New insights on the *Taenia solium* tapeworm using molecular tools: age-based human definitive host prevalence and deliberation on parasite life span

Tiaoying Li\(^ a \), Xingwang Chen\(^ b \), Christine M. Budke\(^ b \), Yuangui Zhou\(^ b \), Mianchuan Duan\(^ b \), Celine Wang\(^ b \), Bo Zhong\(^ b \), Yang Liu\(^ b \), Jianying Luo\(^ b \), Wei He\(^ b \), Jingye Shang\(^ b \) and Akira Ito\(^ c \)

\(^a\)Institute of Parasitic Diseases, Sichuan Center for Disease Control and Prevention, Chengdu, China; \(^b\)Department of Veterinary Integrative Biosciences, College of Veterinary Medicine & Biomedical Sciences, Texas A&M University, College Station, Texas, USA; \(^c\)Department of Endemic Diseases, Mull County Center for Disease Control and Prevention, Mul, China; \(^d\)Williamsville School District, Buffalo, NY, USA; \(^e\)Department of Parasitology, Asahikawa Medical University, Asahikawa, Japan

**ABSTRACT**

Information on age-based *Taenia solium* taeniasis prevalence is crucial for control of cysticercosis. *T. solium* taeniasis prevalence was determined for a village in Liangshan Prefecture, Sichuan Province, China that was co-endemic for *T. solium*, *Taenia saginata asiatica*, and *Taenia saginata*. Individuals who were *Taenia* egg-positive by stool microscopy and/or expelled tapeworms or proglottids post-treatment were diagnosed as having taeniasis. Infecting species was identified via multiplex PCR on tapeworm specimens or coproPCR followed by sequencing. In addition, initial stool samples from 10 children with taeniasis suspected of having spontaneous expulsion of tapeworms within the period between diagnosis and treatment were subject to species confirmation via coproPCR and sequencing. Of the 389 study subjects, 194 (49.9%) were diagnosed with taeniasis. Children (< 16 years of age) had a higher *T. solium* taeniasis prevalence (8.8%) than older individuals (2.5%) (*P* = 0.0127). Molecular analysis of initial stool samples from 7 of 10 children suspected of spontaneously passing tapeworms indicated 6 infections due to *T. solium* and 1 infection due to *T. saginata*. This study found that young children had a higher *T. solium* taeniasis prevalence than older individuals, providing additional support for the belief that adult *T. solium* has a relatively short lifespan compared to other *Taenia* species with human definitive hosts.

**KEYWORDS**

*Taenia solium*; taeniasis; prevalence; lifespan; polymerase chain reaction; sequence

1.  Introduction

Humans act as definitive hosts for the *Taenia solium* tapeworm after consuming undercooked pork containing the larval form of the parasite (cysticerci) [1]. In addition, humans can act as aberrant intermediate hosts, resulting in cysticercosis and neurocysticercosis. Cysticercosis remains a neglected disease of public health importance in numerous countries in Latin America, sub-Saharan Africa, and Asia [2,3]. Cases of cysticercosis are also found in non-endemic areas due to global travel [4]. In 1992, the International Task Force for Disease Eradication declared that human infection with the adult (taeniasis) and larval (cysticercosis) forms of this tapeworm are potentially eradicable [5].

Understanding how human host age impacts *T. solium* taeniasis status is crucial for the control of human cysticercosis. Several studies have been undertaken on the epidemiology of taeniasis at the community level [6–16]. However, it remains unclear how host age affects infection status. Our recent study revealed that children 6–15 years of age from an ethnic minority community in Liangshan Prefecture, Sichuan Province, China had a high (6.7%) prevalence of infection with adult *T. solium* tapeworms [17]. However, additional studies are needed to evaluate how infection frequency changes across the human life span.

In addition to host age, information on parasite life span is important for implementation of effective control practices. In Liangshan Prefecture, we found that some individuals appeared to have spontaneously expelled their tapeworms during the time between case identification via microscopy performed on stool samples and treatment to eliminate the adult worms [17]. For one study site, where treatment was delayed 8 months due to the remote location, 10 of the 23 individuals with *Taenia* eggs initially seen on microscopy did not expel proglottids or tapeworms upon treatment. The remainder discharged worms belonging to *T. solium* (11 cases) and *Taenia saginata* (2 cases). Due to the high efficacy of the purgative agent used [17,18], it is believed that most adult tapeworms present would have been expelled at the time of treatment. In contrast, at another study site where treatment was delivered 3 months after microscopic examination for eggs, all *Taenia* egg-positive children (8 cases) released tapeworms (all were later confirmed as *T. solium*). Based on these observations, it was
hypothesized that some, if not all, of the worms likely expelled within the 8 months between diagnosis and treatment of the children in Liangshan Prefecture were *T. solium*. If this was the case, it would provide insight into the adult *T. solium* lifespan within the human host [9,17,19,20].

Several studies have hypothesized that adult *T. solium* tapeworms have a shorter life span than *T. saginata* and *Taenia saginata asiatica* (an intraspecies variant of *T. saginata*) tapeworms and are naturally expelled within several years [6,9,21–24]. In the present study, we further assessed the initial stool samples from the 10 children who were suspected to have spontaneously expelled their tapeworms over an 8-month period in order to evaluate the hypothesis that the expelled tapeworms were *T. solium*. In addition, human age-based taeniasis prevalence of the three locally recognized human *Taenia* (*T. solium*, *T. saginata*, and *T. saginata asiatica*) was evaluated for samples collected in a village located in Liangshan Prefecture.

### 2. Materials and methods

#### 2.1. Study site and population

This study was conducted in Shuiluo Township located in Muli County of Liangshan Prefecture, Sichuan Province, China (Figure 1). The township has a population of approximately 5,000 and a population density of about 4 persons per km², with inhabitants belonging to the Tibetan, Naxi, and Mongolian ethnic groups. Previous studies have shown that residents of this township are infected with *T. solium*, *T. saginata*, and *T. saginata asiatica* [17,25]. Most township inhabitants own pigs that readily forage in the surrounding environment and have easy access to human feces due to the practice of open defecation. Cattle and yak ownership is more limited, with yaks raised away from the village. Undercooked pork/pork liver and raw yak meat are popular dishes consumed by both adults and children during the slaughtering season (November to January).

#### 2.1.1 Village prevalence study

The study was carried out from December 2017 to May 2018. In cooperation with the Muli County Center for Disease Control and Prevention (CDC) and the Shuiluo Township medical clinic, residents of the village of Guni (population 695) that were 5–75 years of age were invited to participate after providing informed consent or assent. Ethical approval for this study was granted by the ethics committee of the Sichuan CDC.

#### 2.1.2 School-based study

The 23 children (age 6–15 years) positive for *Taenia* eggs who attended Shuiluo Primary School in 2016 [17] were also evaluated as part of this study.

#### 2.2. Data and sample collection

##### 2.2.1. Village prevalence study

Study staff administered a questionnaire to each participant or to a participant’s parent (or with the assistance of a teacher) for children under 15 years of age to obtain basic demographic information and any history of expelling tapeworm proglottids within the previous year. A naturally passed fecal sample was requested from all study participants. Approximately 20 g of stool was collected in a 50 ml tube for microscopy to look for the presence of *Taenia* eggs, and for copro-polymerase chain reaction (coproPCR) if indicated. Direct smears were prepared from stool samples and examined for the presence of *Taenia* eggs.

Individuals positive for *Taenia* eggs and/or reporting segment expulsion were treated within several days of fecal sample collection with a combination of pumpkin seeds and areca nut extract, as previously reported, to expel any adult tapeworms [17,18,26]. Expelled tapeworms or proglottids were kept in 75%
ethanol for species confirmation by multiplex PCR (see below). For stool samples that were positive for *Taenia* eggs, but did not contain tapeworm segments, coproPCR followed by sequencing (see below) was employed to identify the infecting species. Individuals who were *Taenia* egg-positive by stool microscopy and/or expelled tapeworms or proglottids post-treatment were diagnosed as having taeniasis.

### 2.2.2. School-based study

Age and segment expulsion history, previously obtained from 23 children (age 6–15 years) positive for *Taenia* eggs who attended Shuiluo Primary School in Shuiluo Township [17], were reviewed. Initial stool samples from 10 of these 23 taeniasis cases who were suspected to have spontaneously expelled tapeworms prior to treatment 8 months later were also subject to species analysis by PCR and sequencing.

### 2.3. Multiplex PCR, coproPCR, and sequencing

The genomic DNA of parasite isolates was extracted using the DNeasy Blood & Tissue Kit (Qiagen) and subsequently used as a template for PCR. For differentiation of *Taenia* species, multiplex PCR was conducted as described previously [10,17,18,26,27], with a minor revision. Briefly, one reverse and three forward primers were applied to amplify 984, 827, and 269 bp amplicons, specific for mitochondrial (mt) gene cox1 sequences of *T. solium* Asian genotype, *T. saginata*, and *T. saginata asiatica*, respectively. The PCR cocktail contained mixed primers, 10 µl of GoTaq® Green Master Mix (Promega), and 1 µl of template in 20 µl of reaction mixture. PCR protocols were composed of 1 cycle of initial denaturation (4 min at 95°C), 35 cycles of denaturation (30 sec at 95°C), annealing (30 sec at 55°C), and extension (90 sec at 72°C), plus 1 cycle of final extension (5 min at 72°C).

The QIAamp Fast DNA Stool Mini Kit was used to extract DNA from fecal samples. Prior to using the kit, glass beads were added to the fecal samples. The samples were then agitated using a tissue breaker to release as much DNA as possible from the parasite eggs. Extracted DNA was used as a template for multiplex PCR to amplify DNA specific for the three *Taenia*, as described previously [10,18,26,27], with a minor amendment. Briefly, double amplifications were performed for coproPCR, using the first PCR product as a template for the second PCR. The same protocols as described for multiplex PCR were used, except that for coproPCR, two µl of template were added to 50 µl of reaction mixture, and annealing was performed at 52°C. Subsequently, the second PCR products were subject to sequencing for reconfirmation of species. All diagnostic testing was conducted at the Sichuan CDC (China).

### 2.4. Statistical analysis

Age was evaluated for normality and presented using means or medians depending on outcome. Infection frequencies by age group were expressed as proportions. Chi-square tests were used to evaluate differences in *T. solium* taeniasis, *T. saginata* taeniasis, *T. saginata asiatica* taeniasis, and overall taeniasis prevalence between children less than 16 years of age and individuals 16 years of age and older (considered adults for this study) from the village of Guni. Fifteen years of age was evaluated as the cutoff for children since this is the oldest age typically attending primary school. The Student’s t-test was employed to assess mean age of children attending Shuiluo Primary School who were positive for *T. solium* by coproPCR, with supposed natural elimination of their tapeworms within 8 months, versus the mean age of *Taenia* egg-positive children with expulsion of *T. solium* tapeworms following treatment 8 months after diagnosis via microscopy. Significance was set at $P < 0.05$.

### 3. Results

#### 3.1. Village prevalence study

A total of 389 subjects were enrolled from the village of Guni, of which 202 (51.9%) were male and 187 (48.1%) were female. Participant age ranged from 5 to 73 years (median of 31 years), with 276 adults (71.0%) and 113 children (29.0%). Of the 389 subjects investigated, 290 (74.6%) reported a history of passing tapeworm segments within the previous year (Figure 2).

#### 3.1.1 Results of stool microscopy

Fecal samples were available from 281 of the 389 enrolled individuals from the village of Guni. Overall, 67.0% ($n = 185$) of adults and 85.0% ($n = 96$) of children provided samples. Of the 281 samples, 94 (33.5%) were *Taenia* egg positive by microscopy, with 47.6% of samples from adults positive and 6.3% of samples from children positive.

#### 3.1.2 Tapeworms expelled post-treatment

A total of 293 residents of the village of Guni received treatment, of which 185 had originally provided a fecal sample. Of these 185 individuals, 84 were positive for *Taenia* eggs and had a history of segment expulsion, 92 were negative for eggs but reported a history of segment expulsion, and 9 were negative for both but had requested treatment. There were no instances where an individual who was positive for *Taenia* eggs did not have a history of segment expulsion. For the remaining 108 treated subjects who did not provide an initial fecal sample, 97 reported expelling proglottids in the past year and 11 did not have a history of segment expulsion but requested treatment (Figure 2). Tapeworms or proglottids were produced in 175
(59.7%) of 293 treated individuals (Figure 2), but parasite material was disposed of by one subject prior to collection.

### 3.1.3. Results of multiplex PCR and coproPCR

Multiplex PCR-determined infecting species for 174 individuals who expelled tapeworms or proglottids post-treatment are shown in Table 1. CoproPCR was performed on samples from 18 of 19 *Taenia* egg-positive cases without tapeworm specimens. Species-specific DNA was successfully amplified for *T. saginata asiatica* in 16 cases and for *T. saginata* in 2 cases. Subsequent sequencing of coproPCR products succeeded in all 18 cases, exhibiting identical results to the multiplex PCR. That is, 15 were 99% genetically similar and 1 was 97% similar to *T. saginata asiatica* [NCBI Accession number KJ187963.1], and the other 2 were 99% similar to a published genetic sequence for *T. saginata* [NCBI Accession number AB984351.1] (Table 1). Species differentiation was not performed for 2 cases, 1 due to lack of an available stool sample and the other due to inadvertent disposal of the parasitic material by the infected individual.

Combining the results of stool microscopy and tapeworm expulsion post-treatment, the overall taeniasis prevalence was 49.9% (194/389) in the village of Guni. Subjects aged 11–15 years were found to have the highest prevalence of *T. solium* taeniasis (6/53; 11.3%), while those in the 41–50 years age group had the highest prevalence of *T. saginata asiatica* (37/62; 59.7%) and those in the 31–40 years age group had the highest prevalence of *T. saginata* (13/71; 18.3%) (Figure 3). The overall prevalence of taeniasis increased with age, reaching a peak in the 21–30-year age group and then remaining stable (Figure 3). This finding was largely driven by *T. saginata asiatica* prevalence in the

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**Table 1. Species-specific distribution of tapeworms in children (n = 113) and adults (n = 276) in the village of Guni, Sichuan Province, China (2017).**

| Infecting species | Number of cases (%) in children | Number of cases (%) in adults |
|-------------------|---------------------------------|-------------------------------|
| *T. solium*       | 7 (6.19)                        | 4 (1.45)                     |
| (A) *T. Saginata asiatica* | 11 (9.73)                  | 132 (47.83)b                 |
| *T. saginata*     | 1 (0.88)                        | 16 (5.80)c                   |
| *T. solium + T. saginata* | 1 (0.88)                   | 2 (0.72)                     |
| *T. solium + T. saginata* | 2 (1.77)                   | 0 (0.00)                     |
| *T. saginata asiatica + T. saginata* | 1 (0.88)                | 14 (5.07)                    |
| *T. solium + T. saginata asiatica + T. saginata* | 0 (0.00)                   | 1 (0.36)                     |
| Species not identified | 1 (0.88)a                   | 1 (0.36)d                    |

a: stool sample not provided and the post-treatment expelled tapeworms not collected
b: 16 cases identified by coproPCR and sequencing
c: 2 cases identified by coproPCR and sequencing
d: positive for *Taenia* eggs, but no stool sample for coproPCR (subject not treated)
older age groups. Children were more likely to have *T. solium* taeniasis than adults (8.8% vs 2.5%) ($\chi^2 = 6.210, P = 0.0127$). In contrast, adults were more likely than children to have *T. saginata asiatica* taeniasis (54.0% vs 11.5%) ($\chi^2 = 57.802, P < 0.001$) or *T. saginata* taeniasis (11.2% vs 3.5%) ($\chi^2 = 4.892, P = 0.0270$). Similarly, the overall prevalence of taeniasis due to any *Taenia* species was significantly different between adults (61.6%) and children (21.2%) ($\chi^2 = 50.626, P < 0.001$).

### 3.2. School-based study

Of the 23 children positive for *Taenia* eggs who attended Shuiluo Primary School, 14 (60.9%) were male and 9 (39.1%) were female. Twenty-one (91.3%) were female.

### Table 2. Demographic information and infecting species for *Taenia* egg-positive children attending Shuiluo Primary School, Sichuan Province in 2016 without tapeworm expulsion (n = 10) and with tapeworm expulsion (n = 13) following treatment 8 months after diagnosis.

| Sex  | Age (years) | History of SE | Duration of SE | Frequency of SE | Infecting species                      | Number of worms expelled |
|------|-------------|---------------|----------------|-----------------|----------------------------------------|--------------------------|
| male | 6           | positive      | unclear        | occasionally    | *T. saginata asiatica*                 | 0                        |
| female | 7          | negative      | ND             | ND              | *T. solium*                            | 0                        |
| male | 11          | positive      | 3 years        | occasionally    | *T. solium*                            | 0                        |
| female | 11         | positive      | 3 years        | occasionally    | *T. solium*                            | 0                        |
| male | 13          | positive      | 1 year         | occasionally    | *T. solium*                            | 0                        |
| female | 13         | positive      | 1 year         | occasionally    | *T. solium*                            | 0                        |
| female | 14         | positive      | 2 years        | occasionally    | *T. solium*                            | 0                        |
| male | 14          | positive      | unclear        | occasionally    | *T. solium*                            | 0                        |
| male | 15          | positive      | 3 years        | occasionally    | *T. solium*                            | 0                        |
| male | 15          | negative      | ND             | ND              | *T. solium*                            | 0                        |
| male | 8           | positive      | 1 year         | occasionally    | *T. solium*                            | 1                        |
| female | 8          | positive      | 1 year         | occasionally    | *T. solium*                            | 1                        |
| male | 9           | positive      | 1 year         | occasionally    | *T. solium + T. saginata*              | 2                        |
| male | 9           | positive      | 1 year         | occasionally    | *T. solium*                            | 1                        |
| male | 11          | positive      | > 2 years      | frequently      | *T. solium*                            | 1                        |
| male | 11          | positive      | 2 years        | frequently      | *T. solium + T. saginata asiatica*     | 2                        |
| male | 11          | positive      | > 2 years      | frequently      | *T. solium*                            | 11                       |
| female | 12         | positive      | 2 years        | frequently      | *T. saginata*                          | 1                        |
| female | 12         | positive      | unclear        | occasionally    | *T. solium*                            | 1                        |
| female | 13         | positive      | 1 year         | frequently      | *T. saginata*                          | 1                        |
| male | 15          | positive      | 3 years        | frequently      | *T. solium*                            | 6                        |

*: identified by coproPCR and sequencing

**: identified by coproPCR only

SE: segment expulsion

ND: not determined
reported passing tapeworm segments within the past year (Table 2). Initial stool samples were available from 7 of the 10 children who were suspected of having passed tapeworms spontaneously. CoproPCR performed on these 7 samples indicated 6 infections due to *T. solium* and 1 infection due to *T. saginata* (Table 2). Six of the PCR products were successfully sequenced. Five of the sequences were over 99% identical to a published genetic sequence for *T. solium* (NCBI Accession number AB984354.1), while the sixth sequence was 99% identical to a published sequence for *T. saginata* (NCBI Accession number AB984351.1) (Table 2). Sequencing findings were consistent with coproPCR findings for all 6 cases. The mean age (13.0 years) of the six cases identified as having possible natural expulsion of *T. solium* tapeworms was significantly older than the mean age (10.3 years) of the 11 individuals who expelled *T. solium* tapeworms following treatment (P = 0.017).

### 4. Discussion

This is the first study to evaluate the relationship between host age and infection with adult *T. solium* tapeworms, with infecting species confirmed in all but two taeniasis carriers. Prior data about age-related tapeworm infections are scarce, primarily due to the low number of tapeworm carriers identified in individual studies [13–16]. The largest previous study, which was conducted in Guatemala, showed infections with *Taenia* spp. tapeworms increasing with age, and peaking in the 30–39 years age group in areas co-endemic for *T. solium* and *T. saginata* [6]. This pattern was similar to the current study’s findings for overall taeniasis prevalence. Unfortunately, in the Guatemala study, infecting species was not available for more than one-third of the individuals with taeniasis. However, it was shown that *T. solium* was the infecting species for more than half of the evaluated tapeworm carriers and that over half of the detected taeniasis cases in these Guatemalan communities were less than 20 years of age [6]. Another study conducted in Ecuador indicated that infection with *Taenia* spp. also increased with age [8]. However, the study was based on individuals who reported passing tapeworms after treatment rather than those shown to be infected parasitologically.

In two previous studies conducted in Peru and the Democratic Republic of the Congo, young children were found to be at greatest risk for tapeworm infections through detection of coproantigen [9,11]. In these two studies, information on infecting species was also not available. However, *T. solium* was known to be locally endemic through reported cases of human and porcine cysticercosis. Studies in other regions have shown that young children, such as a 4-year-old girl from Bali, can harbor *T. solium* tapeworms [28]. In the study village of Guni, adults and children likely had similar access to infected pork within the household, while the pork provided to children by the school was confirmed to have been purchased from areas in China considered non-endemic for *T. solium*. Although pork consumed at school was likely safe, it was very common for children to grill pork for their own consumption when a pig was slaughtered at home. Lack of sufficient oversight creates the opportunity for children to consume improperly cooked pork. The method by which children incompletely chew food has also been proposed as a risk for acquiring taeniasis [19,29].

Detection of concurrent infection with two or three tapeworm species is not uncommon in co-endemic areas [17,18], and co-infections were also found in the village of Guni. Therefore, there is no reason to think that the low number of *T. solium* infections in adults, in the current study, was due to existing *T. saginata asiatica* or *T. saginata* infections. The larger numbers of *T. saginata asiatica* and *T. saginata* infections in adults compared to children are likely due to adults having accumulated the parasites over a longer duration in combination with the long life span (up to decades) of these parasites. The relatively low prevalence of *T. saginata* in all age groups most likely reflects a diet that more commonly includes pork rather than yak meat or beef.

It has previously been speculated that *T. solium* has a shorter life span than *T. saginata* and *T. saginata asiatica*. Lightowers estimated the average longevity of *T. solium* adults to be less than 3 years by evaluating the rate of taeniasis reoccurrence in populations after mass treatment with cestocidal drugs [20]. [9,11], both inferred that intestinal infection with *T. solium* is short lived since the highest infection prevalence was found in the 5–10 years age group in both Peru and Congo and estimated that the lifespan of *T. solium* tapeworms was most likely less than 5 years. Very early studies with experimental infection of adult Japanese volunteers with *T. solium* cysticerci in the 1930s found that tapeworms produced the greatest number of gravid proglottids during the first month of patency, suggesting a relatively short reproductive life span [19,29].

Other case reports have reinforced the belief that the *T. solium* life span is no longer than several years. In two instances, travelers from Japan and Mongolia both visited India where they reported consuming local pork dishes and spontaneous expelled their tapeworms within 3 years of returning home [30,31]. The current study suggests that a large number of *T. solium* parasites were expelled from children within an 8-month time period. However, the authors acknowledge that stool-based evidence was absent after treatment to ensure worm removal. Overall, the spontaneous passing of *T. solium* tapeworms in children was estimated to occur within 5 years of infection
based on children with naturally expelled worms reporting segment expulsion for no longer than three years.

The possibility of reinfestation is high in highly endemic areas, such as the village of Guni. However, it is not clear if repeated infections impact immunogenicity and worm expulsion. In the current study, all T. solium carriers reported segment expulsion for no longer than five years. Preliminary work done by our research group has shown similar findings in low endemic areas (prevalence less than 1%) of China (unpublished), suggesting that repeated infection likely has no or only a limited effect on parasite longevity. Moreover, new infections are likely deterred through concomitant immunity caused by infection with the same species of tapeworm. This hypothesis is supported by the observation of cases infected with multiple tapeworms of the same species that likely have the same source of infection based on worm morphology and size [32].

Multiple programs have been carried out in an attempt to control T. solium taeniasis and cysticercosis at a regional scale [8,33–36]. However, success has been limited. While the current study does have the limitation of using a convenience sample, 56.0% (389/695) of villagers were evaluated in Guni. Overall, children less than 16 years of age were shown to have a higher T. solium taeniasis prevalence than older individuals. Therefore, children need to be included in T. solium control programs carried out in endemic settings. This study also provided additional support for the hypothesis that T. solium likely has a relatively short life span based on the large number of T. solium parasites that were apparently expelled within an 8-month time period. Information obtained in this study allows us to further understand local T. solium transmission dynamics and develop effective intervention measures.

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Disclosure of potential conflicts of interest

No potential conflict of interest was reported by the author(s).

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