Unique Bacteria in the Coelomic Fluid

**Microbial Communities Associated with Holothurians: Presence of Unique Bacteria in the Coelomic Fluid**

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Marine invertebrates interact with various microorganisms ranging from pathogens to symbionts. One-to-one symbiosis between a single microbial species and a single host animal has served as a model for the study of host-microbe interactions. In addition, increasing attention has recently been focused on the complex symbiotic associations, e.g., associations between sponges and their symbionts, due to their biotechnological potential; however, relatively little is known about the microbial diversity associated with members of the phylum **Echinodermata**. Here, for the first time, we investigated microbial communities associated with a commercially important holothurian species, *Apostichopus japonicus*, using culture-dependent and -independent methods. Diverse and abundant heterotrophs, mostly **Gammaproteobacteria** members, were cultured semi-quantitatively. Using the cloning and sequencing technique, different microbial communities were found in different holothurian tissues. In the holothurian coelomic fluid, potentially metabolically active and phylogenetically unique members of **Epsilonproteobacteria** and **Rickettsiales** were discovered. This study suggests that coelomic fluids of marine invertebrates, at least those inhabiting intertidal areas where physical and chemical conditions fluctuate, provide microbes with unique and stable habitats.

**Key words:** holothurians, microbial diversity, coelomic fluid, host-microbe association, fisheries implications

Marine invertebrates harbor a wide array of ecto- and endo-symbiotic microbes. One-to-one symbiosis between a single microbial species and a single host animal has served as a model for the study of host-microbe interactions (7, 22, 37, 38, 39). Recently, increasing attention has been focused on complex symbiotic associations between a single host species and multiple microbial species/populations, e.g., sponge symbiosis due to their biotechnological potential (57, 58). Many marine invertebrates, particularly sessile to nearly sessile, are rich sources of valuable metabolites (9, 14). An individual sponge harbors dozens of different microbial species, in which bacteria comprise up to 40% of total sponge volume (59). Sponge-dwelling bacteria or the sponge itself produce a variety of bioactive compounds showing antiviral, antibiotic, and antimalarial activities (45, 58). From most phyla of marine invertebrates, host-associated microbial communities have been characterized; however, relatively little is known about microbial diversity within and on **Echinodermata** members such as sea stars, sea urchins, and sea cucumbers (19).

Holothurians are a group of marine invertebrates belonging to **Echinodermata**, and are distributed widely throughout the world from intertidal zones to the deep sea. Some holothurian species have a high market value and thus are heavily overexploited (18). Holothurians are model organisms for studying processes related to organ regeneration and organogenesis since they regenerate their digestive tract within a few weeks after self-evisceration (12, 42). Their body wall and coelomic fluid contain bioactive compounds with antibacterial, antifungal, anticoagulant, and antitumor activities (8, 17, 29). Although some microbes, e.g., members of the genera **Pseudoalteromonas** and **Salegentibacter**, have been isolated in pure cultures from holothurians (1, 23, 34, 48), little is known about the associated microbial diversity. In this study, we investigated microbial communities associated with a commercially important holothurian, *Apostichopus japonicus* (**Echinodermata**: **Holothuroidea**). We quantified culturable microorganisms in major microbial habitats in the holothurian body, including the body surface, digestive tract, and coelomic fluid. In addition, microbial diversity and niche segregation were comprehensively and quantitatively evaluated using culture-independent approaches.

**Materials and Methods**

**Sampling and processing**

Individuals of natural holothurians, *A. japonicus* (**Echinodermata**: **Holothuroidea**), were collected from the Funka Bay and Ainuma fishing port (42°08′N, 140°06′E), Hokkaido, Japan. Individuals (15.0–22.1 cm in length, 195–315.5 g) were rinsed with sterile 75% ASW (NaCl, 22.5 g; KCl, 0.5 g; MgSO₄·7H₂O, 4.0 g; MgCl₂·6H₂O, 8.1 g; CaSO₄·2H₂O, 1.0 g per liter) and were aseptically dissected into the small intestine, large intestine, body surface, and coelomic fluid. Briefly, outer surfaces of the body wall (12 cm² per individual) were sampled using sterile cotton-tipped swabs (Men-tip; J.C.B, Industry, Tokyo, Japan). After wiping the body surface with 70% ethanol, the body wall was cut with a disposable scalpel blade to collect the coelomic fluid using a sterile syringe. Finally, small and large intestines were sampled. The cotton-tipped swabs and digestive tracts were homogenized in sterile 75% artificial seawater (ASW) using a stomacher (Pro-media, SH-IIM; ELMEX, Tokyo, Japan) (49). Samples were serially diluted in sterile 75% ASW for the cultivation test.
Cell counting and cultivation test

Total cell counts were obtained by direct cell counting of the glutaraldehyde-fixed cells with DAPI (4,6-diamidino-2-phenylindole) using epifluorescence microscopy (47). The abundance and diversity of culturable heterotrophs were estimated as previously described (56). After 7 days’ cultivation under 20°C, about thirty colonies were randomly picked up from plates. Isolates were physiologically characterized as previously described (49). In addition, 16S RNA gene sequences of isolates were determined using the primers 24F and 1509R (Table 1). The PCR conditions were as follows: 35 cycles of 94°C for 1 min, 55°C for 1 min, and 72°C for 1.5 min. GoTaq Green Master mix (Promega) was used for the PCR reactions.

16S RNA gene clone library

Microbial diversity was assessed for holothurians collected from the Ainuma fishing port. Microbial DNA was directly extracted from the microbial community in each homogenate sample or coelomic fluid with a Wizard Genomic DNA Purification Kit (Promega). DNA extracted from seawater surrounding the holothurian or the large intestine homogenate. The bacterial community structures were compared by cluster analysis based on the clonal frequency of each representative phylotype. The square distance was determined by the Ward method (41).

Construction of phylogenetic tree

The 16S RNA gene sequences of representative clones and isolates were aligned with ARB software (25). Alignments were manually verified with known secondary structure regions. Phylogenetic analyses were restricted to nucleotide positions that could be unambiguously aligned. Phylogenetic trees were generated by a distance method using PAUP* 4.0b (54) and ARB. Distances were estimated with the Jukes-Cantor correction. Bootstrap analyses with 100 trial replications were used to obtain confidence estimates for the tree topologies.

FISH analysis

Fluorescent in-situ hybridization was performed as described elsewhere (3, 21, 40, 51). In brief, cells were hybridized with the Cy3-labeled probes (Table 2) for 4.5 h at 46°C. The percentage of fluorescently-labeled cells to DAPI-stained cells was determined using epifluorescence microscopy.

Nucleotide sequence accession numbers

The 16S RNA gene sequences of the representative isolates and clones obtained in this study have been deposited in DDBJ/EMBL/GenBank under Accession No. AB550432 to AB550557.

Results and Discussion

Total cell counts and cultivation test

Total cell densities were estimated for the holothurian body surface, large and small intestines, and coelomic fluid from two individuals (Table 3). Unexpectedly, many prokaryotic cells were observed even in the coelomic fluid, which varied in the range of 4.8×10⁵ to 9.9×10⁵ cells mL⁻¹. Cells observed in the coelomic fluid were mostly twisted rods (Fig. S1). The culturable heterotrophic populations generally accounted for less than 5% of the total cell counts (Table 3). In the sample Li-2 (the large intestine from the Ainuma individual), the population of culturable heterotrophs was probably overestimated, potentially due to the uneven distribution of cells in the large intestine homogenate.

Table 2. Oligonucleotide probes and competitors used in this study

| Probe or competitor | Specificity         | Target RNA | Reference |
|---------------------|---------------------|------------|-----------|
| Probe ARCH915       | Archaea             | GTGCTccccCGCCAATTcCT | 16S | 915–935 | 20 (53) |
| Probe EUB338        | Bacteria            | GCTGCCTccccCGTcAGAGT | 16S | 338–355 | 20 (2) |
| Probe ALF968        | Alphaproteobacteria | GGTAGGCTTCTGCGGTTT | 16S | 998–986 | 20 (35) |
| Probe GAM42a        | Gammaproteobacteria | GCCTccccCATCGTTT | 23S | 1,027–1,043 | 35 (26) |
| Probe BET42a        | Betaproteobacteria | GCCTccccACTCGTTT | 23S | 1,027–1,043 | 35 (26) |
| Probe EP402-423     | Epsilonproteobacteria | GAAAGCCTCACCTCCACG | 16S | 402–423 | 30 (55) |
| Probe CFB560        | Bacteroidetes       | WCGCTTTAAACCcART | 16S | 560–575 | 30 (43) |
| Probe PLA886        | Planctomycetes      | GCCTTGcGACcATcCTcC | 16S | 886–904 | 35 (36) |
| Competitor GAM42a   | Gammaproteobacteria | GCCTccccACTCGTTT | 23S | 1,027–1,043 | 26 (26) |
| Competitor BET42a   | Betaproteobacteria | GCCTccccACtCGTTT | 23S | 1,027–1,043 | 26 (26) |
Table 3. Total cell counts and culturable population from the holothurian tissues

| Sample code | Total cell counts | Culturable cell counts | Percentage of culturable cells (%) |
|-------------|-------------------|------------------------|-----------------------------------|
| Si-1        | $4.1 \times 10^5$ | $7.6 \times 10^4$      | 1.9                               |
| Si-2        | $6.1 \times 10^5$ | $1.8 \times 10^4$      | 3.0                               |
| Li-1        | $3.6 \times 10^6$ | $8.8 \times 10^4$      | 2.4                               |
| Li-2        | $3.3 \times 10^3$ | $2.6 \times 10^3$      | 78.8                              |
| U-1         | $1.2 \times 10^4$ | $1.1 \times 10^3$      | 0.9                               |
| U-2         | $3.8 \times 10^3$ | $1.7 \times 10^3$      | 0.5                               |
| C-1         | $9.9 \times 10^3$ | $8.4 \times 10^3$      | 0.9                               |
| C-2         | $4.8 \times 10^3$ | $1.3 \times 10^3$      | 0.3                               |

* Codes denote holothurian body parts and geographical origin. Si, Small intestine; Li, Large intestine; U, Surface; C, Coelomic fluid; 1, Funka Bay; 2, Ainuma Fishing Port.
* Si-1, Si-2, Li-1, and Li-2, cells g$^{-1}$; Li-1 and Li-2, cells cm$^{-2}$; C-1 and C-2, CFU mL$^{-1}$.
* Si-1, Si-2, Li-1, and Li-2, cells g$^{-1}$; U-1 and U-2, cells cm$^{-2}$; C-1 and C-2, CFU mL$^{-1}$.

Microbial communities evaluated by 16S rRNA gene library

Bacterial 16S rRNA gene clone libraries were successfully constructed using two universal primer sets from all holothurian samples (Table 1). The archaeal 16S rRNA gene was not amplified from any samples used in this study, although archaean diversity was previously assessed for the midgut contents of a deep-sea holothurian species (27). A total of 90 different bacterial phylotypes were identified from the 8 libraries on the basis of classification with ≥94% identity (Table S2). The coverage values were 58.3 and 70.0% (small intestine), 61.5 and 65.2% (large intestine), 86.4 and 95.7% (coelomic fluid), and 61.5 and 86.4% (body surface). A distinctive bacterial community was detected in each holothurian niche (Fig. 2).

The rRNA gene clones affiliated to the class Alphaproteobacteria were dominantly detected in all holothurian samples (13.0–54.5% in clonal frequencies). Alphaproteobacterial clones mainly belonged to three different subgroups (Fig. 3A). Roseobacter clade was mainly detected in small and large intestines (up to 43.5% in clonal frequency). These clones were closely related to clones or isolates previously retrieved from various marine environments including coastal and pelagic seawater, sediments, and algae- and invertebrates-associated habitats (4, 5) (Table S2). Members of the genus *Deflavicoccus* were mainly detected from the body surface and the coelomic fluid (up to 54.5% in clonal frequency) (Table S2). Many alphaproteobacterial sequences from the coelomic fluid (28 and 37.5% in clonal frequencies; approximately half of the alphaproteobacterial

Culturable heterotrophs

There were marked differences in the composition of culturable microbial community in each sample (Fig. 1). Heterotrophic microorganisms associated with holothurians were represented by members of eight different phylotic groups within the phyla *Proteobacteria* and *Bacteroidetes*: *Roseobacter* clade, *Shewanella* spp., *Pseudoalteromonas* spp., *Vibrio* spp., and family *Flavobacteriaceae*. The most frequently recovered population consisted of members of the genus *Pseudoalteromonas* (Fig. 1 and Table S1), which accounted for up to 52% of total culturable heterotrophs. The physiological tests of culturable populations indicated that bacteria with the ability to utilize alginate are more frequently recovered from digestive tracts than from other body parts (Table S1). Abundant and diverse heterotrophs were recovered from holothurian coelomic fluids. In the coelomic fluid of Funka Bay individual (C-1), *Flavobacteriaceae* members comprised 27.6% of isolates. In contrast, *Vibrio* members represented the second most frequently recovered population in the Ainuma coelomic fluid (C-2). Considering the total cell counts and the composition of culturable heterotrophs, microbial communities found in the coelomic fluids cannot be regarded as contaminants from other body parts.

Table 4. FISH-direct count analysis of metabolically active microbial populations associated with holothurians

| Samples          | Total cell counts$\pm$SD (10$^5$ cells unit$^{-1}$) | % of DAPI stained cells$\pm$SD |
|------------------|---------------------------------|-------------------------------|
|                  | EUB338 | ARCH915 | ALF968 | BET42a | GAM42a | EP402-423 | CFB560 | PLA886 |
| Small intestine  | 22.1±3.0 | 72.6±9.2 | ND | 27.8±3.9 | 7.0±0.8 | 2.5±0.4 | ND | 4.0±1.2 | 12.2±0.2 |
| Large intestine  | 23.9±0.9 | 74.4±9.8 | ND | 13.9±2.1 | 3.6±2.2 | 3.4±1.4 | ND | 3.0±0.3 | 7.8±1.7 |
| Surface          | 2.2±0.3 | 77.9±6.5 | ND | 46.9±7.5 | 3.9±3.5 | 9.6±2.3 | ND | ND | ND |
| Coelomic fluid   | 0.4±0.1 | 76.3±4.4 | ND | 45.2±2.2 | 2.5±0.3 | 21.3±3.1 | ND | ND | ND |

SD, standard deviation (n=4).
ND, not detected.
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cloned) formed a novel cluster within the order Rickettsiales (Fig. 3A). Most Rickettsiales members have been recognized as obligate intracellular parasites of arthropods with the ability to infect vertebrates (44, 60). Recently, Rickettsiales have also been found in the cells of leeches and marine ciliates (20, 60, 62). This study confirmed that Rickettsiales members could associate with more diverse animals than previously recognized. Clones affiliated to Planctomycetes and Fusobacteria were dominantly detected from holothurian digestive tracts (Fig. 2). These bacteria are frequently retrieved from marine snow and sediments (15), and thus may be transported with sediment particles into holothurian digestive tracts. Clones of the class Epsilonproteobacteria were detected only from the coelomic fluid (8.3 and 52.0% in clonal frequencies) (Fig. 2 and Table S2). As in the case of Rickettsiales clones, epsilonproteobacterial clones formed a novel clade which distantly related to members of the genus Sulfurospirillum (up to 92% sequence similarity) (Fig. 3B). Members of the genus Sulfurospirillum are microaerobic sulfur reducers, which include two validly described marine species, i.e., S. carboxydovorans and S. arcachonense (11, 16). Known marine habitats from which Sulfurospirillum relatives were frequently recovered are restricted to sulfidic environments such as deep-sea vents and whale falls (32). Although the symbiotic ability of the Sulfurospirillum members is unclear, deep-sea vent Epsilonproteobacteria members have the ability to associate with various invertebrates (30, 31, 33, 61).

Whole-cell FISH analysis

Microorganisms with high fluorescence emission after FISH hybridization have high growth rates and high metabolic activity (3). Metabolically active microbial populations were quantified using eight different fluorescent probes (Table 2). In general, FISH analysis confirmed the clone library data. Bacteria detected by the EUB338 probe accounted for approximately three fourths of total cells in all holothurian samples (Table 4). Epsilonproteobacteria specific probe-binding cells were observed only in the coelomic fluid (Fig. 4). ALF968 probe-binding cells were abundantly found in all holothurian samples. These cells existed freely even in the coelomic fluid where Rickettsiales members accounted...
for half of the alphaproteobacterial clones, suggesting that Rickettsiales members detected in the coelomic fluid lack the ability to inhabit holothurian cells.

Conclusion

This study revealed for the first time the diverse and abundant microbial community associated with the holothurian body. In particular, it was remarkable that probably metabolically active and phylogenetically unique Epsilonproteobacteria and Rickettsiales members were discovered in the holothurian coelomic fluid, which contains antibacterial compounds (8). In addition, Echinodermata immunocytes produce reactive oxygen species (ROS) when in contact with bacteria or bacterial cell wall proteins (28). Nevertheless, potentially pathogenic microbes and fungi were repeatedly isolated from the holothurian coelomic fluid (1, 46). In addition, the coelomic fluids of oysters and abalones were used as inocula for isolating novel microorganisms (10, 24, 52). Echinoderms, at least those inhabiting intertidal areas where physical and chemical conditions fluctuate, have increased ability to maintain homeostasis in coelomic fluid (6). These suggest that coelomic fluid of marine invertebrates provides various oceanic microorganisms with unique, stable habitats. Focal points raised by this study for future research include: time course and evolution of symbiont diversity, microbial metabolism, host-microbe interactions and potential biotechnological and fisheries implications.

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