Spontaneous Larval *Gnathostoma nipponicum* Infection in Frogs

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**ABSTRACT.** From June 1993 to September 1997, a survey was carried out for the prevalence of larval *Gnathostoma nipponicum* infection in several kinds of frogs, toads, and their tadpoles collected from an endemic area of this nematode in Aomori Prefecture. Two frog species, *Rana nigromaculata* and *R. catesbeiana*, were infected, and a total of 446 advanced third-stage larvae (AdL3) of *G. nipponicum* were recovered. These results confirmed that two frog species which can serve as the second intermediate and/or paratenic hosts in the life cycle of *G. nipponicum* exist in nature. This report is the first record of spontaneous infection of frogs with AdL3 of *G. nipponicum*. — KEY WORDS: advanced third-stage larva, *Gnathostoma nipponicum*, intermediate host.


In Japan, *Gnathostoma nipponicum*, which is a common parasite found in weasels [14, 17], is a zoonotic parasite [3, 16]. Although the life cycle of this nematode is not completely cleared, it has been confirmed experimentally that two intermediate hosts are required, i.e., some species of copepods can serve as the first intermediate host and various species of animals including frogs can serve as the second intermediate and/or paratenic hosts [2, 7, 9]. In the past, spontaneous infection of advanced third-stage larvae (AdL3) of this nematode have been recorded from freshwater fish (five species), reptiles (two species) and small mammals (two species), and these animals were regarded as the second intermediate and/or paratenic hosts in nature [3, 7, 10–15]. Up to now, no amphibians including frog species naturally infected with AdL3 of *G. nipponicum* have been found. However, some frogs of the genus *Rana* naturally infected with larvae have been recorded from *G. spinigerum* [9], i.e., *R. nigromaculata* and *R. catesbeiana*, and from *G. doloresi* [4, 5], i.e., *R. (Babina) subaspera*, *R. narina*, *R. ishikawae*, and *R. nanyei*. In addition, experimental studies had revealed that several species of frogs and toads had larval infection of *G. spinigerum* [6], i.e., *R. nigromaculata*, *R. catesbeiana*, *R. rugosa*, *Hyla arvorea japonica*, and *Bufo vulgaris formosus*, and of *G. hispidum* [1, 8], i.e., *R. nigromaculata* and *R. limnocharis*. Based on these reports and the results of our previous surveys, we hypothesised that weasels, the final host, might also be infected with *G. nipponicum* by preying on amphibians, such as frogs.

In the present study, we carried out a survey for the prevalence of larval *G. nipponicum* infection in frogs, toads, and their tadpoles collected from eastern Aomori Prefecture [10–15]. From June 1993 to September 1997, a total of 7,169 frogs, toads, and their tadpoles including the following six species (Table 1), *Hyla japonica*, *Rana ornativentris*, *R. nigromaculata*, *R. rugosa*, *Rana catesbeiana*, and *Bufo japonicus formosus* were examined.
R. catesbeiana (common name: bullfrog). Taxonomically, all larvae obtained from the positive two frog species were larvae from species, infection of gnathostome larvae were found in two frog postinoculation. larvae to mammals. All animals were examined at 30 days obtained from were inoculated orally with larvae (10 or 20 per animal) were fixed with 10% hot formalin, cleared and mounted with lactophenol for laboratory mammals) were readily fixed with 10% hot hypodermis, head, visceral organs, and three portions of musculature (frontal legs, posterior legs, and thoraco-abdominal wall) to determine the location of the larvae. In addition, muscle tissues of legs and thoraco-abdominal walls in three R. catesbeiana were examined after pressing between two thick glass plates before digestion. The recovered larvae (except for the experimental infection to laboratory mammals) were readily fixed with 10% hot formalin, cleared and mounted with lactophenol for morphological observation. Three rats (Wistar: male), four mice (ddY: male), and three golden hamsters (Syrian: male) were inoculated orally with larvae (10 or 20 per animal) obtained from R. catesbeiana, to find the infectivity of the larvae to mammals. All animals were examined at 30 days postinoculation.

In the field survey, as shown in Table 1, spontaneous infection of gnathostome larvae were found in two frog species, R. nigromaculata (common name: pond frog) and R. catesbeiana (common name: bullfrog). Taxonomically, all larvae obtained from the positive two frog species were identified as AdL3 of G. nipponicum. The bodies of the larvae from R. nigromaculata and R. catesbeiana were 1.10 × 0.11 mm and 0.72–2.03 × 0.08–0.27 (mean: 1.45 × 0.16) mm, respectively. The mean body sizes of the larvae from R. catesbeiana were larger than those of larvae (1.16 × 0.14 mm) from naturally infected loaches (Misgurnus anguillicaudatus) in the same areas [13]. They had three rows of hooklets on the head-bulb, and the mean number of hooklets of each row from 1st to 3rd was 31, 34, and 39. Other morphological features of the present larvae coincided well with the descriptions of AdL3 of G. nipponicum from loaches reported by previous investigators [3, 9, 13]. Before our present study, spontaneous infection of larval G. nipponicum had not been reported from frog species. Thus, this is the first record of naturally infected R. nigromaculata and R. catesbeiana with this larvae.

The infection rate was 0.2% (1/436) in R. nigromaculata and was 34.7% (51/147) in R. catesbeiana. The prevalence of larvae in R. catesbeiana was higher. The number of larvae per infected R. catesbeiana were 1 to 95 (mean: 8.7), and a total of 445 larvae were recovered (Table 1). We reported previously the infection rates of larval G. nipponicum in several kinds of freshwater fish and small wild mammals collected from the same areas in Aomori Prefecture, i.e., 0.18% in Tribolodon hakonensis, 1.0% in Misgurnus anguillicaudatus, 3.4% in Chaenogobius urotaenia, 2.75% in Oncorhynchus masou, 46.7% in Sillurus asotus, 27.3% in Rattus norvegicus, and 72.5% in Chimarrrogale himalayica [10–13, 15]. In comparison with those data, it seemed that the prevalence in R. catesbeiana in this study was relatively higher level.

The relationship between the number of larvae and body length of R. catesbeiana is summarized in Table 2. Infected frogs were found to be longer than 8.0 cm, and as the body length increased, so did the infection rate and mean number of larvae per frog. The prevalence of frogs larger than 14.1 cm in body length showed extremely higher levels, which is attributed to the feeding habits of the frog. In general, the young-adult and adult stages of R. catesbeiana are essentially carnivorous and they prey chiefly on insect, fish, amphibians, reptiles, and small mammals such as voles, although the neonatal and juvenile stage (tadpole) are usually herbivorous. In fact, by examination of the stomach in this study, various food such as insects, insect larvae, and their body components were found commonly in 56 frogs. Additionally, ingested frogs (22 cases), small fishes (5 cases), loaches (3 cases) (Fig. 1), crawfish (3 cases), voles (3 cases), and a small sized snake (1 case) were also seen in frogs longer than 14.1 cm in body length. Therefore, we speculated that the most important direct sources to R. catesbeiana might be freshwater fish, such as loaches (Misgurnus anguillicaudatus) and gobiid fish (Chaenogobius urotaenia), based on the results of our previous surveys in the same localities. From this viewpoint, it was considered that R. catesbeiana was one of the paratenic hosts infected secondarily with AdL3 by eating the second intermediate hosts. In addition, it also suggested that R. catesbeiana has characteristics suitable to be the host to AdL3 of this nematode. While only one positive

### Table 2. Relationship between the body length of bullfrog (Rana catesbeiana) and prevalence of Gnathostoma nipponicum larvae

<table>
<thead>
<tr>
<th>Range of frog body length (cm)</th>
<th>No. frogs examined</th>
<th>No. frogs infected</th>
<th>Prevalence (%)</th>
<th>No. larvae recovered</th>
<th>No. larvae per infected frog (mean)</th>
</tr>
</thead>
<tbody>
<tr>
<td>⁣</td>
<td>8.0</td>
<td>20</td>
<td>1</td>
<td>5.0</td>
<td>1</td>
</tr>
<tr>
<td>⁣</td>
<td>8.1–10.0</td>
<td>18</td>
<td>2</td>
<td>11.1</td>
<td>2</td>
</tr>
<tr>
<td>⁣</td>
<td>10.1–12.0</td>
<td>38</td>
<td>7</td>
<td>18.4</td>
<td>19</td>
</tr>
<tr>
<td>⁣</td>
<td>12.1–14.0</td>
<td>28</td>
<td>8</td>
<td>28.6</td>
<td>39</td>
</tr>
<tr>
<td>⁣</td>
<td>14.1</td>
<td>43</td>
<td>33</td>
<td>76.7</td>
<td>384</td>
</tr>
<tr>
<td>Total</td>
<td>147</td>
<td>51</td>
<td>34.7</td>
<td>445</td>
<td>1–95 (8.7)</td>
</tr>
</tbody>
</table>
case of *R. nigromaculata* was small, measuring 3.0 cm in body length, the direct source was unclear whether the first or the second intermediate host. However, it seemed that this frog was infected accidentally with larva by eating the first intermediate host such as copepods than the second intermediate host, owing to recovered larva size, body size of infected frog, and feeding habits of this frog species.

In the examinations to determine the location of the larvae in *R. catesbeiana*, the vast majority of the larvae, 386 of 445 (86.7%) were recovered from the musculature. By pressing examination of the three positive *R. catesbeiana*, four encapsulated larvae were clearly found within the musculature (Fig. 2).

The other three species of frogs, one of toad, and tadpoles in this study were all negative. It is unknown whether such negative data result from their non-susceptibility to this larval infection or their feeding habits. Further detailed studies are necessary to clarify the potential characteristics of these frog species and tadpoles to larval *G. nipponicum* infection.

The experimental study revealed that three species of small laboratory mammals were susceptible to AdL3 of *G. nipponicum* from *R. catesbeiana*. All 10 animals inoculated with AdL3 became positive, and recovery rates of the larvae were 57.5% in mouse, 35.0% in rat, and 20.0% in golden hamster at 30 days postinoculation. Most of the larvae (47/50: 94.0%) were recovered from the body muscles. The morphological features of the larvae was similar to those of AdL3 before inoculation. However, body size of the larvae from three mammal species showed slightly development, and they measured 1.60–1.66 x 0.16–0.18 mm. From these results, it seemed that the AdL3 in *R. catesbeiana* were able to infect weasels and also to develop into the adult worms in their final host.

The present study confirmed that two frog species, *R. nigromaculata* and *R. catesbeiana*, infected with *G. nipponicum* larvae exist in nature. In addition, it also suggests that *R. catesbeiana* may be important sources of infection to the weasels in conjunction with their feeding habits, and they may serve as the plausible second intermediate and/or paratenic hosts in the natural life cycle of this nematode.

REFERENCES