First report of segmented filamentous bacteria associated with *Rhigonema* sp. (Nematoda: Rhigonematidae) dwelling in hindgut of *Riukiaria* sp. (Diplopoda: Xystodesmidae)

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Summary

We morphologically and molecularly characterized segmented filamentous bacteria (SFB) associated with *Rhigonema* sp. nematodes in millipede hindguts. Seventy-three *Riukiaria* sp. millipedes were collected from a broad-leaf forest in Japan, and nematodes were excised from the millipede’s hindguts. The occurrence rate of SFB associated with nematodes was 24 % (10/41) for males, 47 % (14/30) for females, and 100 % (2/2) for juveniles. Genomic DNA was extracted from four SFB-rich nematode heads, and we obtained 40 bacterial clones via analysis of nearly full-length 16S rDNA gene sequences. At the phylum level, Firmicutes, Proteobacteria, and Verrucomicrobia accounted for 55 %, 40 %, and 5 % of SFB, respectively. In Firmicutes, Clostridiaceae (28 %) and Lachnospiraceae (15 %) were the dominant groups. Our sequences were divided into seven and three sub-clades between Firmicutes and Proteobacteria in the phylogenetic tree. In the Firmicutes clade, eight sequenced were classified as Lachnospiraceae with a bootstrap value >83 %. A phylogenetic tree involving known uncultured Lachnospiraceae sequences characterized the phylogenetic position of SFB associated with nematodes. Our results suggest that the association of SFB with nematode bodies was probably incidental and that SFB are not always present in millipede hindguts. Our bacterial groups corresponded to those of arthropod hindgut, and SFB associated with nematodes were inferred to belong to Lachnospiraceae. Because the Lachnospiraceae sequences obtained in this study showed specific lineages that differed from all the known deposited sequence data, these groups may be unique to *Riukiaria* sp.

Keywords: Firmicutes; Lachnospiraceae; phylogenetic analysis; morphology; 16S rDNA

Introduction

Guts of many arthropods (e.g., termites, cockroaches, and millipedes) are inhabited by segmented filamentous bacteria (SFB), which are also referred to as “long segmented filamentous structures” (Margulis et al., 1998; Thompson et al., 2012). In the guts, these Gram-positive endospore-forming microbes are attached to the epithelial walls (Klaasen et al., 1992). Although SFB are difficult to culture in vitro, this group is well known as common constituents of gut microbes (Krecek et al., 1987; Brune & Dietrich, 2015). Moreover, SFB were suggested to play a crucial role in host immune function through the coordination of T-cell responses (Ivanov et al., 2009). Among soil-dwelling arthropods, nematodes have frequently been isolated from the guts of termites (Carta & Osbrink, 2005), cockroaches (Ozawa et al., 2014), and millipedes (Hunt & Moore, 1995; Morffe & Hasegawa, 2017). SFB-like organisms were reported to be associated with the gut parasite nematodes *Rhigonema* spp.
in millipede hindgut as early as the 19th century (Leidy, 1853). The SFB associated with Rhigonema spp. were also confirmed in hindguts of Riukiaria spp. (Xystodesmidae) in Japan (Kanzaki et al., 2016). However, the frequency of occurrence and phylogenetic position of SFB associated with Rhigonema sp. in millipede hindguts has not been investigated in detail.

The purpose of this study was to clarify the occurrence rate and phylogenetic position of SFB associated with nematodes. For this, we collected millipedes in a broad-leaf forest and characterized SFB associated with hindgut-dwelling nematodes by both morphological and molecular approaches.

Materials and Methods

Adults and juveniles of Riukiaria sp. were collected at Hinokuma Park in Saga Prefecture (34°33′N, 130°35′E), Japan, in June 2018 (Fig. 1a). At the site, Cinnamomum camphora, Quercus aliena, Quercus serrata, and Schoepfia jasminodora of various ages were patchily distributed, and their broad-leaf litter covered the ground (Fig. 1b). Annual precipitation and annual mean temperature in June 2018, as recorded at the nearest weather station in Saga (8 km from the site), were 291 mm and 24.1 °C, respectively. The millipedes were brought back to the laboratory alive and stored at 20 °C for less than a month, until extraction of nematodes. The hypopygium of Riukiaria sp. was dissected with fine tweezers, and the hindgut was pulled out and excised. Rhigonema sp. individuals were isolated from the hindgut of juvenile, male, and female millipedes in distilled water. All the Rhigonema nematodes retrieved from each millipede’s hindgut were transferred into a Petri dish filled with distilled water. For each millipede, the numbers of Rhigonema sp. with and without SFB were counted under a stereomicroscope (SZX16, Olympus, Tokyo, Japan; maximum magnification ×115).

We performed a χ²-test to ascertain whether the frequencies of occurrence of nematodes and SFB associated with nematodes in guts differed significantly between male and female millipedes. A Pearson’s correlation test was also performed to assess the relationship between the number of nematodes per millipede and

| Sex       | Observed numbers | Body length (mm) | Occurrence frequency of nematodes ‡ | SFB associated with nematodes (%) |
|-----------|------------------|------------------|-------------------------------------|-----------------------------------|
|           |                  |                  | 0  | 1 – 10 | 11 – 20 | 21 – 30 | >30 |                  |
| Male      | 41               | 34.0 ± 0.4       | 4  | 10 (%) | 17 (41 %) | 13 (32 %) | 6 (15 %) | 1 (2 %) | 24 (6 %) |
| Female    | 30               | 36.5 ± 0.5       | 0  | 3 (10 %) | 7 (23 %) | 19 (64 %) | 1 (3 %) | 47 (15 %) |
| Juvenile  | 2                | 19.5 ± 2.5       | 0  | 0      | 1 (50 %) | 1 (50 %) | 0      | 100 (100 %) |

‡ Data shown are mean ± SE.

* Numbers in parentheses indicate relative abundance. The frequency distribution of nematode abundances was significantly different between male and female millipedes (χ²-test, χ² = 21.164, df = 4, P < 0.001), but there was no significant difference between the sexes with regard to occurrence rate of SFB associated with nematodes (χ²-test, χ² = 2.911, df = 1, P = 0.08). Data on juveniles were not used in these analyses owing to the small sample size.

Table 1. Numbers of millipede Riukiaria sp., occurrence frequency of nematode Rhigonema sp., and segmented filamentous bacteria (SFB) associated with nematodes in millipede hindguts.

Fig. 1. Location of sampling site in Japan and a collected millipede. (a) The gray circle indicates the sampling location. Bar = 100 km. (b) View of a sampled broad-leaf forest stand. (c) Riukiaria sp. collected beneath the litter layer on the ground.
the number of nematodes associated with SFB. All data analyses were performed in R 3.3.2 (R Core Team, 2016), and the significance level was set at $P < 0.05$. Extracted nematodes were observed under a light microscope (BX53, Olympus; ×100–400) for detailed observation of SFB. The nematodes were Gram stained with crystal violet (Wako, Osaka, Japan) for the presence of SFB, then photographed (TIFF format, 1360 × 1024 pixels) with a digital camera (DP70, Olympus) connected to the microscope.

Genomic DNA was extracted from four SFB-rich nematode heads using the DNeasy Plant Mini Kit (Qiagen, Valencia, CA, USA). To obtain nearly the full length of 16S rDNA, the extracted DNAs were amplified using Tks Gflex (Takara, Shiga, Japan) with the primer pair 9f (GAGTTTGATCCTGGCTCAG; Yoon et al., 1997) and 1541r (AAGGAGGTGATCCAGCCG; Sato et al., 2004). Thermal conditions were one cycle of 94 °C for 1 min, followed by 30 cycles of 98 °C for 10 sec, 55 °C for 30 sec, and 72 °C for 90 sec. Positive PCR products were cloned using the TA-Enhancer Cloning Kit (Nippon Gene, Toyama, Japan) according to the manufacturer’s instructions. Sixteen successfully inserted white colonies were picked up per nematode head part. Selected colonies were further amplified with Tks Gflex and the primer pair 9f and 1541r. The thermal cycle program comprised 25 cycles of 98 °C for 10 sec, 55 °C for 30 sec, and 72.0 °C for 90 sec. When PCR amplicons were successfully produced, they were purified by using ExoSAP-IT PCR Product Cleanup Reagent (Affymetrix, Thermo Fisher Scientific, Waltham, MA, USA) in accordance with the manufacturer’s instructions. Purified DNAs were labeled by using a BigDye Terminator v3.1 Cycle Sequencing Kit (Applied Biosystems, Carlsbad, CA, USA) and then sequenced by using an ABI 3730 sequencer (Applied Biosystems) with one of the three primers – 9f, 1541r, or 533f (GTGCCAGCAGCCGCGGTAA; Weisburg et al., 1991) – to obtain full-length 16S rDNAs.

Obtained sequences were adjusted manually in MEGA v. 7 software (Kumar et al., 2016; https://www.megasoftware.net/), and sequences of the same sample read by different primers were assembled into a longer sequence whenever possible. Forty sequences were deposited in the DDBJ under accession numbers LC462722–LC462761. Sequences were compared with deposited sequences and identified to the nearest taxon using pairwise searches with the basic local alignment search tool (BLAST) (Altschul et al., 1997; https://blast.ncbi.nlm.nih.gov/Blast.cgi).

To estimate the phylogenetic positions of the bacteria, we constructed phylogenetic trees by using the maximum likelihood

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**Fig. 2.** Micrographs of *Rhigonema* sp. isolated from millipede guts. Images show whole individual and enlarged head part. (a) *Rhigonema* sp. crawl out from a millipede hindgut. (b) *Rhigonema* sp. without segmented filamentous bacteria (SFB). (c) *Rhigonema* sp. with SFB (arrow). (d) Enlarged head part of *Rhigonema* sp. with associated SFB. (e) Gram staining with crystal violet of associated SFB at head part. (f) SFB attached to nematode body surface. Bars represent (a) 1 mm, (b) 500 μm, and (f) 100 μm.
method in MEGA. For the default substitution model, we selected General Time Reversible (GTR) with Gamma-distributed rates plus Invariant sites (G+I), which had the smallest Akaike’s information criterion (AIC) and Bayesian information criterion (BIC) based on the model selection implemented in MEGA. To draw the trees, we referred to the clone library of bacterial sequences derived from the cockroach *Shefordella lateralis* (Schauer et al., 2012), the termite *Reticulitermes santonensis*, and the millipede *Tachypodoiulus niger* (Thompson et al., 2012) deposited in GenBank. The sequences were aligned in MAFFT v. 7 software (Katoh & Standley, 2013; http://mafft.cbrc.jp/alignment/software/) with default settings. In addition, we used the closest sequences of named family

![Graph showing correlation between number of Rhigonema sp. nematodes per Riukiaria sp. millipede hindgut and number of nematodes associated with segmented filamentous bacteria (SFB), for those millipedes that had at least one nematode associated with SFB (n = 25, Pearson’s correlation test, r = 0.593, P = 0.0017).]

**Fig. 3.** Correlation between number of Rhigonema sp. nematodes per Riukiaria sp. millipede hindgut and number of nematodes associated with segmented filamentous bacteria (SFB), for those millipedes that had at least one nematode associated with SFB (n = 25, Pearson’s correlation test, r = 0.593, P = 0.0017).

![Pie chart showing occurrence rates (%) of bacterial taxa detected from segmented filamentous bacteria–enriched Rhigonema sp. heads in hindgut of Riukiaria sp. (40 clones).]

**Fig. 4.** Occurrence rates (%) of bacterial taxa detected from segmented filamentous bacteria–enriched Rhigonema sp. heads in hindgut of Riukiaria sp. (40 clones).
Fig. 5. A maximum likelihood phylogenetic tree of segmented filamentous bacteria (SFB) associated with Rhigema sp. based on partial 16S rDNA sequences. The substitution model was GTR with G+I sites (AIC = 33224.486, BIC = 35454.324). We used clone libraries of 76 bacterial sequences as ingroups derived from the cockroach Shefordella lateralis (Schauer et al., 2012), termite Reticulitermes santonensis, and millipede Tachypodoiulus nigri (Thompson et al., 2012) deposited in GenBank. No outgroups were used. Values ≥80% on branches indicate confidence limits estimated by bootstrap analysis with 1000 replicates.
Fig. 6. A maximum likelihood phylogenetic tree of associated Lachnospiraceae groups of subclade S1 based on partial 16S rDNA sequences. The substitution model was GTR with G+I sites (AIC = 53520.724, BIC = 57505.700). Known sequences of 181 uncultured Lachnospiraceae with nearly the full length of 16S rDNA deposited in GenBank were used as ingroup sequences. Clostridiaceae (JN653039), Eubacteriaceae (JN680665), Ruminococcaceae (JN680615), and Veillonellaceae (JN680647) were used as outgroups. Values ≥80% on branches indicate confidence limits estimated by bootstrap analysis with 1000 replicates.
or genus in GenBank identified by using pairwise BLAST searches as in-group sequences. To increase the resolution of phylogenetic positions, the sequences that were tentatively assigned to Lachnospiraceae (see Results) were re-analyzed by the maximum likelihood method as above involving all the known uncultured Lachnospiraceae sequences with nearly the full length of 16S rDNA deposited in GenBank. For the default substitution model, we selected GTR with G+I, which had the smallest AIC and BIC based on the model selection in MEGA. For the Lachnospiraceae tree, Clostridiaceae (JN653039), Eubacteriaceae (JN680665), Ruminococcaceae (JN680615), and Veillonellaceae (JN680647) in subclade S1 (see Fig. 6), Ruminococcus sp. (JX560551) in subclade S4 (see Fig. 7) were used as outgroups. The reliability of all tree topologies was evaluated by 1000 bootstrap resamplings (Felsenstein, 1985).

**Results**

The microscopy of Rhigonema sp. demonstrated that SFB attached to the nematode body (Fig. 2a–d), and the parts associated with SFB were stained purple with the crystal violet (Fig. 2e, f). The occurrence rates of Rhigonema sp. in millipede hindguts were 90% (37/41) in males, 100% (30/30) in females, and 100% (2/2) in juveniles (Table 1). The highest frequency of Rhigonema sp. per hindgut was 41% (37/41) in males, 100% (30/30) in females, and 100% (2/2) in juveniles (Table 1). The relative abundance of nematodes was significantly different between male and female millipedes (Table 1, χ²-test, χ² = 21.164, df = 4, P < 0.001). The occurrence rate of SFB associated with nematodes was 24% (10/41) in males, 47% (14/30) in females, and 100% (2/2) in juveniles, and there was no significant difference between sexes (Table 1, χ²-test, χ² = 2.911, df = 1, P = 0.08). We noted a significant positive correlation between the number of nematodes per millipede and the number of nematodes associated with SFB (Fig. 3, Pearson’s correlation test, n = 25, r = 0.593, P = 0.0017). Since 6 clones were identified as *Escherichia coli* (negative con-
trol) by BLAST search, they were excluded. DNA sequences were obtained from 69 % (40/58) of clones. At the phylum level, Firmicutes, Proteobacteria and Verrucomicrobia accounted for 55 %, 40 %, and 5 %, respectively (Fig. 4). In the Firmicutes, Clostridiaceae (28 %) and Lachnospiraceae (15 %) were the dominant groups, whereas Desulfovibrionaceae (18 %) dominated the Proteobacteria. These clones were used for the construction of a maximum likelihood tree with known, named sequences (Fig. 5). Our sequences were affiliated with taxonomic clades (Firmicutes, Proteobacteria, or Verrucomicrobia) in the tree. Among them, Firmicutes and Proteobacteria harbored seven and three subclades, respectively, with bootstrap (BS) values greater than 83 % (Fig. 5). The subclade S1 formed with Lachnospiraceae lineages, and two of five sequences (Rh25_5, Rh25_22) were close to uncultured Lachnospiraceae derived from S. lateralis (JN680655), with a 93 % BS value (Fig. 6). On the other hand, clones of Rh28_2, Rh28_10, and Rh29_11 formed a clade with an 88 % BS value. In a tree retrieved from the S4 clade, three of our sequences were positioned differently from most Lachnospiraceae sequences, with 100 % BS values (Fig. 7). The tree was an admix lineage of Clostridiaceae and Lachnospiraceae, and three sequences (Rh23_5, Rh23_23, Rh28_9) were nested within Clostridiaceae groups neighboring Lachnospiraceae clones, with a 99 % BS value.

Discussion

Our results clearly showed that most nematodes in millipede hind-guts were associated with SFB. This study is the first to reveal the identity of SFB associated with nematodes by using molecular methods. Moreover, our bacterial sequences were clustered with known data derived from arthropod guts at high BS values. On the basis of these results, we discuss the occurrence frequency and phylogenetic position of SFB associated with nematodes. Rhigonema spp. are assumed to be common in hindguts of R. ukiaria sp. in Japan (Kanzaki et al., 2016), and the frequency of occurrence of the nematode in millipede guts was high in both sexes (Table 1). However, the number of nematodes per millipede gut was significantly greater in females than males. Females feed more actively than males in order to obtain more nutrients for oviposition (Boggs, 1981), so nutritional levels likely differ between sexes, as reflected in the population size of nematodes in their guts. The occurrence rate of SFB associated with nematodes was less than 50 % for both sexes. Moreover, the number of nematodes per millipede hindgut and the number of nematodes associated with SFB were significantly and positively correlated. Thus, the SFB density in millipede guts might depend on the population size of parasitizing nematodes.

Microscopy of Rhigonema sp. revealed that associating SFB were Gram-positive, formed endospores, and attached to the body of nematodes. These morphological characters agree well with previous reports (Klaasen et al., 1992). Leidy (1853) observed a SFB associated with Rhigonema infectum, Thelastoma attenuatum and Aorurus agilis in the hindgut of the millipede Narceus annularis (summarized in Sayre & Starr, 1988). Further, more recently, Blatta orientalis, Leidyinema spp. nematodes were found to be covered with SFB in the hindgut of black cockroach, Blatta orientalis (Dr. Sergei Spiridonov, pers. comm.). Thus, SFB might generally associate with nematodes harbored within arthropod hindguts.

The most representative phyla in this study were Firmicutes and Proteobacteria, which were also dominant in the hindgut of Japanese cockroach, Periplaneta japonica (Vicente et al., 2018). The most frequent taxa were Clostridiaceae and Lachnospiraceae, which are anaerobic and unculturable taxa known to be an abundant group in arthropod guts (Engel & Moran, 2013; Vicente et al., 2018). Our sequence data were nested with known anaerobic bacteria in gut conditions of millipedes, termites, and cockroaches with a high BS probability (>80 % BS, Fig. 5). Common bacterial groups in arthropods’ hindguts were attached to nematode head parts. The SFB in the hindgut of the termite R. santonensis and the millipede T. niger were classified phylogenetically to a group of Lachnospiraceae (Thompson et al., 2012). Lachnospiraceae were common in the hindgut of arthropods, such as termites, cockroaches, and millipedes, and were attached to their gut walls (Thompson et al., 2012). In the Lachnospiraceae tree in subclade S1, our sequences were clearly separated from other known sequences (100 % BS, Fig. 6). Moreover, the refined tree of subclade S4 was constructed by admixed sequences of both Clostridiaceae and Lachnospiraceae; our sequences were nested within the former taxon but clearly differed from most members of the latter taxon (Fig. 7). The phylum Firmicutes included representatives of common gut bacteria, with most belonging to the family Lachnospiraceae, and highly specific lineages associated with the hindgut cuticle of arthropods (Brune & Dietrich, 2015). Since the sequences obtained in this study also showed specific lineages differing from all the known deposited sequence data, these groups might be unique to Rukiaria sp. In addition, one of our obtained sequences was not placed with any known taxa, implying that the sequence is from an as-yet-unknown taxon. The Lachnospiraceae lineages were distributed among various locations and suggested to include cryptic species, indicating that the phylogenetic relationship within this group should be further investigated.

Our morphological and molecular analyses helped to characterize the SFB associated with nematodes within millipede hindguts in Japan. The association of SFB with nematode bodies was probably incidental and did not occur in all millipede hindguts. While the function of SFB in arthropod guts remains unknown, Ivanov et al. (2009) reported their crucial role in a host immune function through the coordination of T-cell responses. The distribution and functional significance of nematode-associated SFB remain unclear. To resolve these issues, in future research we plan to observe the ultrafine structure of SFB-associated parts in nematodes.
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Conflict of Interest

The authors declare that there are no conflicts of interest.

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