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Morphological and Ecological Adaptation of *Basterotia* Bivalves (Galeommatoidea: Sportellidae) to Symbiotic Association with Burrowing Echiuran Worms

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The burrows created by benthos in tidal flats provide various habitats to other organisms. Echiuran burrows are unique among these in being persistently disturbed by the host's undulating activity, but little is known on how symbionts adapt to such a unique habitat. We report here the morphological and ecological adaptation by two bivalve species of *Basterotia* (Sportellidae), including one new species, which are commensals with burrowing echiuran worms. The burrows of *Ikedosoma gogoshimense* were inhabited by *Basterotia gouldi* at intertidal gravelly mud flats in the central Seto Inland Sea, whereas those of *Ochetostoma erythrogrammon* were inhabited by *Basterotia carinata* n. sp. at an intertidal gravelly coral-sand flat at Amami-Ohshima Island. Both bivalve species were found embedded in the burrow wall with their posterior inhalant and exhalant apertures gaping to the burrow lumen, suggesting that they utilize the water currents created by host echiurans. The posteriorly robust, laterally inflated shell with developed carina is considered an adaptation to symbiotic life, as it is exposed to pressure caused by the host's persistent undulating activity. Females of *Basterotia* bivalves were larger than males, suggesting size-dependent sex change, and possessed brooded veligers in the ctenidium. Our findings suggest that species-specific intimate association with echiurans may be widespread among the Sportellidae bivalves, whose biology remains poorly understood.

Key words: adaptation, burrow, commensalism, ecosystem engineer, symbiosis, Echiura

INTRODUCTION

The echiuran worms (Phylum Echiura), commonly called "spoon worms" or "innkeeper worms", are marine invertebrates with sac- or sausage-like, non-segmented soft bodies and highly extensible proboscides (Ikeda, 1904; Stephen and Edmonds, 1972). Most species of echiuran worms excavate deep tunnels or burrows in soft sediments (Stephen and Edmonds, 1972; Nickell et al., 1995; Rogers and Nash, 1996). Because they pump water that is rich in food and oxygen through their burrows, which are protected from epibenthic predators, these are inhabited by various symbiotic organisms, including bivalves, polychaetes, crabs, shrimps, flatworms, nemerteans, nematodes, gastropods, copepods, entoprocts, and gobiid fishes (Anker et al., 2005; MacGinitie, 1935; Morton and Morton, 1983; Schembri and Jaccarini, 1978). Thus, it is thought that echiuran worms influence the abundance and distribution of diverse invertebrates in local habitats and function as ecosystem engineers, as do other burrowing animals in marine sedimentary environments, such as polychaete worms, sipunculid worms,

and thalassinidean shrimps (Jones et al., 1994, 1997). The inside of an echiuran burrow is a unique environment, persistently disturbed by the peristaltic activity of the host

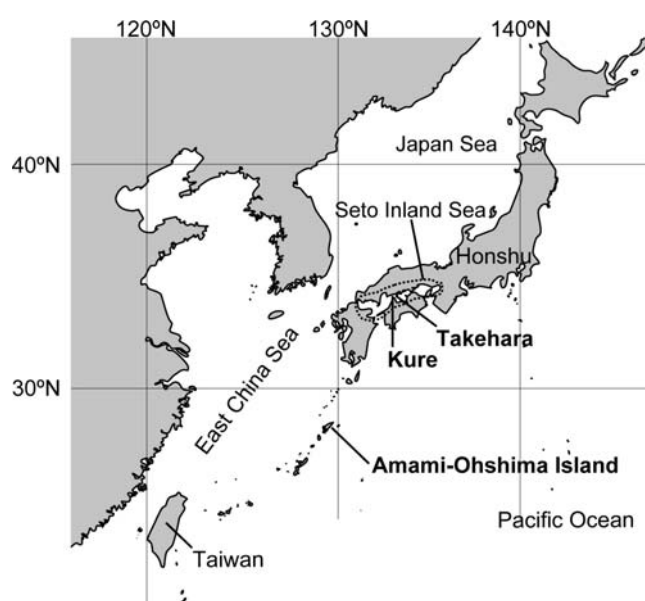


Fig. 1. Map showing the study sites: Kure, Takehara, and Amami-Ohshima Island.

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(Anker et al., 2005). Therefore, to survive in echiuran burrows, symbionts may have to adapt morphologically or behaviorally to the pressure of the undulating movement of the host.

Among the diverse dwellers of echiuran burrows, only a few are obligate symbionts (Anker et al., 2005). In Bivalvia, only two galeommatoidean bivalve species, *Pseudopythina ochestostomae* (Lasaeidae) and *Scintilla thalassemicola* (Galeommatidae), are known to have strong associations with echiurans (Habe, 1962; Morton and Morton, 1983). The former is a small bivalve dwelling in the burrows of *Ochetostoma erythrogrammon* and *Listriolobius sorbillans* in Hong Kong and the Ryukyu Islands (Kosuge et al., 2003; Morton and Scott, 1989). The latter is a small bivalve that attaches to the body surface of *Anelassorhynchus mucosus* in the rocky intertidal zone in Amakusa, Japan (Habe, 1962). Identification of a new close association between echiuran worms and bivalves would provide a good opportunity for further understanding the evolution, adaptation, and specialization of bivalves to symbiotic association with echiuran worms.

Sportellidae is a small bivalve family, recently moved from Cyamioida to Galeommatoidea owing to its molecular phylogeny (Campbell, 2000; Gribet and Distel, 2003; Taylor et al., 2007), and characterized by posterior inhalant and exhalant openings with very short siphons (Coan, 1999; Ponder, 1971). As living Sportellidae specimens previously have been rarely collected, their ecology and anatomy are not well known (Coan, 1999). However, several previous studies have suggested that some *Basterotia* species may have associations with other invertebrates. Kay (1979) briefly mentioned in her work that *Basterotia angulata* was found in the mud tubes of polychaete worms in tide pools in Hawaii, and Anker and colleagues (2005) reported that one specimen of an unidentified sportellid bivalve, possibly *Basterotia* sp., was collected from the burrow of an echiuran, *O. erythrogrammon*, in Venezuela. These fragmentary observations suggest that *Basterotia* may inhabit the burrows of annelid or echiuran worms.

Recently, we found two *Basterotia* species inhabiting echiuran burrows in the Japan Archipelago. To detect

symbiotic associations with echiurans, we investigated their habitat, distribution, morphology, growth patterns, and reproductive biology. Their shell morphology and anatomical structure were very different from those of other galeommatoidean bivalves that are symbiotic with various benthic invertebrates. By comparing the characteristics of *Basterotia* bivalves with those of other galeommatoidean bivalves, we discuss their morphological and ecological adaptations to symbiotic asso-



Fig. 2. Habitats, habits, and morphologies of the echiuran *Ikedosoma gogoshimense* (A–D) and its symbiotic bivalve *Basterotia gouldi* (E–K). (A, B) intertidal gravelly mud flats at Kure and Takehara; (C) a living echiuran; (D) entrance of an echiuran burrow around which mud dung is scattered; (E) symbiotic bivalves with their posterior parts gaping to the burrow lumen (white arrows and broken lines suggests bivalves and the echiuran burrow, respectively); (F) left and right valves illuminated by transmitted light (the shell of the posterior shaded part of both valves is thick); (G) an internal view of the right valve of a female brooding veligers in the ctenidium; (H) veliger larvae harbored in the ctenidium of a female; (I) juveniles attached to the inner ventral margin of the mantle of a female (white arrows); (J) posterior view of a living bivalve; (K) a creeping bivalve using its developed foot with its incurant aperture expanded. Abbreviations: exs, exhalant siphon; ins, inhalant siphon. Scale bar: 1 cm (C), 5 mm (E), 1 mm (F, G, I, J, K), 0.1 mm (H).

ciation with echiurans.

MATERIALS AND METHODS

Study site and study species

From June 2008 to June 2009, we sampled the sportellid bivalves of the intertidal zone at Kure (34°12'N, 132°40'E) and Takehara (34°19'N, 132°53'E) in the Seto Inland Sea and at Amami-Ohshima Island (28°22'N, 129°25'E) in the East China Sea, Japan (Fig. 1). On the intertidal gravelly mud flats at Kure (Fig. 2A) and Takehara (Fig. 2B) are numerous burrows of two echiuran species, *Ikedosoma gogoshimense* (Echiuridae) (Fig. 2C) and *Thalassema fuscum* (Thalassematidae), with the former being more abundant than the latter. *Ikedosoma gogoshimense* is a species endemic to Japan and mainly occurs in the intertidal gravelly mud flats of the Seto Inland Sea. It is a deposit feeder that produces an L-shaped burrow with a vertical and a horizontal branch (Fig. 4A; Kawaguti, 1971). Around the entrance of the burrow, the echiurans scatter ellipsoidal mud particles (Fig. 2D; Supplemental material online). The scatterings of mud particles and the blind burrow suggest that the echiuran is a deposit feeder. The intertidal gravelly coral-sand flats of Amami-Ohshima Island harbored *O. erythrogrammon* (Thalassematidae) (Fig. 3A, B), which is a deposit feeder producing a U-shaped burrow very similar to those reported from Hong Kong (Morton and Morton, 1983). However, the echiuran burrows at our site were somewhat more horizontal than vertical, due to the presence of gravel in the sediment (Fig. 4B).

Method

At Kure and Takehara, we first searched for *I. gogoshimense* burrow entrances by examining the centrifuged scatterings of the ellipsoidal mud particles excreted by the echiurans (see Method in Kawaguti, 1971; Fig. 2D). After finding a burrow entrance, we carefully excavated the burrow and investigated the presence or absence of symbionts; if they were present, we recorded the number, position, and posture of each. *Ikedosoma gogoshimense* burrows collapsed easily because they were constructed in muddy sediments. To prevent bivalves from being overlooked, we sieved the mud collected from alongside the burrow using a 2-mm-mesh sieve.

At Amami-Ohshima Island, *O. erythrogrammon* usually produces burrows under rocks embedded in the sediment (Fig. 4B). Therefore we searched for their burrows by turning over rocks. When we found a burrow, we collected the echiurans with their symbionts. We also checked the number, position, and posture of any symbionts in each burrow. At all sites, we measured the volume of host

echiurans using a graduated cylinder in the field.

To determine whether *Basterotia* bivalves live only in echiuran burrows, we surveyed the burrows of other sympatric invertebrates at Kure and Amami-Ohshima Island. At Kure, the burrows of *I. gogoshimense* were produced in the muddy sand or mud, while at Amami-Ohshima Island, those of *O. erythrogrammon* were pro-



Fig. 3. Habitats, habits, and morphologies of the echiuran *Ochetostoma erythrogrammon* (A, B) and its symbiotic bivalve *Basterotia carinata* (C–L). (A) an intertidal gravelly, coral-sand flat at the type locality of *Basterotia carinata* at Amami-Ohshima Island; (B) an echiuran in its burrow; (C) a bivalve found in an echiuran burrow (a black arrow and white broken lines suggest a bivalve and the echiuran burrow, respectively); (D) the posture of a bivalve with its posterior part opening into the burrow lumen (white broken line suggests the echiuran burrow); (E, F) the hinge structure of the right and left valves; (G, H) internal and external views of the right valves; (I, J) internal and external views of the left valves; (K) ventral view of a living bivalve with juveniles (a black arrow) attached around the internal aperture of the mantle of a female bivalve; (L) posterior view of the bivalve. The *Basterotia carinata* in D, E–J and L is the holotype (NSMT-Mo 77014). Scales: 1 cm (B, C), 5 mm (D), 1 mm (E–L).

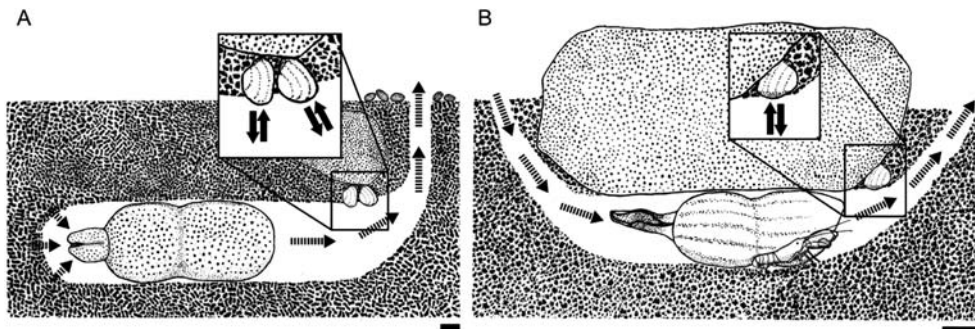


Fig. 4. Schematic representations of the habits of burrow-dwelling echiurans and their symbiotic bivalves (and a shrimp): **(A)** *Ikedosoma gogoshimense* occupied by *Basterotia gouldi*; **(B)** *Ochetostoma erythrogrammon* occupied by *Basterotia carinata* and a shrimp, *Alpheus barbatus*. Solid and broken arrows represent water currents created by the bivalves and their host echiurans, respectively. Scales: 1 cm (A, B).

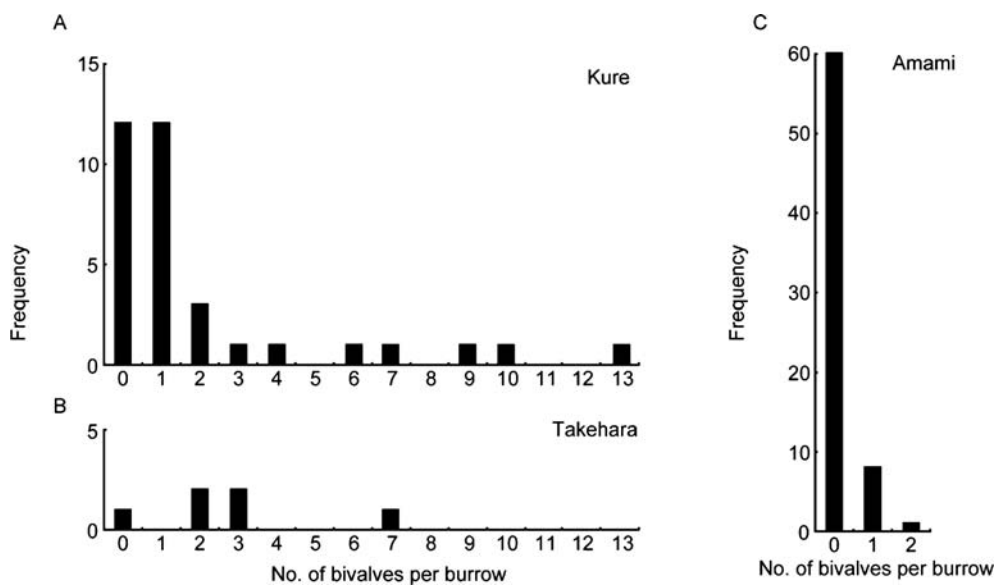


Fig. 5. Frequency distributions of symbiotic bivalves among echiuran burrows; *Basterotia gouldi* at Kure **(A)** and Takehara **(B)** and *Basterotia carinata* at Amami-Ohshima Island **(C)**.

duced in the coral sand which is composed primarily of foraminiferal shells mixed with small fragments of seashells and dead corals. Therefore, we investigated the burrows of invertebrates produced in each substrate type. In the field, we first searched for burrow entrances on the surface of the tidal flat or under rocks. When burrows were found, we carefully dug each burrow and sieved the sand or mud collected from alongside the burrow using a 2-mm-mesh sieve.

To analyze *Basterotia* growth patterns, we measured shell length (antero-posterior), shell height (ventro-dorsal), and shell width (left-right). In addition, to understand the reproductive biology of these bivalves, we dissected *B. gouldi*, which were sampled during the period of 23–25 June 2009 at Kure, under a microscope to determine whether they were brooding veligers or juveniles, and whether their sexual organs contained eggs or sperm. Furthermore, to investigate the interaction between the echiurans and their symbiotic bivalves, we analyzed the relationships between echiuran volume and the number of bivalves per burrow, and that between echiuran volume and bivalve shell length.

RESULTS

Host burrow use by symbiotic bivalves

Twenty-two (64.7%) of 34 *Ikedosoma gogoshimense* burrows in the gravelly mud flats at Kure harbored symbiotic

Basterotia gouldi, and five (83.3%) of the six burrows at Takehara harbored bivalves (Fig. 5A, B). The burrows contained 2.06 ± 0.55 (average \pm SE; range, 0–13) and 2.83 ± 0.95 (0–7) bivalves at Kure and Takehara, respectively. *Ikedosoma gogoshimense* constructed L-shaped burrows in the shallow rock-mud bottom of the mid- to lower intertidal zone (Fig. 4A). *Basterotia gouldi* bivalves were found in both vertical and horizontal portions of the burrow (Figs. 2E, 4A). The bivalves were sometimes found aggregated within the burrow (Fig. 2E). None of the bivalves was attached directly to the bodies of the echiurans. The posterior aperture of the bivalve with inhalant and exhalant siphons always opened into the burrow lumen (Figs. 2E, 4A), whereas the anterior parts were embedded in the burrow wall and attached by byssal threads to mud-embedded pebbles (Fig. 4A). Despite their sessile habit, the bivalves, if detached from the burrow rock and kept in aquaria, actively moved using their developed foot (Fig. 2K). Intensive searches for the bivalves by turning over rocks (for 20 h at Kure and 4 h at Takehara) revealed that the bivalves did not live in habitats other than *I. gogoshimense* (and, very rarely, *Thalassema fuscum*) burrows. The bivalve species were not found in the burrows of other sympatric invertebrates at the study sites (Table 1).

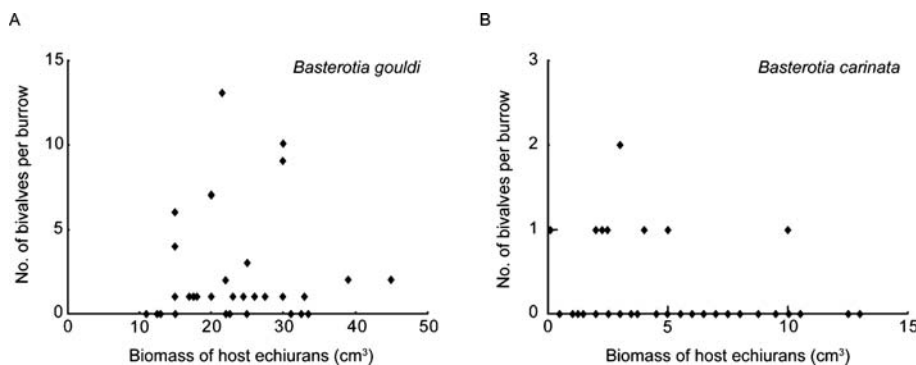
Among the 69 burrows of *O. erythrogrammon* observed at Amami-Ohshima Island, nine (13.0%) harbored *B. carinata* (Fig. 5C). Of these nine burrows, eight harbored one bivalve, and one harbored two bivalves whose shells touched each other. On average, one burrow harbored 0.14 ± 0.05 bivalves (range, 0–2). All bivalves were attached by byssal threads to pebbles embedded in the burrow ceilings (Figs. 3C, D, 4B). Almost all parts of the bivalves were embedded in the burrow wall, with the posterior end opening into the burrow lumen (Figs. 3D, 4B). The results of an intensive search for the bivalves while turning over rocks and digging the burrows of other sympatric benthic invertebrates for a total of 18 hours strongly suggested that the bivalves do not live outside the echiuran burrows (Table 1).

Interaction between echiurans and bivalves

Figure 6 shows relationships between the biomass volume of echiurans and the number of symbiotic bivalves per

Table 1. List of invertebrates excavated from burrows at the study sites, showing substratum type, numbers of burrows observed, and proportions of burrows containing *Basterotia* bivalves.

Locality	Species	Class: Family	Substratum type	No. of burrows observed	Prop. of burrows containing <i>Basterotia</i> bivalves (%)
Kure	Unidentified polychaete	Polychaeta: Terebellidae	muddy sand, mud	40	0
	<i>Sipunculus nudus</i>	Sipunculidea: Sipunculidae	muddy sand	1	0
	<i>Siphonosoma cumanense</i>	Sipunculidea: Sipunculidae	muddy sand	20	0
	<i>Ikedosoma gogoshimense</i>	Echiura: Echiuridae	muddy sand, mud	34	64.7
	<i>Thalassema fuscum</i>	Echiura: Thalassematidae	muddy sand, mud	2	50.0
	<i>Patinapta ooplax</i>	Holothuroidea: Synaptidae	muddy sand	40	0
	<i>Protankyra bidentata</i>	Holothuroidea: Synaptidae	mud	40	0
	<i>Nihonotrypaea japonica</i>	Malacostraca: Callianassidae	muddy sand, mud	20	0
	<i>Nihonotrypaea harmandii</i>	Malacostraca: Callianassidae	mud	2	0
	<i>Upogebia major</i>	Malacostraca: Upogebiidae	mud	1	0
Amami-Ohshima Is.	Unidentified polychaete	Polychaeta: Terebellidae	coral sand	20	0
	<i>Sipunculus nudus</i>	Sipunculidea: Sipunculidae	coral sand	2	0
	<i>Siphonosoma cumanense</i>	Sipunculidea: Sipunculidae	coral sand	20	0
	<i>Phascolosoma scolops</i>	Phascolosomatidea:	coral sand	5	0
		Phascolosomatidae			
	<i>Ochetostoma erythrogrammon</i>	Echiura: Thalassematidae	coral sand	69	13.0
	<i>Patinapta ooplax</i>	Holothuroidea: Synaptidae	coral sand	50	0
	<i>Polycheira rufescens</i>	Holothuroidea: Chiridotidae	coral sand	500	0
	<i>Holothuria paradalis</i>	Holothuroidea: Holothuriidae	coral sand	10	0
	<i>Nihonotrypaea japonica</i>	Malacostraca: Callianassidae	coral sand	20	0
	<i>Upogebia carinicauda</i>	Malacostraca: Upogebiidae	coral sand	3	0

**Fig. 6.** Relationships between the biomass volume of host echiurans and the number of symbiotic bivalves per burrow. *Basterotia gouldi* symbiotic with *Ikedosoma gogoshimense* at Kure (A) and *Basterotia carinata* symbiotic with *Ochetostoma erythrogrammon* at Amami-Ohshima Island (B).

burrow (Fig. 6A: *I. gogoshimense* and *B. gouldi* at Kure; Fig. 6B: *O. erythrogrammon* and *B. carinata* at Amami-Ohshima Island). No significant correlations were detected between parameters (Spearman's rank correlation test: *B. gouldi*, $\rho = 0.107$, $p = 0.554$; *B. carinata*, $\rho = 0.123$, $p = 0.315$).

Figure 7 shows relationships between the biomass volumes of echiurans and the average shell length of symbiotic bivalves in each burrow, and that between the biomass volumes of echiurans and the shell length of the largest bivalve in each burrow (Fig. 7A, C: *I. gogoshimense* and *B. gouldi*; Fig. 7B, D: *O. erythrogrammon* and *B. carinata*). No significant correlations were detected between parameters (Spearman's rank correlation test: host volume–average shell length, *B. gouldi*, $\rho = 0.246$, $p = 0.270$; *B. carinata*, $\rho =$

0.352 , $p = 0.356$; host volume–shell length of the largest bivalve, *B. gouldi*, $\rho = 0.325$, $p = 0.140$; *B. carinata*, $\rho = 0.067$, $p = 0.865$).

Morphology, morphometry, and anatomy of bivalves

B. gouldi have ovate shells with a very thick posterior part (Fig. 2F), whereas *B. carinata* have ovate-triangular shells with a strong carina running from the umbo to the postero-ventral margin (Fig. 3G–J). The shells of both species were equivale and had large posterior and small anterior gapes, even when the valves were tightly closed (Figs. 2J, 3L). The posterior gapes of *B. gouldi*

were larger than those of *B. carinata* (Figs. 2J, 3L). The ratios of shell height to shell length and shell width to shell length were plotted against shell length (Fig. 8), and the results showed that the symbiotic bivalves grew with their lateral shell morphometry unchanged (Spearman's rank correlation test: *B. gouldi*, $\rho = 0.018$, $p = 0.884$; *B. carinata*, $\rho = 0.246$, $p = 0.489$; Fig. 8A, C), whereas their relative shell width increased ontogenetically (*B. gouldi*, $\rho = 0.725$, $p < 0.0001$; *B. carinata*, $\rho = 0.673$, $p = 0.033$; Fig. 8B, D).

The internal anatomy of *B. gouldi* and *B. carinata* is shown in Fig. 9. Although the shell shape of these species is quite different, their internal anatomy is very similar. Both bivalve species have very short inhalant and exhalant siphons at their posterior ends. The posterior mantle has numerous small, well-developed tentacles around the inhal-

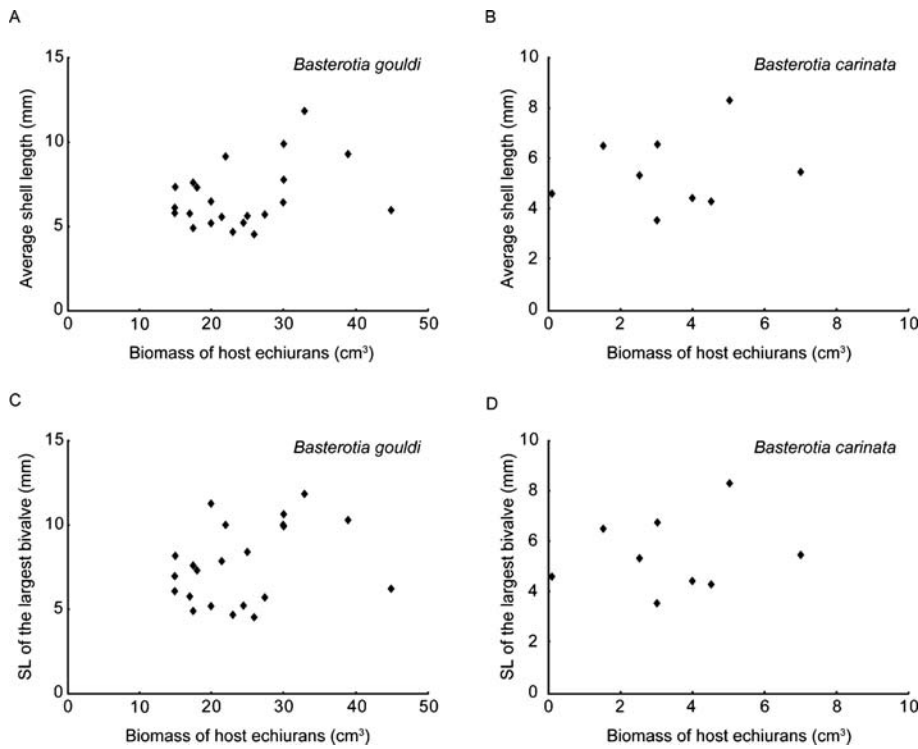


Fig. 7. Relationships of the biomass volume of host echiurans to average shell length of symbiotic bivalves dwelling in each burrow (A, B) and to shell lengths (SL) of the largest bivalves (C, D). *Basterotia gouldi* at Kure (A, C) and *Basterotia carinata* at Amami-Ohshima Island (B, D).

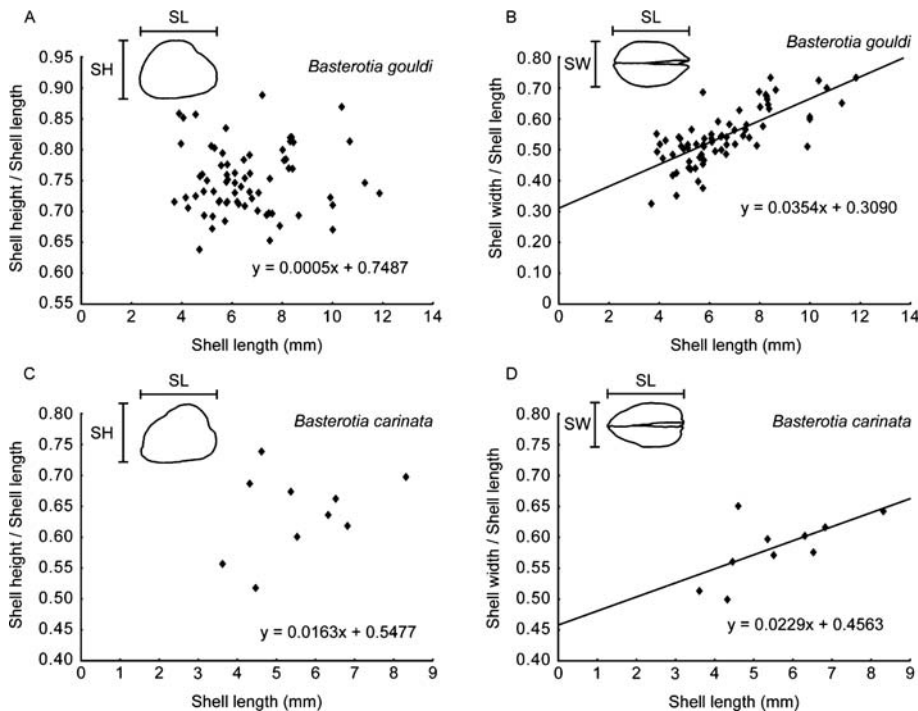


Fig. 8. Relationships of shell length to shell height/shell length ratio (A, C) and to shell width/shell length ratio (B, D). *Basterotia gouldi* at Kure (A, B), and *Basterotia carinata* at Amami-Ohshima Island (C, D).

ant and exhalant pallial apertures. The foot has byssal glands with fine, transparent byssal threads. The anteroventral side of the mantle has an aperture through which the

foot is extended. The outer demi-branch is smaller than the inner. The anterior and posterior adductor muscles are ovate and subequal, with the anterior being slightly smaller than the posterior. The digestive gland is brown, and is located anterior to the large, white gonad.

Reproductive biology and sexual composition of bivalves among burrows

In total, we dissected 40 *B. gouldi* bivalves collected from 12 burrows at Kure. Of these 40 individuals, eight were females, 14 were males, and 18 were immature or intermediate (i.e., individuals that did not exhibit any gonad development) (Fig. 10). Mature sperm were detected in smaller individuals (shell length; 7.0 ± 0.3 mm, range, 6–10 mm), whereas mature ova were detected in larger specimens (shell length; 9.2 ± 0.4 mm, range, 7–12 mm). The shell lengths of mature females were significantly greater than those of mature males (*t*-test, $P < 0.0001$; Fig. 9). Some females were brooding veliger larvae in the ctenidium (Fig. 2G, H), and juveniles with shell lengths of 1–2 mm were found in the ventral margins of the mantle cavities (Figs. 2I, 3K, 9). These small bivalves contained neither sperm nor ova. Of the eight females, one was brooding hundreds of veliger larvae and five juveniles, three brooded a single juvenile, and four brooded neither veliger larva nor juveniles. We also found one juvenile attached to the ventral mantle margin of a male.

Of 12 burrows, five harbored males and females, one harbored only males, and six harbored only immature individuals (Fig. 11). Among the burrows that harbored at least one male and one female, the male-to-female ratio was 1.86 ± 0.79 males to 1 female on average.

The symbiont community in echiuran burrows

A list of symbionts dwelling in the echiuran burrows is provided in Table 2. In addition to *B. gouldi*, we found a small gastropod species *Macromphalus tornatilis* (Vanikoridae) and a small crab species *Pinnixa* sp. (Pinnotheridae) inhabiting the burrow of *I. gogoshimense*.

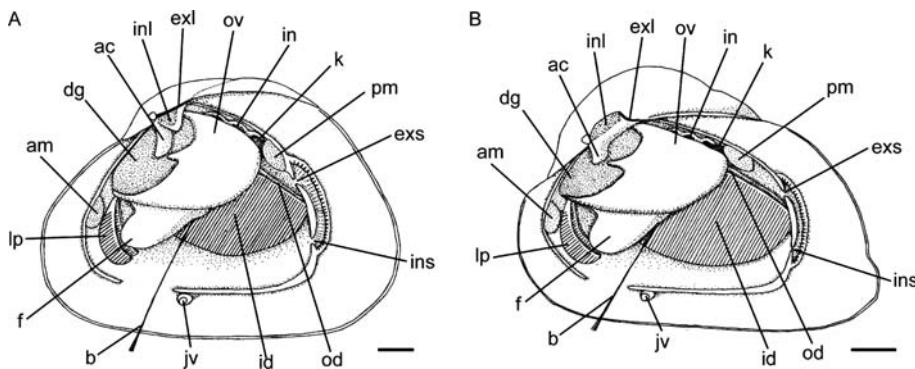


Fig. 9. Internal anatomy of the right parts of females of *Basterotia gouldi* (A) and *B. carinata* (B). Abbreviations: ac, anterior cardinal tooth; am, anterior adductor muscle; b, byssus; dg, digestive gland; exl, external ligament; exs, exhalant siphon; f, foot; id, inner demibranch; in, intestine; inl, internal ligament; ins, inhalant siphon; jv, juvenile; lp, labial palp; od, outer demibranch; ov, ovary; pm, posterior adductor muscle; k, kidney. Scales: 1 mm (A, B).

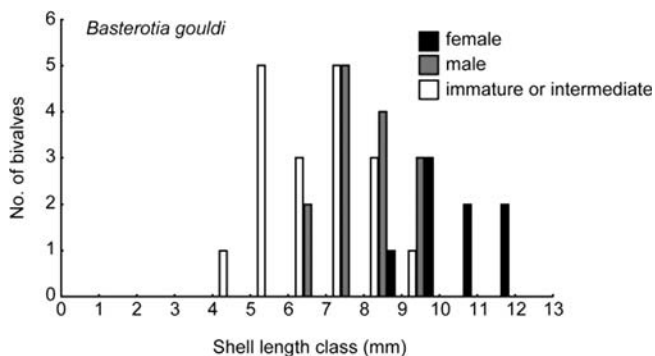


Fig. 10. Size distribution of *Basterotia gouldi* bivalves sorted by gravity and sex.

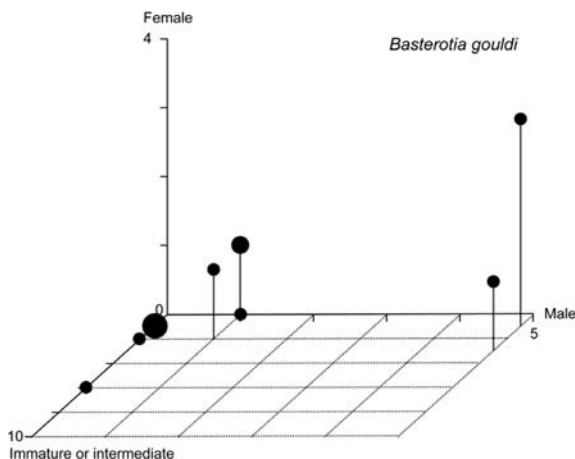


Fig. 11. Composition of sexes of *Basterotia gouldi* in each echiuran burrow. Symbol sizes are proportional to sample size.

M. tornatilis attached to the pebbles embedded in the burrow wall, and one specimen was found on the shell of a living *B. gouldi*. In *O. erythrogrammon* burrows, a small snapping shrimp species, *Alpheus barbatus*, was almost always found, and unidentified scale worms and phenacolepadid snails were sometimes found. *A. barbatus* and the scale worms were usually positioned behind the echiuran

body, whereas the phenacolepadid snails attached to rock embedded in the burrow wall. None of these symbiotic organisms was found outside the host echiuran burrows.

Species description

Family Sportellidae Dall, 1899
Genus *Basterotia* Mayer, in Hörnes, 1859

Basterotia carinata Goto, Hamamura and Kato n. sp.

[New Japanese name: Hainumikazegai]

Holotype: NSMT-Mo 77014, Shell length = 8.3 mm (Figs. 3D–J, L, 9B). The holotype specimen is

deposited with its host echiuran worm in the National Museum of Nature and Science, Japan.

Paratypes: Paratype specimens are deposited with their host echiuran worms in the National Museum of Nature and Science, Japan (NSMT-Mo 77015, Shell length = 6.8 mm; NSMT-Mo 77016, Shell length = 6.3 mm) and in the Smithsonian National Museum of Natural History, USA (USNM 1140542, Shell length = 5.5 mm, Host: USNM 1140543), respectively.

Type locality: Intertidal gravelly coral-sand flat at Nezebu, Amami-Oshima Island, Kagoshima Prefecture, Japan, 28°22'N, 129°25'E.

Diagnosis: Shell small, ovate-trigonal, equivalve, laterally inflated, with strong keel running from beak to postero-ventral corner. Shell having small gape at antero-ventral corner and large slit-like gape at postero-dorsal margin.

Description: Shell length to 8.3 mm. Shell relatively thick, ovate-trigonal, equivalve, laterally inflated, length/height about 1.4, length/width about 1.6 (holotype); beak prosogyrous, situated at about 20% from anterior end; anterior end rounded; posterior end subtruncate; ventral margin straight; external surface of shell with coarse pustules and irregular, coarse, commarginal growth checks; shell having a strong carina running from beak to the antero-ventral corner; angle of a carina most prominent dorsally; carina stout, weakly nodulous; shell having small gape at antero-ventral corner and large slit-like gape at postero-dorsal margin, even when valves are tightly closed; external ligament thin and very short on a short sturdy nymph; internal ligament thick, seated in an oblong, horizontal groove posterior of the beak; right valve with one prominent, antero-ventrally projecting cardinal (Fig. 3E); left valve with one ventrally projecting cardinal (Fig. 3F); both right and left valves having relatively wide escutcheon (Fig. 3L).

Anatomy: Both anterior and posterior adductor muscles ovate, subequal (Fig. 9B); the posterior adductor slightly larger than the anterior; the posterior part of the mantle having very short inhalant and exhalant siphons; both outer and inner demibranches present; the outer demibranch smaller than the inner; ctenidia joined anteriorly to inner and outer labial palps. Mantle edge smooth anteriorly and ventrally but with numerous, small, well-developed tentacles posteriorly

Table 2. List of symbiotic organisms observed in echiuran burrows, showing host echiuran species, number of burrow observed, symbiotic rate, and mean number (with standard error) of individuals per burrow.

Locality	Host echiuran species	No. of burrows observed	Symbiont species	Class: Family	Symbiotic rate (%)	Mean (\pm SE) per burrow
Kure	<i>Ikedosoma gogoshimense</i>	34	<i>Macromphalus tornatilis</i>	Gastropoda: Vanikoridae	26.5	0.47 \pm 0.16
			<i>Basterotia gouldi</i>	Bivalvia: Sportellidae	64.7	2.06 \pm 0.55
			<i>Pinnixa</i> sp.	Brachyura: Pinnotheridae	2.9	0.03 \pm 0.03
	<i>Thalassema fuscum</i>	2	<i>Basterotia gouldi</i>	Bivalvia: Sportellidae	50.0	0.50 \pm 0.50
Amami-Ohshima Is.	<i>Ochetostoma erythrogrammon</i>	69	Unidentified limpet	Gastropoda: Phenacolepadidae	2.9	0.03 \pm 0.02
			<i>Basterotia carinata</i>	Bivalvia: Sportellidae	13.0	0.14 \pm 0.05
			Unidentified scale worm	Polychaeta: Polynoidae	4.3	0.04 \pm 0.02
			<i>Alpheus barbatus</i>	Caridea: Alpheidae	80.0	0.82 \pm 0.05

around the inhalant and exhalant pallial apertures. Digestive gland brown, anterior to white gonad which occupies the middle of the visceral mass; antero-ventral side of the mantle having one aperture through which the foot is extended; foot having byssal gland with fine, transparent byssal threads. Rectum with several compact faecal pellets; the ventral mantle margin attached by several small juveniles.

Biology: Symbiotic in the burrow of *Ochetostoma erythrogrammon*.

Distribution: Only known from Amami-Ohshima Island, southern Japan.

Etymology: From the Latin *carina* meaning keel, *carinata* meaning keeled in reference to the prominent keel running along the central slope of the posterior part of bivalves.

Remarks: The holotype of *Basterotia carinata* resembles that of *Basterotia (Anisodonta) caledonica* in having an equivalve, wedge-shaped shell. However, the holotype of *B. carinata* remarkably differs from that of *B. caledonica* in having an extremely prominent carina running along the central slope of both the right and left valves (Fischer, 1886). This does not indicate that the holotype of *B. caledonica* is ontogenetically less developed than that of *B. carinata*, because the former is 1.5 times larger than the latter (Fischer, 1886). In addition, the anterior adductor muscle of *B. carinata* is ovate, whereas that of *B. caledonica* is rounded (Fischer, 1886). On the other hand, the prominent carinae are similar to those of *B. quadorata*. However, *B. quadorata* have escutcheon almost only on the left valve (Coan, 1999), whereas *B. carinata* are equivalve, having equally-sized escutcheons on both the right and left valves. In addition, *B. carinata* is relatively horizontally longer than *B. quadorata* (the height/length ratio of *B. quadorata* is about 1.1:1, whereas that of *B. carinata* is 1.4:1) (Coan, 1999; this study).

DISCUSSION

Commensalism of *Basterotia* bivalves with echiuran worms

Our field surveys demonstrated that two species of *Basterotia* were symbiotic with specific echiuran species, which were burrow-dwelling deposit-feeders found in gravely mud or coral-sand flats. Although the association of two other *Basterotia* species with annelids or echiurans has

been briefly suggested (Kay, 1979; Anker et al., 2005), to our knowledge the present study is the first report of tight association between *Basterotia* bivalves and burrow-dwelling echiurans.

In our surveys, *Basterotia* bivalves were found embedded in burrow walls with their posterior aperture gaping to the burrow lumen (Figs. 2E, 3D, 4). The postures of the bivalves suggested that they utilized the water pumped through the tunnels by the echiuran hosts. In addition, the burrows of echiurans provide the bivalves with safe sites protected from predation pressure and environmental stresses, such as desiccation and severe fluctuations in temperature and salinity. Conversely, the effect of bivalves on host echiurans is little known. Considering the *Basterotia* bivalve is a filter feeder and was nearly entirely embedded in the burrow wall, any deleterious effect of the bivalves on the hosts would seem to be small. At least, negative effects of the number and size of symbiotic bivalves on the biomass volume of hosts were not detected (Figs. 6, 7). These results suggest that the relationship between echiurans and bivalves may be commensal.

Morphological and ecological adaptations to symbiotic association with echiurans

Recent molecular phylogenetic analysis suggests that *Basterotia* belongs to Galeommatoidea, most members of which have commensal relationships with various invertebrates (Campbell, 2000; Taylor et al., 2007; Gribet and Distel, 2003). However, *Basterotia* bivalves have several morphological characters quite different from those of other galeommatoidean bivalves. Perhaps, these characters may be the result of adaptation or specialization to their unique symbiotic associations with echiuran worms. Here, we discuss the relationships between such morphological characters of *Basterotia* bivalves and their symbiotic association with echiuran worms.

For symbiotic organisms in echiuran burrows, position and posture in the burrow are important because the inside of the burrow is persistently disturbed by the host's peristaltic activity (Anker et al., 2005). *Basterotia* bivalves are almost completely embedded into the burrow wall, and their posterior apertures gape into the burrow lumen (Fig. 4). This posture makes it possible for them to safely utilize the water currents created by the host by reducing the exposure of

their body and friction with host echiurans. To achieve this posture, the posterior position of the inhalant siphon is necessary. *Basterotia* differs from other galeommatoidean species in having a posterior inhalant siphon (Coan, 1999). This morphological trait is considered as a pre- or post-adaptation to life in echiuran burrows. The symbiotic life of *Basterotia* is very similar to that of *Cryptomya californica* (Myidae), which is a small bivalve distributed along the Pacific coast of the Americas from Alaska to Peru. This bivalve species also lives embedded in the tunnel wall of other burrowing animals, such as thalassinidean shrimps, annelids, and echiurans (*Urechis caupo*), and protrudes its extremely short siphon into the burrow cavity for feeding and respiration (Lawly, 1978; MacGinite, 1935; Peterson, 1977; Wicksten, 1978; Yonge, 1951). The extremely short siphon of *Cryptomya* is a unique trait among the close relatives of Myidae and is considered as an adaptation to symbiotic life in the burrow (Lawly, 1978; Yonge, 1951). Similarly, *Basterotia* also have very short siphons (Figs. 2J, 3L, 9). However, the *Basterotia* siphons may have been pre-adapted to symbiotic life, as most members of Galeommatoidea have very short siphons, or lack the siphon entirely (Ponder, 1998).

Other galeommatoideans that live in echiuran burrows, e.g., *Pseudopythina ochetostomae* and *Scintilla thalassemicola*, are characterized by their very flattened shells (Habe, 1962; Morton and Scott, 1989). Given that they cling closely to the burrow wall or host's body, their flattened shell may be an adaptation to reduce the friction between the burrow wall and the host's body. In contrast, the shells of *Basterotia* become laterally inflated as they grow (Fig. 8B, D). This is probably because *Basterotia* bivalves are embedded in the narrow interspace between mud-embedded pebbles and the burrow lumen. Such a posture in the burrow causes morphological constraint, as the bivalves must grow not proportionately, but only laterally. Contrasting differences in shell morphology between *Basterotia* and other galeommatoideans living in the echiuran burrow may reflect the result of differential adaptation to symbiotic association with echiuran worms.

In addition, the posterior parts of the shells in *B. gouldi* are remarkably thick (Fig. 2F) and are reinforced by developed thick carina in *B. carinata* (Fig. 3H, J, L). In the present study, these parts were seen to slightly protrude into the burrow lumen and to interface with the host echiurans (Figs. 2E, 3D, 4). Therefore, these morphological traits are considered to be adaptations for withstanding the pressure caused by the host's undulating movement.

Host preference

Symbiotic association with echiurans is also known in *Pseudopythina ochetostomae*, a small bivalve species living in the burrow of *O. erythrogrammon* and *L. sorbillans* in Hong Kong and the Ryukyu Islands (Kosuge et al., 2003; Morton and Scott, 1989). This bivalve species belongs to Galeommatoidea and has completely lost its siphon (Morton and Scott, 1989). Although *Basterotia* and *P. ochetostomae* occupy similar ecological niches, they have different host preferences. *P. ochetostomae* is frequently found in the burrow of *L. sorbillans* (Kosuge et al., 2003), whereas *Basterotia* has never been found in burrows of *L. sorbillans*

(Kosuge et al., 2003; Goto, unpublished data). *L. sorbillans* extends its proboscis a great distance from the burrow entrance onto the surface of tidal flats and transports minute organic particles into its mouth through ciliate movements on the proboscis surface. The feeding habits of this echiuran suggest that the water current created by the host in the burrow is weak. Conversely, *O. erythrogrammon* and *I. gogoshimesne* powerfully inhale and exhale water in the burrow, thereby creating strong water currents through the burrow (Supplemental material online). Perhaps such strong water currents are necessary for the filter feeding of *Basterotia*, but not for that of *P. ochetostomae*.

Reproductive biology

Some members of Galeommatoidea show size-dependent sex change or protandrous consecutive hermaphroditism (Fox et al., 2007; Goto et al., 2007). Considering the size distribution of male and female bivalves recorded in this study (Fig. 10), *Basterotia* appears to change sex. *B. gouldi* changed sex from male to female when 8–10 mm long (Fig. 10). Some individuals at intermediate sizes between males and females did not exhibit any gonadal development. Such individuals may be intermediates in the process of changing sex from male to female.

Some galeommatoidean species are known to brood veligers or juveniles in the mantle cavity (Kato and Itani, 1995; Lützen and Nielsen 2005). In this study, we found that the largest *B. gouldi* female harbored hundreds of veligers in the ctenidium (Fig. 2G, H), and that mature individuals of *B. gouldi* and *B. carinata* brooded several juveniles in the ventral mantle margin (Figs. 2I, 3K, 9). Although the juvenile-brooding habit is known in two *Basterotia* species (Coan, 1999; Ponder, 1971), the veliger-brooding habit of *Basterotia* is reported for the first time in this study. In addition, we collected juveniles from the ventral margin of a mature male. This suggests that brooded juveniles may contain individuals that reattach to mature individuals after being released from the brooding female.

The symbiont community in echiuran burrows

To date, the only known commensal of *Ikedosoma* has been a snapping shrimp, *Alpheus barubatus*, collected from *Ikedosoma* sp. in Taiwan (Tzeng and Cheng, 1992), whereas no commensals were found in the burrows of *T. fuscum* in Hong Kong (Morton and Morton, 1983). In the present study, we recorded *B. gouldi*, *M. tornatilis*, and *Pinnixa* sp. as symbionts of *I. gogoshimensis*, and *B. gouldi* as a symbiont of *T. fuscum* (Table 2). The small gastropod *M. tornatilis* was often found on pebbles embedded in burrow walls, suggesting that it occupies a relatively similar ecological niche to that of *B. gouldi*. The body of *Pinnixa* sp. is horizontally elongated, a body shape typically found in symbiotic crabs dwelling in echiuran burrows (Anker et al., 2005). In the Seto Inland Sea, three species of crabs, *Sestrostoma balssi*, *Sestrostoma toriumii*, and *Pseudopinnixa carinata*, were collected from the U-shaped burrows of *Urechis unicinctus* (Itani et al., 2005). However, we did not find these symbiotic organisms in *Ikedosoma* or *Thalassema* burrows.

The burrows of *O. erythrogrammon* have been surveyed in Hong Kong, the Ryukyu Islands, and Venezuela (Anker et

al., 2005; Kosuge et al., 2003; Morton and Morton, 1983; Nomura, 2000). The burrows are inhabited by various organisms, such as the small bivalve *P. ochetostomae*, the small crab *Mortensenella forceps*, the small snail *Sigaretornus planus*, the snapping shrimp *A. barbatus*, by an unidentified scale worm in Hong Kong and the Ryukyu Islands (Kosuge et al., 2003; Morton and Morton, 1983; Nomura, 2000), and by the small crab, *Pinnixa* sp. aff. *chaetoptera*, *Pinnixa* sp. aff. *monodactyla*, and the small bivalve, *Basterotia* sp., in Venezuela (Anker et al., 2005). In this study, we found a snapping shrimp, *A. barbatus*, the small bivalve *B. carinata*, an unidentified scale worm, and a phenacolepadid limpet in the burrows of *O. erythrogrammon* at Amami-Oshima Island (Table 2). An extensive geographic comparison of symbiont communities may reveal whether these associations are maintained throughout the tropical waters of the world.

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