Paragonimus westermani metacercariae in two freshwater crab species in Kagoshima Prefecture, Japan, as a possible source of infection in wild boars and sika deer

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ABSTRACT. Paragonimiasis is a particular foodborne parasitic disease that is endemic to southern Kyushu, including Kagoshima Prefecture, Japan. We previously detected Paragonimus westermani triploid larvae in meat samples obtained from wild boars and sika deer hunted in Akune City, Kagoshima Prefecture. These mammals act as paratenic hosts and their meat is a source of human paragonimiasis. Paratenic host mammals and humans become infected with the lung fluke, P. westermani, following consumption of second intermediate hosts, freshwater crab species, namely, Geothelphusa dehaani or Sawagani in Japanese, and Eriocheir japonica or Mokuzugani in Japanese. Therefore, this study aimed to investigate the current infection status of P. westermani in freshwater crabs in Akune City. We collected freshwater crabs from 15 locations and found that the prevalence of P. westermani metacercariae was 1.6% for Sawagani (15 of 941 examined) and 22.1% for Mokuzugani (21 of 95 examined). Based on the morphological characterization of metacercariae and molecular analyses of the internal transcribed spacer 2 region and mitochondrial 16S rRNA gene region using PCR-restriction fragment length polymorphism and sequencing, all metacercariae were identified as the triploid form of P. westermani. These results indicate that Sawagani and Mokuzugani serve as second intermediate hosts to maintain the life cycle of triploid P. westermani. Further, infection in crabs potentially leads to subsequent P. westermani infections in wild mammals, including wild boars and sika deer, both of which are considered important types of game meat in Japan.

KEY WORDS: foodborne disease, freshwater crab, game meat, lung fluke, Paragonimus westermani, triploid form

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Paragonimus westermani is one of the most critical helminths affecting both humans and carnivorous/omnivorous mammals. In Japan, this lung fluke species is present in 2 forms concerning chromosomal structure: the diploid and triploid forms [1]. Both forms infect humans; however, the resulting major respiratory symptoms developed in patients differ. Among patients infected with the triploid form, as adult flukes inhabit worm cysts formed in the lungs, chronic cough with rust-colored sputum is the most common symptom. Conversely, the diploid form generally causes pleural effusion without remarkable lesions in the lung parenchyma [12]. In mammals such as wild boars, which serve as paratenic hosts of both forms, most flukes remain in the larval stage within muscle tissues for an extended period while on their way to the pleural cavities [15]. Therefore, meat from paratenic hosts can be a source of infection in humans. Infection in humans through this route predominantly occurs in southern Kyushu due to the local food habit of consuming uncooked wild boar meat [13, 27]. We previously investigated the contamination status of P. westermani in meat samples from regionally-hunted wild boars and sika deer obtained from a slaughtering and processing facility in Akune City, Kagoshima Prefecture, in southern Kyushu. Larvae of the P. westermani triploid form were detected in samples of sika deer as well as wild boars [26, 27]. Recently, the demand for such wild game meats has increased nationwide [10], which are referred to as “Jibie” in Japanese, based on the French word, “gibier”.

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Paratenic host mammals and humans are infected with the *P. westermani* triploid form even through consumption of the second intermediate crab hosts, namely, the Japanese freshwater crab *Geothelphusa dehaani*, or Sawagani in Japanese, and the Japanese mitten crab *Eriocheir japonica*, or Mokuzugani in Japanese. Therefore, assessing the contamination status of these second intermediate hosts in Akune City is necessary to ensure food safety. Between 1960 and 1964, 41 Mokuzugani were captured, and 4 tested positive for *Paragonimus* metacercariae [8]. However, no epidemiological surveys for *Paragonimus* metacercariae have been conducted in intermediate host crabs thereafter. In this study, we collected Sawagani and Mokuzugani from rivers in Akune City and identified the isolated metacercariae at the species and form levels in accordance with their morphological and molecular characteristics. This study aimed to clarify the *P. westermani* metacercariae infection status of freshwater crabs, the consumption of which, as second intermediate hosts, potentially leads to infection in paratenic hosts.

**MATERIALS AND METHODS**

**Isolation and examination of Paragonimus metacercariae**

We manually collected Sawagani from 15 rivers in the central and southern parts of Akune City, Kagoshima Prefecture, from October 2014 to September 2019. We also captured Mokuzugani from 7 of the rivers, using crab traps, from August 2015 to September 2019 (Fig. 1). Crabs were individually examined using a previously described method [23] to obtain precise data regarding the prevalence and burden of *Paragonimus* metacercariae. Crabs were first immersed in ice-cold water for deep anesthesia through hypothermia. The carapace was then detached from the body. The gills and the internal organs, including the heart, midgut gland, gonad, and intestine, were dissected. These organs were compressed between 2 glass plates (10 cm × 6 cm) and examined for *Paragonimus* metacercariae using a stereoscopic microscope (SZX7; Olympus, Tokyo, Japan). To isolate metacercariae from the muscle tissues of the cephalothorax and legs, the tissues were minced in a meat grinder. Minced crab tissues were incubated with artificial gastric juice (0.1% pepsin, proteolytic activity 1:10,000, Nacalai Tesque, Inc., Kyoto, Japan, plus 0.7% HCl), and placed in a shaking incubator (100 rpm) at 37°C for 3 hr. The digested material was filtered through a metal mesh sieve with a 1-mm pore size. The filtrate was transferred to a 1-l measuring cylinder and allowed to stand for 30 min. The supernatant was aspirated out, and the sediment was transferred to a Petri dish to examine *Paragonimus* metacercariae under a stereoscopic microscope. Isolated metacercariae were placed on glass slides and pressed lightly under a coverslip for morphological observation and measurement using an optical microscope with a digital camera DP71 and imaging software cellSens (standard version 1.12; Olympus). Morphometric differences between the host groups, Sawagani and Mokuzugani, were compared using the Mann-Whitney *U* test, and *P*-values less than 0.05 were considered statistically significant. Statistical analyses were performed using R (version 4.0.3 for Windows).

**Fig. 1.** Map of Japan (A) and Kagoshima Prefecture (B), showing the location where the present study was carried out. Map of Akune City (C) showing 15 locations where crabs were collected in the present study (●, Locations where both Sawagani and Mokuzugani were collected; ○, Locations where only Sawagani were collected). Numbers of the locations correspond to those shown in Tables 1 and 2.
DNA isolation, PCR-restriction fragment length polymorphism (RFLP) analysis, and sequencing

Molecular typing of the isolated metacercariae was performed according to a previous report [25]. Briefly, DNA samples were extracted from individual metacercariae using the QIAamp DNA Micro Kit (Qiagen, Valencia, CA, USA) in accordance with the manufacturer’s instructions. The internal transcribed spacer 2 (ITS2) region of the nuclear ribosomal RNA (rRNA) gene and the mitochondrial 16S rRNA gene region were then amplified via PCR using primer pairs 3S (forward: 5′-GGTACCGGTGGATCACTCGGCTCGTG-3′) and A28 (reverse: 5′-GGGATCCTGGTTAGTTTCTTTTCTCCGC-3′) and T7-1 (forward: 5′-ATTTCATCATGGGGCCGTC-3′) with SP6-1 (reverse: 5′-GATCCTTTACTAGTGAAAAGC-3′), respectively. PCR amplification was conducted at a final volume of 50 µl containing 2.5 µl of template DNA, 2.5 U of Phusion High-Fidelity DNA Polymerase (Thermo Fisher Scientific, Waltham, MA, USA), 0.25 µM of each primer, and the manufacturer’s HF reaction buffer. PCR amplification was performed using a thermal cycler (Takara PCR Thermal Cycler Dice Gradient; Takara Bio, Kusatsu, Japan), with 30 cycles at 98°C for 10 sec, 55°C for 10 sec, and 72°C for 15 sec. Initial denaturation was carried out at 98°C for 30 sec, and final extension was carried out at 72°C for 7 min. For RFLP analyses, we selected the restriction enzymes SnaBI and BsrDI. The former enzyme cleaved the ITS2 PCR products from *P. westermani* and the latter cleaved the PCR products of the mitochondrial 16S rRNA gene region from the *P. westermani* triploid form [25]. The digested samples were electrophoresed on agarose gel (3%, w/v) and visualized using ethidium bromide to examine the cleavage pattern.

Undigested amplicons were sequenced using the corresponding primers to verify identification via RFLP analyses. The dye terminator method was performed using fluorescently-labeled di-deoxynucleotide triphosphates with the BigDye Terminator v 3.1 Cycle Sequencing Kit (Thermo Fisher Scientific) in accordance with the manufacturer’s instructions. Nucleotide sequences were determined using a DNA sequencer (3730xl DNA Analyzer, Thermo Fisher Scientific), and their alignments were analyzed using GENETYX-Win, ver. 13 (GENETYX Co., Tokyo, Japan).

RESULTS

Prevalence and intensity of *Paragonimus* metacercariae in crab hosts

We collected Sawagani from 15 rivers in Akune City, whereas Mokuzugani were collected from 7 of the 15 rivers (Fig. 1). Both Sawagani and Mokuzugani tested positive for *Paragonimus* metacercariae from the Fujigadan, Yumikino, and Nomoto rivers (Tables 1 and 2). From the Kometsugi River and Yamashita River, we collected both Sawagani and Mokuzugani; however, only Mokuzugani tested positive for *Paragonimus* metacercariae. From Katahara River, we collected only Sawagani, which tested positive for *Paragonimus* metacercariae.

Among 941 Sawagani, 15 (1.6%) tested positive for *Paragonimus* metacercariae, from which 24 metacercariae were detected (Table 1). The infection prevalence was the highest at 10.1% in Nomoto River (12 out of 119 Sawagani were positive for metacercariae); the lowest prevalence of 0.8% was observed in Sawagani from the Fujigadan River (1 out of 118 Sawagani was positive). The burden (total number of metacercariae detected/number of Sawagani infected with *Paragonimus* metacercariae) was between 1 and 2 with a mean intensity of 1.6.

Of a total of 95 Mokuzugani examined, 21 (22.1%) tested positive for *Paragonimus* metacercariae, which tested positive for *Paragonimus* metacercariae (Table 1).

| Localities | No. of Sawagani | % infected | Total no. of mc | Intensity |
|------------|-----------------|------------|-----------------|-----------|
| 1. Kometsugi | 84               | 0          |                 |           |
| 2. Souate   | 32               | 0          |                 |           |
| 3. Ikenodan | 27               | 0          |                 |           |
| 4. Takamatsu| 13               | 0          |                 |           |
| 5. Kinzan   | 44               | 0          |                 |           |
| 6. Yokote   | 34               | 0          |                 |           |
| 7. Fujigadan| 118              | 1          | 0.8             | 1         |
| 8. Chaengadan| 68              | 0          |                 |           |
| 9. Jiijiro  | 65               | 0          |                 |           |
| 10. Sashiki | 45               | 0          |                 |           |
| 11. Yamashita| 103             | 0          |                 |           |
| 12. Yumikino| 112              | 1          | 0.9             | 1         |
| 13. Takashu | 29               | 0          |                 |           |
| 14. Nomoto  | 119              | 12         | 10.1            | 21        |
| 15. Katahara| 48               | 1          | 2.1             | 1         |
| Total      | 941              | 15         | 1.6             | 24        |

1) Numbers of the localities correspond to those shown in Figs. 1 and 4. 2) mc: metacercariae that were identified as the triploid form of *Paragonimus westermani*. 

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metacercariae were detected (Table 2). The metacercariae-positive Mokuzugani were obtained from 5 of the 7 rivers. The highest prevalence was recorded in Nomoto River (100%; only 1 Mokuzugani was captured, and it was positive), and the lowest prevalence was observed in Kometsugi river (3.6%; 1 out of 28 Mokuzugani was positive). The burden was between 1 and 38 with a mean intensity of 4.0.

**Morphology of the metacercariae detected in freshwater crabs**

The metacercariae isolated from the 2 species of freshwater crabs were almost spherical (Fig. 2). A large I-shaped excretory bladder filled with excretory granules was observed in the center of their bodies and a convoluted intestine surrounded both sides of the excretory bladder. A stylet was observed along the dorsal edge of the oral sucker. The diameter of the inner cyst wall of metacercariae from Sawagani (n=6) was 393 × 398 to 499 × 499 µm (mean: 435 × 440 µm) and the thickness of the inner cyst wall was 12 to 20 µm (mean: 16 µm; Fig. 2). The diameter of the inner cyst wall of metacercariae from Mokuzugani (n=50) was from 378 × 382 to 502 × 522 µm (mean: 412 × 414 µm) and the thickness of the inner cyst wall was from 12 to 22 µm (mean: 16 µm; Fig. 2). The differences in the mean size of metacercariae from the 2 crab species were not statistically significant (all \( P > 0.05 \)). Based on these key morphological features and the morphometric data, the metacercariae detected in the 2 species of freshwater crabs were presumed to be *P. westermani*. The only difference between metacercariae from the crab species was the presence or absence of red granules in the metacercarial body. Granules were observed in almost all metacercariae from Mokuzugani; however, red granules were never detected in those from Sawagani (Fig. 2).

**Molecular identification/typing of the metacercariae**

Species and form were identified using RFLP analysis of the PCR products amplified from metacercariae isolated from either Sawagani or Mokuzugani. ITS2 products of approximately 520 bp were generated through PCR from all metacercarial DNA samples irrespective of host crab species (Fig. 3; lane 1). The amplicons were cleaved into 2 fragments (expected sizes for *P. westermani* of approximately 420 and 100 bp; Fig. 3; lane 2) through SnaBI digestion. The PCR products for the mitochondrial 16S rRNA gene region of approximately 840 bp were also amplified from all metacercarial DNA samples irrespective of host crab species (Fig. 3; lane

### Table 2. Prevalence and number of *Paragonimus* metacercariae in Mokuzugani from Akune City, Kagoshima Prefecture

| Localities1) | No. of Mokuzugani | % infected | Total no. of mc2) detected | Intensity |
|-------------|-------------------|------------|---------------------------|-----------|
|             | Examined | Infected |               | Range | Average |
| 1. Kometsugi | 28       | 1        | 3.6 | 2 | 2 | 2.0 |
| 3. Ikenodan | 15       | 0        |          |       |     |     |
| 6. Yokote   | 1        | 0        |          |       |     |     |
| 7. Fujigadan| 12       | 3        | 25 | 5 | 1–3 | 1.7 |
| 11. Yamashita| 28      | 15       | 53.6 | 38 | 1–14 | 2.5 |
| 12. Yumikino| 10       | 1        | 10 | 1 | 1 | 1.0 |
| 14. Nomoto  | 1        | 1        | 100 | 38 | 38 | 38 |
| Total       | 95       | 21       | 22.1 | 84 | 1–38 | 4.0 |

1) Numbers of the localities correspond to those shown in Figs. 1 and 4. 2) mc: metacercariae that were identified as the triploid form of *Paragonimus westermani*.

Fig. 2. *Paragonimus* metacercariae isolated from Sawagani (a) and Mokuzugani (b) captured in Akune City, Kagoshima Prefecture. Numerous minute red granules were evenly distributed throughout the bodies of the metacercariae isolated from Mokuzugani.
Restriction fragment length polymorphism (RFLP) the major cause of human paragonimiasis in this region.

either Sawagani or Mokuzugani, or both, as the second intermediate hosts, with the triploid form of Paragonimus cases were also reported following venison consumption in accordance with the dietary history revealed by patients processed in a facility in Akune City [26]. Moreover, Sawagani exoskeleton has been reported in the stomach contents of sika deer.

metacercariae were detected in only Sawagani in 1 river. It is plausible that the life cycle of triploid with a few exceptional locations in Kagoshima Prefecture, where the triploid form of metacercariae was detected in Sawagani [5], detected [20]. Miyazaki [12] reported that the triploid form of Paragonimus was used to estimate the size of fragments (lane 5). Based on PCR-RFLP patterns, we identified metacercariae as the triploid form of Paragonimus westermani. Metacercariae from Mokuzugani exhibited the identical cleavage pattern as those from Sawagani.

We confirmed no variation in the RFLP patterns for any of the PCR amplicons. Furthermore, we sequenced the undigested amplicons using the corresponding primers and revealed that the amplified products were of 463 bp and 805 bp for the ITS2 region and 16S rRNA gene region, respectively; primer sequences were not included. The sequences obtained in this study were identical to previously deposited sequences for the P. westermani triploid form (DDBJ/EMBL/GenBank accession numbers: LC577864 for ITS2 region and LC577863 for 16S rRNA gene region) from Japan. Therefore, the nucleotide sequences obtained in this study have been deposited in the DDBJ/EMBL/GenBank database as the metacercarial stage of the triploid form of P. westermani. The ITS2 region and 16S rRNA gene region sequences from Sawagani isolates were deposited under accession numbers LC578473 and LC578471, whereas those from Mokuzugani isolates were deposited under accession numbers LC578472 and LC578315.

DISCUSSION

The lung fluke, P. westermani, is a medically and veterinarily crucial foodborne helminth. This lung fluke species is widely distributed in East, Southeast, and South Asian countries, and comprises a species complex with extremely high genetic diversity in the ITS2 region of rRNA gene, in particular [23, 31]. However, P. westermani, or its species complex, can be readily identified through patterns of lobulation of ovary and the arrangement of cuticular spines on adult flukes. Recently, more localities with freshwater crabs positive for P. westermani metacercariae have been discovered, e.g., in Thailand [24] and Vietnam [3]; the morphological/molecular features of the isolated metacercariae were highly variable. These findings imply that existing molecular markers may not be versatile for identifying Paragonimus metacercariae as P. westermani. Under such circumstances, experimental infection studies are required to inoculate metacercariae in plausible definitive hosts, such as dogs or cats, so as to obtain gravid adults, which can then be used to identify the species as P. westermani or its species complex by anatomical characterization with solid taxonomic evidence [12]. In Japan, however, comparative studies have established associations between adult morphology and genetic characteristics observed at various life cycle stages among the lung flukes of the genus Paragonimus at the species/subspecies and form levels [16, 25]. Therefore, we identified the isolated metacercariae from both Sawagani and Mokuzugani collected from rivers in the central and southern parts of Akune City, Kagoshima Prefecture, as the triploid form of P. westermani based on their morphological and molecular characteristics. Red granules observed in the bodies of the metacercariae isolated from Mokuzugani. These granules probably developed during the growth process of larvae in the second intermediate crab host, Mokuzugani [12].

Akune City is in the Hokusatsu region in the northern part of the Satsuma Peninsula, along with 5 other neighboring municipalities, namely, Isa City, Izumi City, Satsumasendai City, Satsuma Town, and Nagashima Town. This region has long been recognized as endemic for human paragonimiasis [8, 19]. Intensive field surveys have been performed to clarify the infectivity and prevalence of P. westermani metacercariae in second intermediate crab hosts in the region. Regarding Mokuzugani, surveys were previously conducted in Akune City [8], Sendai City (currently Satsumasendai City, the city/town name was changed following a municipal merger) [18], and Izumi City [19]. These field surveys all detected metacercariae of P. westermani. Sawagani were also previously collected in Kedouin Town (currently Satsumasendai City), in which the triploid P. westermani metacercariae were detected [20]. Miyazaki [12] reported that the triploid form of P. westermani metacercariae was primarily detected in Mokuzugani with a few exceptional locations in Kagoshima Prefecture, where the triploid form of metacercariae was detected in Sawagani [5, 20]. In the current study, we detected metacercariae of triploid P. westermani in both Sawagani and Mokuzugani in 3 rivers, and metacercariae were detected in only Sawagani in 1 river. It is plausible that the life cycle of triploid P. westermani is maintained in either Sawagani or Mokuzugani, or both, as the second intermediate hosts, with the triploid form of P. westermani accounting for the major cause of human paragonimiasis in this region.

Wild boars harbor P. westermani larvae in their muscles and play an important role as a source of infection in humans [13, 15]. Paragonimiasis cases were also reported following venison consumption in accordance with the dietary history revealed by patients [13, 17, 32, 33]. The triploid form P. westermani larvae have been previously detected in the muscles of sika deer hunted and processed in a facility in Akune City [26]. Moreover, Sawagani exoskeleton has been reported in the stomach contents of sika deer in central Japan [9]. Therefore, it is reasonable to consider sika deer as a paratenic host of P. westermani. Indeed, the importance of

Fig. 3. Restriction fragment length polymorphism (RFLP) patterns of PCR products amplified from the DNA of Paragonimus metacercariae from Sawagani. The internal transcribed spacer 2 region of nuclear ribosomal DNA was amplified using PCR (lane 1) and treated with the restriction enzyme SnaBI (lane 2). The mitochondrial 16S ribosomal RNA gene region was amplified (lane 3) and treated with the restriction enzyme BsrDI (lane 4). A 100-bp DNA ladder marker was used to estimate the size of fragments (lane 5). Based on PCR-RFLP patterns, we identified metacercariae as the triploid form of Paragonimus westermani. Metacercariae from Mokuzugani exhibited the identical cleavage pattern as those from Sawagani.

3). Amplicons were cleaved into 2 fragments through BsrDI digestion (expected sizes for the P. westermani triploid form of approximately 560 and 280 bp; Fig. 3, lane 4). The digestion pattern was consistent with that of the triploid form of P. westermani. Accordingly, we used 5 metacercariae from Sawagani and 7 from Mokuzugani collected from all rivers where positive crabs were detected. We confirmed no variation in the RFLP patterns for any of the PCR amplicons.
Fig. 4. The 2 km × 3 km mesh map of the central and southern parts of Akune City showing the locations where Paragonimus westermani-positive Sawagani were collected; ■, Locations where triploid P. westermani-positive Mokuzugani were collected. Numbers of the locations correspond to those shown in Tables 1 and 2. The original mesh map encompassing Japan was drawn by Dainihon Ryoyukai, the Japan hunters’ association, and we were approved to trace the map of Akune City for this manuscript. Red-bordered squares on this map designate areas where P. westermani-positive wild boars and sika deer were hunted [26, 27].

In Japan, the former Cabinet implemented the “Growth Strategy 2017”, which proposed the effective use of meat from wild birds and beasts as game meat to support local economies [2]. The Ministry of Agriculture, Forestry and Fisheries supports the Cabinet policy and has selected several areas as Japan’s pioneer models to facilitate the provision of safer and higher-quality game meat [10]. The Ministry of Health, Labour and Welfare has established guidelines for food safety management throughout the entire process from hunting to the consumption of game meat, to prevent food poisoning [11]. In brief, the initiative to secure safe game meat is gaining momentum and has currently received increasing attention in Japan. Traditionally, meat from wild birds and beasts has been consumed primarily by hunters and their families. However, more recently, opportunities for the consumption and distribution of game meat over a wider area have increased through promoting campaigns and online sales [6, 10, 28]. To prevent paragonimiasis outbreaks on the national level, governmental agencies for public health should accelerate the spread of information describing the need to cook wild boar and sika deer meat sufficiently before consumption, e.g., at 75°C for 1 min in the innermost region [11].

In conclusion, we identified the metacecarias as the triploid form of P. westermani that were isolated from Sawagani and Mokuzugani collected in Akune City, Kagoshima Prefecture. The triploid form of P. westermani larvae have been detected in the muscles of wild boars and sika deer hunted and processed in this city [26, 27]. These paratenic host mammals might become infected with triploid P. westermani through consumption of infected Sawagani or Mokuzugani in Akune City or the surrounding area. Studies are, therefore, required to assess the contamination status of Sawagani and Mokuzugani in Japan, as crab infection potentially leads to subsequent P. westermani infection of wild mammals, such as wild boars and sika deer. Furthermore, security safeguards should be constituted against P. westermani infections through the persistent health education campaigns for people who consume uncooked game meats or freshwater crabs.

POTENTIAL CONFLICT OF INTEREST. The authors have nothing to disclose.

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