2644-2754	7	09°43.5'S	10°57.0'E	22.5.68	ES
	201	1964-2031	41	09°25.0'S	11°35.0'E	23.5.68	ES
	202	1427-1643	21	08°56.0'S	12°15.0'E	15.5.68	ES
Walda	DS20	2514	4	02°32.0'S	08°18.1'E	--.71	DS
	DS25	2470	31	02°19.8'S	07°49.2'E	--.71	DS

In addition to the material listed above we have examined the following specimens from the Porcupine and Valorous Expeditions housed in the BMNH: -85.11.5.397-402; 6.9.27,28; 85.11.5.494-5; 89.11.11.10-13; 89.9.5.26-9; 77.11.28.24. We have also examined specimens from the Galathea Expedition housed in the ZMUC, and specimens mostly from the U.S. Fish Commission expeditions housed in the USNM and MCZ.

**DISTRIBUTION.** Common throughout the Atlantic north of the Argentine and Cape Basins and south of the Norwegian and Arctic Basins. It is also found in the southwestern Pacific. It mainly occurs at slope depths, occasionally at abyssal depths (West European Basin), from 508m to 4829m.



**Fig. 3.** *Neilonella salicensis*. **a**, internal inclined ventral view of the hinge plate and teeth of a left valve; **b**, dorsal view of shell to show the extent of the external ligament and detail of sculpture. Both specimens taken from Walda station DS 25 in the Angola Basin. Scale = 1mm.

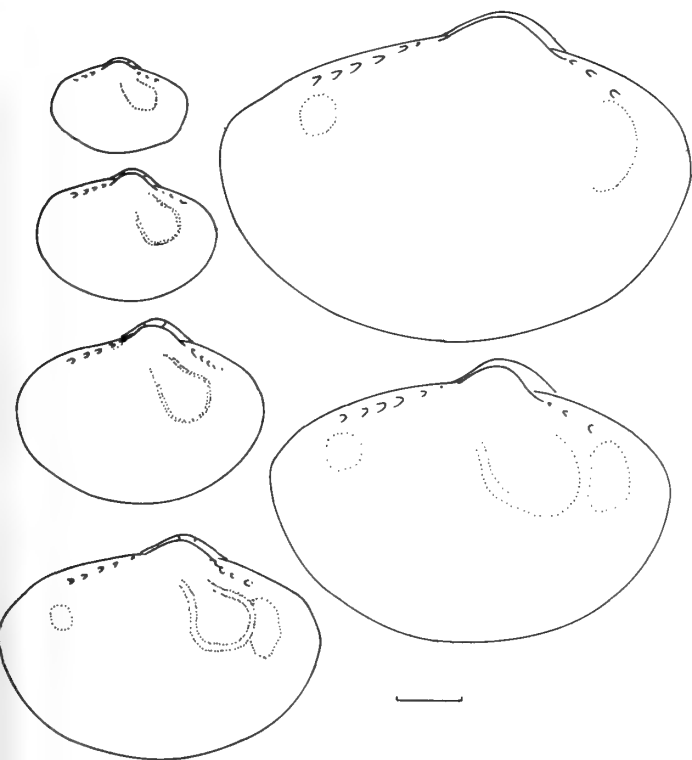


Fig. 4. *Neilonella salicensis*. A series of shells, from Atlantis II station 73 from the North Atlantic Basin, in lateral view from right side to show change in shape with increasing size. Scale = 1mm.

#### SHELL DESCRIPTION (Figs 2-4)

Shell robust, moderately elongate, moderately wide, ornamented

with concentric ridges sometimes in groups of three to four well-defined ridges alternating with two to three slightly less well-defined ridges slightly wider spaced than the former, light to dark straw-coloured periostracum; umbos moderately prominent, somewhat anterior to mid-line (postumbonal length *c.* 54% of total length), inward and slightly anterior facing; dorsal shell margins deep set close to umbo but no lunule or true escutcheon, antero-dorsal margin slopes down to join anterior and ventral margins in smooth curve, postero-dorsal margin almost straight, may be slightly angled at posterior limit of hinge short of acutely rounded posterior margin; posterior limit of shell somewhat dorsal to mid horizontal line, anterior limit of shell at mid line; posterior margin of smaller specimens less acutely rounded; hinge plates elongate, moderately broad, increasing slightly in width distally, anterior and posterior hinge teeth series meet below umbo with no edentulous space between, teeth numerous, up to 19 in posterior series and 14 in anterior series depending on size of specimen, obtuse chevron-shape, proximal hinge teeth very small, those more distal elongate and so closely articulated that it may be impossible to separate valves without damage to them; internal ligament (resilium) microscopic, present ventral to umbo at outer margin of hinge plate, external ligament opisthodontic, relatively short, short anterior part present. Maximum total length of present specimens 8.0mm.

As in most deep-sea protobranchs, there is a change in the shape of the shell outline as growth occurs (Figs 4 & 5), although this is not as marked as in many protobranch species (e.g. *Ledella* spp. Allen and Hannah, 1989). The ratio of postero-umbonal length to total length increases and at the same time the height to total length decreases so that larger shells (>3mm total length) are more posteriorly attenuate than smaller specimens. There is a fair degree of variation in the ratios which is not related to the different populations sampled. Unusually in some samples the length frequency histograms are skewed to the right (Fig. 6) and with clearly

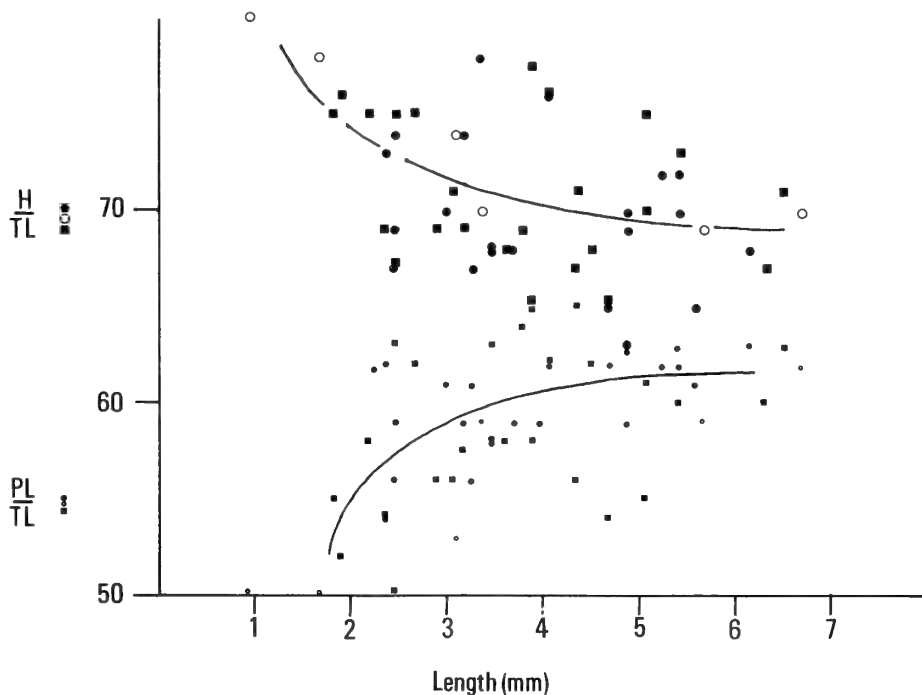


Fig. 5. *Neilonella salicensis*. The ratios height to total length (H/TL)(large symbols) and post-umbonal length to total length (PL/TL)(small symbols) plotted against total length to show changes in shell proportions with increasing length. Open circles are specimens from Biogas III station DS 49 West European Basin, closed circles from Sarsia Station S44 West European Basin and closed squares, from Chain 58 station 103 North America Basin.

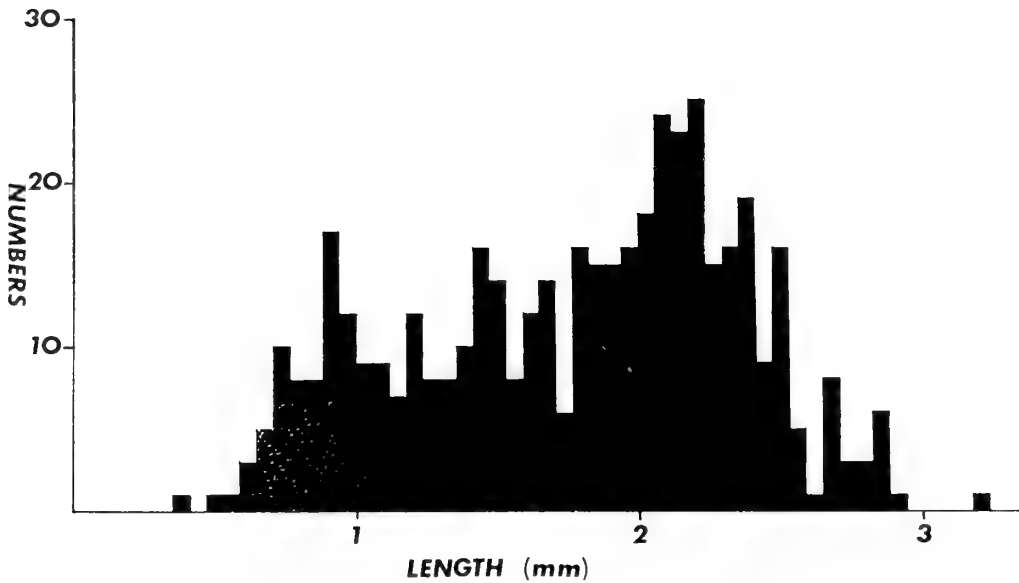


Fig. 6. *Neilonella salicensis*. Length/frequency histogram of a sample taken from Atlantis II station 73 from the North America Basin.

marked size groups. It is unclear whether these latter relate to annual settlements or to fortuitous periodic settlements.

#### INTERNAL MORPHOLOGY (Figs 7 & 8)

This was one of the few deep-sea species that were kept alive for a few weeks following the processing of the sample. The external drab olive/yellow of the shell is in marked contrast to the spectacularly vivid array of colours of the body organs. The stomach is a bright emerald green, the digestive gland a bright olive green and the gills are bright red. The gill plates are particularly vivid below the frontal cilia and dorso-laterally, in addition the mantle overlying the gills is also red. The margins of the palps are a diffuse pink as too is a band underlying the crest of the inner folds. The red and pink pigments are probably a cytochrome. The blood is not red in colour and probably contains a haemocyanin as do other protobranchs (Taylor, Davenport and Allen, 1995). Elsewhere the palps are pale yellow and the yellow/pink effect is in marked contrast to the palp proboscides which are pure white.

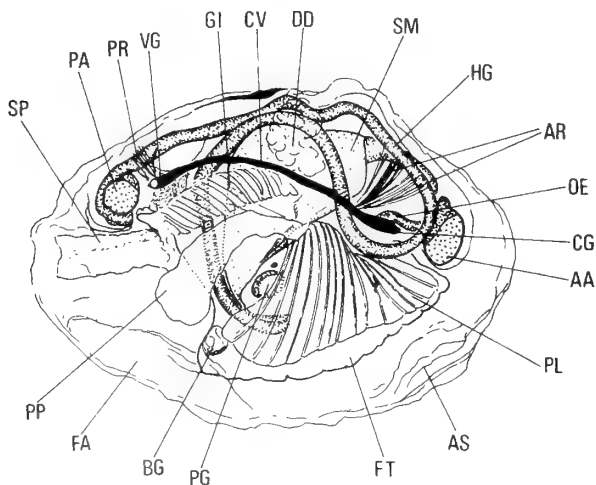


Fig. 7. *Neilonella salicensis*. Semidiagrammatic view of the internal morphology as seen from the right side. See abbreviations to text-figures on p. 102. Scale = 1mm.

The mantle has three marginal folds of which the inner is fused at the posterior end to form the siphon. Fusion is minimal, restricted to the dorsal margin of the siphon (Fig. 8). Ventrally the siphonal margins are unfused as too are the longitudinal ridges that mark the division between the dorsal exhalent and ventral inhalent channels. The gill axes attach to the inner ends of the longitudinal ridges. There is a shallow siphonal embayment and attached to its inner limit on the left or the right side is a siphonal tentacle which in living specimens extends beyond the shell margin at distance equal to a third of the shell length. Ventral to the anterior adductor muscle there is an elongate anterior sense organ, which consists of a flap of tissue derived from the middle sensory fold. A major ciliated rejection tract is present on the inner face of the inner mantle fold. The adductor muscles are relatively small, unequal in size, with the posterior the smaller. The posterior muscle is round in cross-section while the anterior is bean-shaped.

The gills, with up to 26 plates, are slung from a horizontal axis extending from the base of the siphon to a point about half way across the body. Individual gill plates are exceptionally elongate for a deep-sea protobranch. The foot is broad and the divided sole has papillate margins, the anterior papillae are the more pronounced. There is a small median papilla present posterior to the aperture to the 'byssal' gland. The latter is extremely large. The pedal muscles consist of a pair of broad posterior retractors, one on each side of the hindgut anterior to the posterior adductor, and a series of four pairs of anterior retractors posterior to the anterior adductor. The two inner muscles of the four lie more close together than to the two outer.

The palps are relatively large with up to 27 inner ridges depending on the size of the specimen. Each bears a long narrow palp proboscis. A ciliated rejection tract traverses the lateral face of the foot at the junction of of the muscular and visceral parts and the posterior ventral point of the palp is positioned at the posterior limit of this tract in the living animal.

The mouth is set far posterior to the anterior adductor. The oesophagus curves first antero-dorsally to meet the inner face of the anterior adductor then postero-dorsally to open on the anterior face of the stomach. The stomach is large and slung diagonally within the visceral mass. The pedal muscles form a ventral 'basket' in which the stomach rests. The stomach, which is similar to that of other

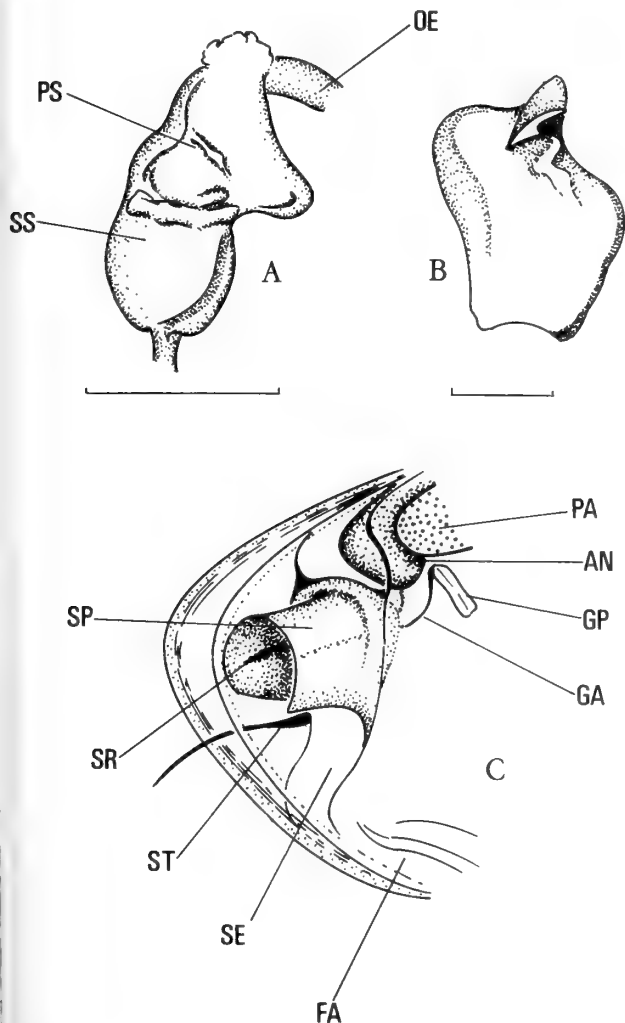


Fig. 8. *Neilonella salicensis*. Detail of internal morphology; a, lateral view from the right side of the stomach and style sac; b, internal view of a dissected gastric shield; c, diagrammatic enlargement of the siphonal region to show the relationship of the parts. See abbreviations to the text-figures p. 102. Scales = 0.5mm (A); 0.1mm (B).

protobranchs (Purchon, 1956), is one third lined with a gastric shield dorsally and to the left (Fig. 8). The unlined right wall bears very few sorting ridges. The style sac is flask-shaped rather than the more usual conical outline of other protobranchs. The combined mid gut is guarded by minor and major typhlosoles. The hind gut penetrates the foot to a point ventral to the pedal ganglia before taking a course posterior to the stomach and thence to form a loop to the right side of the body. This single loop extends anteriorly as far as the anterior adductor muscle and where it then takes a dorsal course along the margin of the body to the anus. The hind gut has a particularly wide diameter compared with most other deep-sea protobranchs and has a well-marked typhlosole present along its entire length. The digestive gland which lies anterior and dorso-lateral to the stomach has three sections each connecting with the stomach via a duct. The apertures of the ducts are ventral, and left latero-ventral to the oesophageal aperture.

The nervous system is of a typical protobranch design with elongate pairs of cerebral and visceral ganglia and large ovoid pedal ganglia. These are connected by very stout cerebro-visceral and cerebro-pedal commissures. A statocyst lies dorsal each pedal gan-

glion. From visceral ganglia three main nerves pass to the gill, hind gut and mantle/siphon, and from the cerebral ganglia nerves pass to the palps and mantle.

The sexes are separate, specimens greater than 4mm total length have gonads. The gonad first develops at the periphery of the visceral mass and as it matures it spreads across the lateral surface of the viscera. On the right side of the body, apart from a small portion postero-dorsal to the loop of the hind gut the gonad does not extend beyond the confines of the of the loop. While different samples show different degrees of maturity, the specimens of each sample appear to be maturing synchronously. The ratio of males to females is approximately 50:50.

### *Neilonella hampsoni* new species

TYPE SPECIMEN. Holotype BMNH 1995049; paratype BMNH 1995050.

TYPE LOCALITY. Atlantis II cruise 31, station 155, Sierra Leone Basin, 00°03.0'S 27°48.0'W, 3730–3783m.

#### MATERIAL:

Cruise	Sta	Depth (m)	No	Lat	Long	Date	Gear
SIERRA LEONE BASIN							
Atlantis 11	155	3730–3783	3	00°03.0'S	27°48.0'W	14.2.67	ES
J. Charcot	DS03	5150	1	10°59.0'N	45°15.0'W	16.11.77	DS Vema

DISTRIBUTION. An abyssal species, to date only found in the Sierra Leone Basin at depths from 3730m to 5150m.

#### SHELL DESCRIPTION (Figs 9 & 10)

Shell robust, oval, wide, ornamented with concentric ridges, straw-coloured periostracum; umbo prominent, anterior to midline (postumbonal length c.60% of total length in large specimens), inward facing; lunule and escutcheon, both well-defined; postero-dorsal margin in lateral view almost straight, slopes relatively steeply from umbo to proximal limit of hinge, rounded angle to posterior margin, anterior and ventral margins form a smooth curve, distally antero-dorsal margin slightly concave then slopes steeply in smooth curve to anterior margin; anterior limit of shell coincides

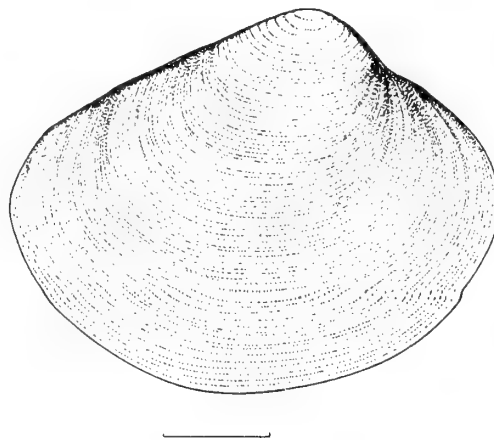


Fig. 9. *Neilonella hampsoni*. Lateral view of the right valve of the holotype, from Atlantis II station 155 from the Sierra Leone Basin. Scale = 1mm.

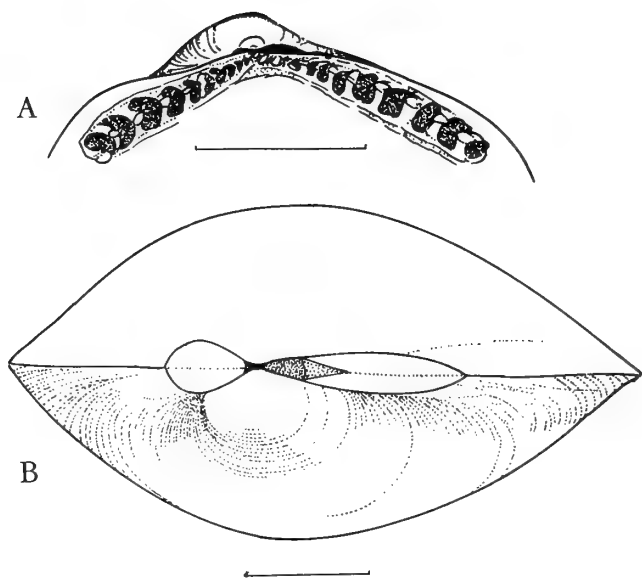


Fig. 10. *Neilonella hampsoni*. a, lateral view of the hinge plate of the right valve of a paratype; b, dorsal view of the shell of the holotype. Both specimens taken from Atlantis II station 155 from the Sierra Leone Basin. Scales = 1mm.

with horizontal midline, posterior limit of shell immediately ventral to mid line; hinge plate stout extending along most of the dorsal shell margin, anterior and posterior tooth series meet below umbo, with faint suggestion of an edentulous space between. 13 posterior and 10 anterior teeth in specimen 3.9mm total length, teeth increase in size distally, teeth chevron-shaped, obtuse, so much so that teeth appear to be a straight line transverse to hinge plate; ligament opisthodontic, external, short, anterior outer layer extremely short, hidden beneath umbo, no resilium.

Maximum total length of present specimens is 7.5mm.

Apart from *N. salicensis*, from which it differs in having a shorter, wider shape, and well-marked lunule and escutcheon, the only other protobranch species with which it has some similarity is '*Leda sericea* var *ovata* Jeffreys 1876 (Jeffreys, 1879). One of us (JAA) has examined specimens of this latter species in the Natural History Museum, London, (BMNH 85 11 5483-84) and find that *L. sericea* is more ovate, with the dorsal margins much less sloping, a more anteriorly positioned umbo and a much more narrow hinge plate.

#### INTERNAL MORPHOLOGY

The morphology is similar to *Neilonella salicensis*. Such differences that do exist include the adductor muscles, both of which are small, similar in size, but with the posterior oval and the anterior round in cross section. The foot is somewhat smaller but with a moderately large byssal gland with a small, hooked, median papilla posterior to its aperture. Except for the posterior quarter of their length, the margins of the divided foot are finely papillate. There are approximately 20 gill plates and 25 ridges on the palps of a specimen 3.9mm total length. The hind gut makes a simple loop to the right side of the body, it has a wide lumen (0.21mm in diameter) with a single well-defined typhlosole running its entire length. The stomach is large and the mouth lies some distance posterior to the inner wall of the anterior adductor. The kidney extends in a narrow band from the posterior margin of the posterior adductor, anteriorly narrowing over the viscera, and terminating at the posterior edge of the digestive diverticula.

It is named after our good friend and colleague George Hampson who accompanied us on so many of our cruises and without whom sampling at abyssal depths would not have been the resounding success that it proved to be.

#### *Neilonella corpulenta* (Dall 1881)

TYPE LOCALITY. Blake station off Havana, 823m (station number not recorded but, only station 51, 23°11.0'N 82°21.0'W, is listed as having a depth of 450fm (823m) (Smith, 1888)).

TYPE SPECIMEN. Holotype, U.S. National Museum 63169. Cited specimen: BMNH 1995048.

*Leda* (*Neilonella*) *corpulenta* Dall, 1881, 125; 1886, 254, pl. 7, figs. 1a, 1b.

*Neilonella* (*N.*) *corpulenta* Laghi 1986, pl.9, figs 1-3.

#### MATERIAL:

Cruise	Sta	Depth (m)	No	Lat	Long	Date	Gear
BRAZIL BASIN							
Atlantis II	162	1493	1	08°02.2'S	34°03.0'W	19.2.67	ES 31

The type specimen has been examined by us.

DISTRIBUTION. An upper slope species previously recorded only from the Gulf of Mexico but here found in the Brazil Basin. It occurs at depths from 347m to 1493m.

#### SHELL DESCRIPTION (Fig. 11)

Dall (1881) gave an adequate description which was later (Dall, 1886) augmented by good internal and external lateral views of the shell. The specimen collected from the Brazil basin differs little from the type (Dall, 1881, 1886)(Fig. 11):-

Shell elongate, solid, oval, ornamented with concentric ridges; umbo not particularly large or elevated, somewhat anterior to the mid line; postero-dorsal margin almost straight, slightly upturned posterior to the distal limit of the hinge, then sharply and smoothly curved to posterior margin, ventral margin shallow smooth curve, not posteriorly sinuous, anterior margin smoothly curved, antero-dorsal margin relatively steeply angled, distally slightly raised; hinge plate elongate, relatively wide, hinge teeth chevron-shaped, 9 in anterior and 12 in posterior series; external ligament slightly opisthodontic, short; resilium small ventral to umbo.

Dall (1881) states that there are an equal number of teeth in the anterior and posterior series (15), however the type specimen which is larger than the present specimen, has 17 anterior and 20 posterior teeth.

The total length of present specimen is 3.1mm.

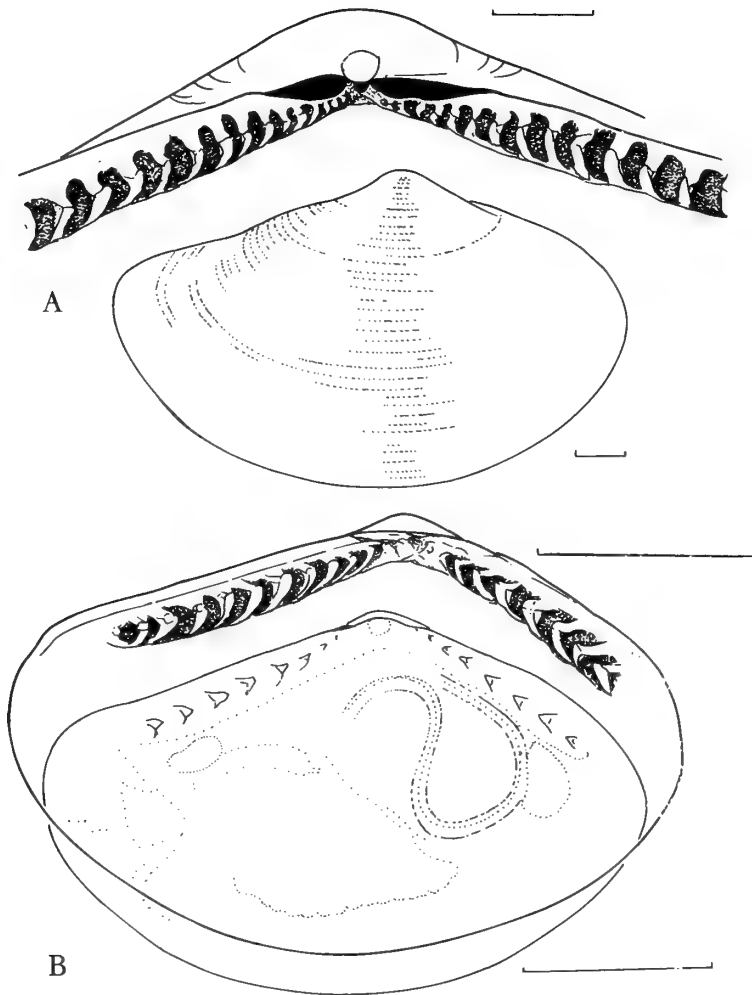
#### INTERNAL MORPHOLOGY

Both adductor muscles are oval in cross section, the anterior is somewhat larger than the posterior but neither is particularly large. The foot is relatively short with large marginal papillae. The palps are relatively short with 7 broad internal ridges. The gill is also short. The siphonal embayment is shallow and the siphons similar to those described for *N. salicensis*. The hind gut forms a single loop to the right side of the body and has a typhlosole along its entire length.

#### *Neilonella whoii* new species

TYPE SPECIMEN. Holotype BMNH 1995052; paratypes BMNH 1995053.





**Fig. 11.** *Neilonella corpulenta*. **a**, an external lateral view of the right valve of specimen No 63169 from the USNM and an internal lateral view of the central region of the hinge of the same valve; **b**, lateral view of the intact shell, from Atlantis II station 162 from the Brazil Basin, and an internal view of the left valve of the same specimen to show detail of the hinge plate. Scales = 1 mm.

CITED SPECIMEN. BMNH 1995054.

TYPE LOCALITY. Chain cruise 50, station 78, North America Basin, 38°00.8'N 69°18.7'W, 3828m.

**MATERIAL:**

Cruise	Sta	Depth (m)	No	Lat	Long	Date	Gear
<b>WEST EUROPEAN BASIN</b>							
Chain 106	323	3356-3338	7	50°08.3'N	13°53.7'W	21.8.72	ES
	326	3859	5	50°04.9'N	14°23.8'W	22.8.72	ES
	328	4426-4435	6	50°04.7'N	15°44.8'W	23.8.72	ES
	330	4632	16	50°43.5'N	17°51.7'W	24.8.72	ES
J.Charcot	DS20	4226	2	47°33.0'N	09°36.7'W	24.10.72	DS
Polygas	DS22	4144	2	47°34.1'N	09°38.4'W	25.10.72	DS
	DS23	4734	18	46°32.8'N	10°21.0'W	26.10.72	DS
	DS28	4413	1	44°23.8'N	04°47.5'W	2.11.72	DS
J.Charcot	CV23	2034	1	47°32.7'N	08°34.2'W	25.8.73	CV
Biogas III	DS41	3548	4	47°28.3'N	09°04.2'W	26.8.73	DS
	CV27	4023	1	47°34.2'N	09°32.4'W	28.8.73	CV
	CV30	4518	53	46°32.8'N	10°20.0'W	28.8.73	CV
	DS47	4230	2	44°26.8'N	04°50.7'W	31.8.73	DS
J.Charcot	CV34	4406	1	44°27.2'N	04°49.1'W	19.2.74	CV

Biogas IV	DS53	4425	30	44°30.4'N	04°56.3'W	19.2.74	DS
	DS54	4659	13	46°31.1'N	10°29.2'W	21.2.74	DS
	DS56	4050	2	47°32.7'N	09°28.2'W	23.2.74	DS
	DS60	3742	2	47°26.8'N	09°07.2'W	24.2.74	DS
	DS66	3480	4	47°28.2'N	09°00.0'W	17.6.74	DS
Cryos							
Biogas V	DS75	3250	4	47°28.1'N	09°07.8'W	22.10.74	DS
J.Charcot	DS76	4228	8	47°34.8'N	09°33.3'W	23.10.74	DS
Biogas VI	CP14	4237	1	47°32.0'N	09°35.9'W	23.10.74	CP
	DS78	4706	29	46°31.2'N	10°23.8'W	25.10.74	DS
	DS79	4715	10	46°30.4'N	10°27.1'W	26.10.74	DS
	CP18	4721	1	46°30.5'N	10°26.0'W	26.10.74	CP
	DS81	4715	1	46°28.3'N	10°24.6'W	27.10.74	DS
	CP19	4434	3	44°24.9'N	04°51.3'W	28.10.74	CP
	DS82	4462	16	44°25.4'N	04°52.8'W	29.10.74	DS
J.Charcot	OS01	2634	3	50°15.2'N	13°11.0'W	30.7.76	OS
Incal	DS11	4823	1	48°18.6'N	15°12.0'W	1.8.76	DS
	CP11	4823	1	48°21.2'N	15°13.7'W	1.8.76	CP
	OS02	4829	6	48°19.1'N	15°15.5'W	2.8.76	OS
	OS05	4296	3	47°32.9'N	09°34.7'W	7.8.76	OS
	OS06	4316	3	47°27.9'N	09°36.0'W	9.8.76	OS
	DS16	4268	1	47°30.3'N	09°33.4'W	9.8.76	DS
	WS09	4277	2	47°27.9'N	09°34.0'W	10.8.76	WS
<b>CANARY BASIN</b>							
Discovery	6711	2988	1	27°14.9'N	15°36.3'W	19.3.68	ES

SIERRA LEONE BASIN						
Atlantis II 31	146	2842-2891	2	10°39.5'N	17°44.5'W	6.2.67 ES
	147	2934	4	10°38.0'N	17°52.0'W	6.2.67 ES
	148	3814-3828	4	10°37.0'N	18°14.0'W	7.2.67 ES
	149	3861	1	10°30.0'N	18°18.0'W	7.2.67 ES
GUINEA BASIN						
J. Charcot Walda	DS28	1261	2	04°21.2'N	04°35.2'E	?8.71 DS
ANGOLA BASIN						
Atlantis II 42	195	3797	46	14°49.0'S	09°56.0'E	19.5.68 ES
	197	4595	25	10°29.0'S	09°04.0'E	21.5.68 ES
	198	4559-4566	20	10°24.0'S	09°09.0'E	21.5.68 ES
	199	3764-3779	3	09°49.0'S	10°33.0'E	22.5.68 ES
	200	2644-2754	8	09°43.5'S	10°57.0'E	22.5.68 ES
NORTH AMERICA BASIN						
Atlantis II 264	2	3752	2	38°05.0'N	69°36.0'W	22.5.61 AD
Atlantis II 12	64	2886	2	38°46.0'N	70°06.0'W	21.8.64 ES
	72	2864	9	38°16.0'N	71°47.0'W	24.8.64 ES
Chain 50	76	2862	3	39°38.3'N	67°57.8'W	29.6.65 ES
	77	3806	752	38°00.7'N	69°16.0'W	30.6.65 ES
	78	3828	199	38°00.8'N	69°18.7'W	30.6.65 ES
	80	4970	1	34°49.8'N	66°34.0'W	2.7.65 ES
	85	3834	1150	37°59.2'N	69°26.2'W	5.7.65 ES
Atlantis II 24	124	4862	1	37°26.0'N	63°59.5'W	22.8.66 ES
	126	3806	48	39°37.0'N	66°47.0'W	24.8.66 ES
Atlantis II 40	175	4667-4693	1	36°36.0'N	68°29.0'W	28.11.67 ES
Chain 106	334	4400	3	40°42.6'N	46°13.8'W	30.8.72 ES
	335	3882-3919	5	40°25.3'N	46°30.0'W	31.8.72 ES
Knorr 35	340	3264-3356	95	38°14.4'N	70°20.3'W	24.11.73 ES
BRAZIL BASIN						
Atlantis II 31	156	3459	6	00°46.0'S	29°28.0'W	14.2.67 ES
GUYANA BASIN						
Knorr 25	287	4980-4934	72	13°16.0'N	54°52.2'W	24.2.72 ES
	288	4417-4429	19	11°02.2'N	55°05.5'W	25.2.72 ES
	291	3859-3868	43	10°06.1'N	55°14.0'W	26.2.72 ES
	301	2487-2500	23	08°12.4'N	55°50.2'W	29.2.72 ES
	303	2842-2853	8	08°28.8'N	56°04.5'W	1.3.72 ES
	307	3862-3835	15	12°34.4'N	58°59.3'W	3.3.72 ES
J. Charcot Biovema	DS05	5100	3	10°46.0'N	42°40.3'W	19.11.77 DS
ARGENTINE BASIN						
Atlantis II 60	242	4382-4402	1	38°16.9'S	51°56.1'W	13.3.71 ES
	243	3815-3822	3	37°36.8'S	52°23.6'W	14.3.71 ES
	247	5208-5223	6	43°33.0'S	48°58.1'W	17.3.71 ES
	256	3906-3917	8	37°40.9'S	52°19.3'W	24.3.71 ES
	259	3305-3317	11	37°13.3'S	52°45.0'W	26.3.71 ES

## SHELL DESCRIPTION (Figs 12 &amp; 13)

Shell robust, ovate, moderately wide, ornamentated with marked concentric ridges, straw-coloured periostracum; umbos prominent, inwardly facing, clearly anterior of vertical midline in specimens >3.0mm, more central in smaller specimens (post-umbonal length *c.* 57% of total length); no clearly marked lunule or escutcheon, but some specimens with faint indications; postero-dorsal margin slight concave curve, very slightly angulate opposite limit of hinge plate then steepening to posterior margin, posterior margin may be somewhat flattened particularly in small specimens, ventral margin moderately curved joining anterior and antero-dorsal margins in a smooth curve, anterior and posterior limits of shell dorsal to mid-horizontal line; hinge plate broad, continuous, elongate, short chevron-shaped hinge teeth, up to 14 in posterior series, 12 in anterior series, numbers varying with size of specimen, edentulous space below umbo very small; external ligament short, opisthodontic, resilium microscopic, close to shell margin, ventral to umbo. Maximum total length of present specimens is 9.8mm.

*Neilonella whoii* most closely resembles *N. salicensis*. It can be separated from the latter species by its more rounded shape and greater height. The posterior shell margin is not as acute and the posterior limit of the shell is more dorsal in position as compared with *N. salicensis*. Furthermore, the post-umbonal length of *N. whoii* is somewhat longer than in *N. salicensis* and the hinge plate is not so broad having a smaller ratio of anterior to posterior teeth.

We name this species in honour of the Woods Hole Oceanographic Institution, through whose auspices these studies were carried out.

## INTERNAL MORPHOLOGY (Figs 14 &amp; 15)

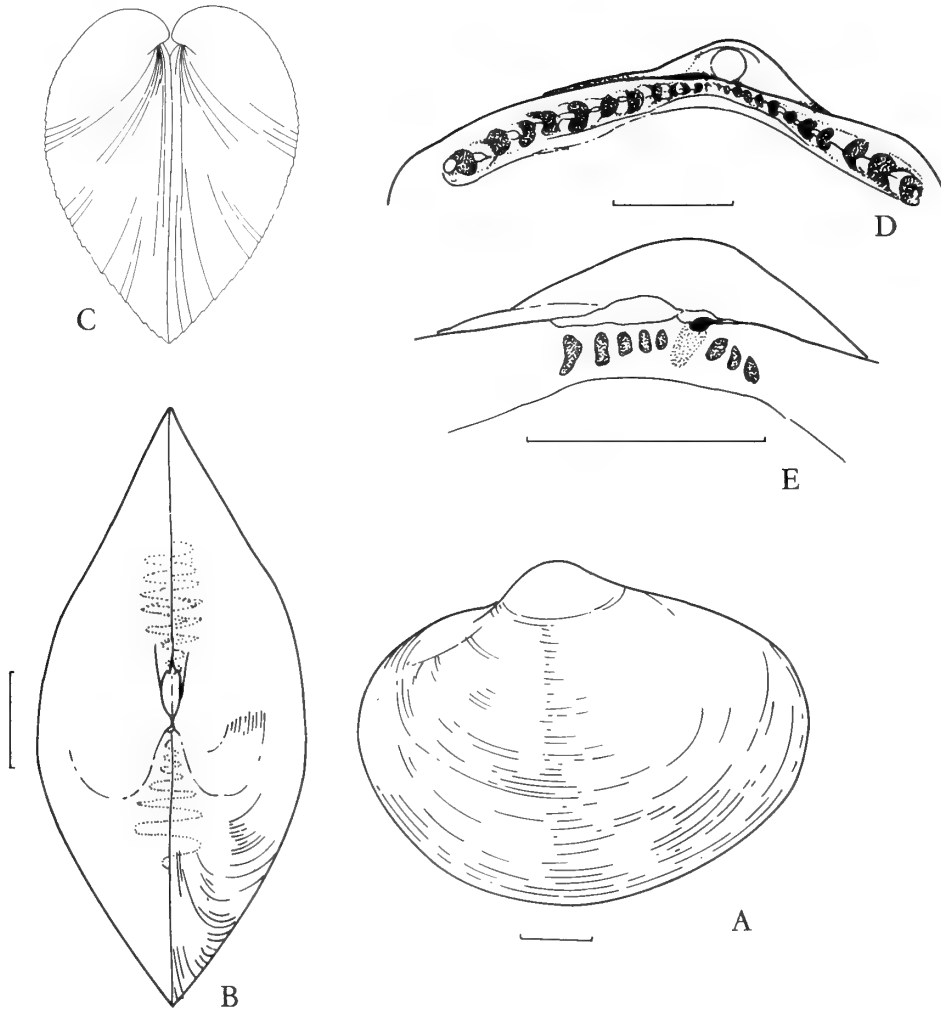
For the most part the morphology of *N. whoii* is similar to that of *N. salicensis*. Of the mantle structures, the construction of the siphon is similar, although the siphonal embayment is less deep than in *N. salicensis*. The posterior adductor muscle is oval in cross-section and not much smaller than the anterior, probably reflecting the more rounded shell outline of *N. whoii*. The anterior sense organ is poorly developed, the least developed of all the deep-sea nulanids that have been described to date. The gills and palps of *N. whoii* are similar in size and form to those in *N. salicensis* with up to a maximum of 18 gill plates and 17 palp ridges.

The foot is large with a few moderately deep papillae at the margin. The mouth lies some distance posterior to the anterior adductor. The stomach is large with 9 or 10 ridges forming the posterior sorting area. The hind gut makes a single loop to the right side of the body, the loop being somewhat larger and more smoothly curved than that in *N. salicensis*. The ganglia and commissures are not so stout as they are in the latter species.

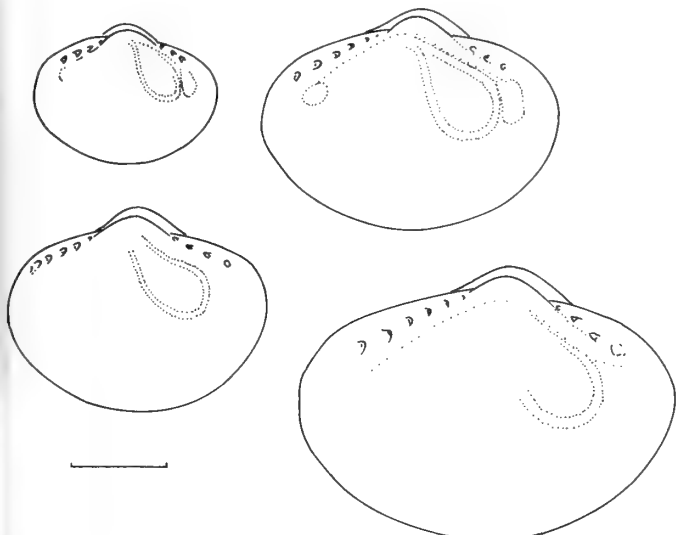
Clearly *Neilonella whoii* is closely related to *N. salicensis*. Although similar in form, they have markedly different depth distributions, *N. salicensis* occurring mainly at lower slope depths and *N. whoii* occurring mainly at abyssal depths. We believe that in the past there may have been misidentifications, and specimens of *N. salicensis* recorded from abyssal depths deserve re-examination.

Specimens which are narrower and relatively smaller in height to length ratio than those described above (Fig. 16) are present in some samples and do not occur other than with typical specimens of *N. whoii*. In other respects they are no different in their morphology to *N. whoii*. We consider them to be variants at the limit of a range of shell outlines and not a subspecies.

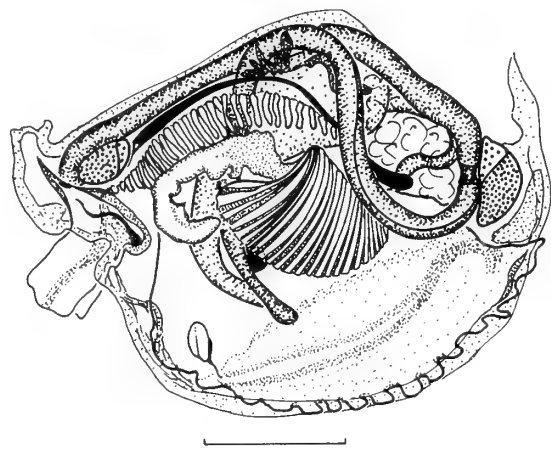
**DISTRIBUTION.** An abyssal species, found widely within temperate and tropical latitudes at depths ranging from 2487m to 5223m.



**Fig. 12.** *Neilonella whoii*. Details of shell form of specimens, from Chain station 78 from the North America Basin, the type locality; **a**, left lateral, **b**, dorsal and **c**, anterior views intact shells; **d**, the hinge plate of a specimen, from Incal station DS05 from the West European Basin; **e**, detail of the umbonal region of the hinge plate of a specimen, from Chain station 78. Scales = 1mm.



**Fig. 13.** *Neilonella whoii*. Four shells in outline, from Atlantis II station 72 from the North America Basin, in left lateral view to illustrate the small changes in shape with increasing size. Scale = 1mm.



**Fig. 14.** *Neilonella whoii*. A semidiagrammatic view of the internal morphology as seen from the right side. For the identification of the parts see text-figure 7, p. 106. Scale = 1mm.

**Family Nuculanidae** Adams and Adams 1858

A recent definition of the family is given by Allen, Sanders and Hannah (1995).

Subfamily **Ledellinae** Allen and Sanders 1982

The subfamily is defined by Allen and Hannah (1989) and comprises two genera, *Ledella* and *Tindariopsis*.

Genus **LEDELLA** Verrill and Bush 1897

TYPE SPECIES. *Ledella bushae* Warén 1978. SD – Warén 1981.

Shell small, short, robust, surface matt, concentric sculpture, in some species scattered incomplete radial striae, usually rostrate, single postero-dorsal ridge in some species, postero-ventral margin very slightly sinuous, ventral margin in older specimens maybe flattened, anterior and posterior hinge teeth series separated by edentulous space bearing short internal amphidetic ligament (resilium) which may be restricted to dorsal portion of hinge plate, outer layer of ligament visible externally and maybe extended anteriorly and posteriorly for a short distance, hind gut with various configurations.

Genus **TINDARIOPSIS** Verrill and Bush 1897

TYPE SPECIES. *Malletia (Tindaria) agatheda* Dall 1889. OD.

Shell veneriform, matt surface, concentric sculpture, umbo large, short rostrum defined by slight radial ridge and furrow, postero-ventral margin slightly sinuous, anterior and posterior hinge teeth separated by very small edentulous space, internal ligament small, close to shell margin, external ligament robust, amphidetic.

***Ledella acinula*** (Dall 1890)

TYPE SPECIMEN. Holotype USNM 95438.

TYPE LOCALITY. U.S. Fish Commission Steamer 'Albatross' Sta. 2754, 11°40'N 58°33'W, East of Tobago, 1609m.

CITED SPECIMENS. BMNH 1995047

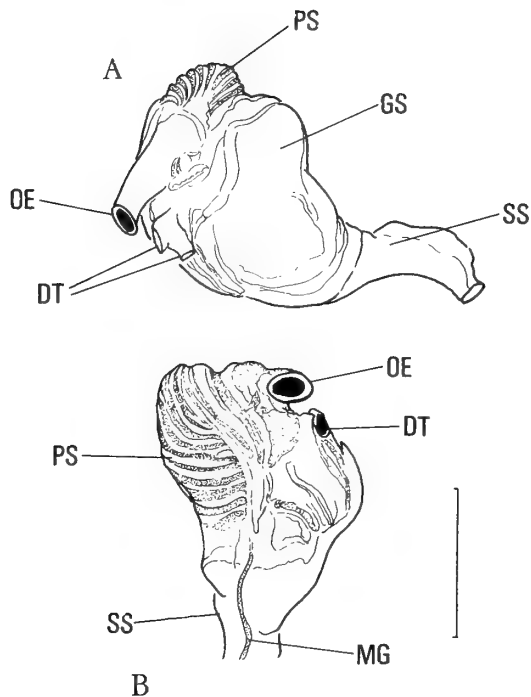
*Malletia (Tindaria) acinula* Dall, 1890, 253, pl.XIII, fig.4.

*Tindaria acinula* Verrill and Bush 1898, 881.

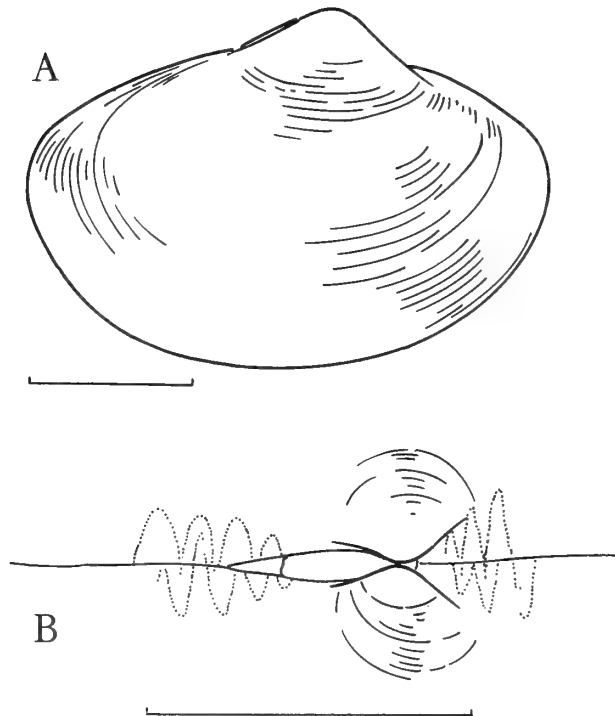
'*Tindaria*' *acinula* Sanders and Allen 1977, 55, figs 44, 45.

MATERIAL:

Cruise	Sta	Depth (m)	No	Lat	Long	Date	Gear	
<b>BRAZIL BASIN</b>								
Atlantis II	167	943–	1	07°58.0'S	34°28.0'W	20.2.67	ES	
	31	1007						
<b>GUYANA BASIN</b>								
Knorr 25	293	1456–	2	08°58.0'N	54°04.3'W	27.2.72	ES	
		1518						
		295	1000–	4	08°01.9'N	54°16.4'W	28.2.72	ES
			1022					
		299	1942–	108	07°55.1'N	55°42.0'W	29.2.72	ES
			2076					
<b>WEST EUROPEAN BASIN</b>								
Chain 106	330	4632	3	50°43.5'N	17°51.7'W	24.8.72	ES	
J.Charcot	DS79	3226	15	46°30.4'N	10°27.1'W	26.10.74	DS	
<b>BIOGAS VI</b>								
<b>GUINEA BASIN</b>								
J. Charcot	DS28	1261	4	04°21.2'N	04°35.2'E	--.71	DS	
Walda								



**Fig. 15.** *Neilonella whoii*. The dissected stomach and style sac as seen from a, in left lateral and b, frontal view. See abbreviations to text figures on p. 102. Scale = 1mm.



**Fig. 16.** *Neilonella whoii*. a, right lateral view of shell of elongate form, from Atlantis II station 242 from the Argentine Basin, in right lateral view and b, detail of the umbonal region of the same shell in dorsal view. Scales = 1 mm.

The type specimen has been examined by us.

**DISTRIBUTION.** Found predominantly in tropical and subtropical latitudes, and southern temperate latitudes in the eastern Atlantic, at mid-slope to abyssal depths ranging from 943m to 4632m.

Many protobranch species with robust shells having subrostrate or ovate outlines and with concentric ridges and external ligaments, in the past have been referred to the genera *Malletia*, *Neilonella* or *Tindaria* (e.g. Dall, 1890). *Ledella acinula* is a case in point. Having addressed this problem in earlier papers (Sanders and Allen, 1977, 1985), and the present, we have been able to define more rigorously the families Tindariidae, Neilonellidae and Malletiidae.

Dall (1890) who described large specimens of *L. acinula* referred them to the subgenus *Tindaria*. Large specimens do show some resemblance to neilonellids and tindariids, but had Dall seen the shells of smaller specimens (Fig. 18), he would have been unlikely to have made the error.

In our studies on *Tindaria* (Sanders and Allen, 1977) we re-examined and briefly redescribed *L. acinula*, and recognized that there was a problem in identification but deferred final judgment until we had made further comparative studies. Now that the Ledellinae have been reported upon (Allen and Hannah, 1989), the taxonomic relationship of this species is clear.

That *L. acinula* is siphonate clearly distinguishes it from members of the family Tindariidae. Although similar to the neilonellids in having an external amphidetic ligament, it differs in having a small but well-defined internal ligament and in being semi-rostrate with a slightly sinuous postero-ventral margin. A further significant difference is the form of the hind gut and the course that it takes within the body (Sanders and Allen, 1977). In *L. acinula* the hind gut is not particularly wide in diameter and is not restricted to the right side of the body (see below) having a configuration only known to occur in species of the subfamily Ledellinae e.g. *Ledella galatheae* Knudsen

1970, *L. oxira* (Dall 1927), *L. acuminata* (Jeffreys 1870) (Allen and Hannah, 1989).

#### SHELL DESCRIPTION (Figs 17 & 18)

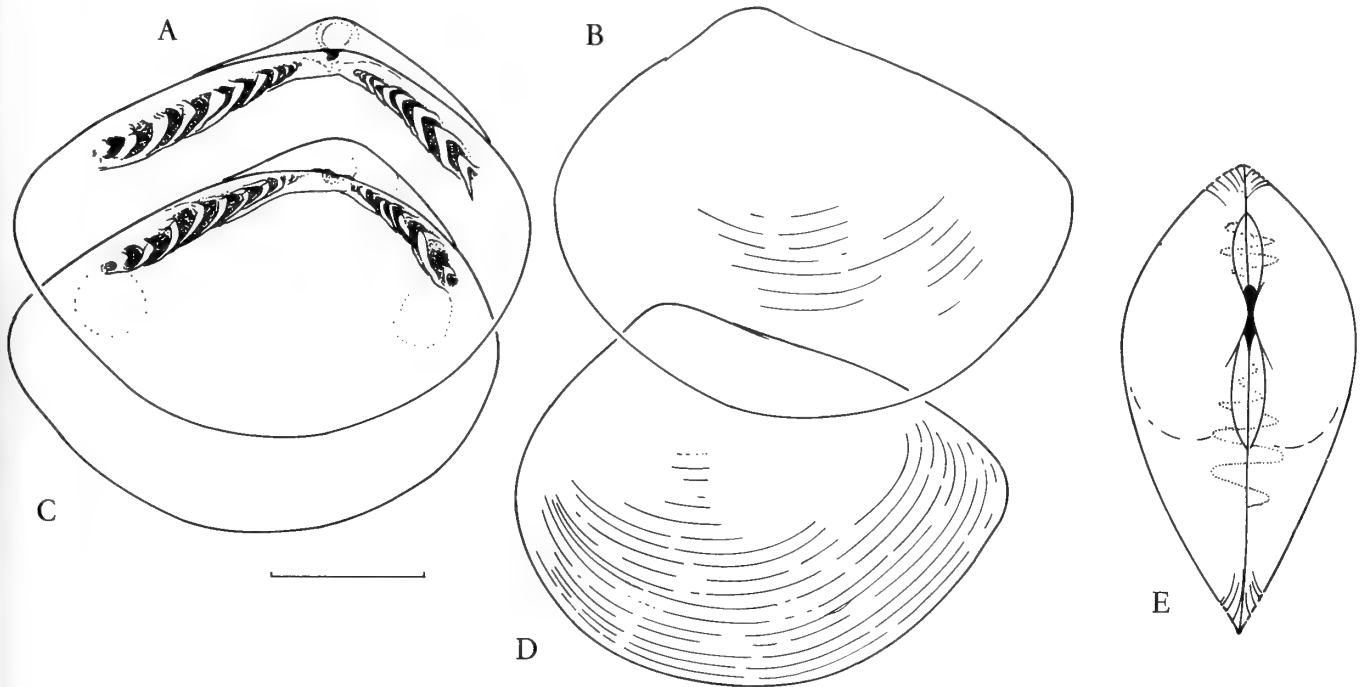
Shell robust, posteriorly angulate, ornamented with concentric ridges particularly well-defined on ventral part of shell, straw-coloured periostracum; umbos relatively low in profile, inward facing, anterior to midline; posterior rostral region characteristically broad and blunt when seen in dorsal view; postero-dorsal shell margin almost straight, angulate at posterior limit of hinge plate - particularly so in smaller specimens, barely so in larger, posterior margin sharply rounded, ventral margin deeply curved, postero-ventral margin slightly sinuous, particularly in smaller specimens, antero-dorsal, anterior and antero-ventral margins form a smooth curve; posterior and anterior limits of shell at or slightly ventral to mid horizontal axis, ventral limit of shell posterior to vertical axis through umbo; hinge plate elongate, broad, anterior and posterior tooth series separated by relatively long edentulous space, chevron-shaped teeth acutely angled, up to 10 anterior and 12 posterior teeth depending on size of specimen; ligament amphidetic, external parts short, internal resilium small, rounded, occupying upper central part of hinge plate below umbo. The maximum total length of the present specimens is 6.0mm.

The shape of the shell changes significantly with growth (Figs 18 & 19). While the ratio of height to length remains more or less the same, the postero-umbonal length increases from 50% to 60% of the total length of the shell. With increasing size the postero-dorsal margin also becomes less angulate at the posterior limit of the hinge, also the postero-ventral margin becomes less sinuous, at most being somewhat flattened.

#### INTERNAL MORPHOLOGY

This has been described and illustrated by us in our earlier studies on the family Tindariidae (Sanders and Allen, 1977). Only essential features relating to the taxonomy need be mentioned.

Combined siphons are present and there is a well-developed



**Fig. 17.** *Ledella acinula*. **a & b**, internal and external views of a left valve, from the type locality Albatross station 2754, USNM 95438; **c**, internal view of left valve of specimen, from Atlantis II station 167 from the Brazil Basin; **d & e**, lateral view of left side and dorsal view of a shell, from Knorr station 299 from the Guyana Basin. Scale = 1mm.

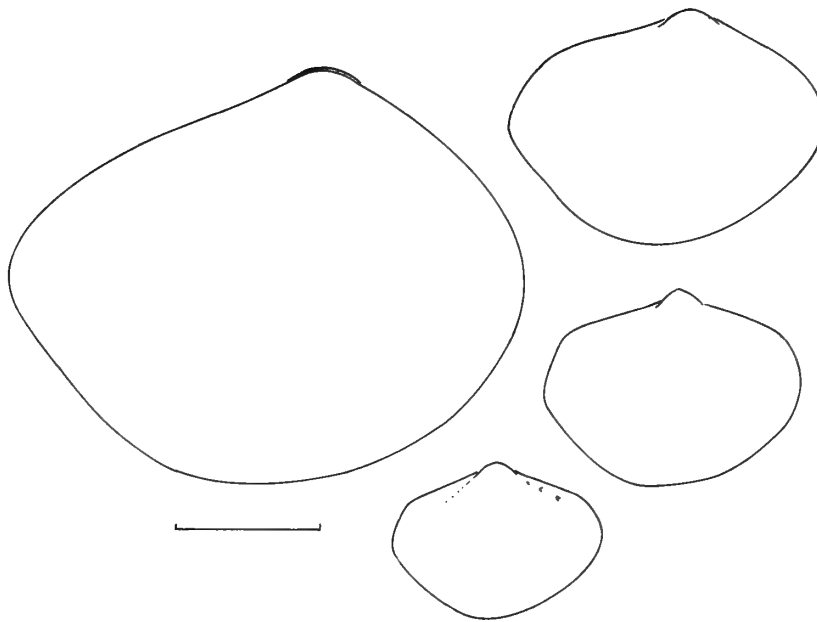


Fig. 18. *Ledella acinula*. Four shells in outline, from Knorr station 299 from the Guyana Basin, in right lateral view to illustrate changes in shape with increasing size. Scale = 1mm.

feeding aperture ventral to the siphonal embayment. The adductor muscles are relatively large, the anterior ovate and the posterior more circular in cross section. The anterior sense organ lies far anterior, ventral to the anterior adductor. The palps and gills are moderate in size with relatively few ridges (up to 15) and plates (up to 17) respectively. The foot has a well-defined neck, this is probably related to the relatively large height of the shell. There is a large 'byssal' gland present in the heel of the foot. The hind gut first makes a single loop on the right side of the body before passing to the left side of the body between the oesophagus and the inner face of the anterior adductor muscle. On the left side of the body the hind gut forms a double coil. Because of the anterior penetration of the gut to the left side, the mouth is displaced some distance posterior to the anterior adductor muscle.

### *Ledella aberrata* (new species)

TYPE SPECIMEN. Holotype BMNH 1995045; paratypes BMNH 1995046.

TYPE LOCALITY. Chain cruise 60, station 247, Argentine Basin, 43°33.0'S 48°58.1'W, 5208–5223m

#### MATERIAL:

Cruise	Sta	Depth (m)	No	Lat	Long	Date	Gear
ARGENTINE BASIN							
Atlantis II 60	242	4382–4402	2	38°16.9'S	51°56.1'W	13.3.71	ES
	247	5208–5223	34	43°33.0'S	48°58.1'W	17.3.71	ES
	252	4435	4	38°29.8'S	52°09.1'W	22.3.71	ES
GUYANA BASIN							
J.Charcot Biovema	KG13	5100	1	10°47.6'N	42°40.4'W	20.11.77	KG
NORTH AMERICA BASIN							
Chain 50	85	3834	1	37°59.2'N	69°26.2'W	5.7.65	ES
WEST EUROPEAN BASIN							
Chain 106	330	4632	3	50°43.4'N	17°51.7'W	24.8.72	ES
J.Charcot Polygas	DS23	4734	5	46°32.8'N	10°21.0'W	1.11.72	DS
Biogas II	DS32	2138	1	47°32.2'N	08°05.3'W	17.4.73	DS
Biogas IV	DS54	4659	7	46°31.1'N	10°29.2'W	21.2.74	DS
Cryos	DS68	4550	2	46°26.7'N	10°23.9'W	19.6.74	DS
Biogas V							
J.Charcot	DS78	4706	18	46°31.2'N	10°23.8'W	25.10.74	DS
Biogas VI	DS79	4715	17	46°30.4'N	10°27.1'W	26.10.74	DS
	DS80	4720	3	46°29.5'N	10°29.5'W	27.10.74	DS
	DS81	4715	2	46°28.3'N	10°24.6'W	27.10.74	DS
INCAL							
CP10	4823	1	48°25.5'N	15°10.7'W	31.7.76	CP	DS
DS11	4823	2	48°18.8'N	15°11.5'W	1.8.76	DS	WS
WS03	4829	18	48°19.2'N	15°23.3'W	1.8.76	WS	CP
CP11	4823	1	48°20.4'N	15°14.6'W	1.8.76	CP	OS
OS02	4829	1	48°19.2'N	15°15.9'W	2.8.76	OS	WS
WS10	4354	1	47°27.3'N	09°39.9'W	11.8.76	WS	
CAPE BASIN							
J.Charcot	DS05	4560	2	33°20.5'S	02°34.9'E	30.12.78	DS
Walvis							

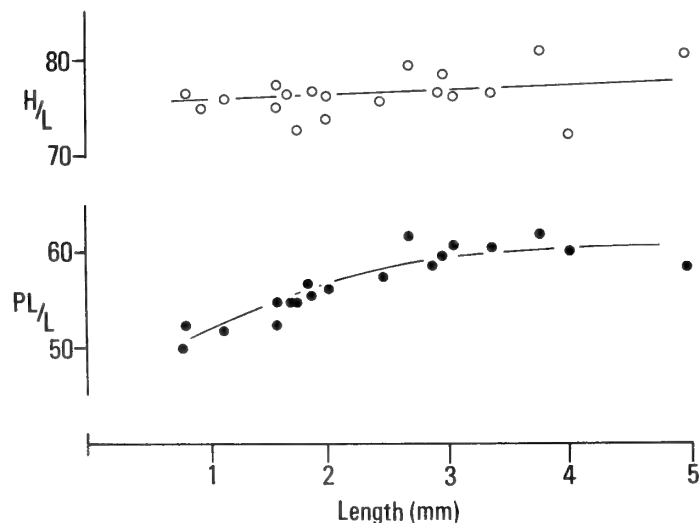
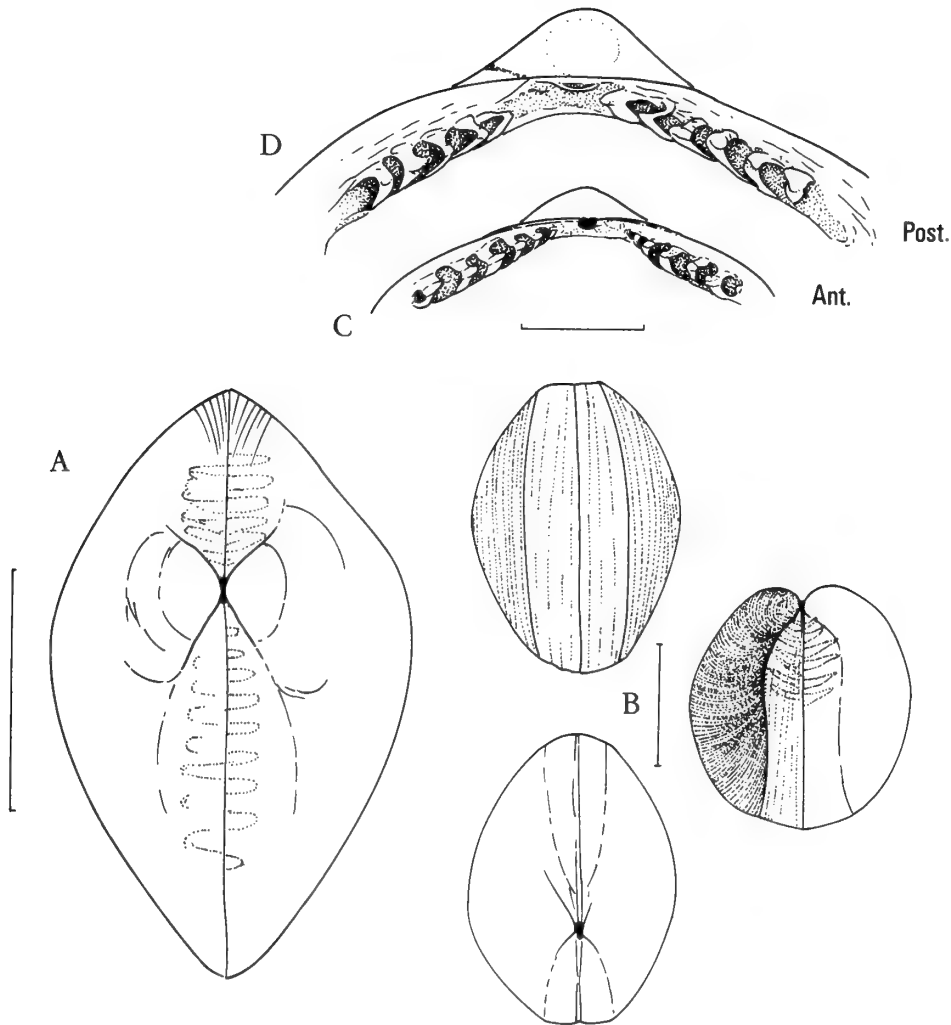


Fig. 19. *Ledella acinula*. The ratios of height to total length (H/L) (open circles) and post-umbonal length to total length (PL/L) (closed circles) plotted against total length to show changes in shell proportions with increasing length. Specimens from Knorr station 299 from the Guyana Basin.



**Fig. 20.** *Ledella aberrata*. **a**, dorsal view of shell; **b**, dorsal, ventral and anterior view of thickened shell; **c**, lateral view of the hinge plate of a left valve, all from Atlantis II station 247 from the Argentine Basin; **d**, lateral view of the hinge plate of a right valve, from Chain station 85. Scales = 1mm.

**DISTRIBUTION.** In temperate and tropical basins of the Atlantic at abyssal depths >4000m, rare in the North America Basin. Depth range, 2138–5223m.

#### SHELL DESCRIPTION (Figs 20 & 21)

Shell small, ovate, relatively high, moderately wide, ornamented with concentric ridges; umbo moderate in size, inwardly turned, anterior to mid-line but less so in juveniles; no lunule or escutcheon; periostracum pale straw colour; postero-dorsal margin slightly convex becoming more straight with increasing size, slightly angulate at posterior limit of hinge plate and at posterior margin, postero-ventral margin very slightly sinuous, otherwise ventral margin deeply curved with ventral limit posterior to vertical axis through umbo, anterior margin sharply curved, antero-dorsal margin slightly convex with slight change in slope at anterior limit to hinge plate; shell outline characteristically asymmetrical, shell margin in larger specimens changes direction of growth producing a broad flattened ventral margin; hinge plate broad, up to 6 chevron teeth in anterior series and 7 in posterior series, edentulous space between series relatively broad; ligament small, amphidetic, internal part restricted to upper part of hinge plate, external part extremely short situated below umbo. The maximum length of the present specimens is 2.6mm.

#### INTERNAL MORPHOLOGY (Figs 22 & 23)

The adductor muscles are moderately large and oval in shape. The combined siphon is relatively short. The dorsal margins of the exhalent part are fused proximally for a short distance and the ventral margins of the inhalent part are not fused but are slightly thickened and probably adhere in living specimens. Internally where inhalent and exhalent parts join, there is a thickened median ridge on each side which together together with the posterior continuation of the gill axes probably act as guides when the faecal pellets are extruded. The siphonal embayment is small and there is a small, slender, tentacle attached to the left side at the inner limit of the embayment. The anterior sense organ is small and is situated ventral to the anterior adductor muscle.

The palps are small with up to 11 broad ridges. The gills are also small each with up to 11 plates the most posterior of which lies some distance from the siphon. The gills are attached to the posterior limits of the median guides by slender extensions of the gill axis.

The hind gut is greatly extended and takes a similar but more complex course to that described for *L. acinula*. Like the latter, the hind gut passes from the right side of the body to the left immediately posterior to the anterior adductor muscle and returns by the

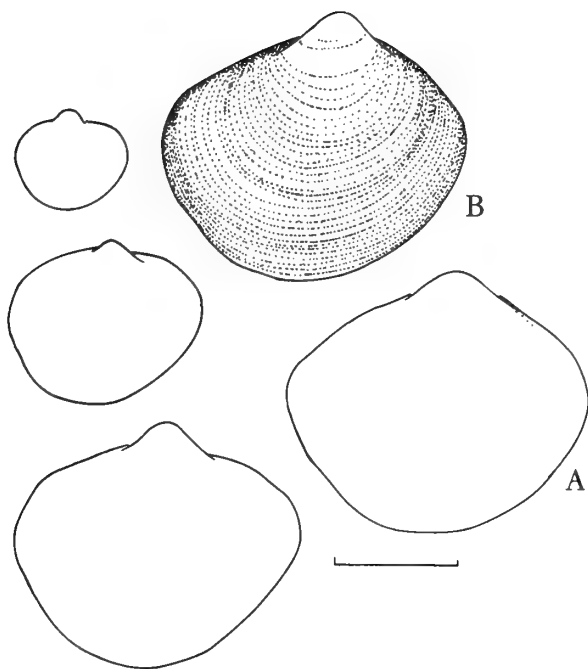


Fig. 21. *Ledella aberrata*. Five shells in outline to show variation in shape with growth. a, is a specimen with a thickened margin; b, has a slightly thickened margin; the remaining three shells are unthickened. Scale = 1mm.

same route. Unlike the latter species, it makes a single coil on the right side as well as a double twinned coil on the left.

The foot is unusual in having a large heel and a narrow muscular anterior part. The marginal papillae are few in number and restricted to the anterior margins of the sole. The sole is less deeply divided as compared with other protobranchs. There is a large 'byssal' gland in the heel of the foot.

Although the shell outline of the smaller specimens is more characteristic of the genus *Ledella* than in larger mature specimens, the general shell outline of *L. aberrata* (and *L. acinula*) is much deeper and more ovate than in other described species (Figs 18 & 21) nor is it markedly rostrate. Despite this, the characters place them in the Ledellinae (Allen and Hannah, 1989) and we see no reason for erecting another genus.

Apart from *L. aberrata*, only two species of protobranchs, have been reported as exhibiting a change in shell growth to produce a flattened shell margin (Fig. 20). Both are ledellids, namely *L. ultima* (Smith 1885) and *L. solidula* (Smith 1885) (Allen and Hannah, 1989). Like *L. aberrata* these two latter species also have elongate hind guts. The hind gut of *L. solidula* is very similar to that of *L. aberrata* in having double twinned coils to the left side of the body (Fig. 22), although it does not have an additional single coil to the right as does the present species. The type of course taken by the hind gut in *L. acinula* is also found in other species of *Ledella* (e.g. *L. oxira*) (Allen and Hannah, 1989).

We named this species after the familiar appellation to which it was referred during our original analysis of the samples.

### *Tindariopsis agatheda* (Dall 1889)

TYPE SPECIMEN. USNM 95437, lectotype here designated.

TYPE LOCALITY. U.S. Fish Commission Sta. 2754, east of Tobago, 11°40'N 58°33'W, 1609m.

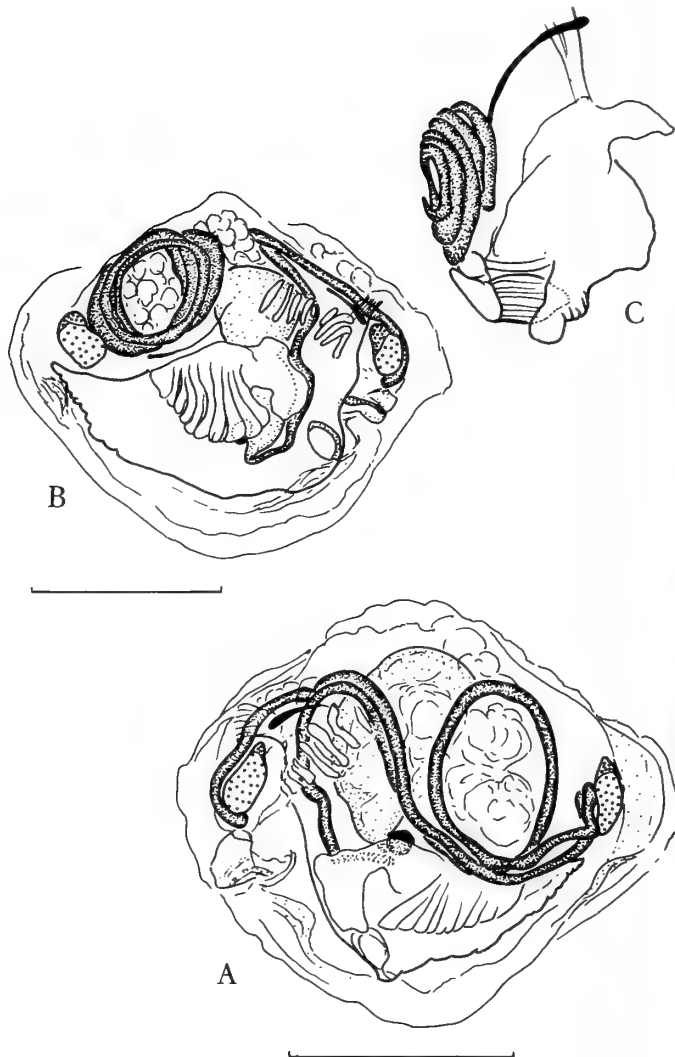


Fig. 22. *Ledella aberrata*. Internal morphology as seen in a, right lateral view, b, left lateral view and c, left ventro-lateral view. For identification of the parts see text-figure 7, p. 106. Scales = 1mm.

CITED SPECIMEN. BMNH 1995062.

*Malletia* (*Tindaria*) *agatheda* Dall 1890, 252, pl. xiii, fig. 10.  
*Tindaria* (*Tindariopsis*) *agatheda* Verrill and Bush 1897, 59.  
*Saturnia* (*Tindariopsis*) *agatheda* McAlester 1969, N235.  
*Tindariopsis agatheda* James 1972, 98, figs 60–62.  
*Neilonella* (*Tindariopsis*) *agatheda* Laghi 1986, 190, pl.8, figs 2–6.

### MATERIAL:

Cruise	Sta	Depth (m)	No	Lat	Long	Date	Gear
BRAZIL BASIN							
Atlantis II	167	943–	1	07°58.0'S	34°28.0'W	20.2.67	ES
	31	1007					
GUYANA BASIN							
Knorr 25	293	1456–	14	08°53.1'N	54°04.3'W	27.2.72	ES
		1518					
	299	1942–	8	07°55.1'N	55°42.0'W	29.2.72	ES
		2067					
	301	2487–	7	08°12.4'N	55°50.2'W	1.3.72	ES
		2500					

The type specimen has been examined by us.



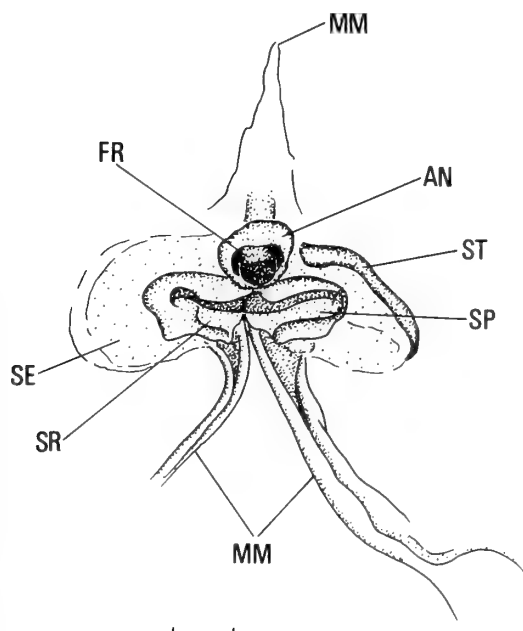


Fig. 23. *Ledella aberrata*. Siphonal region, as seen from the inside of the mantle cavity. See abbreviations to the text-figures p. 102. Scale = 0.1mm.

This species occurs at upper to mid slope depths in the tropical western Atlantic in the Brazil, Guyana, Caribbean and Gulf of Mexico Basins (Dall, 1889; James 1972). Depth range; 943–2500m.

The holotype for *T. agatheda* was not designated by Dall (1890), nevertheless he did illustrate the left valve from Albatross Sta. 2754, 11°40'N 58°33'W (USNM 95437) which we have here nominated as

lectotype. We have redrawn the shell and added detail of the hinge plate (Figs 24 and 25). In addition, James (1972) reported that two valves of *T. agatheda* were included in USNM 63149 from Blake sta. 236, 2909m off Bequia, furthermore USNM 94326 from Blake stas 26 and 30 between Cuba and Yucatan, identified by Dall as *Leda pusio*, are examples of *T. agatheda*.

#### SHELL DESCRIPTION (Figs 24 & 25)

Shell small, sub-ovate, wide, sub-rostrate, ornamented with concentric ridges, postero-lateral furrow ventral to sub-rostrum, ill-defined lunule outlined by obscure ridge, escutcheon similarly ill-defined and bounded by faint ridge; umbo prominent, posterior to mid-line, inwardly directed; antero-dorsal margin concave, with change in slope at anterior limit of hinge plate, slightly flattened anteriorly dorsal to the anterior limit of shell, antero-ventral and ventral margins smoothly curved, postero-ventral margin slightly sinuate, posterior margin acutely angled, postero-dorsal margin slightly concave, marked angle at posterior limit of hinge plate and thereafter almost straight to form sub-rostrate posterior margin; anterior and posterior limits of shell are ventral to horizontal mid-line; hinge plate strong, with edentulous space ventral to umbo, hinge teeth stout, chevron-shaped, up to 12 in each series in shell 6mm total length; internal ligament small, close to shell margin, external ligament amphidetic, moderately short, stout.

Young shells are less rostrate, with the posterior and anterior limits of the shell more dorsal in position.

Although there is variation in the shape of the shell, the height/length and the post-umbonal length/total length ratios increase slightly with increasing size (Table 1). The maximum total length of the present specimens is 6.0mm.

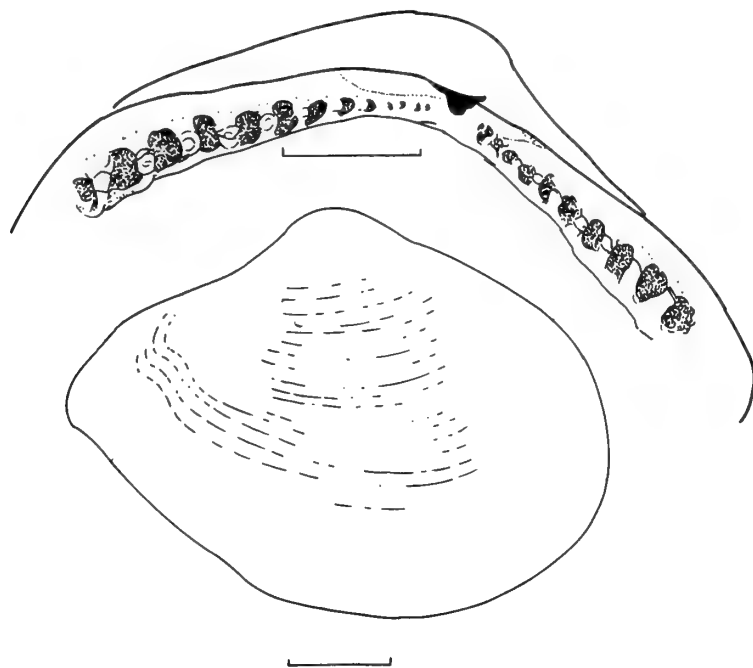
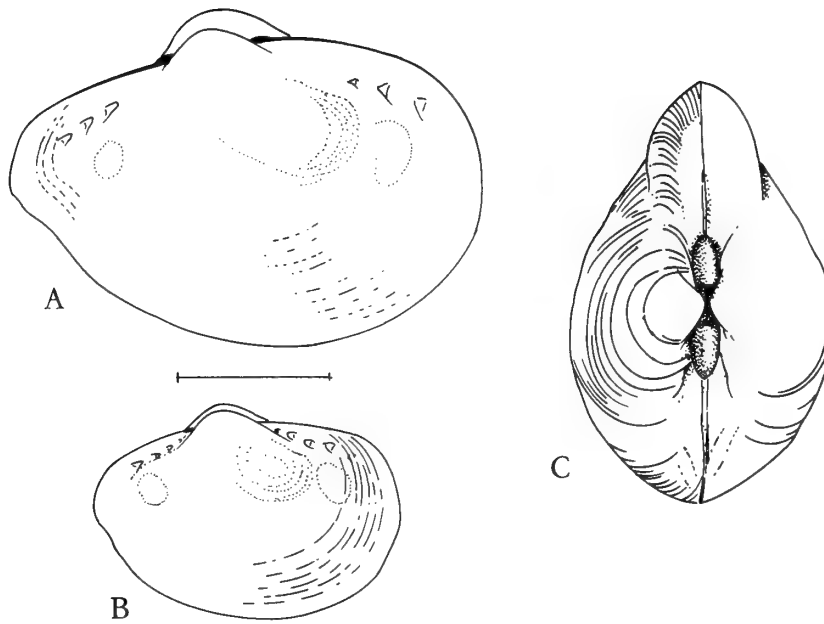


Fig. 24. *Tindariopsis agatheda*. External lateral view of the right valve of the lectotype and an internal view of the hinge plate of the same valve, from U.S.Fish Commission station 2754 East of Tobago. USNM 95437. Scales = 1mm.



**Fig. 25.** *Tindariopsis agatheda*. External lateral views of right side of two specimens of differing size to show change in shape with growth. a, from Knorr station 301 and b, from Knorr station 293, both from the Guyana Basin; c, external dorsal view of a shell also from station 293. Scale = 1mm.

**Table 1.** Measurements and ratios of shell parameters of the sample from Knorr sta. 293.

Length (L) (mm)	Height (H) (mm)	Post-umbonal (PL) length (mm)	PL/L	H/L
6.00	4.95	3.00	0.50	0.83
5.60	3.95	2.65	0.47	0.71
5.00	3.65	2.20	0.44	0.73
4.90	3.65	2.05	0.42	0.75
4.70	3.15	1.95	0.42	0.67
4.50	3.10	1.85	0.41	0.69
4.45	3.20	1.85	0.42	0.72
3.70	2.55	1.70	0.46	0.69
3.45	2.50	1.45	0.42	0.73
2.35	1.58	0.93	0.39	0.67
2.05	1.43	0.93	0.45	0.70
1.95	1.30	0.88	0.45	0.67
1.83	1.28	0.83	0.45	0.70
1.10	0.88	0.48	0.43	0.80

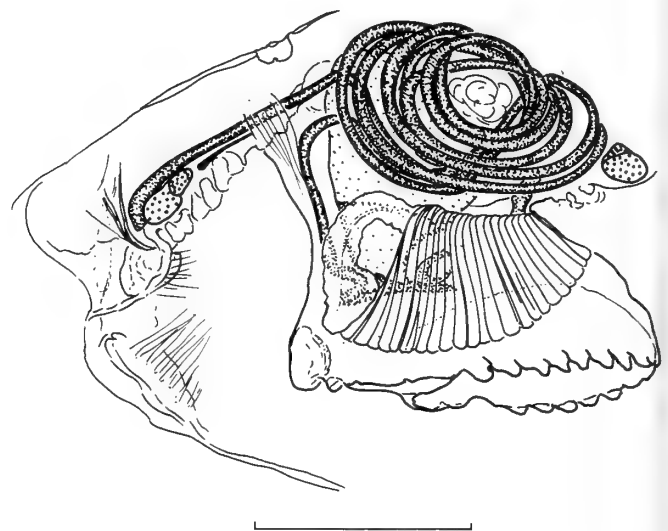
#### INTERNAL MORPHOLOGY (Fig. 26)

The siphonal embayment is shallow and dorso-ventrally narrow. In contrast, the feeding aperture is broad and well-supplied with radial pallial muscles. The adductor muscles are small, the posterior muscle is oval and the anterior muscle is circular in cross section. The foot is large with a well-developed heel containing a large 'byssal' gland. The gland opens into the posterior limit of the pedal groove via a small papilla. The anterior two-thirds of the margins of the divided sole are broadly papillate. The palps are large with up to 25 ridges in the largest specimens. The gills are narrow ill-defined with about 16 plates in the largest specimens.

The mouth lies close to the posterior face of the anterior adductor muscle. The oesophagus opens into a large stomach and style sac. The hind gut passes posterior to the style sac and stomach to the dorsal side of the viscera and thence across the right side of the body where it forms 8 coils before returning along the same path to the dorsal side of the viscera and from there through the heart and then dorsal to the posterior adductor muscle to the anus.

Usually not more than six coils are visible, the others being overlain by those to the outside of them. The form of the hind gut is derived by 4 complete turns of the closely parallel anterior and posterior lengths of the hind gut on the right of the body. This particular disposition of the hind-gut is to be found in other ledellids (e.g. *L. ultima*) and yoldiellids (e.g. *Y. ella* Allen, Sanders and Hannah 1995) (Allen and Hannah, 1989; Allen, 1992; Allen, Sanders and Hannah, 1995).

The nervous system is similar in its arrangement to that of other deep-sea protobranchs, however the cerebral and visceral ganglia are noticeably smaller and the commissures much finer than observed in other species.



**Fig. 26.** *Tindariopsis agatheda*. Internal morphology as seen from the right side of a specimen from Knorr station 293. For identification of the parts see text-figure 7, p. 106. Scale = 1mm.

*Tindariopsis aeolata* (Dall 1890)

TYPE SPECIMEN. Holotype, USNM 95436.

TYPE LOCALITY. U.S. Fish Commission Sta. 2754, East of Tobago, 11°40'N 58°33'W, 1609m.

CITED SPECIMEN. BMNH 1995061.

*Malletia* (*Tindaria*?) *aeolata* Dall 1890, 252.

*Tindaria* (*Tindariopsis*) *aeolata* Dall 1898, 582.

*Tindariopsis aeolata* James 1972, 97, figs 57–59.

## MATERIAL:

Cruise	Sta	Depth (m)	No	Lat	Long	Date	Gear
GUYANA BASIN							
Knorr 25	299	1942–	4	07°55.1'N	55°42.0'W	29.2.72	ES
				2076			
	301	2487–	5	08°12.4'N	55°50.2'W	29.2.72	ES
		2500					

The type specimen has been examined by us.

This species occurs from mid to lower slope depths in the tropical western Atlantic in the Guyana and Caribbean Basins and the Gulf of Mexico. Depth range: 1609–3466m.

## SHELL DESCRIPTION (Figs 27 &amp; 28)

Shell small, subquadrate, rostrate, ornamented with marked concentric ridges; periostracum pale yellow; umbos moderately large, posterior to midline (post-umbonal length 45–48% of total length), facing inwards, slightly separated by external ligament; distally antero-dorsal shell margin horizontal, then curves smoothly and steeply to anterior margin, postero-ventral margin sinuous, ventral margin somewhat flattened, postero-dorsal distal margin slopes gently to limit of hinge plate then curves sharply to rostral point, latter rounded and somewhat eroded in large specimens, more pointed in smaller, rostrum in mid horizontal plane in small specimens and dorsal to it in large specimens, limit of anterior margin

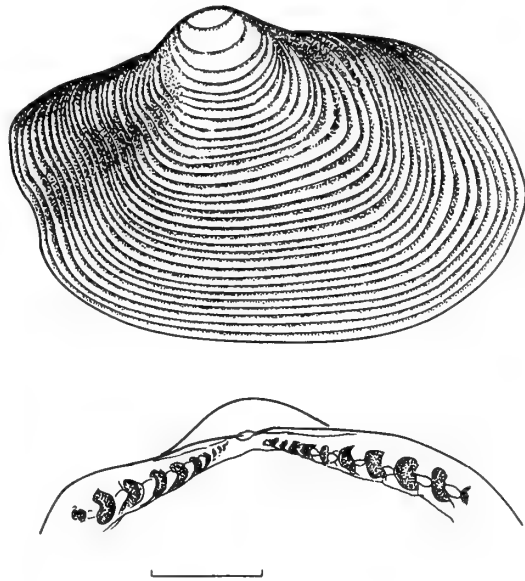


Fig. 27. *Tindariopsis aeolata*. External lateral view of the right valve and the hinge plate of the left valve of the holotype, from U.S. Fish Commission station 2754, USNM 95436. Scale = 1mm.

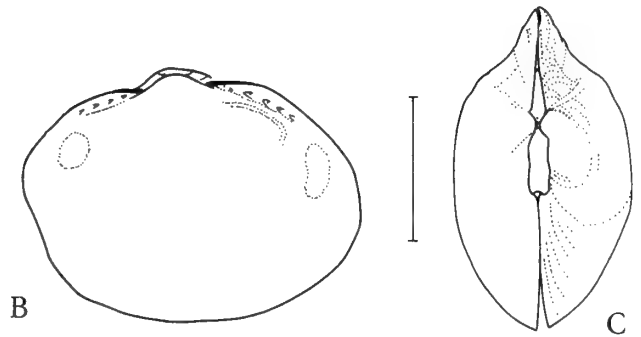
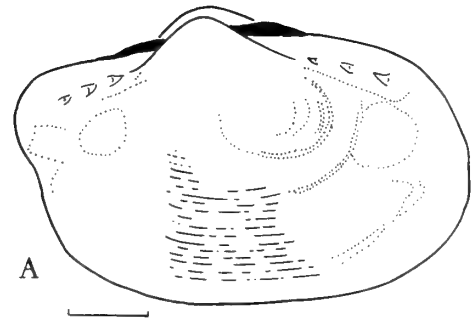


Fig. 28. *Tindariopsis aeolata*. External lateral views of the right side of shells of differing size to change in shape with growth. a, from Knorr station 301 and b, from Knorr station 299 from the Guyana Basin; c, external dorsal view of a shell also from Knorr station 299. Scales = 1mm.

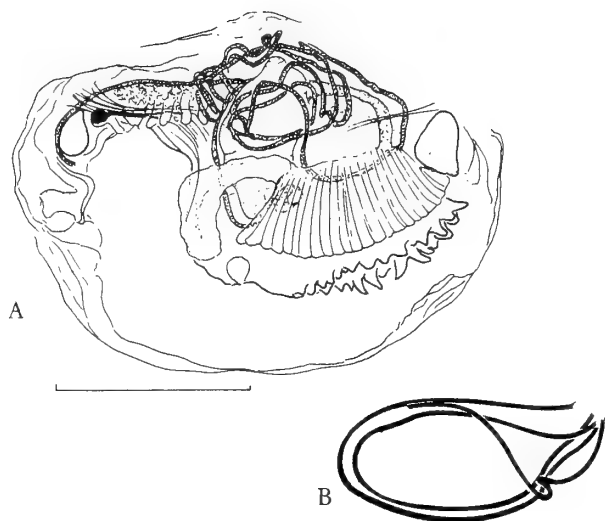
ventral to the mid horizontal plane; hinge plate stout, small edentulous space between tooth series, 9 chevron-shaped teeth in anterior series and 10 in posterior series in largest specimens; ligament amphidetic, external except for small resilifer at margin ventral to umbo, external part thickened, particularly so in large specimens. The maximum length of the present specimens is 5.8mm.

In lateral view the rostrum, although dorsal to mid horizontal line, is reminiscent of *Ledella*, while the robust external ligament is more reminiscent of *Spinula*.

## INTERNAL MORPHOLOGY (Fig.29)

The siphonal embayment is relatively shallow and the contained siphon is similar to that of *T. acinula*. The adductor muscles are moderately small, ovate in cross-section and equal in size. The foot is large, with a divided sole fringed with large papillae. The 'byssal' gland is moderate in size. The palps are very large with many ridges (c 26 in the largest specimen) and the palp proboscides are broad. The gills are small with 11 gill plates in the largest specimen. The kidney is long and narrow. The nervous system is of typical protobranch design. The ganglia are relatively large and, in contrast, the commissures are unusually slender.

The mouth lies some distance posterior to the anterior adductor muscle (see below). The oesophagus opens on to the anterior face of a moderately large stomach. The latter lies almost vertical within the posterior part of the visceral mass. The hind gut is very small in diameter and takes an extraordinarily complex course through the body. There are two loops to the left side of the body (Fig.29B) and one major loop to the right side of the body, all three pass from one side to the other ventral to the umbo. There is also a complex series of loops anterior and to the right of the stomach. This morphology has not been encountered before in the protobranch bivalves and is very different from that seen in *T. agatheda*. Yet, it is debateable



**Fig. 29.** *Tindariopsis aeolata*. a, internal morphology as seen from the right side of a specimen from Knorr station 301 from the Guyana Basin; b, the form of the hind gut on the left side of the body. For identification of the parts see text-figure 8, p. Scale = 1mm.

whether the difference warrants generic status. Other protobranch genera show an array of hind gut morphologies (e.g. Yoldiellidae, Allen, 1992; Allen, Sanders and Hannah, 1995) which we believe relate to changes in the benthic food resource as depth increases. For this reason we are reluctant to erect a new genus when in other respects *T. aeolata* is clearly within the genus *Tindariopsis*.

#### Subfamily **Nuculaninae** Allen and Sanders 1982

The subfamily is defined by Allen and Hannah (1986) and comprises three genera *Nuculana*, *Propeleda* and *Adrana*.

#### Genus **NUCULANA** Link 1807

**TYPE SPECIES.** *Arca rostrata* Gmelin 1791 = *Arca pernula* Müller 1779. OD.

Shell robust, moderately elongate, concentric sculpture, occasionally with radial ribs, slightly rostrate, usually bicarinate; umbo anterior; postero-dorsal margin straight or somewhat concave, posterior margin may be slightly sinuous; escutcheon present; no internal ridge from umbo to posterior margin; hinge moderately robust, teeth chevron-shaped; ligament small, for most part internal, usually amphidetic and vertical, sometimes posteriorly oblique.

#### Genus **PROPELEDA** Iredale 1924

**TYPE SPECIES.** *Leda ensicula* Angas 1877. OD.

Shell very elongate, thin, glossy, concentric sculpture may be ill-defined, 3/4 shell post-umbonal, usually with two marked carinae from umbo to upper and lower limit of rostrum, posteriorly truncate; umbo small; postero-dorsal margin concave, postero-ventral margin not sinuous; internal ridge usually from umbo, skirts ventral margin of posterior adductor to posterior margin, second ridge may be present from hinge plate to rostral margin; hinge plate slender, hinge

teeth chevron-shaped, one or both arms of the chevron may be elongate, anterior tooth series curve round the outer margin of the anterior adductor, posterior series extends posterior to adductor; ligament in large part internal, opisthodontic and oblique.

#### Genus **ADRANA** Adams & Adams 1858

**TYPE SPECIES.** *Nucula lanceolata* Lamarck 1819. SD Stoliczka 1871.

Shell extremely elongate, slender, lanceolate, fragile, smooth or with fine concentric and sometimes oblique sculpture, without carinae, glossy; umbo almost central, barely raised; escutcheon elongate, flattened, narrow; postero-dorsal margin straight, antero-dorsal margin slightly convex, postero-ventral margin sinuous; hinge plate slender, hinge teeth fine, obtuse, chevron-shaped; chondrophore present; ligament internal, amphidetic.

#### *Nuculana acuta* (Conrad 1831)

**TYPE SPECIMEN.** Lectotype here designated, chosen from ANSP 30613, remainder of lot designated paralectotypes.

**CITED SPECIMEN.** BMNH 1995055.

**TYPE LOCALITY.** Tertiary fossil beds, near Suffolk, Virginia.

*Nucula acuta* Conrad 1831, 32, pl.6, fig.1.

*Nucula cuneata* Sowerby 1833, 198.

*Nucula carinata* H.C. Lea 1843, 163, (non M' Coy 1844).

?*Leda jamaicensis* d'Orbigny 1846, 263, pl. XXIV, figs 30–32.

?*Leda inornata* A. Adams 1856, 48.

*Leda unca* Verrill 1880, 401, (?non Gould 1862).

*Leda acuta* Dall 1886, 251, pl.7, figs 3a, 3b and 8.

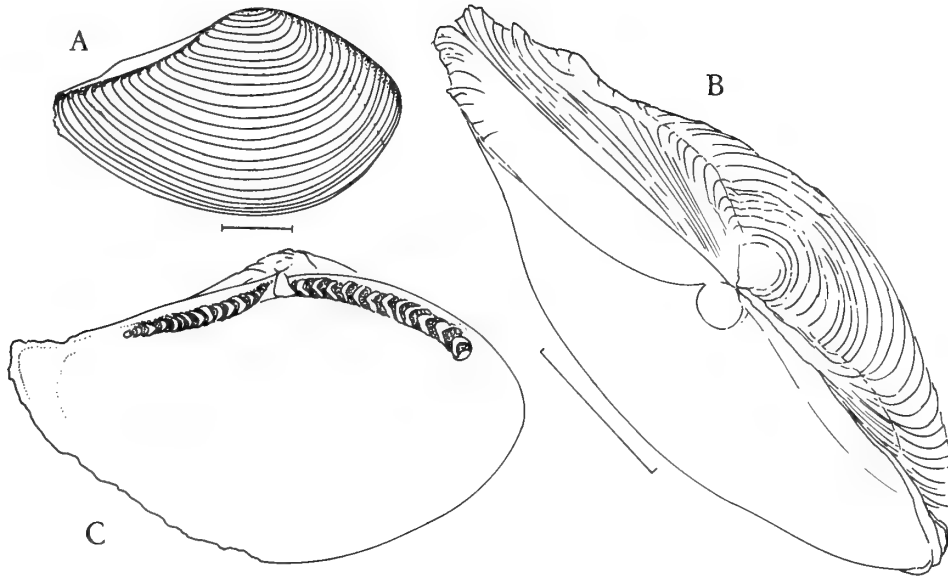
*Nuculana acuta* Morris 1951, 7, pl.6, fig.2.

#### **MATERIAL:**

Cruise	Sta	Depth (m)	No	Lat	Long	Date	Gear
NORTH AMERICA BASIN							
Atlantis 264	C1	97	2+2v	40°20.5'N	70°47.0'W	25.5.61	AD
Atlantis 283	Slope sta.2	200	8+6v	40°01.8'N	70°42.0'W	28.8.62	AD
Atlantis II 24	114	197	8	40°04.1'N	70°27.8'W	15.8.66	ES
Atlantis II 40	172	119	7+2v	40°12.3'N	70°44.7'W	27.11.67	ES
	173	123	4	40°10.8'N	70°43.6'W	28.11.67	ES

The type specimens have been examined by JAA.

Conrad (1831) described this species from fossils obtained from the Miocene beds near Suffolk and the banks of the James and York rivers, Virginia. He later redescribed the species (Conrad, 1845) adding that he had found Recent specimens in deep water in the Gulf of Mexico. His first account refers to specimens being in 'Cabinet of the Acad. Nat. Sciences, No.1738.' This reference does not correspond with any lot of *N. acuta*, Recent or fossil, in the Academy of Natural Sciences, Philadelphia today. There are specimens in the Invertebrate Paleontology collection of the Academy (catalogue number 30613) that up to now have been considered as possible syntypes of the species. The lot comprises 4 right valves, 3 left valves 1 left and 1 right broken valve, 1 intact shell, 1 shell with rostrum tip missing and 3 fragments. These specimens, labelled by



**Fig. 30.** *Nuculana acuta*. a & b, lateral and dorsal external views of the lectotype, ANSP 30613; c, lateral internal view of a paralectotype from the same lot. Scales = 1mm.

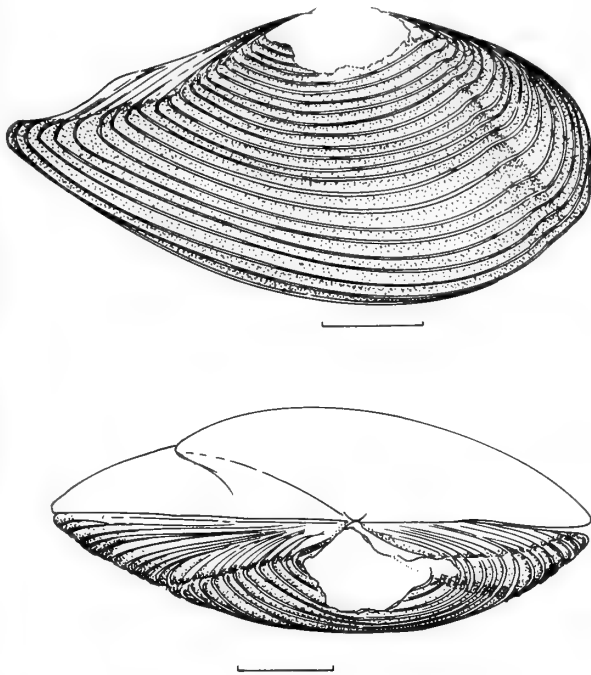
Conrad, may include those figured by him. Comparison with the figures (Conrad, 1831 and 1845) shows that it is impossible to say which, if any one, was figured nor is it possible to be absolutely certain that these are the specimens from which the original description was made, although we believe that they are. Thus, the intact shell from lot 30613 has been chosen as the lectotype, the remainder being paralectotypes.

Campbell (1993) listed *Leda jamaicensis* d'Orbigny 1846, *Leda*

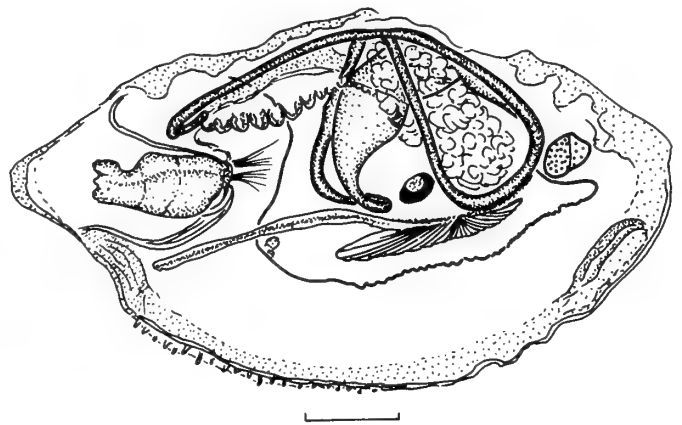
*inornata* A.Adams 1856 and *L.unca* Gould 1862 as synonyms of *N. acuta*. Verrill (1882) describes in detail differences that he found between *N. acuta* and *L.jamaicensis* and *L.unca* which cast doubt as to the synonymy, although Dall (1886) maintains the synonymy of *L. jamaicensis*. Similarly, we have doubts as to synonymy with *L.inornata* A.Adams which is a 'gibbose', 'fuscous', 'sulcate' species from New Guinea. Thus, although *Nuculana acuta* is a well-described species (e.g. Verrill 1882, 1884; Dall, 1886; Abbott, 1974) because there are closely related species in the Atlantic and elsewhere, we include a description here. It occurs off the East coast of North America, in the Caribbean Sea and off Brazil at depths from the outer shelf to lower slope depths, 97–2909m (James, 1972).

**SHELL DESCRIPTION (Figs 30, 31)**

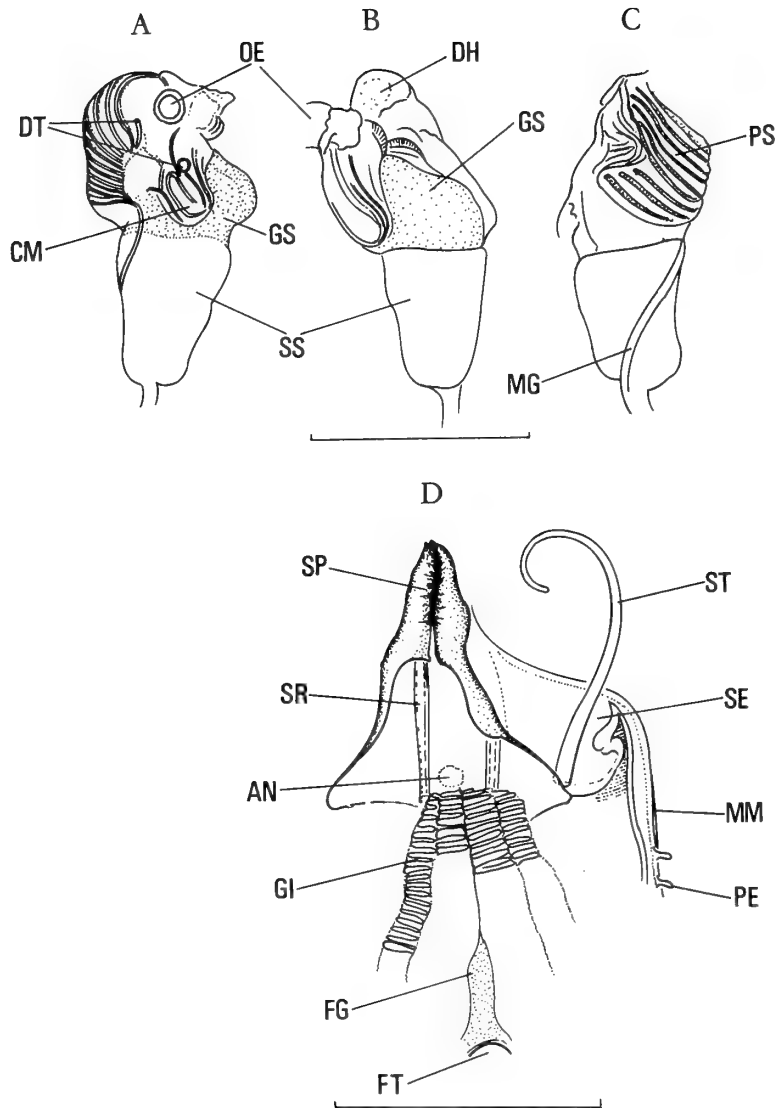
Shell moderately large, elongate, somewhat inflated, rostrate, ornamented with deep concentric ridges flattened at the apex, rostral ridge from umbo to ventral limit of rostrum, very faint radial ridge from umbo to antero-ventral margin, yellow periostracum; lunule



**Fig. 31.** *Nuculana acuta*. Lateral and dorsal external views of a shell, from Atlantis 283 station 2 from the North America Basin. Scales = 1mm.



**Fig. 32.** *Nuculana acuta*. Internal morphology as seen from the right side of a specimen from Atlantis II station 197 from the North America Basin. For identification of the parts see text-figure 7, p. 106. Scale = 1mm.



**Fig. 33.** *Nuculana acuta*. Internal morphology, a, anterior, b, left lateral & c, posterior views of a dissected stomach and combined style sac; d, ventral view of siphonal region. See abbreviations to text-figures on p. 102. Scales = 1 mm.

broad, elongate, outlined with faint ridge, escutcheon broad, outlined by rostral ridge; umbos small, inwardly directed, anterior to mid-line; antero-dorsal margin broadly concave, anterior, antero-ventral, ventral and postero-ventral margins form smooth curve, postero-dorsal margin raised, distally straight, proximally – posterior to hinge plate – slightly concave, may be slightly upturned in larger specimens; hinge plate elongate, relatively broad, hinge teeth chevron-shaped, up to 18 teeth in each series depending on size of specimen; ligament small, amphidetic, internal pear-shaped in sagittal section, extends slightly ventral to hinge plate. The maximum length of the present specimens is 9.2mm.

#### INTERNAL MORPHOLOGY (Figs 32, 33)

The siphonal embayment is deep, with an elongate tentacle attached to the inner right or left side. The siphons are elongate, combined and except anteriorly, the ventral margins are fused. In the contracted state the line of fusion is marked by deep ventral furrow. The anterior sense organ is far anterior, situated at the point where the radial ridge meets with the shell margin. Between the feeding (ventral to the siphonal embayment) and the pedal gape, the inner

folks of the ventral margin are applied to each other. In this section of mantle margin approximately 30 small sensory papillae are attached to each middle sensory fold in a specimen 6.5mm total length. The adductor muscles are small the anterior unusually so. The anterior adductor is circular in cross-section and the posterior elongate-oval.

The foot is moderately large, elongate, the sole with papillate margins. The heel is not marked but there is a large 'byssal' gland present internal to the posterior limit of the sole. The gills are well-developed with up to 48 alternating gill plates. The dorsal margins of the left and right inner demibranchs are fused. In life the gills are a bright orange-red colour. In contrast the palps are cream. The latter are relatively small, elongate and dorso-ventrally narrow and, for the most part, hidden under a fold of the body wall. This latter is more pronounced on the right side of the body where the hind gut loop meets the ventral margin of the visceral wall. Each palp has up to 24 ridges on the inner face. The palp proboscides are also long and tapering.

The digestive gland is bright orange in life. As in other species of *Nuculana*, the course of the hind gut describes a single loop to the

right side of the viscera. There is a single typhlosole present along its entire length. The stomach is of moderate size and internally is similar in form to that of shallow water species of *Nuculana* (Yonge, 1939). The gastric shield lines much of the left wall of the stomach. To the right there is a large posterior sorting area with 13 ciliated ridges. A deep caecum is ventral to the oesophageal aperture. Two digestive ducts open close to the antero-dorsal margin of the posterior sorting area and a single duct opens antero-dorsally close to the oesophageal opening.

As will be seen *Nuculana acuta* is remarkably similar in its shell features and anatomy to *Nuculana commutata*. This similarity is discussed under the latter species (p. 123).

### *Nuculana commutata* (Philippi 1844)

TYPE SPECIMEN. ZMHU.

TYPE LOCALITY. Pliocene, Palermo, Sicily.

CITED SPECIMEN. BMNH 1995212

*Arca fragilis* Chemnitz 1784, 199, pl.LV, fig.546.

*Arca pella* Gmelin 1790 (non Linné), 3307.

*Arca minuta* Brocchi 1814 (non Fabricius), 482, pl.XI,fig.4.

*Nucula pella* Payraudeau 1826 (non Linné), 64.

*Lembulus deltoideus* Risso 1826 (non Lamarck), 320, pl. XI, fig. 164.

*Nucula minuta* Scacchi 1836 (non Fabricius), 4.

*Nucula striata* Philippi 1836 (non Lamarck), 64.

*Nucula commutata* Philippi 1844, 101.

*Leda fragilis* Jeffreys 1879, 575.

*Leda minuta* Jeffreys 1856 (non Fabricius), 25.

*Leda commutata* Hanley 1863, pl.CCXVIII, figs, 80, 81.

*Lembulus commutatus* Monterosato 1878, 6.

*Leda (Portlandia) tenuis* Sturany 1896, 6.

*Nuculana (Jupiteria) fragilis* Nordsieck 1969, 9, pl.I, fig.02.25.

*Nuculana (Jupiteria) commutata* Smith and Heppell 1991, 56.

#### MATERIAL:

Cruise	Sta	Depth (m)	No	Lat	Long	Date	Gear
WEST EUROPEAN BASIN							
Sarsia	S29	119	16	47°40.0'N	05°00.0'W	12.8.67	ES

#### SHELL DESCRIPTION (Fig.34)

Shell moderately large, elongate, slightly inflated, rostrate, ornamented with concentric ridges, pale straw-coloured periostracum; radial ridge from umbo to antero-ventral margin; rostral ridge well-defined, delimits posterior dorsal area, within this area a faintly outlined escutcheon extending half the length of postero-dorsal margin; lunule elongate, defined by fine ridge; less well-defined ventral rostral ridge extends from umbo to postero-ventral margin; umbos anterior to midline, inwardly directed; antero-dorsal margin proximally straight, distally slightly concave merging with rounded anterior margin to where it meets with ventral limit of anterior radial ridge, ventral margin broadly concave, postero-ventral margin sinuate where ventral rostral ridge meets margin, posterior margin acute, slightly upturned, postero-dorsal margin slightly raised with shallow angulation at limit of hinge plate; hinge plate elongate, relatively broad, acute chevron teeth, 16 on both anterior and posterior hinge plates of specimen 8.3mm total length; ligament internal, amphidetic, triangular, extends slightly ventral to hinge plate.

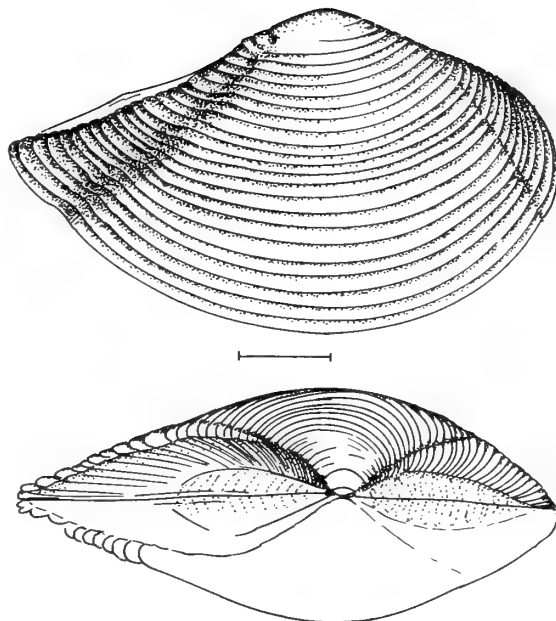


Fig. 34. *Nuculana commutata*. Lateral and dorsal views of a shell, from Sarsia station S29 from the West European Basin. Scale = 1mm. Maximum length of present specimens is 8.3mm.

#### INTERNAL MORPHOLOGY (Fig.35)

The internal morphology differs little from that of *N. acuta* (Fig.32). The most noticeable differences are that *N. commutata* has less attenuate palps with fewer palp ridges and larger adductor muscles than does *N. acuta*.

Other differences between the two species are that in *N. commutata* the ridge from the umbo to the antero-ventral margin is more marked, the apices of the concentric ridges are less broad, the postero-dorsal margin is not so raised and the shell is somewhat less elongate in relation to its height.

These differences are of degree and at that point where separation into species rather than subspecies is a subjective judgement. Nevertheless, these differences are more marked than those between *N. commutata* and *N. illirica* Carrozza 1987 (paratypes BMNH 1995213 examined by JAA), a species that has been recently described from the northern Adriatic Sea (Carrozza, 1987). In contrast *N. commutata*

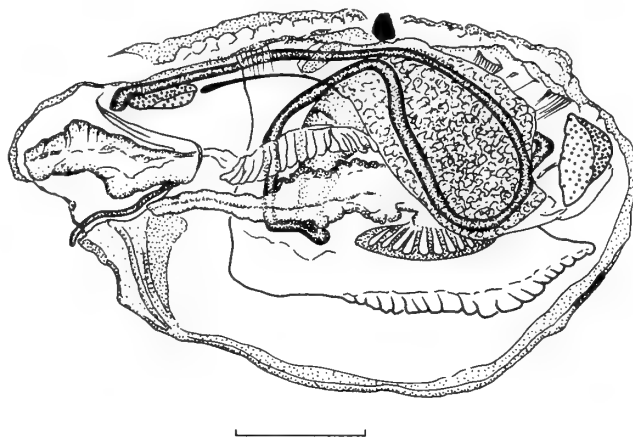


Fig. 35. *Nuculana commutata*. Internal morphology as seen from the right side of a specimen from Sarsia station S29 from the West European Basin. For identification of the parts see text-figure 7, p. 106. Scale = 1mm.

is widely distributed throughout the Mediterranean and Lusitanian Atlantic. Prior to the paper by Carrozza (1987) there had been debate as to whether *N. fragilis* and *N. commutata* were the same species (Locard, 1891, 1898; Bucquoy *et al.*, 1887–98). It is not possible to determine whether this earlier debate was a presage to the study of Carrozza (1987). In contrast *N. acuta* is even more widely distributed off the eastern North America, West Indies and off Brazil (Abbott, 1974). It must be assumed that these are three sibling species.

### *Nuculana vestita* (Locard 1898)

TYPE SPECIMEN. MNHN

TYPE LOCALITY. Talisman stas 96–98 & 101, West of Senegal, 2324–3200m, 19°12'N 17°57'W – 16°38'N 18°24'W

CITED SPECIMENS. BMNH 1995056 and 1995211

*Leda vestita* Locard 1898, 340, pl.XIV, figs 12–18

*Nuculana vestita* Clarke 1962, 53.

*Leda macella* Barnard 1963, 448, fig.11d; type locality: West off Cape Point, S.W.Africa, Africana II stas A190, A192, A317, A319, 2268–3200m, SAM (not seen).

#### MATERIAL:

Cruise	Sta	Depth (m)	No	Lat	Long	Date	Gear
SIERRA LEONE BASIN							
Atlantis II 31	146	2842 –2891	1	10°39.5'N	17°44.5'W	6.2.67	ES
ANGOLA BASIN							
Atlantis II 42	201	1964 –2031	1	09°29.0'S	11°34.0'E	23.5.68	ES
	203	527 –542	742	08°46.0'S	12°47.0'E	23.5.68	ES

Specimens taken by the Galathea Expedition described by Knudsen (1970) examined by JAA, ZMUC. Knudsen (1970) following examination of specimens synonymized *L. macella* with *N. vestita*.

*Nuculana vestita* is a well-described species (Locard, 1898; Theile and Jaekel, 1931; Knudsen, 1970) occurring off West and South-west Africa at lower slope depths (715–2891m) in the Sierra Leone, Guinea and Angola basins.

#### SHELL DESCRIPTION (Figs 36 & 37)

Shell moderately large, inflated, somewhat elongate, rostrate, ornamented with concentric ridges, pale brownish-yellow periostracum; umbos large, inwardly directed, anterior to midline; antero-dorsal distal margin horizontal for short distance, proximal margin broadly convex forming smooth curve with anterior margin, ventral margin broadly convex to rostrum, postero-dorsal margin, raised in small specimens but less so in large, proximally straight or slightly concave, in small specimens angulate at limit of posterior hinge plate, distally slightly concave to posterior limit of rostrum; broad ridge extends from umbo to rostrum forming outer limit of escutcheon; anterior and posterior hinge plates broad, meet shell margin ventral to umbo, hinge teeth broad chevrons, up to 19 and 16 in anterior and posterior series respectively in specimen 8.3mm total length; ligament internal, amphidetic, pear-shaped in sagittal vertical section, extends ventral to hinge plate. The maximum length of the present specimens is 13.8mm.

#### INTERNAL MORPHOLOGY (Fig. 38)

The adductor muscles are moderately large and oval. The siphonal embayment is deep with the siphonal tentacle to the right side. The siphons are entire. The feeding aperture is small but well-defined with the mantle surface ridged internally. Radial mantle muscles are well-developed forming a broad band internal to the inner lobe

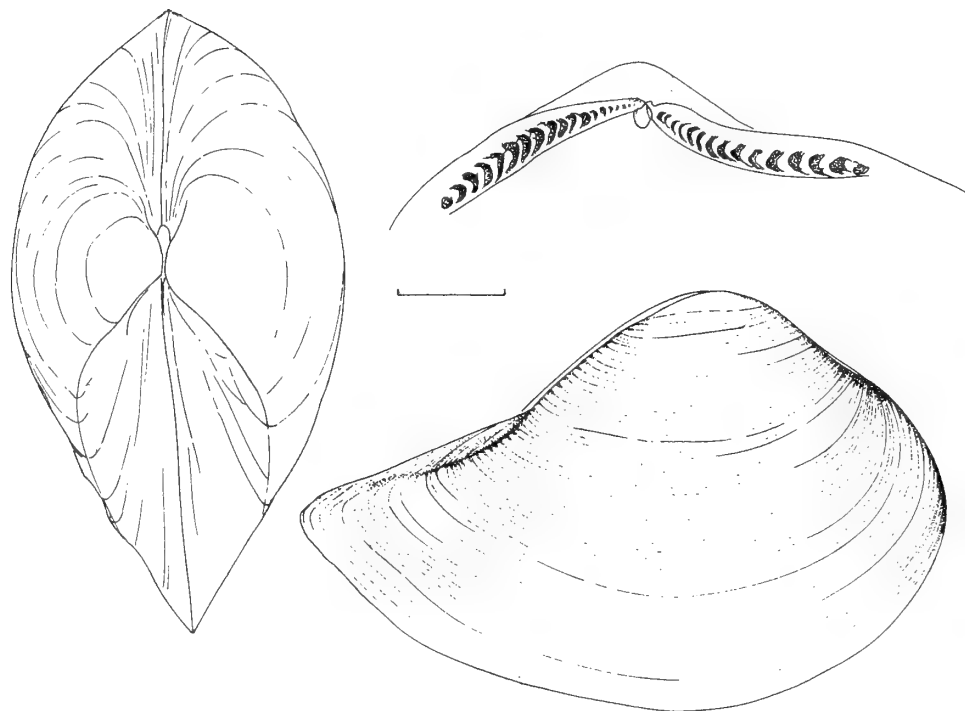
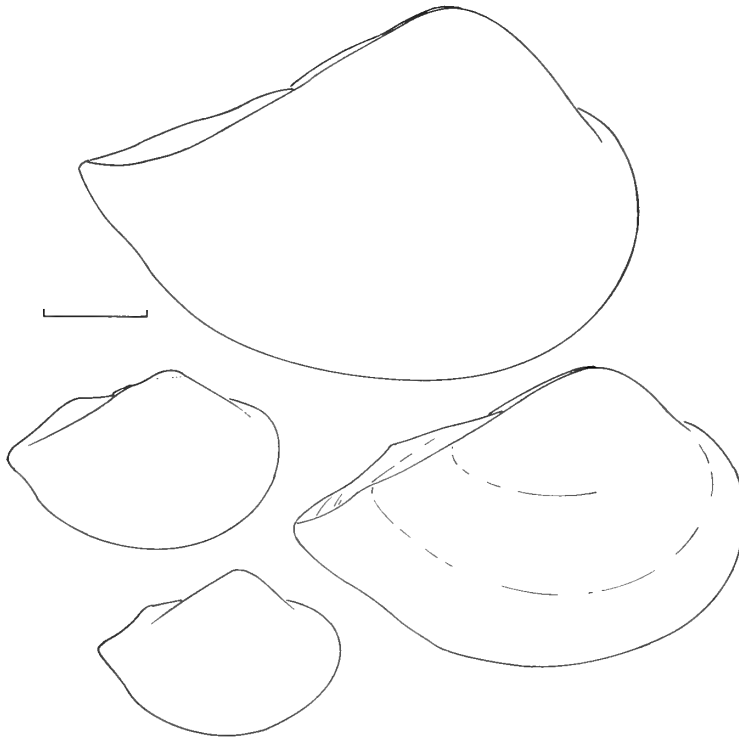


Fig. 36. *Nuculana vestita*. External right lateral and dorsal view of a large adult shell and a lateral view of the hinge plate of a right valve. Specimens are from Atlantis II station 203 from the Angola Basin. Scale = 1mm.





**Fig. 37.** *Nuculana vestita*. Lateral views of right side of four shells in outline to show differences in shape with increasing size. Specimens from Atlantis II station 203 from the Angola Basin. Scale = 1mm.

of the mantle edge. The anterior mantle sense organ is well-developed.

The foot is large with the margins of the sole finely papillate. The palps are very small with up to 12 broad folds. The palp proboscides are stout and elongate, even in the contracted state. The gills are elongate and broad with up to 36 gill plates.

### *Propeleda carpenteri* (Dall 1881)

TYPE SPECIMEN. Syntypes, USNM 63151 and MCZ 7936–7938.

TYPE LOCALITY. Off Barbados, 'Blake' stations 5, 9, and 21, 100fm–287fm.

CITED SPECIMENS. BMNH 1995057

*Leda carpenteri* Dall 1881, 125; 1886, 249, pl.8, fig.10, pl.9, fig.3.

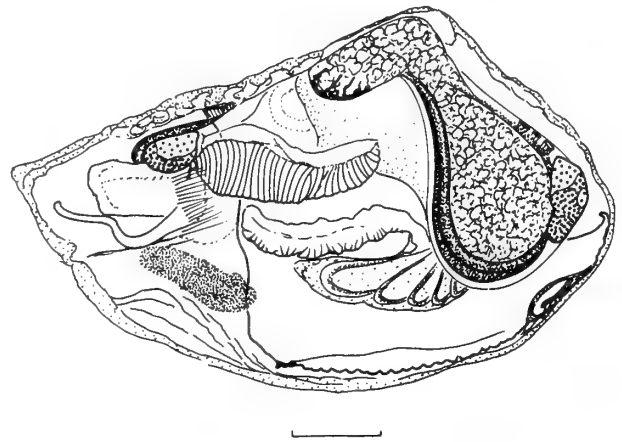
*Nuculana carpenteri* Johnson 1934, 16.

### MATERIAL:

Cruise	Sta	Depth (m)	No	Lat	Long	Date	Gear
ARGENTINE BASIN							
Atlantis II 60	237	993– 1011	194	36°32.6'S	53°23.0'W	11.3.71	ES
	239	1661– 1679	8	36°49.0'S	53°15.4'W	11.3.71	ES
	240	2195– 2323	8	36°53.4'S	53°10.2'W	12.3.71	ES

Specimen USNM 63151 examined by JAA.

Previously reported off N. Carolina, Gulf of Mexico and Eastern Caribbean (Dall, 1889; Rice and Kornicker, 1965; James, 1970), the present specimens are from the Argentine Basin. This species has a



**Fig. 38.** *Nuculana vestita*. Internal morphology as seen from the right side of a specimen from Atlantis II station 203 from the Angola Basin. For identification of the parts see text-figure 7, p. 106. Scale = 1mm.

somewhat unusual distribution from shelf to lower slope depths, 200–2323m

### SHELL DESCRIPTION (Figs 39–41)

Shell fragile, semi-transparent, slender, moderately elongate, rostrate with two post umbonal carinae, ornamented with faint concentric ridges; periostracum pale straw colour; umbos small, far anterior (postumbonal length 60–68% of total length), inwardly facing; antero-dorsal margin slightly flattened, ventral margin smoothly curved, postero-dorsal margin raised, slightly sinuous, concave proximally, convex distally, meets posterior margin at limit of dorsal post-umbonal carina, posterior margin concave between posterior

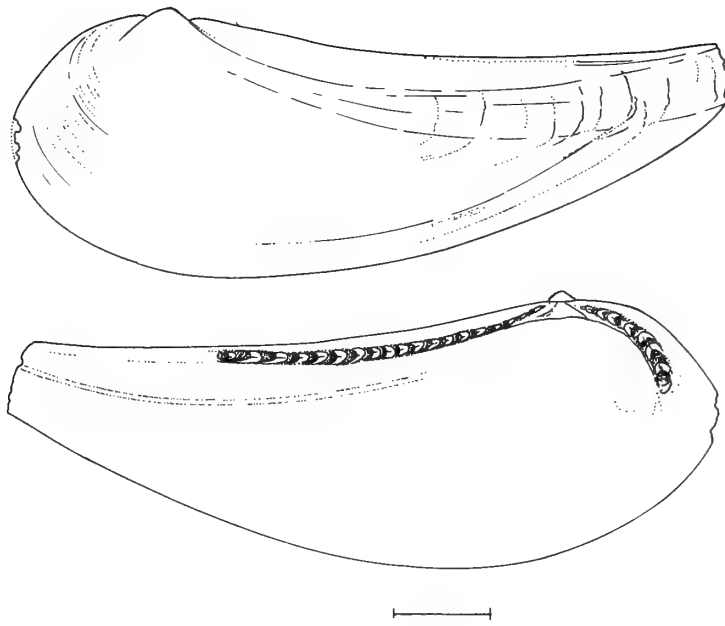


Fig. 39. *Propeleda carpenteri*. External lateral view of the left valve and an internal view of the same valve of a syntype USNM 63151, from off Barbados in 100fms. Scale = 1mm.

limits of dorsal and ventral carinae; escutcheon lanceolate; hinge plates relatively broad, posterior plate short, occupying little more than half of postero-dorsal margin, anterior plate approximately half length of posterior, hinge teeth acute chevron-shape, relatively few in number, up to 18 in posterior series and 16 in anterior; ligament internal, oblique, posterior to umbo; internal shell ridge extends from mid posterior margin to approximately opposite the mid-point of the posterior hinge plate between and parallel to the lines of the

post-umbonal carinae. The maximum length of the present specimens is 15.3mm.

In most specimens the posterior shell margin is damaged, often being markedly shortened and specimens frequently show regeneration of the shell posterior to the posterior adductor. The repaired shell is without concentric ornamentation. It is possible that extended siphons are predated upon and that the shell is damaged when this occurs.

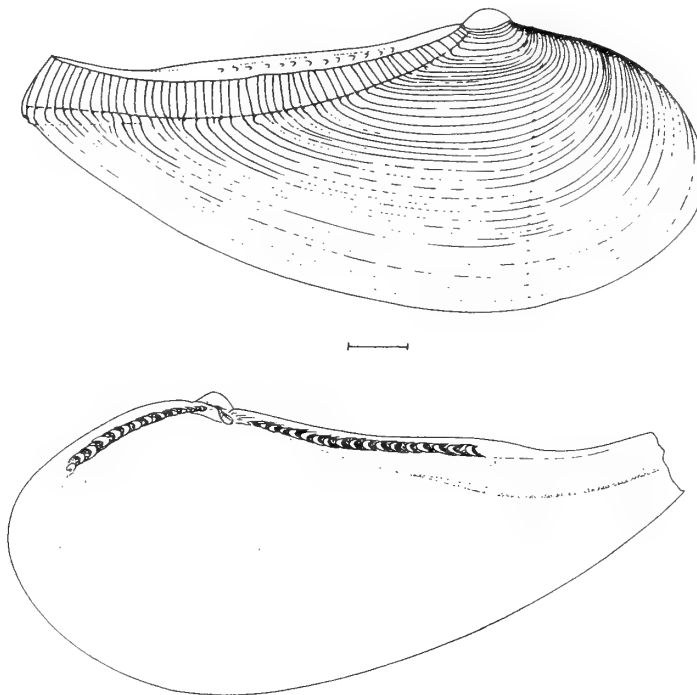
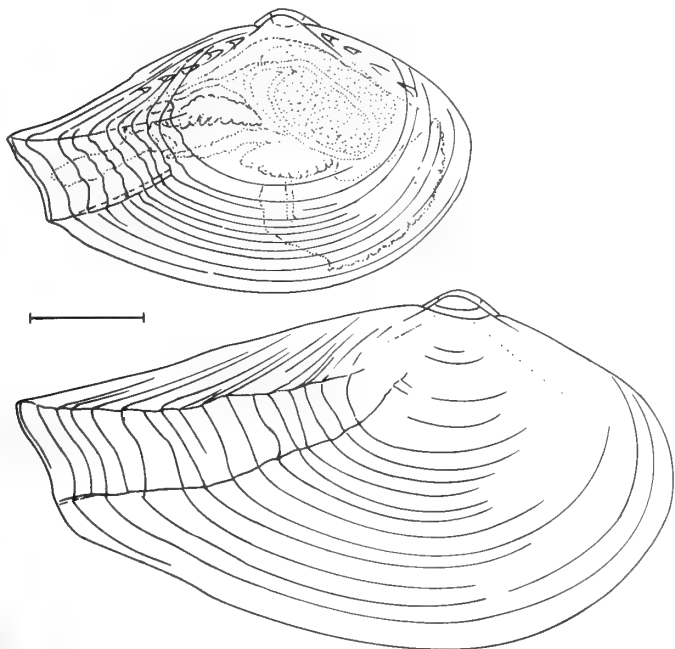


Fig. 40. *Propeleda carpenteri*. External lateral view of the right side of a shell and an internal lateral view of a right valve from Atlantis II station 237 from the Argentine Basin. Scale = 1mm.



**Fig. 41.** *Propeleda carpenteri*. External lateral views of the right sides of two small shells to show differences in shape from the specimen illustrated in text-figure 40. Specimens taken from Atlantis II station 239 from the Argentine Basin. Scale = 1mm.

The concentric ridges on the shell of the present specimens while not particularly marked are more so than those described by Dall, though the syntypes that we have examined are dead valves that are somewhat worn (Fig.39). Our specimens also are very slightly more anteriorly extended than the syntypes, though the characteristic antero-dorsal curvature of the shell margin is the same. Our specimens correspond well with the figures given by James (1972, figs 67 and 68). These latter come are from similar depths (2340–2627m) to some of our own. It would appear that specimens from mid to lower slope differ slightly from those taken at shallower depths however, we regard the differences as being at most infrasubspecific.

There is a marked change in shape during growth. Juveniles are much shorter than the adults and subsequent growth involves increasing elongation of the post-umbonal shell. The prodissoconch is extremely large measuring 630µm in length.

**INTERNAL MORPHOLOGY (Fig.42)**

The adductor muscles are oval in cross-section, the posterior being the more fusiform, both are set some distance in from the shell margin. The posterior adductor muscle lies opposite the limit of the

posterior hinge plate, the anterior muscle is attached to the shell just dorsal of the mid-horizontal shell axis. The siphonal embayment is elongate, the siphons are slender and entire. The anterior sense organ is small in size.

The foot lies in the anterior half of the mantle cavity in preserved specimens, it is relatively elongate and has numerous small papillae present along the margins of the sole. The palps are small, each bearing an extremely long, narrow, palp proboscis. Depending on the size of the specimen there are up to 17 palp ridges. The gills are elongate, slender, and have up to 17 plates.

The course of the hind gut is similar to that in *Nuculana* in that it passes to the right side of the body where it forms a broad loop that passes close to the posterior wall of the anterior adductor muscle. The stomach is large and occupies a vertical position in the posterior part of the visceral mass. The digestive gland is extensive occupying much of the antero-dorsal visceral space.

***Propeleda louiseae* (Clarke 1961)**

**TYPE SPECIMEN.** Holotype, MCZ 224958.

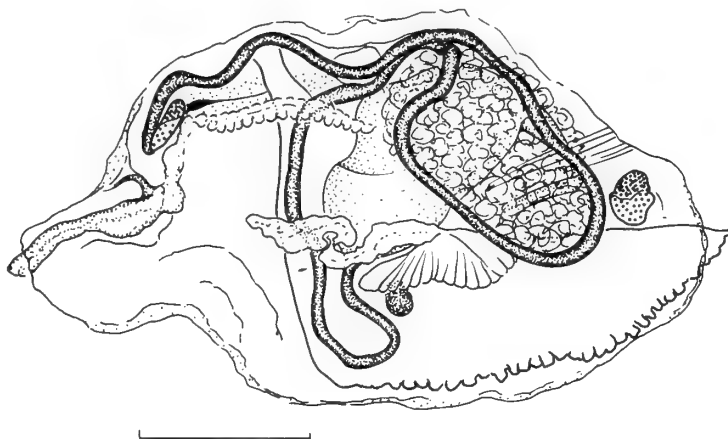
**TYPE LOCALITY.** R.V.Vema biology station 121, Argentine Basin, 1000 miles ESE of Buenos Aries, 5105 metres.

**CITED SPECIMENS.** BMNH 1995058

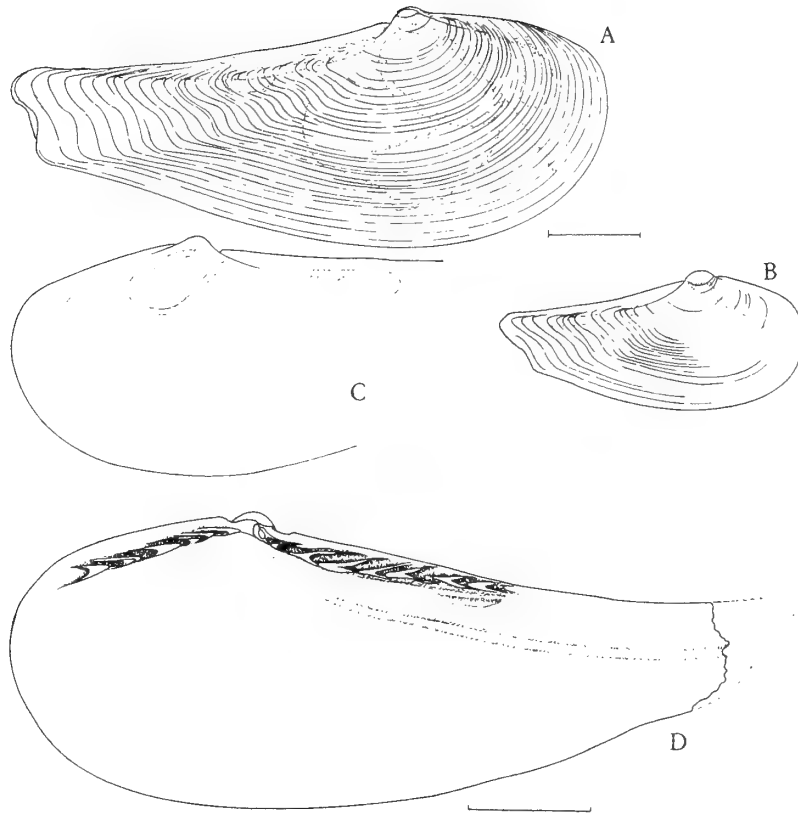
*Nuculana (Thestyleda) louiseae* Clarke 1961, 375, pl.1, fig. 7.

**MATERIAL:**

Cruise	Sta	Depth (m)	No	Lat	Long	Date	Gear
<b>ARGENTINE BASIN</b>							
Atlantis II	242	4382–	25	38°16.9'S	51°56.1'W	13.3.71	ES
60		4402					
	243	3815–	2	37°36.8'S	52°23.6'W	14.3.71	ES
		3822					
	247	5208–	2	43°33.0'S	48°58.1'W	17.3.71	ES
		5223					
	256	3906–	3	37°40.9'S	52°19.3'W	24.3.71	ES
		3917					



**Fig. 42.** *Propeleda carpenteri*. Internal morphology as seen from the right side of a specimen taken from Atlantis II station 239 from the Argentine Basin. For identification of the parts see text-figure 7, p. 106. Scale = 0.5mm.



**Fig. 43.** *Propeleda louiseae*. a & b, external lateral views of the right sides of shells of differing size to show differences in shell proportions with growth; note outline internal morphology through semi-transparent shells, in particular the form of the hind gut and position of the adductor muscles; c, outline of shell from the left side showing the outline of hind gut and adductor muscles; d, internal view of a left valve. All specimens taken from Atlantis II station 242 from the Argentine Basin. Scales = 1mm.

Type specimen examined by HLS.

Distributed at abyssal depths in the Argentine Basin; depth range: 3815–5223 metres.

#### SHELL DESCRIPTION (Fig.43)

Shell extremely elongate, slender, fragile, semi-transparent, ornamented with moderately spaced concentric ridges, two post-umbonal rounded ridges, one dorsal and one ventral at posterior shell margin and crossed vertically by wavy continuations of the concentric ridges, faint anterior radial ridge from umbo to antero-ventral margin; umbo slightly raised, far anterior (post-umbonal length 65–70% of total length), inwardly facing; antero-dorsal margin with short proximal notch, distally margin raised and slightly concave, faint angulation before anterior margin, anterior margin joins with ventral margin in smooth curve, postero-ventral margin very slightly sinuate, posterior margin angled and sinuate, postero-dorsal margin notched at umbo, distally somewhat raised and keeled, concave overall but slightly sinuous dorsal to hinge plate; hinge plate relatively broad, posterior plate short occupying approximately half the postero-dorsal shell margin, hinge teeth elongate, acute chevron shape, up to 12 in anterior and 20 in posterior series; ligament small, internal, oblique, pear-shaped; rounded internal ridge extends from umbo to posterior margin and marks junction between inhalant and exhalant siphons. The maximum length of the present specimens is 20.3mm. The prodissoconch is large measuring 300µm in length.

Clarke (1961) records a long, thin, external ligament in the type

specimen – the latter being a single large valve. We find no evidence of an external part to the ligament and believe that Clarke mistakenly confused periostracum along the postero-dorsal margin for ligament.

#### INTERNAL MORPHOLOGY

With one notable exception the internal morphology of *P. louiseae* differs little from that of *P. carpenteri*.

Unlike the latter species the hind gut of *P. louiseae* first passes to the left side of the body where it forms a relatively small loop immediately ventral to the umbo (Fig.43). From there it passes to the right of the body and forms a loop that is considerably larger than that on the left although not as extensive as that in *P. carpenteri* (Figs 42 & 43).

The adductor muscles are relatively large, the elongate posterior muscle is situated at the distal limit of the posterior hinge plate. The gill is very short and slender with few (c. 13) gill-plates.

#### *Propeleda paucistriata* (new species)

TYPE SPECIMEN. Holotype BMNH 1995059; Paratypes BMNH 1995060.

TYPE LOCALITY. Atlantis II station 203, Angola Basin, 08°48.00'S 12°52.00'E, 527–542m.

## MATERIAL:

Cruise	Sta	Depth (m)	No	Lat	Long	Date	Gear
ANGOLA BASIN							
Atlantis II	203	527–	31	08°48.00'S	12°52.00'E	23.5.68	ES
42		542					

**DISTRIBUTION.** Restricted to the Angola Basin at upper slope depths, 527–542 metres.

**SHELL DESCRIPTION** (Fig.44)

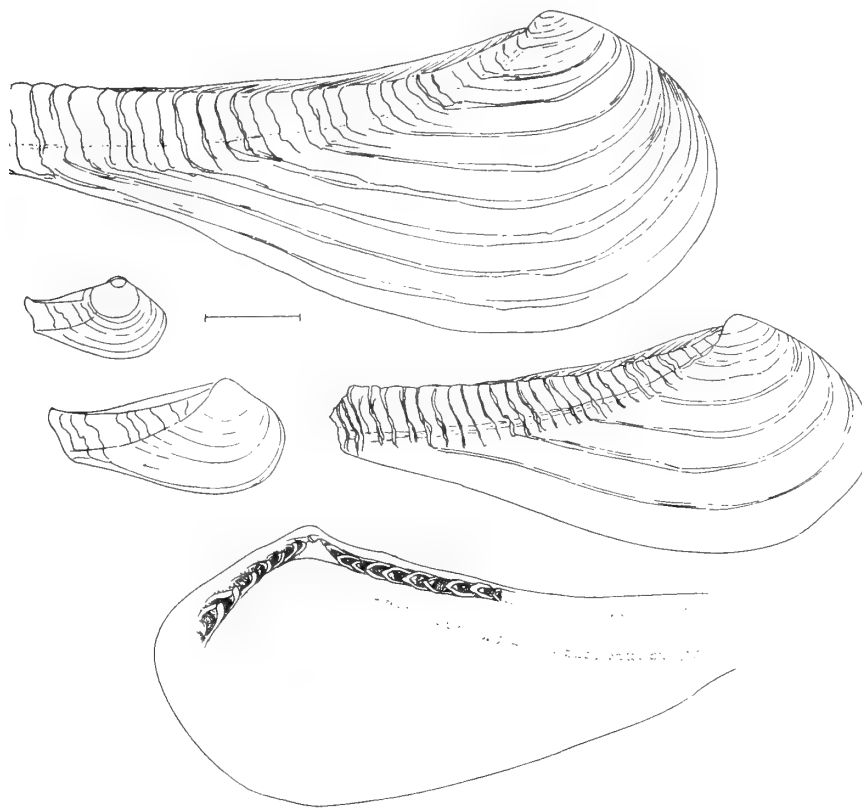
Elongate, fragile, translucent shell, moderately slender, two carinate ribs from umbo to posterior margin, widely spaced prominent, relatively broad, concentric ribs with overhanging ventral margin, 2–4 fine concentric lines between ribs, between carinae vertical ribs and lines equally prominent; umbo moderately raised, far anterior in largest specimens (post-umbonal length 79% of total length) but less so in smaller specimens, beaks inwardly facing; antero-dorsal margin sloping, proximally convex but almost straight in small specimens, joins with anterior and antero-ventral margins in smooth curve; postero-ventral margin very slightly sinuous, posterior margin usually damaged in large specimens, intact margin angled and sinuate, forming a hook dorsally where postero-dorsal margin and dorsal carina meet, postero-dorsal margin concave, proximally raised, elongate escutcheon outlined by dorsal carina; hinge elongate, moderately broad, large, acute chevron-shaped teeth up to 16 in anterior series and up to 28 in posterior series, anterior series extends to anterior limit of anterior adductor muscle, posterior series extends approximately half length of postero-dorsal margin to anterior limit

of posterior adductor muscle, ventral margin of hinge plate corresponds to line of dorsal carina, ventral to umbo teeth approach shell margin, anterior and posterior hinge plates continuous; ligament internal, ventral to umbo and close to shell margin, slightly inclined posteriorly; rounded internal ridge extends from umbo to posterior margin. The maximum recorded shell length is 14.1mm. The prodissoconch is very large and measuring 560µm in length.

Juvenile shells are more ovate and, before posterior elongation occurs, could be mistaken for a yoldiellid (Fig.44).

**INTERNAL MORPHOLOGY** (Fig.45)

The anterior adductor muscle is oval in cross section, while the posterior adductor is smaller and more elongate. Both are set in from the shell margin, the posterior is positioned at approximately two-thirds the distance between the umbo and the posterior limit of the shell. There is a small anterior sense organ formed from the sensory fold of the mantle, ventral to the anterior adductor. The siphons are joined with their ventral margins fused to form entire lumina. They are slender and particularly elongate and when contracted are contained in the elongate siphonal embayment. The foot and viscera lie in the anterior half of the mantle cavity. The foot is elongate and directed anteriorly. In most preserved specimens the tip of the foot lies between the anterior adductor and the shell margin. The margins of the sole are fringed with numerous relatively small papillae. There are three anterior and two posterior pedal retractor muscles. The palps and gills are markedly narrow and elongate. The are at least 22 palp ridges in the largest specimens and the palp proboscides are attenuate each with a straight dorsal margin and a papillate ventral margin. In a few preserved specimens the palp proboscides extend from the feeding aperture. The gills are similarly attenuate



**Fig. 44.** *Propeleda paucistriata*. External lateral views of the right sides of four shells of differing size to show change of shape with growth and an internal view of a right valve. All specimens taken from Atlantis II station 203 from the Angola Basin. Scale = 1mm.

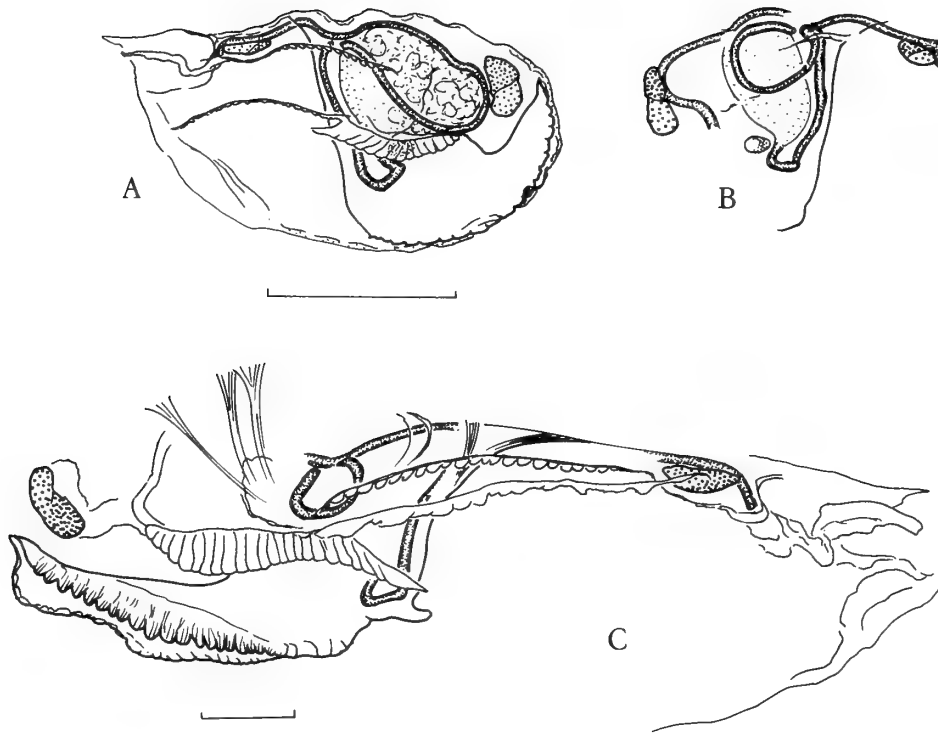


Fig. 45. *Propeleda paucistriata*. a, internal morphology as seen from the right side; b, part of the left side of the same specimen to show details of the course of the gut; c, the internal morphology as seen from the left side of a much larger specimen. All specimens taken from Atlantis II station 203 from the Angola Basin. For identification of the parts see text-figure 7, p. 106. Scales = 1mm.

and extend from the posterior visceral mass to the anterior limit of the posterior adductor. There are at least 22 gill plates in larger specimens. A slender extension of the axis extends from each gill from ventral to the posterior adductor to the inner junction between inhalent and exhalent siphons.

From the large combined stomach and style sac the course of the hind gut takes it first dorsal and posterior to the stomach and then to the left side of the body where it makes a small loop. From there it passes ventral to the umbo to the right side of the body where it makes a much larger loop at the perimeter of the viscera and passing close to the inner face of the anterior adductor. From there it passes mid-dorsally to the anus. The mouth is set some distance posterior to the anterior adductor muscle. The oesophagus is broad and elongate and the combined stomach and style sac is positioned vertically in the posterior part of the visceral mass. The pedal ganglion is large and lies immediately anterior to the junction of mid gut and hind gut.

The shell surface in some larger specimens is covered with epifaunal solitary hydroids. This would indicate that *P. paucistriata* lives close to the surface of the sediment. This is also suggested by the fact that the posterior tips of the shells of larger specimens are broken. We believe that this is the result of predation on the siphons.

The extreme post-adductor elongation of the shell is advantageous in that it provides distance between predator and the more vulnerable viscera with damage being restricted to more easily generated tissue.

The shell of this species differs from others described by the small number of pronounced concentric shell ridges and we name it with reference to this characteristic feature. This is the first species of *Propeleda* to be recorded off the south-west coast of Africa. At approximately the same latitude off the east coast of Natal a species named *Leda lanceta* by Boshoff (1968) occurs at upper slope depths. Nijssen-Meyer (1972) believes that this latter species is a *Propeleda*,

and we concur with her. *P. lanceta* is more robust, more arcuate and with far more numerous concentric ridges than is the case in *P. paucistriata*.

## DISCUSSION

The major point of interest in this particular account of deep-sea protobranch bivalves is the evidence it provides to further our understanding of the evolution of the nuculanoid protobranchs. In our earlier studies on the Tindariidae we speculated as to how the nuculanoids could have evolved from the nuculoids (Sanders and Allen, 1977). In functional terms, this involved a change in the inflow of water into the mantle cavity from an antero-ventral position to a posterior position and the beginning of specialization of the posterior mantle edge, a view also expressed by Yonge (1939). In the tindariids this latter involves the development of sensory papillae from the sensory fold of the mantle at the points of ingress and exit of the circulatory water. Although infaunal and deposit feeding, the tindariids, like the nuculids, live close to the surface and, like many other bivalves that occupy this position, they are ovate and robust. Many of these subsurface dwelling bivalves, including the tindariids, have stout external ligaments.

The development of siphons was the next step in the evolutionary process and the neilonellids are illustrative of this. The shell form and ligament as seen in the tindariids is largely retained in the neilonellids, but short siphons, as yet only fused dorsally, are now present and these are contained in a shallow siphonal embayment. Although the shell remains stout and ovate there is some posterior elongation and an area ventral to the inhalent siphon from where the palp proboscides are extended is more defined. Like the tindariids the neilonellids are deposit feeders living close to the surface of the sediment.

The hind gut in *Neilonella*, like that in *Tindaria*, has a wide lumen and single pronounced typhlosole. Although the course that the hind gut takes in neilonellids makes a single loop on the right side of the body, it does not penetrate mantle space as it does in tindariids (Sanders and Allen, 1977). In this respect the hind gut of neilonellids probably represents the more primitive condition. We have argued elsewhere (Allen, 1992) that elongation and the complexity of form of hind gut configuration are related to food procurement at great depths, and this applies to the tindariids (Sanders and Allen, 1977). The neilonellids are for the most part upper slope species and the hind gut would be expected to be less specialized and less elongate.

In *Nuculana* posterior elongation becomes more extreme and the ventral margins of the combined siphons are fused such that the exhalent and inhalent lumina are separate and entire. The shell remains robust, but is more slender. The ligament is restricted to a small internal structure separating elongated anterior and posterior series of hinge teeth. We believe that elongation is correlated to the almost vertical orientation of the animal in the sediment but which retains contact with the surface via the extended posterior body and siphons. The genus *Nuculana* is found mainly in shelf and upper slope sediments and as such the available food resources are relatively abundant. The hind gut is not greatly extended and remains as a single loop to the right side of the body.

In *Propeleda* the evolutionary trend of posterior body elongation seen in *Nuculana* becomes more extreme, particularly posterior to the posterior adductor muscle. The posterior adductor muscle is more elongate and dorso-ventrally narrow, and the gill axes, siphons and the palp proboscides are exceptionally long and slender. The shell of *Propeleda*, particularly in abyssal species, is much more fragile and is further specialized in that it possesses an internal posterior longitudinal ridge. The function of this ridge is not entirely clear and has await examination of the living animal but, possibly, it is involved in the separation of excretion, feeding and respiratory functions in the extremely elongate posterior mantle cavity. It may also help to strengthen the otherwise very fragile shell and assist in predation damage limitation. In *Propeleda* post-adductor elongation involves body tissues that can be relatively easily regenerated, much in the same way as has been reported in deposit feeding tellinids (Edwards, Steele and Trevallion, 1970). Specimens showing shell repair posterior to the posterior adductor are present in our samples.

The evolution of the Ledellinae and an assessment of their functional morphology was discussed earlier (Allen and Hannah, 1989). In respect of the species of *Ledella* and *Tindariopsis* described here, little needs to be added to that account other than to note, again, that the hind gut in these abyssal protobranchs is extraordinarily lengthened and takes the most complex courses within the visceral mass.

The other item of note is the description of yet another ledellid in which the shell, after reaching a certain length, changes its direction of growth. In *Ledella aberrata* as in *L. ultima* the result of this change is to produce a broad shell margin and lateral expansion of the shell cavity. This adaptation has been construed as possibly providing more space for the gonads that begin to develop at about the time the change in direction occurs.

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