Platycopia compacta n. sp., the Second Species of Platycopioida (Crustacea: Copepoda) in the Indo-Pacific Region, with Remarks on Development, Feeding, Swimming, and Zoogeography

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Platycopia compacta n. sp., the Second Species of Platycopioida (Crustacea: Copepoda) in the Indo-Pacific Region, with Remarks on Development, Feeding, Swimming, and Zoogeography

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ABSTRACT—Platycopia compacta new species (Copepoda: Platycopioida) is described from the hyperbenthic zone in the Tokara Islands, southern Japan. This is the second species of the order in the Indo-Pacific region, and the eighth species of the genus. The present new species is distinguishable from the closely related P. inornata from the Bahamas by (1) the formation of allobasis in the antenna; (2) two unequal setae on the basal exite of the maxillule; (3) the intermaxillipedal process with only 3 pairs of prominences at tip; (4) the fusion of the coxa and basis in leg 3; and (5) the fusion of the second and third endopod segments in female legs 3-5. Copepodid stages III, V, and VI are compared with those of P. orientalis collected from Okinawa, and differ in the developmental patterns of the antennule and legs. The presence of an intermaxillipedal process and gut content analysis revealed that Platycopia is carnivorous. The armature of legs 2-5 of the hyperbenthic genus Platycopia is stouter than that of cavernicolous platycopioids, which indicates that Platycopia may penetrate into loose sediment, whereas the latter group might be more adapted to pelagic life. The horizontal distribution of the genus Platycopia apparently exhibits a full Tethyan track.

INTRODUCTION

Hyperbenthic and cavernicolous copepods have provided significant new information about the morphology, zoogeography, ecology and phylogeny of the maxillopodan subclass Copepoda. Some calanoid genera such as Exumella, Metacalanus, Paramisophria, and Stephos are broadly distributed in both the hyperbentic zone and marine caves and indicate faunistic similarities between the two realms (cf. Boxshall et al., 1990; Jaume and Boxshall, 1995a; Ohtsuka et al., 1993b,1994; Riera et al., 1991). Recently numerous new families, genera, and species of copepods have been discovered from anchialine caves (e.g., Fosshagen and Iliffe, 1985, 1988, 1989, 1991; Jaume and Boxshall, 1995a,b; Ohtsuka et al., 1993a,b; Rocha and Iliffe, 1991). These have shed new light on phylogenetic relationships within each copepod order (Boxshall, 1989; Ho, 1994) and brought into question the totality of the Messinian Salinity Crisis in the Mediterranean Sea about 5.5 Myr BP (Jaume and Boxshall, 1996b).

During our investigation of hyperbenthic copepods we have found another species of Platycopia, which belongs to the hyperbenthic/cavernicolous copepod order Platycopioida, in the Nansei Islands, southern Japan. This is the second species of the order in the Indo-Pacific region and the eighth species of the genus worldwide. The present paper describes the new species with remarks on its development, feeding, swimming, and zoogeography.

MATERIAL AND METHODS

Copepods were collected with a dredge (mouth 50 cm wide, 15 cm high; mesh size 5 mm) at 67 m depth off Kuchino Island, Tokara Islands, south of Kyushu, Japan (30° 00.82’ N, 129° 53.92’ E), on June 1, 1997 (local time 0815-0847). Sediment was stirred in sea-water several times and the supernatant was filtered with a plankton net (mesh size: 0.1 mm). Copepods were fixed in 10% neutralized formalin/sea-water immediately after filtering.

The type specimens are deposited in the Natural History Museum and Institute, Chiba. Terminology follows Huys and Boxshall (1991).

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DESCRIPTION
Order Platycopioida Fosshagen, 1985
Family Platycopiidae Sars, 1911
Genus Platycopia Sars, 1911
Platycopia compacta new species
(Figs. 1-6)

Material examined.—Holotype: adult female (body length 0.76 mm), dissected and mounted on 14 glass slides, CBM ZC 4035. Paratype: adult female (body length 0.81 mm), whole specimen, CBM ZC 4036. Non-types: copepodid V female (body length 0.74 mm), dissected and mounted on 4 glass slides; copepodid V male (body length 0.62 mm), dissected and mounted on 2 glass slides; copepodid III (body length 0.47 mm), dissected and mounted on 1 glass slide.

Description.—Adult female (holotype). Body (Fig. 1A,B) compact. Prosome about 2.8 times as long as urosome; cephalosome and first pedigerous somite incompletely fused, without distinct arthrodial membrane; fifth pedigerous somite produced posteriorly into round lobe reaching beyond middle of first abdominal somite; several developing ova observed in posterior part of prosome in dorsal and lateral view. Rostrum (Fig. 1C) sharply pointed, ventrally directed, without filaments at tip. Urosome (Fig. 1D) 5-segmented; genital somite small, almost concealed beneath dorsal extension of fifth pedigerous somite; paired slit-like gonopores (arrowed in Fig. 1D) located ventrally, slit-like; first to third abdominal somites with posterior margin finely striated; third abdominal somite protruded dorsoposteriorly into w-shaped plate partly covering anus; fourth abdominal somite prominent ventroposteriorly. Caudal rami (Fig. 1D,E) symmetrical, each with triangular dorsal process; seta I rudimentary, seta II slender, setae III-VI plumeose; setae IV and V with finely punctate proximal surface.

Antennule (Fig. 2A) 23-segmented. First segment with two longitudinal ridges and elongate, heavily chitinized anterior seta. Armature and fusion pattern as follows: I-III may be incorporated in first segment, which is not traceable based on elements; IV-VI—6 + ae; VII—2 + ae; VIII—1 short spiniform seta; IX—2 + ae; X—1 + 1 short spiniform seta; XI—2 + ae; XII—1; XIII—1 + 1 short spiniform seta; XIV—2 + ae; XV—1; XVI—1 + ae; XVII—0; XVIII—1; XIX—0; XX—1; XXI—1 + 1; XXII—0; XXIII—1; XXIV—1; XXV—1 + ae; XXVI—2; XXVII—1 + 1; XXVIII—4 + ae.

Antenna (Fig. 2B) with unarmed coxa and basis. Endopod with first segment completely fused to basis to form allobasis; second and third segments incompletely coalesced, bearing 3 and 7 setae, respectively. Exopod indistinctly 7-segmented; setal formula 1, 0, 1, 1, 2, 1, 4.

Mandibular gnathobase (Fig. 2C) bearing 1 monocusped tooth and 4 bicusped teeth. Mandibular palp (Fig. 2D) with unarmed basis. Endopod unisegmented, with 7 terminal setae. Exopod 4-segmented, first to fourth segments, respectively, with 0, 2, 1, and 2 setae.

Maxillule (Fig. 2E) well developed; praecoaxal arthrite expanded, bearing 6 stout, multicurved spines and 6 setae, 2 setae of which being originated from center of anterior surface; coxal endite with 3 setae; basal exite (arrowed in Fig. 2E) lobate, bearing 2 setae of unequal length; proximal basal endite elongate, bearing 3 terminal spinulose setae; distal basal endite probably represented by a knob with 2 unequal setae; both rami completely incorporated into basis, bearing 4 setae in total.

Maxilla (Fig. 2F) stout; 2 praecoaxal, 2 coxal, and 1 basal endites bearing 4, 1, 3, 3, and 3 setae, respectively, one setae on each second coxal endite and on basal endite fused to segment. First endopod segment completely fused to basis, represented by 1 spiniform seta; second to fourth segments each bearing 1 spiniform seta; third and fourth segments with 1 and 2 setae, respectively, in addition to spiniform seta.

Maxilliped (Fig. 3A) with praecoaxa and coxa incompletely coalesced; coxa with 2 spinulose, subterminal setae; basis bearing 2 inner setae, one of them being rudimentary; endopod 6-segmented, setal formula 1, 2, 2, 1, 1, 4. Intermaxillipedal process (Fig. 1B,F) with 3 pairs of anteriorly directed, acute prominences at tip.

Leg 1 (Fig. 3B) with 2-segmented rami; coxa unarmored, produced into acute outer process subterminally; basis expanded, bearing plumose setae near base of endopod. First endopod segment with longitudinal groove on anterior surface along basal seta, segment produced at outer distal corner. First exopod segment unarmored, prominent at outer distal corner; second with acute prominence and row of setules along outer margin and 6 setae, outermost more heavily chitinized than others. Leg 2 (Fig. 3C) with 3-segmented rami. Legs 3 (Fig. 4A) and 4 (Fig. 4B) with second and third endopod segments not fully separate on posterior surface; both legs similar, but differing in: (1) coxa and basis separate in leg 4 and incompletely fused in leg 3; (2) third endopod segment bearing 5 spiniform setae in leg 3 but 4 in leg 4; (3) second exopod segment lacking inner seta in leg 3. Leg 5 (Fig. 4C) with expansion along inner margin of basis; second and third endopod segments almost fused, with suture visible on both surfaces. Spine and seta formula of legs 1 to 5 shown in Table 1.

Male unknown.

Copepodid III (CIII) (Although legs 3-5 are damaged in the only individual, it was identified as CIII on the basis of the number of urosomal somites, the number of ramal segments of leg 2 and the presence of leg 5. See Ohtsuka and Boxshall, 1994).—Urosome 2-segmented. Antennule 15-segmented (Fig. 5): armature as follows: 1—4 + ae (Fig. 6A); 2—0; 3—1 + ae; 4—0; 5—1; 6—0; 7—1; 8—1 + ae; 9—0; 10—1; 11—1; 12—1 + ae; 13—2; 14—1 + 1; 15—4 + ae (see Development in Discussion). Antenna with basis completely fused to first endopod to form allobasis. Basal exite of maxillule with 2 setae of unequal length. Maxilliped with only single seta on coxa; basis as in adult female; endopod 4-segmented; setal formula of 1, 1, 1, 4. Intermaxillipedal process with 2 pairs of prominences at tip. Leg 1 with 2-segmented rami; second exopod segment bearing only 5 setae. Leg 2 (Fig. 6C) with 2-segmented rami; first exopod segment with single outer spine. Legs 3-5 missing.

Copepodid V (CV) female.—Urosome 4-segmented. An-
tennule 22-segmented (Fig. 5); armature as follows: 1—8 + 2 ae (Fig. 6B); 2—1 short spiniform seta; 3—2; 4—1 short spiniform seta; 5—2 + aesthetasc; 6—1; 7—1 + 1 short spiniform seta; 8—2; 9—1; 10—1 + ae; 11—0; 12—1; 13—0; 14—1; 15—1 + ae; 16—0; 17—1; 18—1; 19—1 + ae; 20—2; 21—1 + 1; 22—4 + ae (see Development in Discussion). Basal exite of maxillule with 2 setae of unequal length. Maxilliped as in adult female. Intermaxillipetal process with 3 pairs of prominences at tip. Legs 1 and 2 as in adult female. Leg 3 with 2-segmented endopod and 3-segmented exopod; armature as

Fig. 1. *Platycopia compacta*, adult female (holotype). (A) Habitus, dorsal; (B) Habitus, lateral, intermaxillipetal process arrowed; (C) Rostrum; (D) Urosome, ventral, gonopores arrowed; (E) Anal somite and caudal rami, dorsal; (F) Intermaxillipetal process, viewed from ventral side. Scale bars in mm.
in adult female, but second and third endopod segments not separated, without arthrodial membrane. Leg 4 with 2-segmented endopod and 3-segmented exopod; armature as in adult female, but second and third endopod segments not separate. Leg 5 (Fig. 6D) with intercoxal sclerite not fully separated from coxa; 2-segmented rami; first endopod unarmed, second bearing 6 spines; first exopod segment with single outer spine, second segment bearing 4 flanged spines and 3 short spines.

Copepodid V (CV) male.—Urosome, antennule (Fig. 6B),

Fig. 2. Platycopeia compacta, adult female (holotype). (A) Antennule; (B) Antenna; (C) Mandibular cutting edge; (D) Mandibular palp; (E) Maxillule, basal exite arrowed; (F) Maxilla. Scale bars in mm.
New Species of Platycopia

basal exite of maxillule, intermaxillipedal process, and legs 1-4 as in CV female. Leg 5 (Fig. 6E) with intercoxal sclerite not fully separated from coxa; 2-segmented rami; second endopod segment with 6 short spines; second exopod segment bearing 4 flanged spines (2 missing) and 2 short spines.

Etymology.—The specific name “compacta” (Latin compactus meaning plump) is derived from the compact prosome of the new species, compared with that of *P. orientalis* first collected from Japan.

Comparison.—The mouthparts and leg 1 of species of the genus *Platycopia* exhibit no remarkable differences while the armature and fusion pattern of legs 2-5 are the important characters for distinguishing between species (Wilson, 1946; Ohtsuka and Boxshall, 1994). The presence or absence of the intermaxillipedal process is also important. Some species such as *P. inornata* Fosshagen, 1972 and *P. perplexa* Sars, 1911 are furnished with it (see Sars, 1911; Huys and Boxshall, 1991), whereas *P. orientalis* Ohtsuka and Boxshall, 1994 lacks a distinct process. Although some other congener may also bear a process, it was not mentioned in their original descriptions.

The present new species closely resembles *Platycopia inornata*, *P. perplexa*, *P. robusta* Andronov, 1985, and *P. sarsi* M.S. Wilson, 1946 in: (1) 3-segmented rami in legs 2-4 (symplesiomorphy); (2) the first exopod segments of legs 3 and 4 of adults bearing 1 inner seta (symplesiomorphy). Female *P. perplexa* and *P. robusta* both lack an inner seta on the first exopod segment of leg 5 (synapomorphy) while *P. inornata* and the new species carry one (symplesiomorphy). Since only males are known in *P. sarsi*, it is impossible to compare the female legs 5. The present new species is distinguishable from *P. inornata* in: (1) the basis and endopod of the antenna being completely fused (separate in *P. inornata*); (2) the 2 setae on the basal exite of the maxillule being of unequal length (almost equal in *P. inornata*); (3) the intermaxillipedal process having 6 spinous prominences at its...
Fig. 4. *Platycopia compacta*, adult female (holotype). (A) Leg 3, anterior; (B) Leg 4, anterior; (C) Leg 5, anterior. Scale bar in mm.

*Platycopia sarsi* was poorly described on the basis of a single adult male and two copepodids by Wilson (1946). The following features differ between *P. sarsi* and the new species: (1) the proximalmost spine on the praecoxal arthrite of maxillule is bifurcate in *P. sarsi* but tricusped in *P. compacta*; (2) the outer subterminal corner of the coxa of leg 1 is round.
Table 1. Seta and spine formula of legs 1 to 5 of *Platycopia compacta* n. sp. Roman and Arabic numerals indicate spines and setae, respectively.

<table>
<thead>
<tr>
<th>coxa</th>
<th>basis</th>
<th>exopod</th>
<th>endopod</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>1 2 3</td>
<td>1 2 3</td>
</tr>
<tr>
<td>Leg 1</td>
<td>0-0</td>
<td>0-1</td>
<td>0-0:1,1,4</td>
</tr>
<tr>
<td>Leg 2</td>
<td>0-0</td>
<td>I-1</td>
<td>II-0:1-0;III,1,2</td>
</tr>
<tr>
<td>Leg 3</td>
<td>0-0</td>
<td>I-1</td>
<td>II-0:1-0;III,1,2</td>
</tr>
<tr>
<td>Leg 4</td>
<td>0-0</td>
<td>I-1</td>
<td>II-1-1;III,1,2</td>
</tr>
<tr>
<td>Leg 5</td>
<td>0-0</td>
<td>I-0</td>
<td>II-1-1;III,1,2</td>
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in *P. sarsi* but acutely pointed in *P. compacta*. The late copepodid female described by Wilson (1946) probably corresponds to the fifth copepodid stage because of the 22-segmented antennule, 3-segmented rami of legs 2-4, and 2-segmented rami of leg 5 (cf., Ohtsuka and Boxshall, 1994). In *P. compacta* the fifth copepodid stage of both sexes bears a 2-segmented endopod in leg 5, in which the second and third segments of the adult are not fully separated. Leg 5 of the CV female of *P. sarsi* differs slightly from that of *P. compacta*: the proximal outer spine on the second endopod segment and the proximal inner spine on the second exopod segment are relatively more elongate than those of *P. sarsi*.

**DISCUSSION**

**Development**

The segmentation and setation of the antennules of *P. compacta* are depicted in Fig. 5. The most proximal antennular segment of female CVI is interpreted as IV-VI based on Huys and Boxshall (1991): “the setation pattern provides no evidence of the fate of the 3 most proximal segments”. Although the fourth copepodid stage (CIV) is unknown, the segmentation of the distal 13 segments are unchanged from CIII to CVI. At the last molt from CV to CVI, an aesthetasc is added into the ancestral segments IV-VI, IX, and XIV; a seta is added to the segment X. The first compound segment of CV consists of the ancestral segments IV-VII, the last of which is separate from the compound segment IV-VI at CVI.

Developmental patterns of antennules, maxillipeds and legs can be compared between *P. compacta* and *P. orientalis* as described by Ohtsuka and Boxshall (1994). In CIII the number of segments of the antennules is 15 in *P. compacta* and 18 in *P. orientalis*. The distal 13 segments (ancestral segments XVI-XXVIII) of *P. compacta* have the same armature as in *P. orientalis*. In CV the segmentation and setation of the antennules are the same in both species except for the presence of an aesthetasc on the ancestral segment XIV and incomplete separation between the first (precursor of III-VII) and second (VIII) segments in *P. orientalis*. In CVI (adult female) the ancestral compound segment III (or IV)-VI and segment VII are separate in *P. compacta* and fused in *P. orientalis*. The differences between the armatures of CVI of both species are as follows: an aesthetasc is present on the compound segment IV-VI of *P. compacta* but absent on the part corresponding to IV-VI on the compound segment III-VII of *P. orientalis*; the ancestral segment VII is separate from the compound segment IV-VI in *P. compacta* but incorporated in *P. orientalis*; the ancestral segment XIII bears 1 long and 1 short spiniform seta in *P. compacta*, but only a single long seta in *P. orientalis*. In addition, the appearance of aesthetascs at the molt from CV to CVI differs: in *P. compacta* an aesthetasc appears on segments IV-VI, IX, and XIV, whereas in *P. orientalis* only one aesthetasc is added on segment IX.

The development of maxillipeds has so far been unknown in the Platycopioida including *P. orientalis* (Ohtsuka and Boxshall, 1994). In *P. compacta* the numbers of the maxilliped endopodal segments in CV and CIII are 6 and 4, respectively. Therefore that in CIV may be 5. If the development follows the pattern proposed for the Calanoida by Ferrari (1995), the penultimate endopod segment of CIII can be identified as a precursor of the third to fifth segments of the adult (although Ferrari (1995) regards the first endopod segment as a lobe of the basis, we follow Huys and Boxshall (1991): maxillipeds primitively bear 6-segmented endopod rather than 5-segmented); at the molt from CIII to CV the third segment of the adult may first appear to result in 5-segmented endopod;

![Fig. 5. Schematic illustration of antennules of Platycopia compacta. (III) third copepodid stage; (V) fifth copepodid stage (both sexes); (VI) adult female. Roman numerals indicate the ancestral segment scheme proposed by Huys and Boxshall (1991).](https://bioone.org/journals/Zoological-Science on 14 Dec 2019 Terms of Use: https://bioone.org/terms-of-use)
the fourth segment of the adult could be added at the molt from CIV to CV. In the Calanoida the adult endopod segmentation is completed at CIV (see Ferrari, 1995; Ferrari and Markhaseva, 1996) while in *P. compacta* presumably at CV. The completion of the endopod segmentation may be accelerated in the Calanoida.

The setation of the maxilliped endopod of *P. compacta* is completed probably at CV as well as the segmentation. Perhaps the setal numbers of the first and sixth endopod segments of the adult are the same from CIII to CVI. Following the pattern of the Calanoida (Ferrari, 1995; Ferrari and Markhaseva, 1996), the setal formula of the endopod of CIV may be 1, 2, 2, 1, 4.

The developmental pattern of legs 1-5 of *P. compacta* is shown in Table 2 to compare it with the common pattern of copepod leg ontogeny proposed by Ferrari (1988). Simultaneous additions are suspended in both rami of leg 1 here as well as in *P. orientalis*. The developmental pattern of legs 2 and 5 of *P. compacta* almost coincides with the common developmental pattern although the second and third endopod segments of legs 3 and 4 are not fully separated. Probably separations of these segments are delayed in legs 3 and 4. In contrast, simultaneous additions during the last molt are suspended in the endopods of legs 2-4 in *P. orientalis*, which results in 2-segmented endopods in these legs. In CIII a proximal outer spine on the first exopod segment of leg 2 appears in *P. orientalis* (Ohtsuka and Boxshall, 1994), but not in *P. compacta*.

**Feeding and swimming**

Little is known about the biology and ecology of the order Platycopioida (Huys and Boxshall, 1991). *Platycopia* bears...
well-developed maxillulary praecoxal arthrites, maxillae with stout spiniform setae and spines, and an intermaxillipedal process in some species, all of which probably play a role in grasping and holding prey. This is supported by gut content analysis which indicated that the CV female of *P. compacta* had preyed upon copepodids of other copepods.

The strong spines and spiniform setae of legs 2-5 of *Platycopia* imply that it may penetrate loose sediment but not interstitial spaces. The leg armature of the hyperbenthic *Platycopia* is much stronger than that of the cavernicolous *Antrisocopia* Fosshagen, 1985 and *Nanocopia* Fosshagen, 1988, which suggests that *Platycopia* is more closely associated with sediment than the cave-dwelling genera (cf., Bowman and González, 1961), which are more adapted to a pelagic life.

**Zoogeography**

The order Platycopiida was the first group to diverge from the main lineage of the subclass Copepoda and retains numerous extremely plesiomorphic characters. Platycopioids are restricted to the hyperbenthic zone and anchialine caves in shallow waters (Huys and Boxshall, 1991). All eight species of the genus *Platycopia* are distributed exclusively in the hyperbenthic zone in the Atlantic and Indo-Pacific Oceans (Huys and Boxshall, 1991; Ohtsuka and Boxshall, 1994; present study), while *Antrisocopia* Fosshagen, 1985 and *Nanocopia* Fosshagen, 1988 are monotypic inhabitants of anchialine caves in Bermuda (Fosshagen and Iliffe, 1985, 1988). Since there are numerous anchialine caves and other types of marine caves in the Nansei Islands, southern Japan (Hayami and Kase, 1993), undescribed taxa of the Platycopioida could be discovered in these regions.

The distribution of the genus *Platycopia* is shown in Fig. 7 on the basis of Huys and Boxshall (1991), Ohtsuka and Boxshall (1994), and the present study. This pattern evidently coincides to a full Tethyan track (circum-tropical through the entire region of the former Tethys Sea) as defined by Stock (1993), who insisted that stygobionts that display such a distributional pattern and have poor dispersal abilities must have existed in place ever since the existence of the Tethys Sea (ca. 120-20 Myr BP). Although the genus *Platycopia* is not a real stygobiont, the distributions of many hyperbenthic copepods seem to coincide with the full Tethyan track. For example, hyperbenthic calanoid copepods such as *Paramisophria*, *Pseudocyclops*, *Stephos*, and the Ridgewayiidae are distributed in the shallow waters in the Caribbean, the eastern Atlantic, and Indo-West Pacific regions (Boxshall et al., 1990; Fosshagen, 1968a,b, 1970a,b; Jaume and Boxshall, 1995a; Ohtsuka and Hiromi, 1987; Ohtsuka et al., 1993a,1996; Riera et al., 1991). In addition, some of these taxa accommodate both hyperbenthic and cavernicolous species (Ohtsuka et al., 1993b,1994; Yeatman, 1969). Such a faunistic similarity between in the hyperbenthic zone and marine caves strongly supports the above assumption.
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