



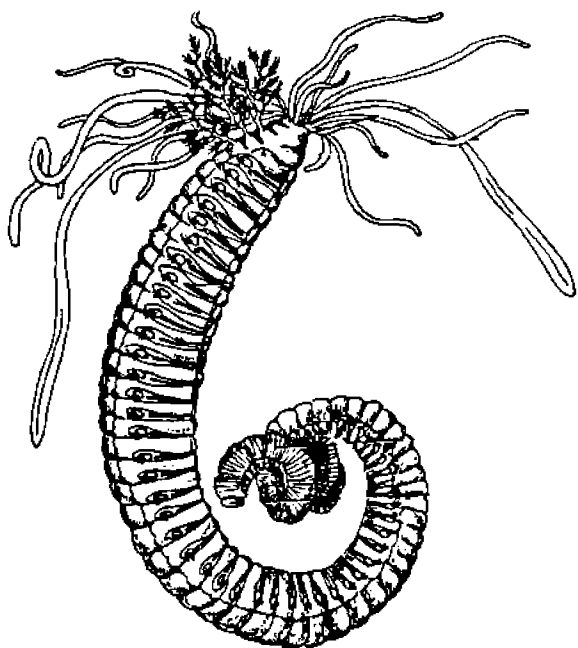
**Southern California Association of
Marine Invertebrate Taxonomists**

3720 Stephen White Drive
San Pedro, California 90731

October, 1993

Vol. 12, No. 6

NEXT MEETING:	Corymorphine Hydroids
GUEST SPEAKER:	John Ljubenkov, MEC Analytical Systems Inc., Carlsbad, CA
DATE:	November 15, 1993 (note third Monday)
TIME:	9:30am-3:00pm
LOCATION:	MEC Analytical Systems Inc., Carlsbad, CA (map is included)



NOVEMBER 15 MEETING

The meeting in November will be on Corymorphine Hydroids of southern California. In addition, some of the unique cnidarians generated from the master species list will be reviewed. Please bring any cnidarians you need to have identified or confirmed. The workshop will be led by John Ljubenkov of MEC Analytical Systems Inc. It will be held at MEC Analytical Systems Inc., Carlsbad, CA.

Amphitrite ornata from Invert. Zoology (Sec. Ed.) by
Paul A. Meglitsch

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Scamit Newsletter is not deemed to be a valid publication for formal taxonomic purposes.

**MINUTES FROM MEETING ON
OCTOBER 19**

The SCAMIT Christmas party has been scheduled for December 11th at the Cabrillo Marine Aquarium, San Pedro, CA. If anyone is interested in organizing the party and coming up with a theme please do not hesitate to do so.

The City of San Diego is pleased to announce four new employees. They are Laura Essex, Ami Groce, Megan Lilly, and Rick Rowe.

Don Cadien (Los Angeles County Sanitation Districts) informed attending members about new literature; the Amphipod Newsletter 19 and Amphipods, a noble obsession: Essays in memory of J. Laurens Barnard (1928-1991), Journal of Natural History 27(4): 723-988.

Included in this newsletter is a list of publications available from De L'Institut Oceanographique in Paris, France.

Also included is a call for abstracts for the 1994 Water Environment Federation 67th Annual Conference and Exposition in Chicago, Ill.

Don Cadien chaired the workshop on Anthurid Isopods. Included in this newsletter is a handout prepared by Don and Richard C. Brusca. If anyone has any comments please send them to Don at LA County Sanitation Districts, Marine Biology Lab., 24501 S. Figueroa St., Carson, Ca 90745, (310) 775-2351 ext. 403. He will be modifying the key for future reference.

FUTURE MEETINGS

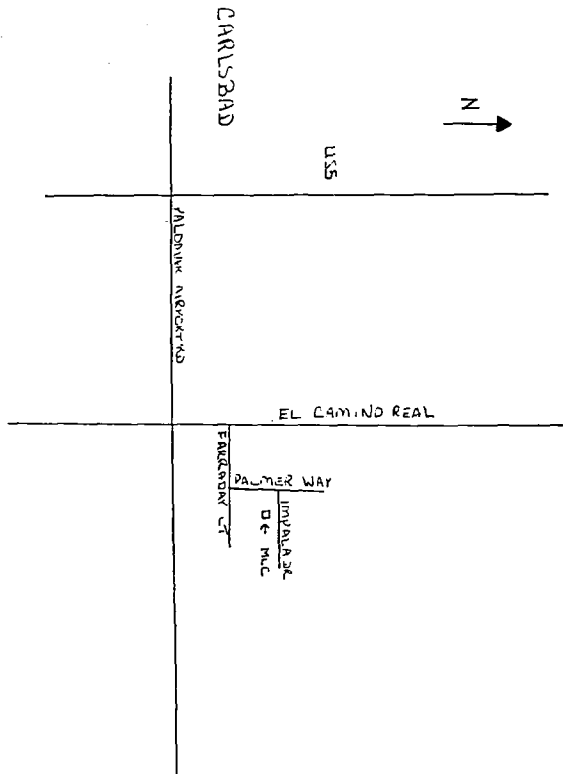
The December 13 meeting will be a show and tell with polychaete specimens that are weird, strange, or rare from the recently generated species list. There will also be some discussion on what SCAMIT's responsibility will be for the species list, how we can use it and whether we can distribute it. Tentatively the meeting will be held at Kirk Fitzhugh's polychaete lab at the Los Angeles Natural History Museum.

SCAMIT OFFICERS:

If you need any other information concerning SCAMIT please feel free to contact any of the officers.

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MAP TO MAP THE ECOLOGICAL CONSULTANTS (MEC)

ANTHURIDEAN ISOPODS (CRUSTACEA) OF CALIFORNIA AND THE TEMPERATE NORTHEAST PACIFIC

Don Cadien and Richard C. Brusca
(presented at the October 19, 1993 meeting of SCAMIT)

I. Introduction

Literature on the anthuridean isopod fauna of California and the northeast Pacific has not recently been synthesized. Since the most recent comprehensive report (Schultz 1977) family and generic level reviews have altered the nomenclature of several species. Environmental survey and monitoring programs have generated many new geographic and bathymetric distributional records for eastern Pacific anthurideans, most as yet unpublished, and have collected several undescribed species. The current review was undertaken to update and standardize anthuridean taxonomy in California, and to disseminate information derived from a variety of unpublished sources.

II. Definition of the Group

Isopods of the suborder Anthuridea are most easily recognized by their slender elongate bodies (usually 7 or more times longer than wide), lateral uropods that curve up and over the pleotelson, and presence of (usually) one or two pleotelsonic statocysts (Fig. 1.1). Unlike most isopods, anthurideans are not much flattened dorso-ventrally and are circular or oval in cross-section. According to Brusca and Wilson (1991), the specific defining synapomorphies of the Anthuridea are: mandible without distinct lacinia mobilis or spine row, instead with a lamina dentata (which may be secondarily lost in some species); maxillae reduced, minute, fused to paragnath (or lost entirely); coxae of maxillipeds fused to head; maxillipedal endite without coupling spines; and uropodal exopod folded dorsally over pleotelson.

Brusca and Wilson (1991) placed this suborder within the "flabelliferan complex" (the Flabellifera *sensu lato*). The suborder contains four families, all now known to occur in the temperate northeast Pacific. Most species achieve a moderate size (8-15mm length), but a few are much smaller (4mm) or larger (45mm). Most anthurideans are marine, but some genera have marine, brackish, and freshwater members (e.g. *Cyathura*), some are exclusively freshwater (e.g. *Cruregens*), some are primarily stygofaunal (e.g. *Stygocyathura*), and some are primarily anchialine or interstitial (e.g. *Curassanthura*). About 200 species have been described, but this is almost certainly only a small percentage (probably less than half) of the world fauna.

III. Aspects of Anthuridean Biology

Reproduction - Mature male and female anthurideans are easily separated by secondary sexual characters, particularly the enlarged, multiarticulate, aesthetasc-fringed flagellum of the male first antenna (Fig. 2). In some cases males and females differ so greatly in gross morphology that they were initially described as separate species. The apparent separation of sexes can, however, be misleading, as protogynous sequential hermaphroditism occurs in many species. In others, as in some tanaids (Buckle-Ramirez 1965), there is also male polymorphy, with some animals always male and some males developing secondarily from post-brood females (Legrand & Juchalt 1963, Burbanck & Burbanck 1974, 1979).

Sex ratio in collections of anthurids is often skewed strongly towards females and juveniles, with few adult males (Kensley & Schotte 1989), although in *Apanthura* the reverse may be true. Examining a collection of several hundred *Apanthura* from tropical Australia, Poore and Lew Ton (1988b) noted no oostegite bearing females, and they suggested reproduction in this genus might deviate from the normal anthuridean pattern. Seasonal fluctuations of sex ratio in some species appear related to protogynous hermaphroditism (Burbanck & Burbanck 1979).

As in other isopods, the gonopores are located ventrally on the sternite of the fifth pereonite of the female and the seventh pereonite of the male. The inner ramus (endopod) of the male second pleopod also bears an appendix masculina as in other isopods. These structures assist in sperm transfer between the penile papillae of the male and the gonopore of the female. Their structure can be useful in anthuridean taxonomy, but details are unknown for most species, and may vary within a species due to male polymorphy.

Fertilization may occur in the ovary as in sphaeromatids (Shuster 1991) or may take place in the oviduct, before the eggs pass out through the gonopore and into the marsupium following molting (while the exoskeleton is still elastic). Eggs in the marsupium are already fertilized (at least in *Cyathura*) since they are encased in a vitelline coat lacking a micropyle for admission of sperm (Strömberg 1972). The marsupium is formed by paired oostegites on pereonites 2 or 3 through 5. Once in the marsupium the young undergo epimorphic development, eventually leaving as manca. Manca exit with only six pairs of pereopods and thus can be differentiated from post-manca juveniles with seven pairs. The genera *Cruregens* and *Colanthura* are neotenous, and have the formation of the seventh pereopod suppressed even in the adult.

Growth - No information is available on growth rates or molting frequencies for any eastern Pacific anthurideans. In the Atlantic species *Cyathura carinata* growth rate is dependent on temperature and food availability, and growth ceases during reproduction (Bamber 1985). Rate of growth declines with age in *Cyathura carinata* (Bamber 1985), but information is lacking on eastern Pacific species.

Feeding - There are two types of mouth parts in anthurids; those modified for piercing and sucking (Fig. 1.2), and those adapted for biting and chewing (Fig. 1.3). Chewing/biting mouth parts are used to feed either on detritus (Schultz 1977) or on living prey (Wägele 1981). Burbanck & Burbanck (1979) reported that while normally feeding on detritus, *Cyathura polita* may also consume both live and dead polychaetes, oligochaetes, amphipods, shrimp, and fish when the opportunity arises. Piercing mouth parts occur only in the family Paranthuridae, and are associated largely with species living among and feeding upon algae (Schultz 1977). Feeding ecology of eastern Pacific species has not been studied.

Habitats - Anthurideans are important and often abundant components of the offshore soft-sediment marine environment. Most live in sediment burrows or tubes, or within algal mats, habits in agreement with their narrow and elongate bodies. They may excavate burrows themselves (Fig. 3.1), or move into tubes or burrows abandoned by other organisms (e.g. *Cyathura polita* - Burbanck & Burbanck 1979). Anthurideans from hard substrates may live in crevices or fissures, in holes formed by other species, or in the attached tubes of other organisms (Wägele 1981). They are often found associated with littoral and sublittoral algae. *Eisothistos* sp. A lives among the incomplete septae at the eroded bases of the colonial coral *Coenocyathus bowersi*. A few other local species live outside of burrows or other shelter, finding adequate concealment among the tangled thalli of filamentous algae (e.g. *Paranthura elegans*), or among the rhizomes of seagrasses.

Family Hyssuridae

Hyssuridae gen. A, sp. A [MBC, 1984] Formerly reported as *Apanthura* sp. A; see comments below.

Family Anthuridae

Amakusanthura californiensis (Schultz, 1964)** Formerly placed in *Apanthura* and *Apanthuretta*; see comments below.

Calathura branchiata (Stimpson, 1855) Formerly placed in *Anthura*; see comments below.

Cyathura carinata (Kroyer, 1849). Originally placed in *Anthura*; see comments below.

Cyathura munda Menzies, 1951*

Eisothistos sp. A [MBC, 1984]. Formerly reported as *Heteranthura* sp. A; see comments below.

Eisothistos sp. B [Cadien, 1990]

Haliophasma geminatum Menzies & Barnard, 1959** Formerly placed in *Silophasma*; see comments below.

Mesanthura occidentalis Menzies & Barnard, 1959**

IV. Comments on Individual Species (listed alphabetically)

Amakusanthura californiensis (Fig. 4). The brief original description (Schultz 1964) was based on a lot of twelve females from "several to 11mm long," taken from black mud at a depth of 80m off Santa Monica, California. Schultz (1977) was aware of no additional records of the species, and we are aware of no other published records since the original description. However, this species has been collected in several environmental monitoring programs from southern California. It also occurred in samples from west Mexico taken during Allan Hancock Foundation cruises, ranging as far south as Isla Guadalupe (pers. obsv., LACMNH collections). The species was transferred from *Apanthura* Stebbing, to *Apanthuretta* Wägele by Poore & Lew Ton (1985), and subsequently to *Amakusanthura* Nunomura when *Apanthuretta* itself was synonymized (Poore & Lew Ton 1988b). The holotype of *A. californiensis* has been reexamined and inaccuracies and omissions in the original description are being corrected (Wetzer & Brusca, in press). Most importantly, pleonites 1-5 are dorsally fused along the midline, and the maxillipedal endite is broad and lobelike.

Ananthura luna (Fig. 5). *Bathura* Schultz was originally differentiated from *Ananthura* Barnard by a low tooth on the palm of the first pereopod, by the characteristic broadly-radiating setal clusters at the distal tips of the uropodal rami and pleotelson, and by the lack of serrations on the outer margins of the uropodal endopods (Schultz 1966). Kensley (1978) deemed these characters insufficient to support separate generic status and synonymized both *Bathura* and *Ananthura* with *Anthelura* Norman and Stebbing. These genera were later reexamined by Poore and Lew Ton (1988d), who separated *Ananthura* and *Anthelura* on the basis of their statocysts. *Bathura*, which was described with two statocysts, was reevaluated as having one central statocyst with a slitlike dorsal pore, as in *Ananthura*. Although this feature was not interpreted as a statocyst by Schultz, it was clearly indicated in his illustration of the holotype. *Ananthura luna* is a large species (to 21mm length; Schultz 1977) that is infrequently encountered in relatively deep water (783-1298m) off the southern California borderland between the Coronado and Santa Monica Submarine Canyons. It may also occur in shallower water around canyon heads, based on a sample from Santa Monica Bay (taken in 78m) in the LACMNH collection.

Calathura branchiata (Fig. 6) was originally described from New Brunswick (eastern Canada) by Stimpson (as *Anthura branchiata*), and has since become the senior synonym for two of G.O. Sars' northeast Atlantic species (*Paranthura norvegica* Sars and *Paranthura arctica* Sars). Gurjanova's (1936) record of *C. branchiata* from north Pacific, from the Sea of Okhotsk and the Bering Sea, and Coyle and

Predators - Many fishes are known to feed on the west Atlantic estuarine species *Cyathura polita*, as do blue crabs (Burbanck and Burbank 1979). Predation by crabs, and other invertebrates is likely for eastern Pacific species, but has not been documented. In an evaluation of trophic relationships between fishes and benthic invertebrates at Catalina Island, Hobson & Chess (Ms.) found 11 fishes feeding on anthuridean isopods. Forty-one anthurids were found in the guts of 28 fish. Most of the isopods were consumed by three species; black surfperch *Embiotoca jacksoni* (6 guts, 10 isopods), blackeye goby *Coryphopterus nicholsi* (5 guts, 8 isopods), and California sheephead *Pimelometopon pulchrum* (5 guts, 10 isopods). Species taking anthurideans at lower frequencies were rock wrasse *Halichoeres semicinctus*, señorita *Oxyjulis californicus*, kelp surfperch *Brachyistius frenatus*, island kelpfish *Alloclinus holderi*, garibaldi *Hypsypops rubicundus*, halfmoon *Medialuna californica*, kelp bass *Paralabrax clathratus*, and blue-banded goby *Lythrypnus dalli*.

Anthurideans are slow compared to many other peracarids, and they swim only clumsily. Outside their refuges their movements are awkward, and they are probably easy prey to predatory nemertean, annelids, and other arthropods. Despite the lack of special protective or offensive structures, some anthurideans respond aggressively to attack. If seized from behind, *Paranthura elegans* will twist around and strike at its attacker (pers. obsv.). Perhaps this aggressive response is sufficient to deter some would-be predators.

The relatively indurated pleotelsonic region of many anthurideans apparently serves as an operculum to block access to certain tube or burrow-dwelling species. Observations on living *Eisothistos* (Wägele 1981) indicated that they adopt a head down position in serpulid worm tubes while feeding on the original occupant. This leaves the ornamented pleotelson and uropods in the position of the worm's operculum (Fig. 3.2). Foraminiferans and sponges observed attached to the tail-fan of *Eisothistos* sp B suggest they may move little once established in a tube, thus minimizing exposure to predators.

IV. Anthuridea of the West Coast of North America (North of Mexico)

Apart from *Edanthura linearis* Boone, 1923, the first anthurideans known from the northeastern temperate Pacific were those described by Menzies (1951). The temperate fauna of the northeast Pacific currently contains at least 15 recognizable species. Three are undescribed species and 12 are nominate species, of which one is a *nomen nudum* (*Paranthura linearis*), one may be a misidentification or incorrect locality record (*Paranthura algicola*), and one is clearly a questionable record (*Cyathura carinata*). Holotypes (***) or paratypes (*) of most of these species are in the collection of the Los Angeles County Museum of Natural History (LACMNH) (Wetzer et al. 1991), as noted below. North of California, anthuridean isopods are both less common and less diverse. No members of this suborder were reported by Richardson (1905) for the northeast Pacific, by Hatch (1947) from Washington, or by George & Strömberg (1968) from Puget Sound. In a detailed environmental analysis of benthic communities in Puget Sound, Lie (1968) reported *Haliophasma geminatum*, and three other species have since been reported from the northeast Pacific: *Cyathura carinata*, *Calathura branchiata*, and *Eisothistos* sp B.

Family Antheluridae

Ananthura luna (Schultz, 1966)** Formerly placed in *Bathura*; see comments below.

Family Paranthuridae

Califanthura squamosissima (Menzies, 1951)* Formerly placed in *Colanthura*; see comments below.

Colanthura bruscai Poore, 1984*

Paranthura algicola Nunomura, 1978 Questionable species; see comments below.

Paranthura elegans Menzies, 1951*

Paranthura linearis *nomen nudum*. Formerly placed in *Edanthura*; see comments below.

Müller's (1981) record from the Gulf of Alaska, establish this species as circum-north Pacific in distribution. It has not been reported south of Alaska, and its reported depth range is 20-1500m

Califanthura squamosissima (Fig. 7). Schultz (1977) sunk *Colanthura* Richardson, on the basis of a supposed synonymy of *Colanthura tenuis* Richardson (the type species) and *Paranthura infundibulata* Richardson, and he erected *Califanthura* as a replacement genus for *Colanthura squamosissima*. Poore (1980), however, resurrected *Colanthura*, declaring both it and *C. tenuis* to be valid taxa. Poore's conclusion was based, in part, on a reexamination of the types of *C. tenuis* and *P. infundibulata* by Kensley, who also did not substantiate their synonymy (in Poore 1980). Although Poore's (1980) move sunk Schultz' *Califanthura* into *Colanthura*, he later (Poore 1984) reestablished it as a valid genus, which now contains six species worldwide. *C. squamosissima* is a small species, reaching only about 5.2mm in length. It occurs in shallow water (18-90m) from Dillon Beach, California (Schultz 1977) to Magdalena Bay, west Baja California (Nunomura 1978), and has also been collected intertidally at Morro Bay and La Jolla.

Colanthura bruscai (Fig. 8) is similar to *C. squamosissima* in general appearance and size. However, it is predominantly Panamic in distribution, with its northernmost occurrence at San Clemente, California (Poore 1984), and from there ranging south to at least Costa Rica. It occurs intertidally at most locations, although some northern records are subtidal to a maximum depth of 27m. The maximum reported length is 5.4mm (Poore 1984).

Cyathura carinata is a northern European species. Bernard's (1978:576) record from the Strait of Georgia (British Columbia, Canada), if accurate, may reflect a relict north Pacific population from a former circumboreal distribution. However, because there are no other reports of this well-known Atlantic species from the Pacific Ocean, this unpublished Pacific record needs confirmation. The record in Austin (1985) presumably is derived from Bernard's report. This species was originally placed in *Anthura*; and transferred to *Cyathura* by Norman and Stebbing (1886). It is not included in our key.

Cyathura munda (Fig. 9) is a moderate size (to 9 mm), narrow (length more than 9 times width) species, usually associated with brown algal holdfasts on hard substrates. The type material from northern California was all taken from the holdfasts of *Egregia* and *Laminaria*. All the subtidal records of Menzies & Barnard (1959) are from stations where the samples were noted to contain either kelp or rocks (Allan Hancock Foundation, 1965). LACMNH material of this species usually indicates collection from kelp or from surfgrass (*Phyllospadix*). This species has been taken from the intertidal zone (Menzies 1951) to 58m (Menzies and Barnard 1959), from Tomales Point to the Mexican border, and in the Gulf of California. More recent collections in the Santa Maria Basin extend the depth range down to 132m on rocks. Brusca and Iverson (1985) described a very similar species from intertidal habitats on the Pacific coast of Costa Rica (*C. guaroensis*).

Eisothistos sp. A. A single juvenile specimen (1.4mm) of this species was taken off Tajiguas, Santa Barbara Co., California at 77m depth, in the washings of rocks retrieved during a submersible dive in 1984. It was initially called by the unpublished name *Heteranthura* sp. A. However, Wägele (1981) synonymized *Heteranthura* Kensley and *Eisothistos* Haswell, hence the generic reassignment. This specimen, while clearly not belonging to any other eastern Pacific anthuridean species, is not sufficiently adult to compare with other species of *Eisothistos*, of which there are over a dozen worldwide. Additional specimens were later taken by Hans Kuck (LACMNH) in 1989, in association with colonies of the coral *Coenocyathus bowersi* collected at 5-8m depth off the eastern shore of Catalina Island. These specimens were larger (2-2.5mm), but still not fully adult. In gross morphology this species is similar to *Eisothistos antarcticus* as described by Wägele (1984b), with serrate uropodal and pleotelsonic margins, and a single row of spines down the middle of the pleotelson. The range of this undescribed California species, as currently known, is 5-77m, Tajiguas to Catalina Island. The genus *Eisothistos* was recently transferred from Hyssuridae to Anthuridae by Poore & Lew Ton (1988c).

Eisothistos sp. B was encountered in environmental monitoring samples from Alaska related to the Exxon Valdez oil spill. Although the exact locations of the sampling sites were unavailable because of litigation, the animals were collected between the intertidal zone and 10m depth somewhere in Prince William Sound. Numerous specimens were taken from the tubes of serpulid polychaetes, a common habitat for members of this genus. This species resembles both *Eisothistos* sp. A and *Eisothistos minutus* (Sivertsen and Holthuis, 1980) of the tropical east Atlantic. Post-brood adult females, which undergo elongation of pereonites 2-6 (Fig. 10) as described for other species (Wägele 1981), may reach 5mm in length. This species has not been recorded from California waters.

Haliophasma geminatum (Fig. 11). Schultz (1977) erected a new genus (*Silophasma*) for this species, which the revision of Poore (1975) had placed beyond the bounds of a redefined *Haliophasma* Haswell. Subsequently, the definition of *Haliophasma* was expanded such that *Silophasma* was no longer needed, and it fell into synonymy with *Haliophasma* (see Negoescu and Wägele 1984 and Poore and Lew Ton 1988a). Poore (1975) changed the spelling of the trivial name from "geminata" to "geminatum" to match the gender of the generic name. Schultz (1977) gave 7mm as maximum size for *Haliophasma geminatum*, but we have seen specimens from California as large as 12mm in length. This species ranges from Monterey, California (Iverson 1974) to San Quintin Bay, Baja California, Mexico (Menzies 1962) over a broad depth range (9-512m). Lie (1968) also recorded it from Puget Sound.

Hyssuridae gen. A sp. A (Fig. 12). Collections made in the western Santa Barbara Channel and in the Santa Maria basin in central California encountered scattered specimens of this small species (5-6mm length). This may be the same as the "Anthurid n. sp. & n. gen." reported but not well described by Menzies (1962) from off San Quintin Bay, Baja California. In his discussion, Menzies indicated a close affinity to *Kupellonura* for his specimens, but felt they might constitute a new genus. The present material matches the characters Menzies noted: indurated pleotelson with a ventral keel, separation of all pleonal segments, antennal flagellum article counts, and details of the uropods. Menzies did not illustrate his material, and nothing in his brief discussion is unique enough to definitely establish identity between his material and our own. Redefinition of the genera of the Hyssuridae by Poore and Lew Ton (1988c) places the current material close to both *Kupellonura* Barnard and *Hyssura* Norman and Stebbing. One might be inclined to assign it to *Kupellonura* because of the presence of lobes on the lateral margins of the uropodal exopods, a unique synapomorphy for this genus (Poore and Lew Ton 1988c). It also possess a triangular carpus on pereopods IV-VII, whereas the carpus of *Hyssura* species is rectangular in shape. However, the mouth parts are more characteristic of *Hyssura* in that the mandibular molar process is acute (not blunt, as is characteristic of *Kupellonura*), and the maxillipedal endite is short, reaching only the second palp article (rather than the third article, as is typical of *Kupellonura*). One of the specimens of this species we examined had a 4-articulate flagellum on the left antenna and an 8-articulate flagellum on the right. Other than our own observations and Menzies' possible record, this species has not been reported from the northeast Pacific. Our material came from a sample taken off the southeast end of San Miguel Island, and from seven MMS sampling stations between Oso Flaco and the north side of Anacapa Island, from 47 to 166m.

Mesanthura occidentalis (Fig. 13). The original description of this species distinguishes it solely on the basis of the dorsal pigmentation pattern. Illustrations of the pleotelson apex, the maxilliped, the antennae, and the last three articles of the first pereopod were provided, but not discussed. This was amplified by description of a paratype, with a more complete illustration of its antennae, mouth parts, and appendages by Wägele (1984a). Although taken subtidally by grab, the 7mm holotype female came from a sample containing kelp fragments and red algae. Menzies and Barnard (1959) recorded this species from two localities (Point Conception and Point Fermin, California), both containing either kelp or rock, and both from shallow water (12-20m). An additional lot was reported by Schultz (1964) from off Palos Verdes, also in shallow water (20m). Schultz (1977) later gave this species' range as "Point Conception to San Quintin Bay, Baja California" and "from shallow water to 55m deep," perhaps a transcription error of

earlier literature. The records of Menzies and Barnard (1959) suggest that this is a shallow-water species, probably associated with either macroalgal holdfasts, or with algal mats or turf. Brusca (1980) reported a similar appearing congener (*Mesanthura* sp.) from intertidal algal mats in the Gulf of California that may, in time, prove to be a variant of *M. occidentalis*. *Mesanthura nubifera* Wägele, 1984, also from intertidal habitats in the Gulf of California, does not match the pigmentation of Brusca's (1980) species.

Paranthura algicola (Fig. 14) was described by Nunomura (1978) on the basis of two female specimens (5.5mm and 10mm in length) sent to him by Waldo Schmitt in the 1970's. The locality was given as simply a "rocky beach in California, washed from algae, 24 November 1916." Judging by Nunomura's illustrations, his animals may have been *Paranthura elegans* showing the effects of long-time preservation. Nunomura stated that *P. algicola* differed from *P. elegans* in having: "eyes with scattered ocelli" [sic], pleonites medially fused, and by the "shape of the posterior border of the sixth pleonal somite." In fact, the eyes of *P. elegans* are large with many ommatidia and could easily appear as figured and described by Nunomura after many years of preservation; the pleonites are free in *P. elegans* but the articulations are very faint and can easily be mistaken as being fused; and, we see no significant differences between these two species in the posterior margin of the sixth pleonite (aside from what could be attributed to poor renditions by both Nunomura and Menzies). Nunomura's description and figures are difficult to interpret, but the type material was reported as being at the USNM and should be reexamined to establish the correct disposition of this species. We did not include this species in the key that follows. Nunomura (1978) also described another species of *Paranthura*, which he gave the unfortunate name of *P. californiae*, from Magdalena Bay (Baja California, Mexico) that closely resembles *P. elegans*.

Paranthura elegans (Fig. 15) ranges from Dillon Beach at least to San Quintin Bay (west coast of Baja California, Mexico), from the intertidal zone to a depth of 55m (Schultz 1977), and also throughout the Gulf of California (Brusca 1980). It frequents algal mats and clumps, mud bottoms, encrusted pier pilings, and rocky low intertidal habitats. Adults reach about 9.5mm in length in California waters, but are larger in the warmer waters of the Gulf of California (8-15mm). Differences in adult size along a latitudinal gradient are not uncommon, and have been reported for idoteid isopods in the eastern Pacific (Brusca and Wallerstein 1979, Wallerstein and Brusca 1982), and for *Cyathura polita* on the east coast of America (Burbanck and Burbanck 1979).

Paranthura linearis has remained enigmatic since its description (as *Edanthura linearis*). Boone (1923) reported this animal from Laguna Beach, California. She described its mouth parts only as "well developed, unique"; perhaps accurate but certainly imprecise. Menzies (1951) considered *Edanthura* Boone a synonym of *Paranthura* Bate and Westwood, and also recommended *E. linearis* be reduced to *nomen nudum* status. Poore (1984) and Negoescu & Wägele (1984) apparently agreed with these assignments. The type has not been found at the USNM (where Boone indicated it had been deposited), and its whereabouts remains unknown. This species is not included in the key that follows.

V. Key to the Species of Anthuridea Known from the Northeast Pacific (North of Mexico)

1. Mouth parts adapted for piercing and sucking, together forming an anteriorly directed cone-like structure under the head; maxillipedal palps long, thin, and tapering; mandibular incisor smooth, styliform, not toothed; mandible without molar process or lamina dentata; 0 or 1 statocyst in pleotelson; first pleopods enlarged and operculate to others 2
- Mouth parts adapted for biting and chewing, not forming a conelike structure; maxillipedal palps broad; mandibular incisor often toothed; mandible usually with molar process and lamina dentata; 0, 1, or 2 pleotelsonic statocysts; first pleopods may or may not be operculate to others 4

2. Pereonite 7 at least 50% as long as 6; seventh pereopods present *Paranthura elegans*
 - Pereonite 7 less than 20% as long as 6; seventh pereopods absent 3
3. Pereonite 1 twice as long as 2; pleonites free, not fused *Colanthura bruscai*
 - Pereonites 1 and 2 subequal; pleonites fused dorsally *Califanthura squamosissima*
4. With no statocysts in pleotelson; first pleopods not enlarged and operculate to others; body extremely elongate, about 15 times longer than wide (Hyssuridae) Hyssuridae gen. A, sp. A
 - With 0, 1 or 2 statocysts in pleotelson; first pleopods always enlarged and operculate to pleopods 2-5; body length 6-10 times width 5
5. With 1 pleotelsonic statocyst; maxillipedal endite and palp very wide; pleonites 1-5 entirely free never fused dorsally (Antheluridae); the only known California anthelurid is blind and its uropodal tips bear radiating setal clusters *Ananthura luna*
 - With 0, 1 or 2 pleotelsonic statocysts; maxillipedal endite and palp normal, not especially broad; pleonites 1-5 free or dorsally fused (Anthuridae) 6
6. Pleotelson with a dorsal median spine row 7
 - Pleotelson smooth or ridged, but without dorsal spines 8
7. Uropodal endopod with distolateral margin more or less evenly serrate *Eisothistos* sp. A
 - Uropodal endopod with distolateral margin divided into two cusps by three prominent denticles, evenly serrate between these points *Eisothistos* sp. B
8. Pleonites 1-5 completely free and separate in both dorsal and lateral view
 *Calathura branchiata*
 - Pleonites 1-5 completely fused or fused mediodorsally, although segments may be visible in lateral view 9
9. Carpus of pereopods 4-7 rectangular; pleotelson with three raised dorsal longitudinal ridges *Haliophasma geminatum*
 - Carpus of pereopods 4-7 triangular; pleotelson without dorsal ridges 10
10. Pleonites 1-5 fused only along dorsal midline, segments free laterally; uropodal endopods narrow (<60% of pleotelson width), exopods much shorter than either pleotelson or endopods
 *Amakusanthura californiensis*
 - Pleonites 1-5 completely fused dorsally, segmentation indicated in lateral view only by faint lines and setal bundles; uropodal endopods subequal in width to pleotelson, exopods nearly as long as pleotelson and endopods 11
11. Maxillipedal palp 3-articulate; pereonites pigmented dorsally, with complete or nearly complete dark ovals on pereonites 2-6 *Mesanthura occidentalis*
 - Maxillipedal palp 2-articulate; pereonites pigmented dorsally with dark splotches, but without pigment rings *Cyathura munda*

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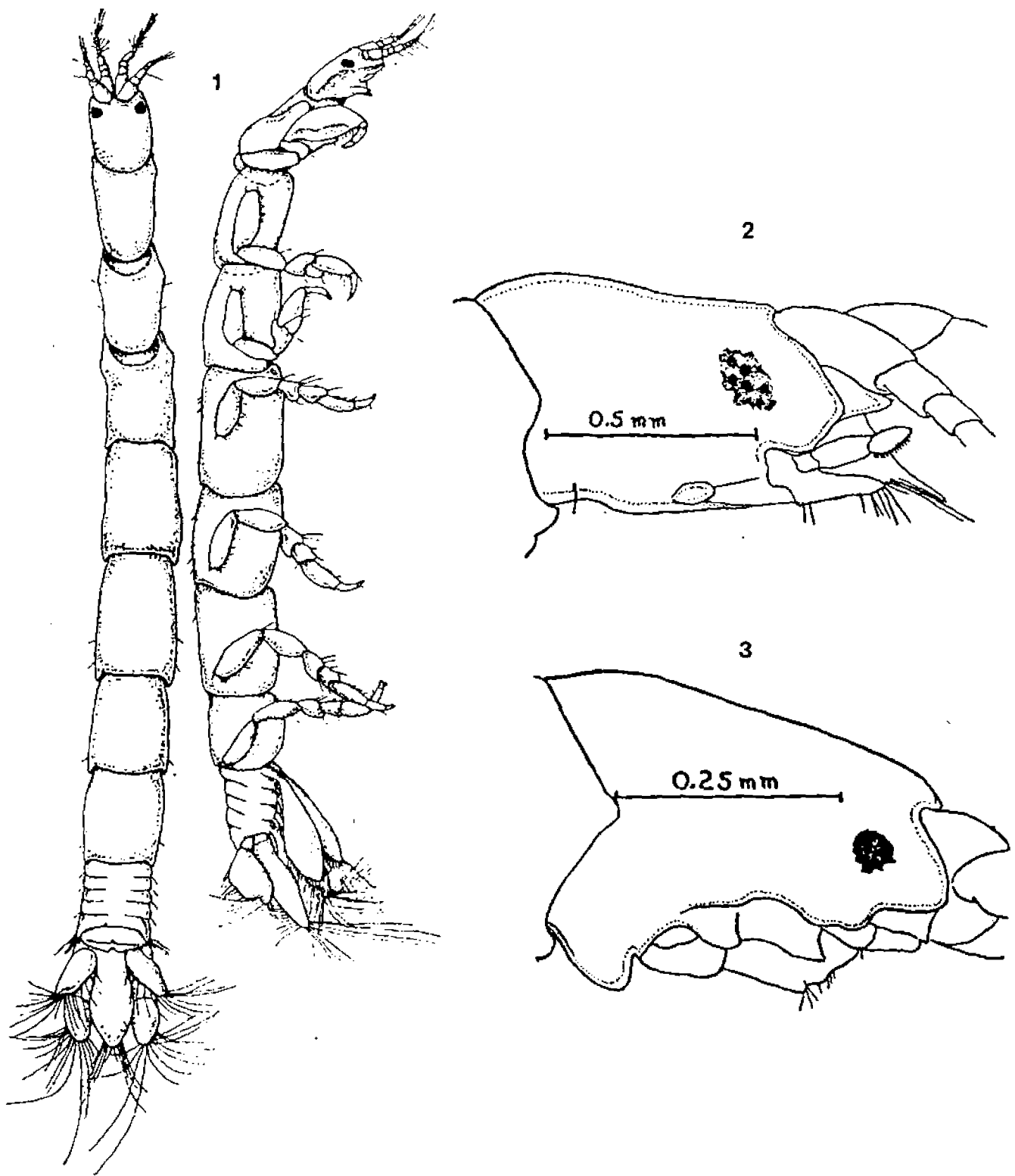


FIGURE 1. 1) Dorsal and lateral views of *Aphantura libyana* ♀ (from Negoescu 1980); 2) piercing mouthparts of *Paranthura elegans* (from Menzies 1951); 3) biting mouthparts of *Cyathura munda* (from Menzies 1951)

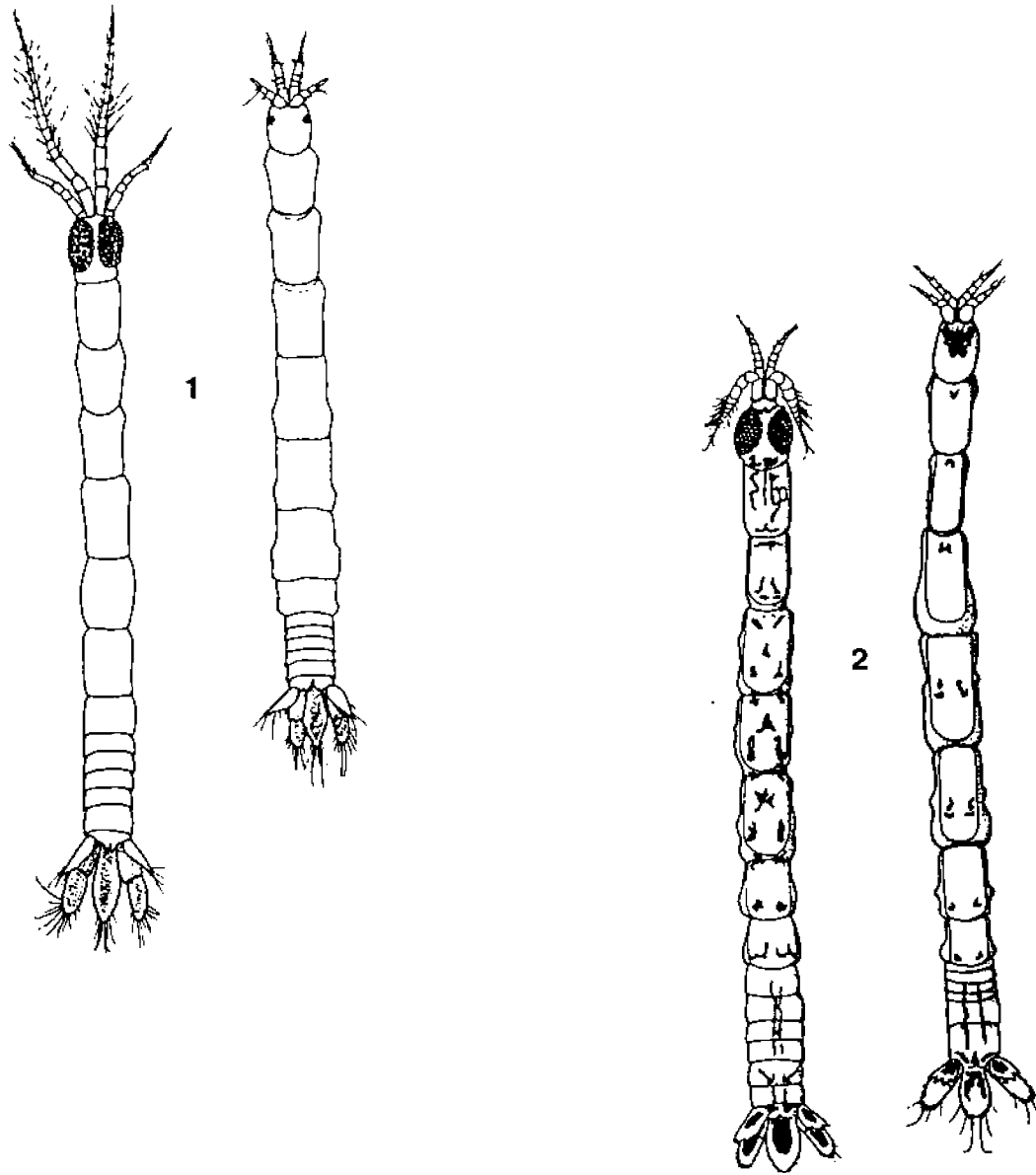
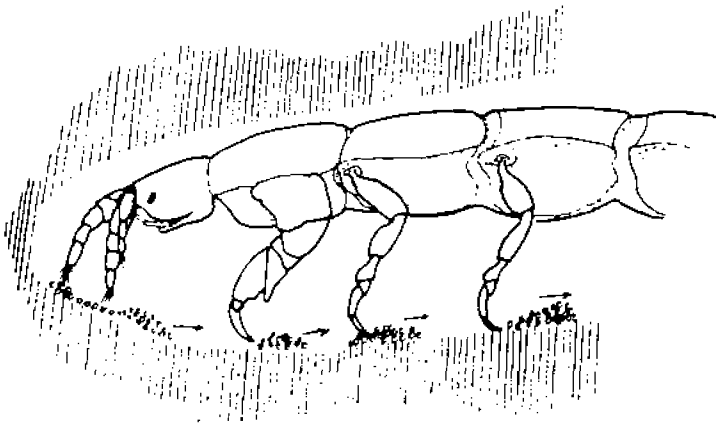


FIGURE 2. Anthurid sexual dimorphism. Male/female pairs of 1) *Chalixanthura lewisi* and 2) *Chalixanthura scopulosa* (male to the left of each pair)(from Kensley & Schotte 1989)



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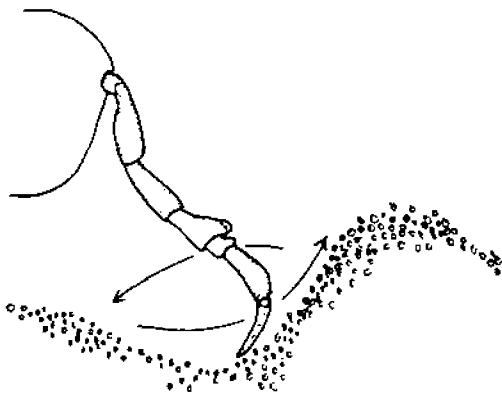
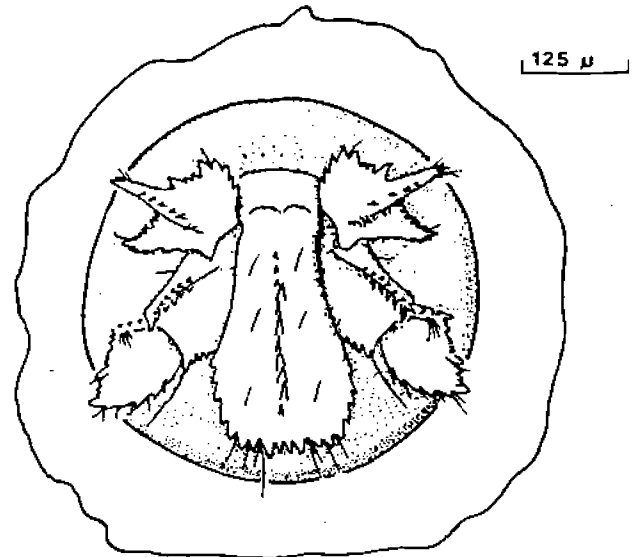
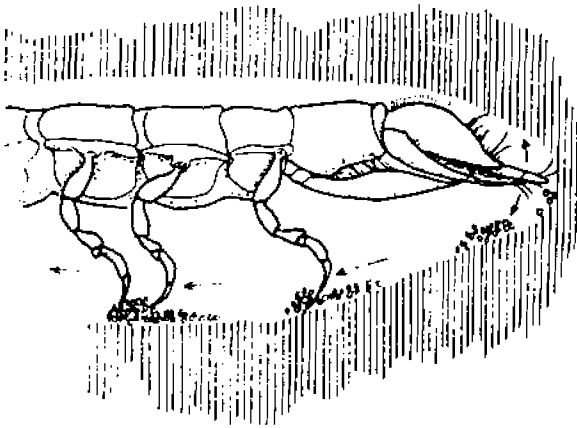


FIGURE 3. 1) *Cyathura carinata* excavating a burrow with anterior appendages, posterior appendages and pleotelson, and pereopods (from Wägele 1981); 2) tail fan of *Eisothisτος macrurus* at the aperture of a serpulid tube (from Wägele 1981)

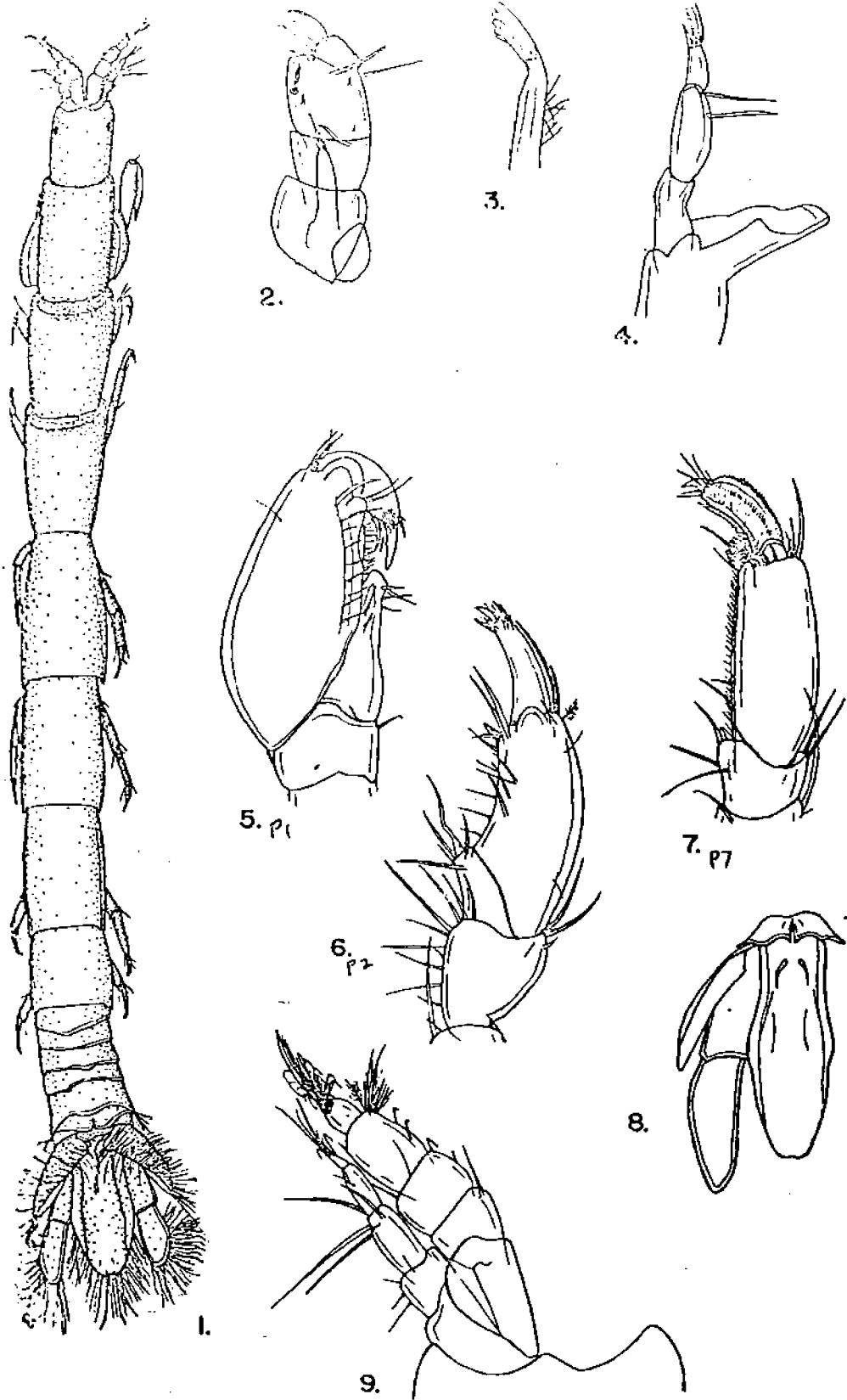


FIGURE 4. *Amakusanthura californiensis* - 1) ♀ whole animal, dorsal view; 2) maxilliped; 3) maxilla 1; 4) mandible; 5) pereopod 1; 6) pereopod 2; 7) pereopod 7; 8) tail fan; 9) antennae (from Schultz 1964)

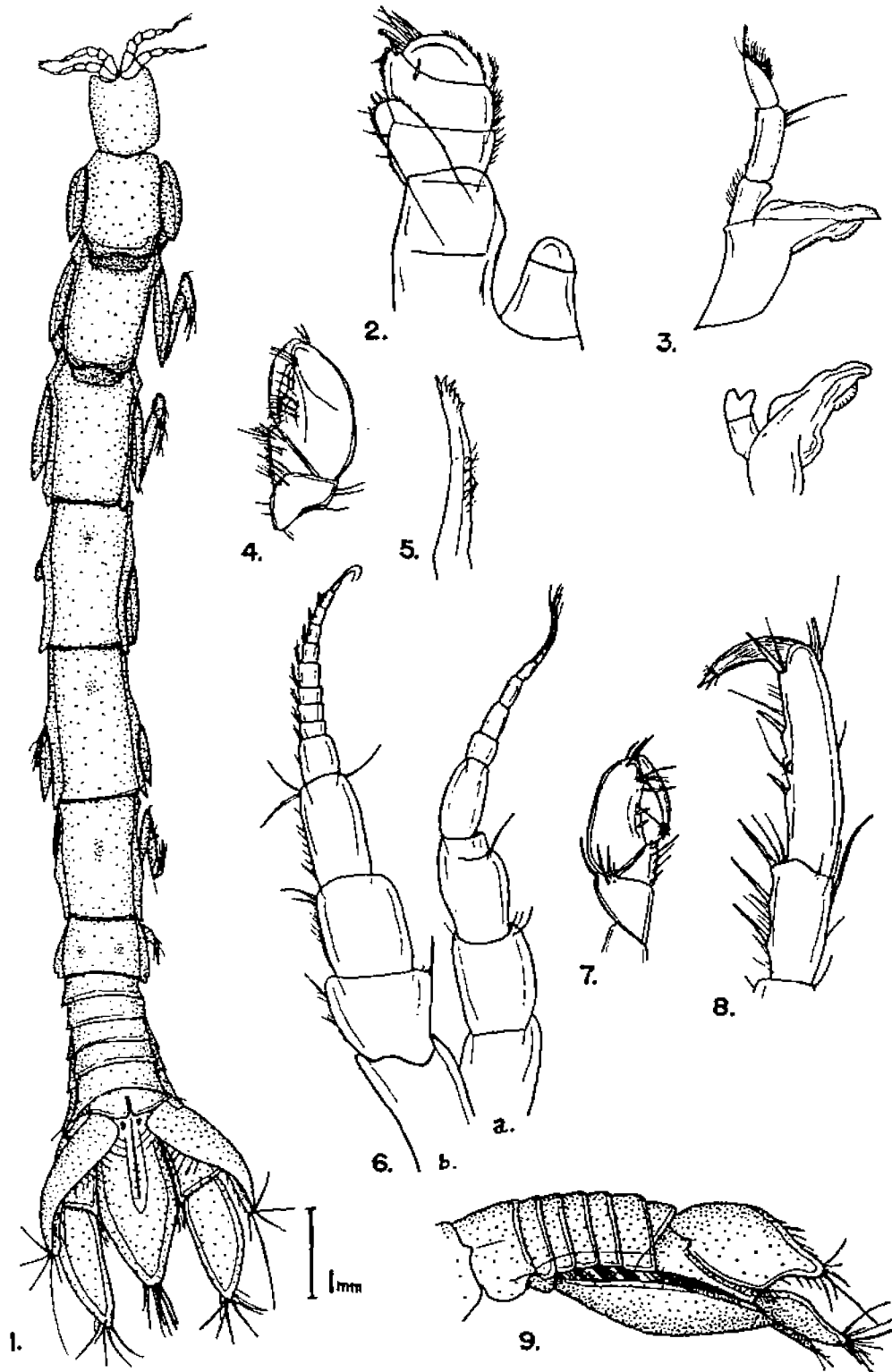


FIGURE 5. *Ananthura luna* - 1) ♀ whole animal, dorsal view; 2) maxilliped; 3) mandible; 4) pereopod 1; 5) maxilla 1; 6) antennae; 7) pereopod 2; 8) pereopod 7; 9) lateral view of pleon and tail fan (from Schultz 1966)

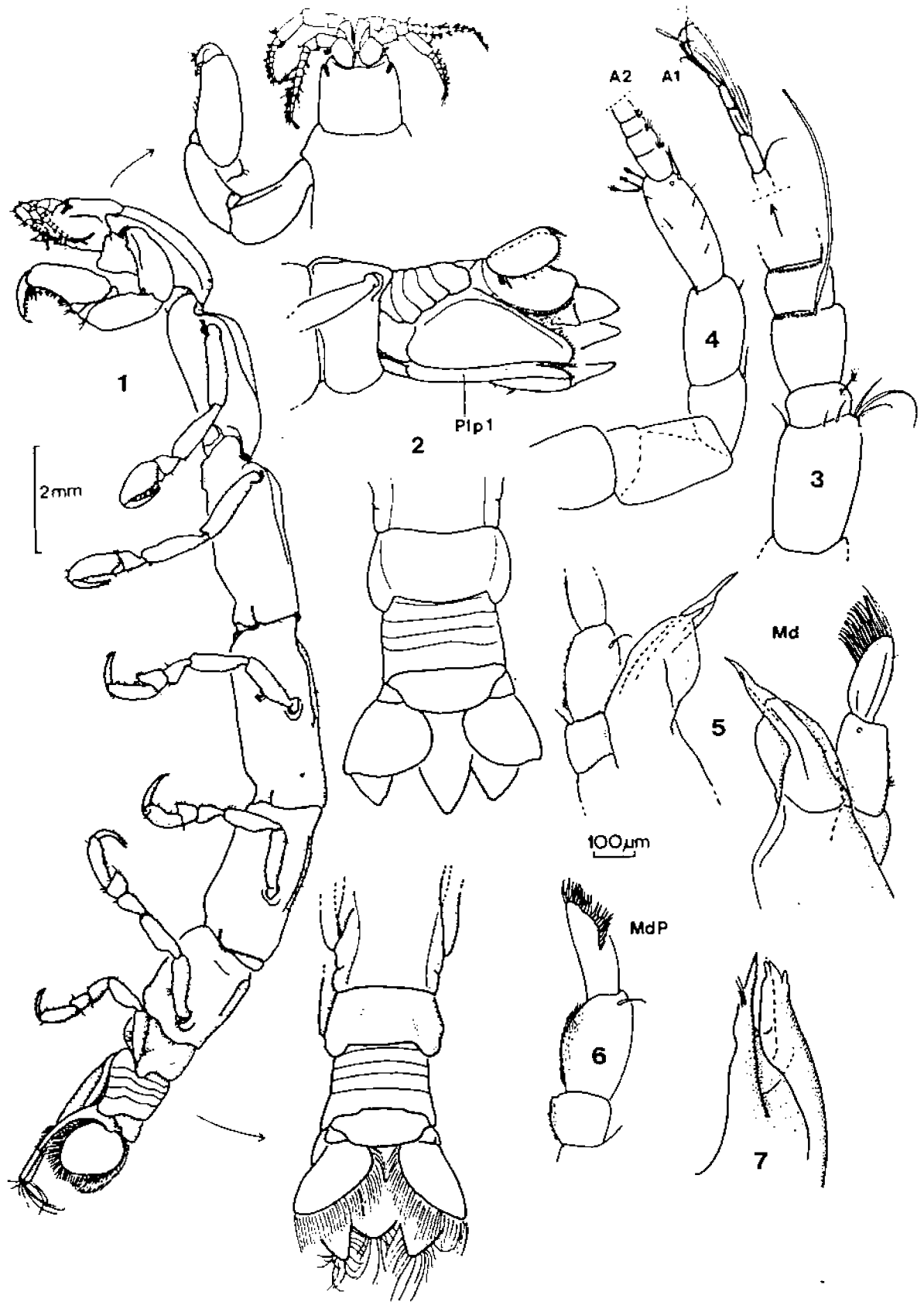


FIGURE 6. *Calathura branchiata* - 1) ♀ whole animal lateral view, with dorsal views of head and pleon/tail fan; 2) ventral oblique and dorsal views of the pleon/tail fan of another specimen; 3) antenna 1; 4) antenna 2; 5) mandible (2 views); 6) mandibular palp; 7) lower lip (labium)(from Wägele 1981)

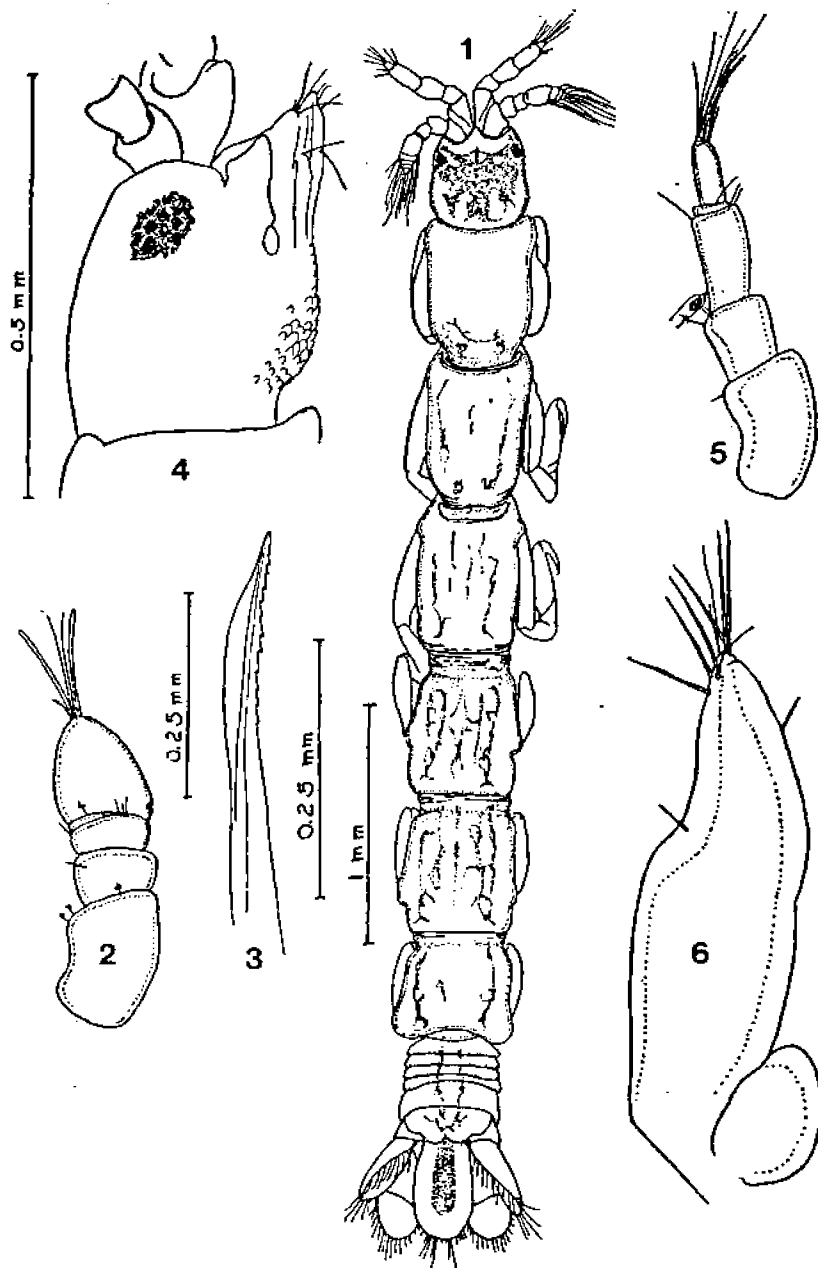


FIGURE 7. *Califanthura squamosissima* - 1) holotype ♂, dorsal view; 2) antenna 1 of juvenile ♂; 3) tip of maxilla 1; 4) later view of head, holotype ♂; 5) antenna 1 of ♀; 6) maxilliped of holotype ♂ (from Menzies 1951)

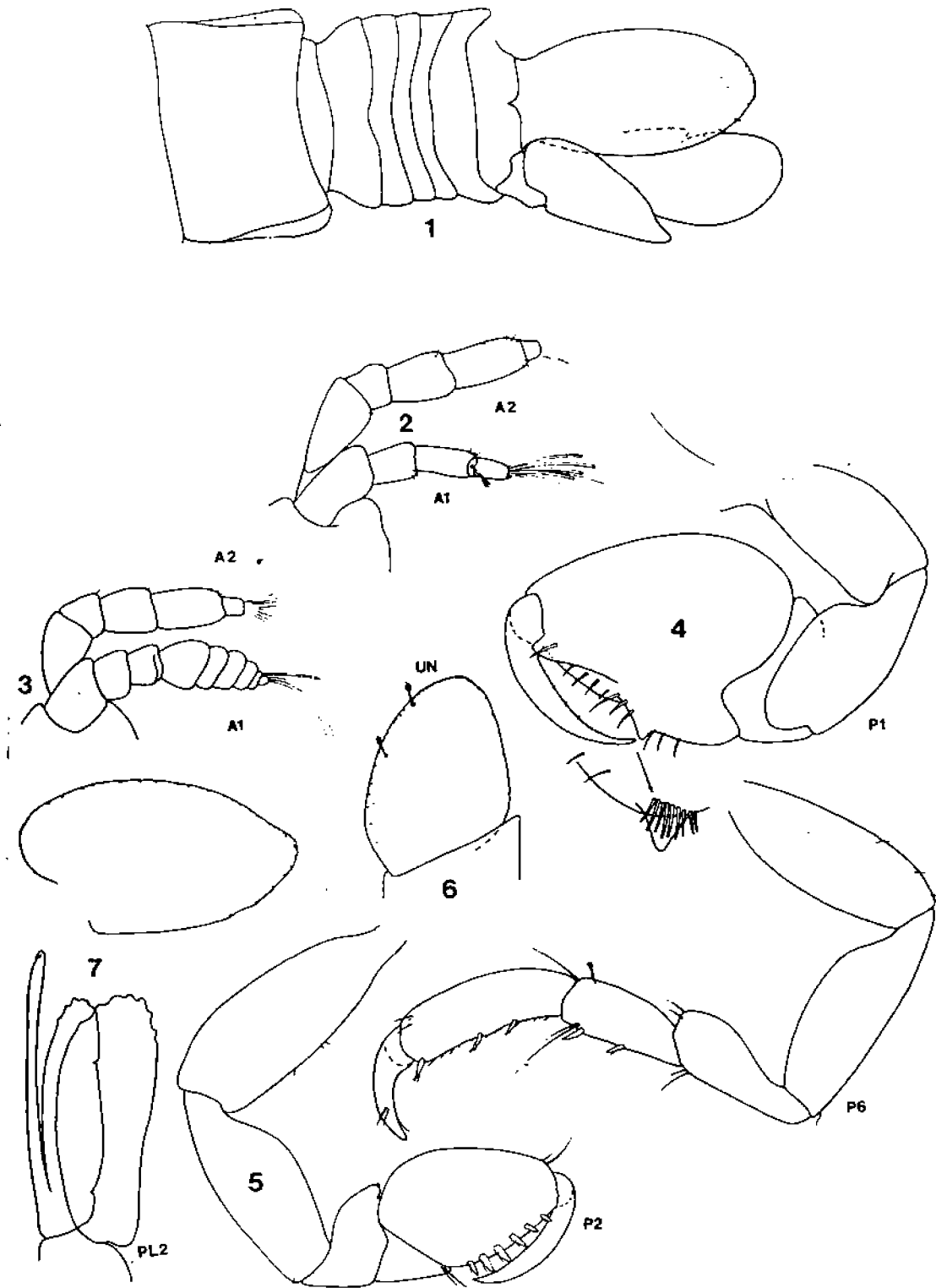


FIGURE 8. *Colanthurus bruscai* - 1) dorsal view of ♀; 2) antennae of ♀; 3) antennae of ♂; 4) pereopod 1 of ♀; 5) pereopod 2 of ♀; 6) uropodal endopod of ♀; 7) uropodal exopod of ♀; pleopod 2 of ♂ (from Poore 1984)

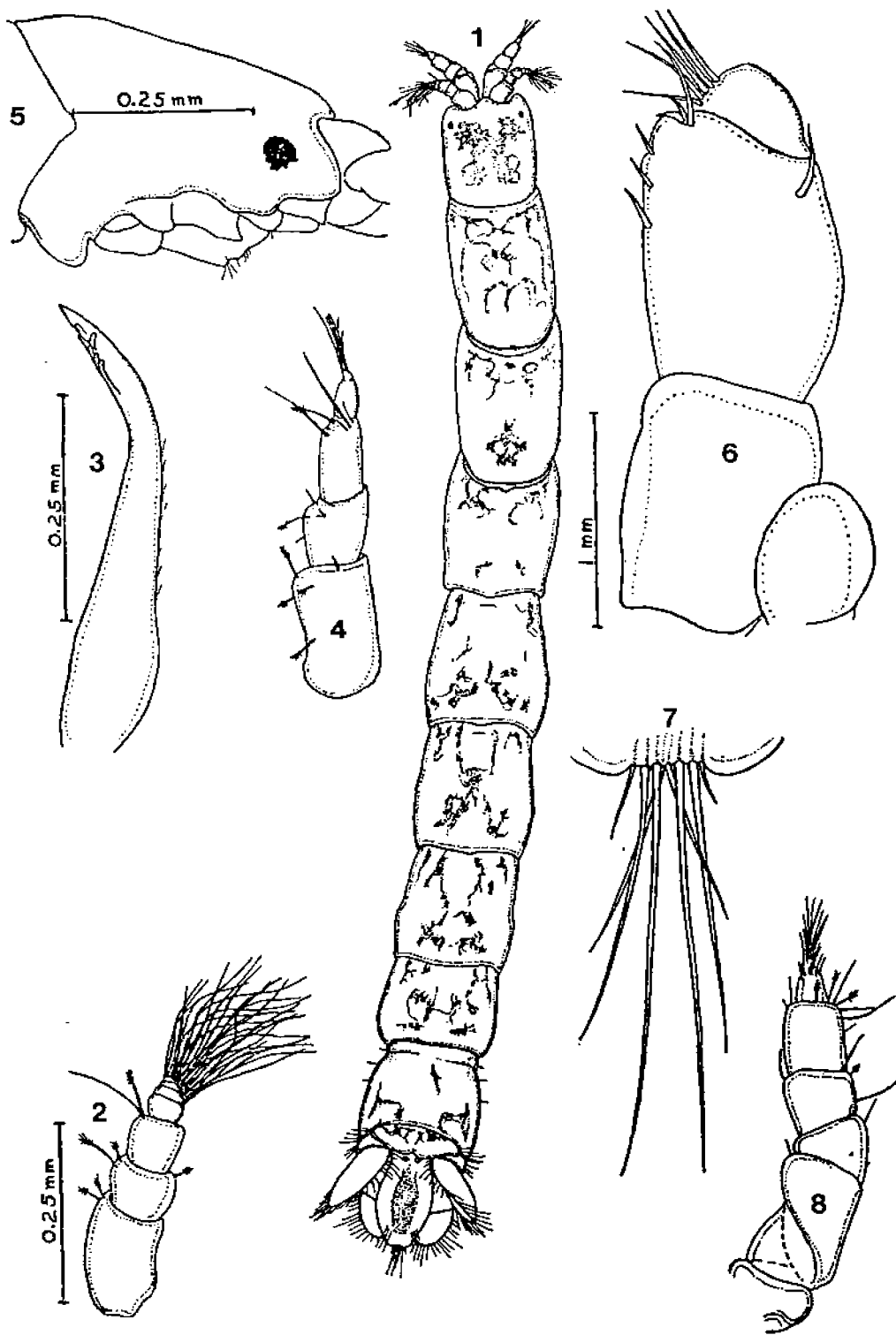


FIGURE 9. *Cyathura munda* - 1) dorsal view of ♂; 2) antenna 1 of ♂; 3) maxilla 1; 4) antenna 1 of ♀; 5) lateral view of ♂ head; 6) maxilliped of ♀; 7) tip of pleotelson of ♂; 8) antenna 2 of ♂ (from Menzies 1951)

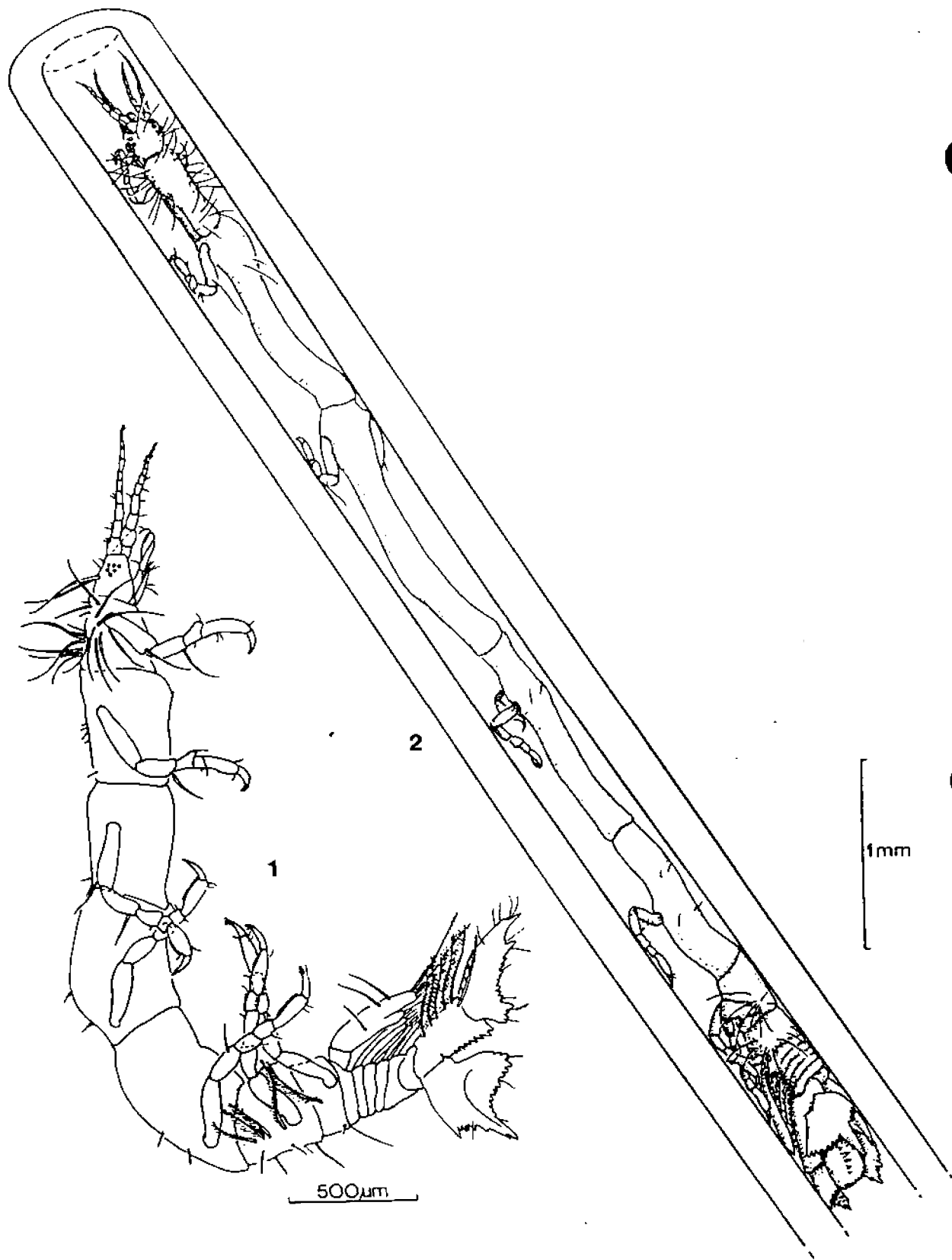


FIGURE 10. *Eisothisτος macrurus* - 1) pre-brood ♀ in lateral view; 2) post-brood female in capillary tube showing elongation of pereonites (from Wägele 1981)

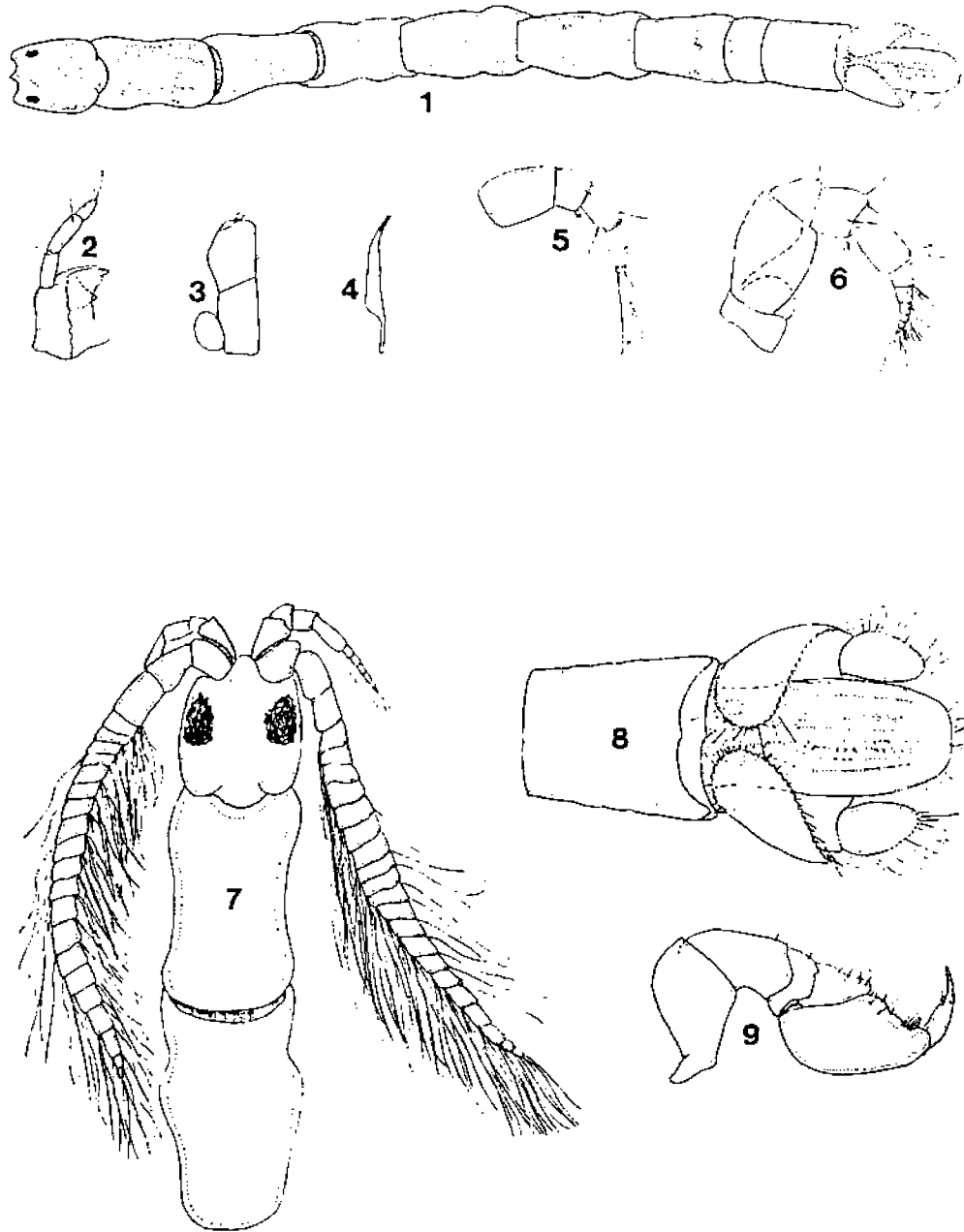


FIGURE 11. *Haliophasma geminatum* - 1) dorsal view of ♀; 2) mandible; 3) maxilliped; 4) maxilla 1; 5) antenna 1 of ♀; 6) antenna 2 of ♀; 7) anterior dorsal view of ♂; 8) pleon & pleotelson of ♂; 9) pereopod 1 of ♂ (from Menzies & Barnard 1959)

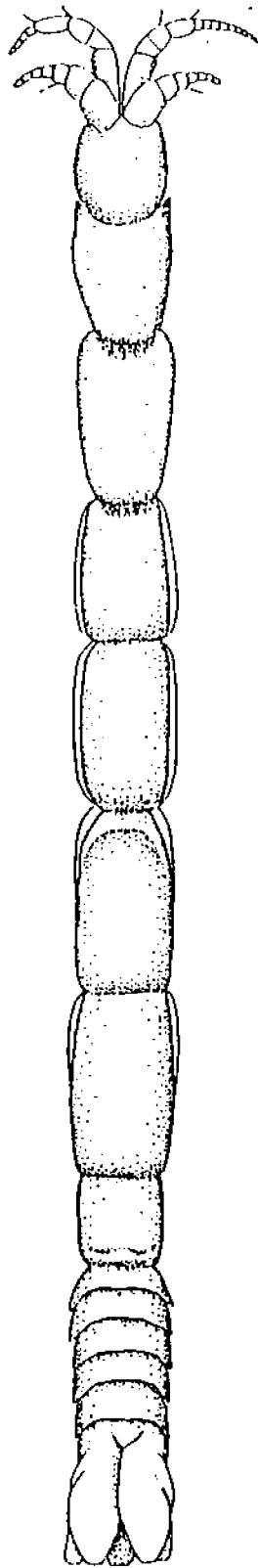


FIGURE 12. Hysurridae gen A sp A - dorsal view (from Wetzer & Brusca, in press)

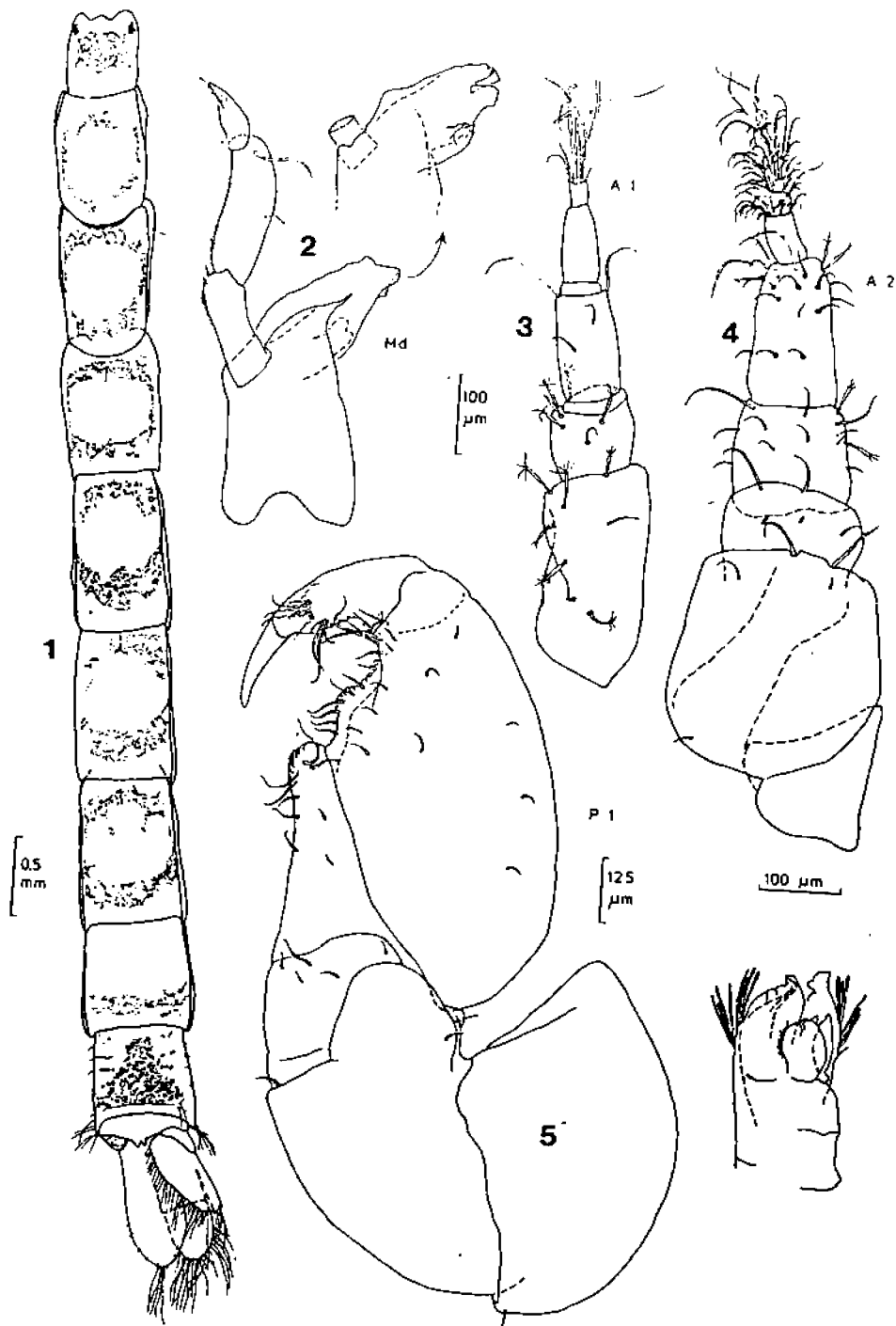


FIGURE 13. *Mesanthura occidentalis* - 1) dorsal view of juvenile; 2) mandible (2 views); 3) antenna 1; 4) antenna 2; 5) pereopod 1 (from Wägele 1984)

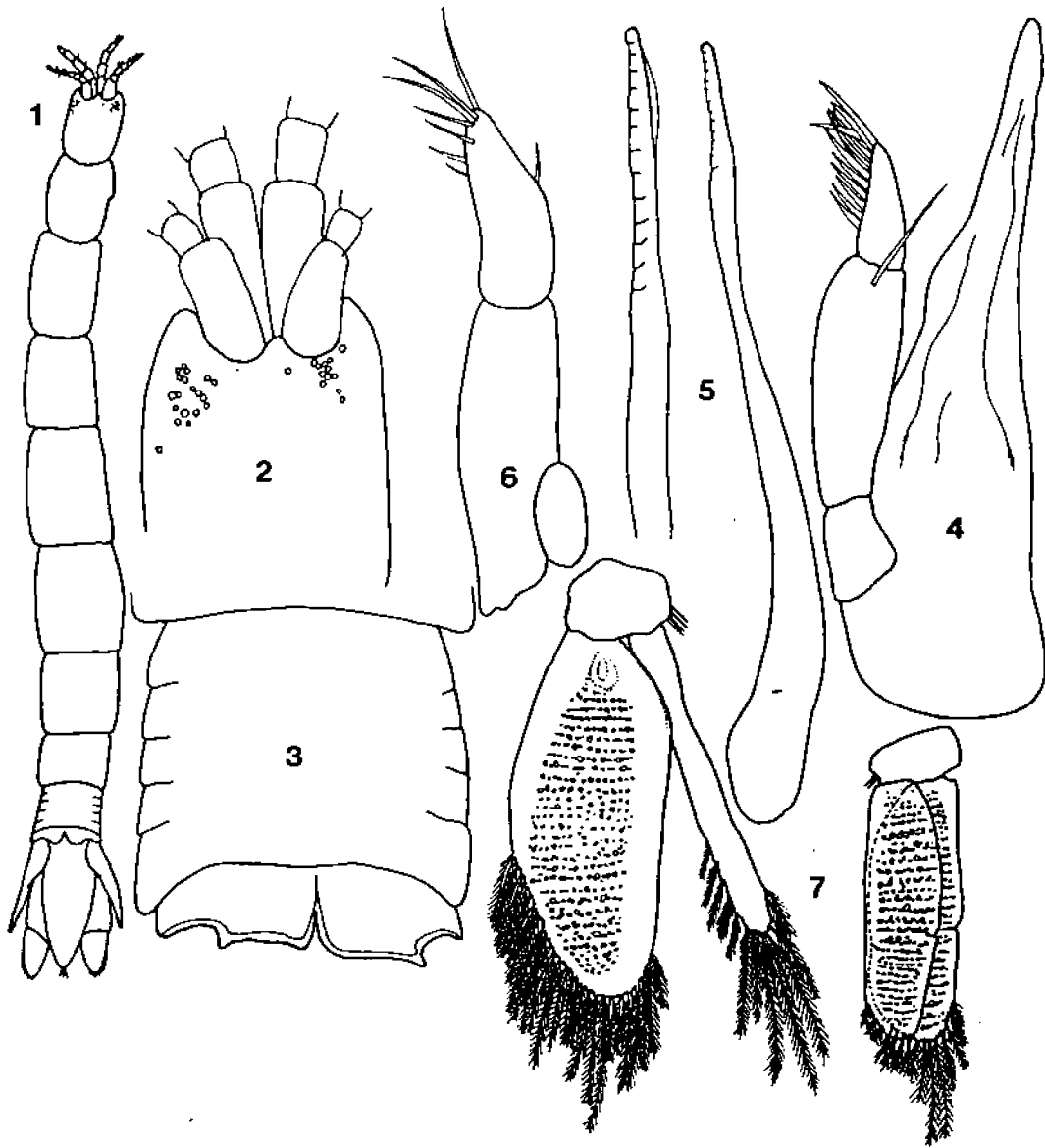


FIGURE 14. *Paranthura algicola* - 1) dorsal view of ♀; 2) head; 3) pleon; 4) mandible; 5) maxilla 1 (2 views); 6) maxilliped; 7) pleopods (2 views) (from Nunomura 1978)

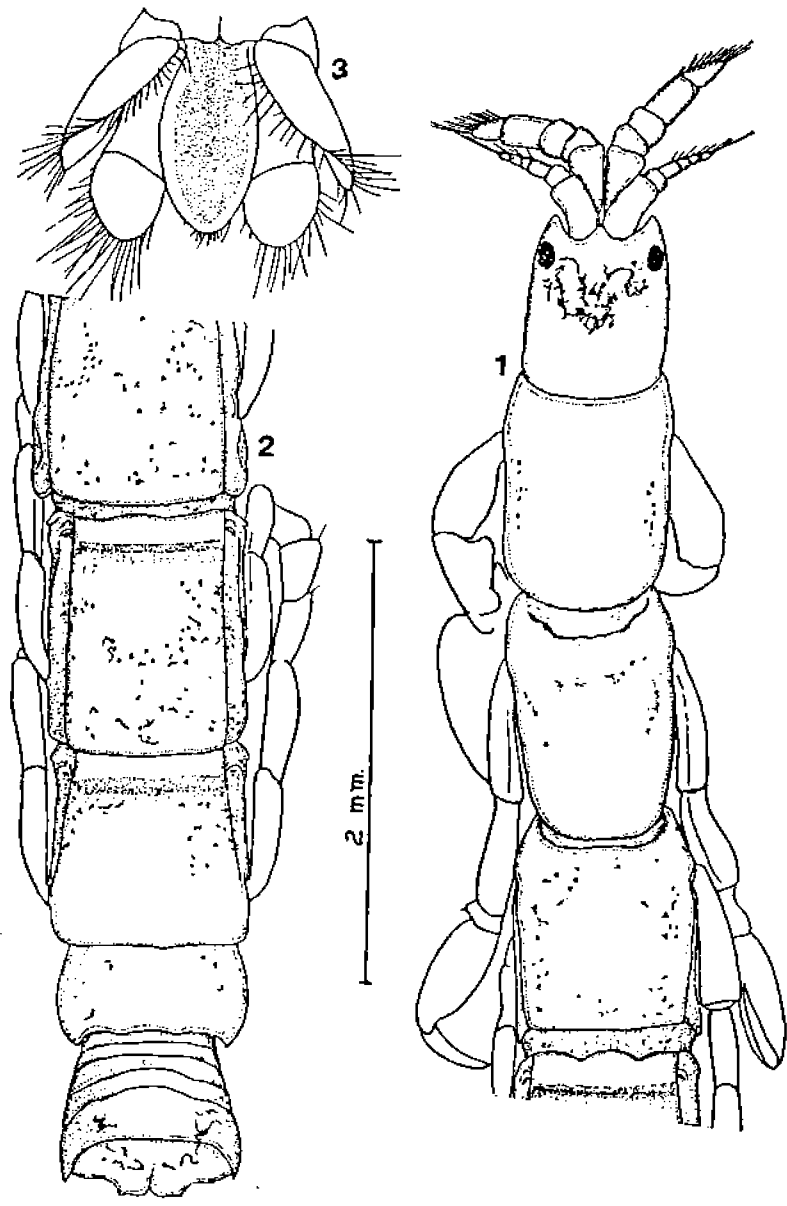


FIGURE 15. *Paranthura elegans* - 1) dorsal view of anterior body of ♀; 2) dorsal view of posterior body of ♀; 3) dorsal view of tail fan of ♀ (from Menzies 1951)

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- . Toxicity Reduction Evaluations
- . Sediment Quality Criteria
- . Assessment of Sediment Contamination (extent & type)
- . Sediment Bioavailability Issues
- . Evaluation of Cumulative Impacts
- . Regional Planning
- . Water Quality Criteria and Standards (including site specific)
- . Freshwater & Marine Water Quality and Ecosystem Issues
- . Bioassessment, Rapid Bioassessment Protocols and Biocriteria
- . Great Lakes: (Development & Implementation of Regulations; Impact on Regulated Communities; Watershed Management; Site-specific Municipal and Industrial Permitting; and Regional Water Quality)

The deadline for submission of abstracts is January 10, 1994. Authors will be notified of tentative selection of abstracts by April 2; final acceptance of papers is contingent on submission of a full manuscript of the selected abstract by July 1, 1994.

Submit abstracts to:
Water Environment Federation
Attn: Maureen Novotne, Technical & Educational Services
601 Wythe Street, Alexandria, VA 22314-1994
(703) 684-2400, ext. 7450



Abstract Submittal FORM

WEF Control No. _____

Water Environment Federation

67th Annual Conference & Exposition

McCormick Place North

Chicago, Illinois October 16-20, 1994

A photocopy of this form must be used as the title page for each copy of the abstract. The session topic for which the abstract is submitted must be identified by letter in the appropriate space on the form. Another platform session or the poster session may be indicated for alternate consideration if the paper is not accepted for the primary session topic.

Send copies of the complete submittal to the Federation office. Sending abstracts to session managers or other members of the Program Committee may delay consideration of the paper. Abstracts must arrive at the Federation office by January 10, 1994. No FAX submissions can be accepted for consideration. Submissions received after this date will receive consideration only after prior submissions have been evaluated, and on a space available basis.

Title of paper: _____

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Will this paper be presented elsewhere before September 1, 1994? Yes No

If so, where? _____

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Enter letter from attached list. If submitted for A-H (Symposia series), submit 15 copies of the abstract; for all other sessions, submit 5 copies. This form must be used as the cover page for each copy of the abstract.

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Submit abstracts to:
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Alexandria, Virginia 22314-1994 USA

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For Committee
use only:

1	2	3	4	5	Total



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Dec. 13th SCAMIT Meeting
on Odd Polychaetes

Polychaetes brought by Larry Lovell

Scoelelepis

sp. B ^{of Rossi} see → tridentate hooks

(*S. bullibranchia*)

multidentate hooks, ^{in key} notosetae absent on 1st setiger

→ see Couplet insert in Lovell + Pasko Key of Spionidae
(tri. vs multidentate hooks) (new version in process)

occipital cirrus absent here ~~in~~ in *S. sp. B*
hooks begin on setiger 15th.

Scoelelepis sp. B. of Rossi

- notosetae absent on 1st setiger
- occipital cirrus absent
- multidentate hooks not tri- as in *S. bullibranchia*

Maciulek uses the shape of the main fang of the hooded hooks to separate Scoelelepis into 2 main groups
one has a sharper main fang and the other is more blunt

Scoelelepis sp. B brought by Larry could be Scoelelepis
sp. 1 of Pt. Loma Larvis from lagoon but Pt. Loma sees ^{them offshore}
→ { has some palpal papillae as drawn in
 { Maciulek 1987 and interramal ventral lamellae

See Maciulek '87 (Pettibone version)

Scotolepis tridentata has notosetae on setiger 1.
unlike Ssp B.

Scotopolos acmeceps (brought by L. Lowell)

Family Orbiniidae

found deep in offshore basins

Larry has seen it much shallower

branchiae begin on setiger 12

* Check for uncinial setae in anterior setigers because
there are fewer capillaries to interfere in view
the capillaries form a U shape around these
uncinial setae

No subpodial lobes found - easier to see if stained
with methyl green

Scotopolos tends to have its anterior end flattened
dorso-ventrally whereas Leitoscoloplos is inflated
and has some sort of ducts that ~~make~~ make a dashed line
running from setae to setae across ventrum. Scotopolos
doesn't have these

Mostly S. armiger is seen especially by Pt. Loma
Hyperion • hasn't reported this as of yet.

Nothua occidentalis

no pigment, large eyes, pigment spot on anterior prost.
first few parapodia point anteriorly

on 1st setiger large presetal rounded lobe

found 60 meters in coarse sediment ^{with} heavily encrusted rock tubes

See Hartman atlas under Nothua conchyliga

for figures that refer to N. occidentalis.

Onuphis pallida - one from Larry and one from us.

Larry's - very different from ours

which keys out to O. graphiliformis according to R. Velarde but should be checked against type.

He thinks it looks close to O. iridescens except that branchiae begin on setiger 4 not 1.

Larry's from Montecito

- has what looks like eye spots on ceratophores
- subcircular hooks present from setiger 10
- Compound spinigers
- branchiae begin at 1 (for Monoonuphis - they begin at 6 or 7) and are bifid
- pseudocompound hooks - tridentate or bidentate?
- pseudocompound hooks in 1st 3 setigers on one side and

1st 4 setigers on other side

Keys out to Oruphis eremita parva if we
consider pseudocompound hooks in 1st 4 setigers
rather than 3

Need to check against Shisto's O. annulata

Syllidae

Sphaerosyllis branhaeroti: (Larva specimen from
look at Banse + Hobson (Orange Co. in 90 meters))

* whether or not dorsal cirrus is present on 2nd setiger

S. branhaeroti has it.

3 pairs of eyes

dorsal cirrus on 1st 2 setigers

simple setae present on setiger 1

no dentitions on edges of compound setae and

the simple setae are smooth

*^a main character to distinguish species with

S. californiensis has no dorsal cirrus on 2nd setiger
and only 2 pair of eyes

Maldanidae

see Hobson + Banse and voucher sheet by Karen Greene

Petaloproctus borealis - 21 setigers + smooth margin
what to look at { pygidial scoop w/ smooth or crenulated margin?
20 or 21 setigers?

Notopectus, Nicomancho, Petaloproctus all have very long setae

all these are quite rare
most aren't found whole

Cirratulidae

Apheleochaeta parva

spinally coiled with a mid dorsum stripe

large like A. multifilis

Tony Phillips will examine it

Moeroruphis stigmatus

branchiae simple + strap like begin at setiger 21
and ^{begin} start on setigera 12-19

peristomium with dark brown pigment
pigment bands across setigers - double bars
small eyes at base of ceratophores

Tell Tom about

Sabellides sp. 1 - Pt. Loma - Common at all depths

11 thoracic unciniger

4 pairs of branchiae

another pair of appendages inserted between
branchiae

See Fauchald, 1972 and Uebelacker - Gulf of Mexico
it differs from Asabellides in having 1 less unciniger
and having palae