Real-Time Polymerase Chain Reaction for Detection of Low-Intensity Schistosoma japonicum Infections in China

Tore Lier,* Gunnar S. Simonsen, Tianping Wang, Dabing Lu, Hanne H. Haukland, Birgitte J. Vennervald, Joachim Hegstad, and Maria V. Johansen

INTRODUCTION

Despite tremendous success in controlling Schistosoma japonicum in the past half century, the parasite is still a major public health concern in schistosomiasis-endemic countries, with China having the most cases.¹² In most of the villages where infection occurs, the prevalence and average infection intensity is now low.¹ This finding creates a diagnostic dilemma because the usefulness of diagnostic tests commonly used in China, such as the Kato-Katz thick smear, the hatching test, and antibody detection, changes with the epidemiology of the disease. When endemicity is high, morbidity control is usually attempted through mass chemotherapy.³⁴ Under such circumstances, it is cost-effective to use a simple and cheap diagnostic test. High sensitivity is less important, and even high specificity is not vital because praziquantel is inexpensive and relatively safe. When the level of endemicity decreases, transmission control and interruption are desirable goals, but these strategies will accordingly require more sensitive and specific diagnostic tests. Inferior test performance will compromise the positive predictive value, which is vital for the design and execution of control efforts.¹⁰

Kato-Katz thick smears are easy, quick, quantitative, inexpensive to perform, and have for several decades been successfully used to diagnose schistosomiasis and other helminth infections in highly endemic areas.¹ However, it has repeatedly been shown that the sensitivity of the Kato-Katz technique decreases with decreasing intensity of S. japonicum infection.⁸⁹ For this reason, antibody detection in serum has been extensively used, especially the indirect hemagglutination assay (IHA) and the enzyme-linked immunosorbent assay with soluble egg antigen.¹⁰–¹² The main problem with this approach has been low specificity because of persistent antibodies from previous infections and cross-reactions with other helminth infections. A commonly used diagnostic strategy has therefore been to screen for antibodies and subsequently examine the stool of seropositive persons by using the Kato-Katz or hatching test.³ Only those persons who have positive results in the latter tests are treated.

The polymerase chain reaction (PCR) has the potential for high sensitivity and specificity. Pontes and others reported the first use of PCR for detection of S. mansoni in a population study.¹³ A few Schistosoma PCR assays for diagnosing human cases have been reported, but with a single exception, they have all been applied only to a limited number of clinical samples.¹⁴ This finding is also true for three PCR assays used for detection of S. japonicum in humans.¹⁵–¹⁸ Thus, the epidemiologic situation makes the search for improved diagnostic tests especially relevant for this species. We evaluated a probe-based real-time PCR as a diagnostic alternative and compared it with the Kato-Katz test, the hatching test, and the IHA with clinical samples from China.

MATERIALS AND METHODS

Study population and sample collection. The study population was recruited from the Guanghui and Heping villages in Laozhou Township, which is located on an island in the Yangtze River in Tongling County, Anhui, China. Serum was collected in capillary tubes from the ear lobe. The Kato-Katz test, the hatching test, and the PCR were performed with one stool sample. The Kato-Katz test, the hatching test, and the IHA were performed by experienced technicians at the Anhui Institute of Parasitic Diseases. Extraction of DNA for the PCR was conducted at the same institute, and PCR amplification was performed at the institution of the first author. Informed consent was obtained from the participants. Ethical approval was provided by the village leadership, the Scientific Committee at Anhui Institute of Parasitic Diseases, and the Tongling County Health Bureau. Participants who were positive were treated with one oral dose of praziquantel (40 mg/kg) (Nanjing Pharmaceutical Factory, Nanjing, China).

Indirect hemagglutination assay. Antibody detection was performed by using the IHA and a kit containing human erythrocytes coated with soluble egg antigen. The kit is commercially available from the Anhui Provincial Institute of
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Parasitic Diseases (Wuhu, China). The test procedure was as described by Zhou and others.22 The test result was considered positive when a positive reaction appeared at a titer ≥ 1:10.

**Real-time PCR.** Plastic spoons were used to measure a stool volume equivalent to one gram. DNA was extracted by the use of rapid one-step extraction buffer and ethanol precipitation as described.13,18 The PCR was based on a previously evaluated assay targeting the mitochondrial NADH dehydrogenase I gene, but with new primers and probe-based detection to maintain high specificity.17,18 The consensus sequence from alignment of NADH dehydrogenase I gene sequences from \( S. \) japonicum (GenBank accession nos. AF215860, AF056291–96, and AF056288–90) covering seven geographic locations in China and The Philippines were used as basis for primer and probe design.9–21 Alignment was performed using SeqMan version II 5.05 Lasergene software (DNASTAR, Madison, WI).

The new primers and probe were designed using Primer Express version 3.0 software (Applied Biosystems, Foster City, CA). The primer set F-SJ-ND1 (forward: 5′-ACTGGTTA TGGTTGTTGATGTTAGT-3′) and R-SJ-ND1 (reverse: 5′-AGGCCACGAAAGACATTAC-3′) synthesize a 75-basepair amplicon. The probe was a minor groove binding probe, MGB-SJ-ND1 (Applied Biosystems): 6-FAM-5′-AGGTTCCTGGAAAAAGTAT-3′-MGBNFQ (MGBNFQ is a minor groove binder/non-fluorescent quencher). The reaction mixture consisted of 5 µL of DNA template, 12.5 µL of TaqMan Universal PCR Master Mixture (Applied Biosystems), 300 nM of each primer, 250 nM of probe, 2 µL of bovine serum albumin (Promega, Madison, WI), and distilled water to give a final volume of 25 µL.

The cycle protocol consisted of incubation at 50°C for 2 minutes, 95°C for 10 minutes, and 50 cycles at 95°C for 15 seconds and 60°C for 1 minute. A 7300 Real Time PCR System thermal cycler with accompanying SDS software (both from Applied Biosystems) was used. The PCRs were carried out in duplicate for every sample. Samples that were positive in only one PCR were tested in triplicate, and the test result was regarded as positive when at least one of these triplicate results was positive. As a result of this modification, 12 samples had to be retested, of which 5 were regarded as having positive results after the second run. We found the PCR amplification curves easy to classify into positive and negative results. Non-template controls, with water instead of DNA template, were included in all runs to check for contamination.

Sensitivity for different egg counts was examined by spiking one-gram stool samples with various numbers of \( S. \) japonicum eggs recovered from mice livers.15 The 46 samples contained 1,000 eggs (n = 2), 100 eggs (n = 8), 50 eggs (n = 13), 10 eggs (n = 7), 5 eggs (n = 7), and 1 egg (n = 9). The PCR result was positive for all the samples except for one sample that contained one egg per gram (epg) of stool.

The PCR specificity was tested in three types of material. The first type was DNA extracted from adult worms of the following species: \( S. \) japonicum (Anhui strain), \( S. \) mansoni, \( S. \) haematobium, \( S. \) bovis, and \( O. \) viverrini. The \( S. \) japonicum strain was PCR positive, and the other Schistosoma species and \( O. \) viverrini were PCR negative. The second type was the study database that was searched for samples that were Kato-Katz positive for other helminth eggs, but negative for \( S. \) japonicum by the Kato-Katz test, the hatching test, and the IHA. We found 6 samples containing *Ascaris lumbricoides* and 34 containing *Trichuris trichiura*. All 40 samples were PCR negative. The third type was DNA extracts from 29 bacterial and yeast species in stool (all from American Type Culture strains; LGC Standards AB, Boras, Sweden) and human DNA from two cell lines (MRC-5 and HE). All 31 samples were PCR negative.

**Hatching test and Kato-Katz thick smear.** The hatching test was conducted by transferring approximately 30 grams of stool into a metal container with a coarse metal mesh. Fine material was flushed into a fine-meshed nylon bag by using flowing water and a stirring stick. Sediment was washed by flushing with water until the water was clear and by stroking the outside of the bag with chopsticks. The content of the bag was then transferred to an Erlenmeyer flask containing 300 mL of non-chlorinated water from the Yangtze River. The flasks were left in a well-lit room at approximately 28°C and checked by two of the staff for swimming miracidia after 4, 6, 8, and 24 hours by using a strong light that illuminated the back or side of the flasks.

The Kato-Katz thick smear was performed as described by using nylon screens and plastic templates (Zhejiang Ningbo Medical Instrument Factory, Ningbo, China). Three slides (41.7 mg each) were prepared from one stool sample from each person and examined within a week.

**Data analysis.** Data were stored in an Excel 2002 spreadsheet (Microsoft Corp., Redmond, WA) and analyzed with SPSS version 14.0 for Windows software (SPSS, Inc., Chicago, IL). Only stool samples with results from all three stool-based tests were selected to examine the agreement between the three tests (n = 1,106; Figure 1). The value of \( \kappa \) was calculated using \( \kappa = (\text{P}(o) - \text{P}(e))/\left(1 - \text{P}(e)\right) \).

To calculate effects on patient treatment by changing from a traditional IHA/Kato-Katz–based algorithm to a strategy based on the IHA and PCR, we selected all samples with available results for IHA, Kato-Katz, and PCR (n = 942).

![Figure 1](image.png)

**Figure 1.** Agreement between polymerase chain reaction, Kato-Katz thick smear, and hatching test for stool-based diagnosis of *Schistosoma japonicum* infections, China. Values indicate the number of positive samples in each group.
RESULTS

Prevalence and egg counts. Samples were collected from 1,727 persons (52% males and 48% females, age range = 6–84 years, mean = 40 years). Results from all diagnostic tests were not available for the whole study population. The total number of samples tested, number of positive samples, and prevalence of *S. japonicum* determined by using the three diagnostic tests are shown in Table 1.

The 46 Kato-Katz–positive samples could be divided into 3 egg count groups: 26 samples (57%) with < 40 epg, 15 samples (33%) with 40–99 epg and 5 samples (11%) with > 99 epg. The lowest possible egg count in this study was 8 epg because this value represents 1 egg per 3 Kato-Katz slides. There were 9 samples with 8 epg and 7 with 16 epg (total of 2 eggs per 3 slides).

Agreement between tests. The degree of agreement between positive results from the three stool-based tests is shown in Figure 1. A total of 52% of the PCR positive samples were positive only by the PCR. The corresponding values for the Kato-Katz and hatching tests were 49% and 33%, respectively. Calculation of the strength of agreement resulted in a value of $\kappa = 0.32$ the between the PCR and Kato-Katz test, 0.43 between the PCR and hatching test, and 0.34 between the Kato-Katz and hatching tests.

Agreement between IHA and stool-based test results is shown in Figure 2. This figure shows the proportion of positive IHA results in samples positive in each of the three tests and the proportion of positive IHA results in samples where one test result was positive and the other test result was negative.

The proportion of positive IHA results increased with increasing egg count. In samples with less than 20 epg (n = 13) 31% were IHA positive, with 20–100 epg (n = 24) 58% were IHA positive, and with > 100 epg (n = 5) 80% were IHA positive.

Treatment algorithm. If the Kato-test had been used in a traditional treatment algorithm (screening with IHA and testing the seropositive samples with the Kato-Katz test), it would have resulted in treatment of 22 persons. Nineteen patients who were Kato-Katz positive but IHA negative would not have received treatment. If the PCR had been used as the stool-based test, 50 persons would have been treated, and 13 persons who were PCR positive but IHA negative would not have been treated.

DISCUSSION

Consistent with the increasing success of the schistosomiasis control program in China, making a reliable diagnosis with the methods used has become increasingly problematic. In search of a diagnostic alternative, we compared a real-time PCR with the Kato-Katz thick smear, hatching test, and IHA, which are diagnostic tests commonly used in the schistosomiasis control program in China. In a large nationwide survey in China during 2004, the average *S. japonicum* prevalence in humans was 2.5% in villages classified as endemic for schistosomiasis. In more than half of the villages, the prevalence was < 1%. The average egg output was 33 epg. Such low prevalence and low egg output is a huge challenge for use of diagnostic tests. Performance of diagnostic tests is often evaluated in populations with higher *S. japonicum* prevalence than what is now most common in China. The predictive value of a test cannot automatically be transferred between two populations with different prevalences.

In the present study, the prevalence ranged from 3.0% to 5.3%, depending on the stool-based test used. The PCR test gave a higher prevalence of *S. japonicum* infections than the Kato-Katz and hatching tests. Pontes and others reported the prevalence of *S. mansoni* to be 30.9% in 3 stool samples examined with duplicate Kato-Katz smears in contrast to 38.1% in one sample examined by PCR. In Senegal, ten Hove and others found that a PCR with two stool samples resulted in lower *S. mansoni* prevalence (73% versus 80%) than the Kato-Katz test with the same two samples (duplicate 25-mg smears from each sample). However, these results cannot be compared directly to the present study because the assay and the epidemiology are different.

A surprising finding in our study was the low overall agreement between the three stool-based tests and their large variation in concordance with the IHA. The total number of positive samples detected by the Kato-Katz and hatching tests was almost the same, but the results corresponded in less than half of all cases. Low agreement between *S. japonicum* diagnostic tests was also evident in earlier studies.

Several explanations may account for the discrepancies in the results among the different test methods. The reproducibility of test results will be compromised when egg content in each sample is close to the detection limit of the tests. In the present study, 35% of the Kato-Katz–positive samples had only one or two eggs on 3 slides (< 20 epg). The PCR uses a larger sample than the Kato-Katz test, but even one gram amounts to only approximately 0.6% of the daily stool output in a person in China. An even larger amount of stool is examined in the hatching test, but hatchability, and thus sensitivity of the assay, may be influenced by a number of ill-defined factors. Finally, the uneven distribution of *S. japonicum* eggs within stool samples may further lead to discrepancies between test results.
An alternative explanation is that IHA has lower sensitivity when egg counts are low.12 Evaluation of stool-based methods is further complicated by the uncertain status of the IHA as a gold standard for past or present Schistosoma infections. Low specificity of the IHA in previous studies may in part be explained by use of a limited number of Kato-Katz smears or hatching of low sensitivity as the gold standard. Yu and others reported an IHA sensitivity and specificity of 80% and 48%, respectively, using a comprehensive stool examination with duplicate Kato-Katz slides from 7 stool samples and hatching as the gold standard.11 The diagnostic strategy for schistosomiasis must change as control of the parasite progresses in an area.10 The two-step algorithm commonly used in China, with an antibody test followed by the Kato-Katz test of samples from patients who were seropositive, may not be appropriate as prevalence continues to decrease. In the present study, nearly half of the Kato-Katz–positive cases would not have received treatment if this traditional algorithm was used. It remains uncertain whether the main problem in this situation is false-positive Kato-Katz results or false-negative IHA results. More than twice as many persons would have received treatment if the PCR had replaced the Kato-Katz test and to a lesser extent the hatching test is dependent upon the experience of the technician, while objective criteria can be made for a positive PCR. We have demonstrated the use of an inexpensive, non-commercial DNA extraction method applicable to local laboratories with limited resources. Amplification and detection can be centralized to larger facilities. As DNA-based diagnostic technologies are developed to become simpler and more affordable, they will be available in many laboratories, even in less affluent countries. However, it still is an expensive diagnostic alternative.

A satisfactory diagnostic algorithm for low-intensity S. japonicum infections in schistosomiasis-endemic areas has yet to be determined. More research in this field is needed to evaluate diagnostic algorithms with traditional tests and search for new methods. Diagnosing such infections will benefit patients and help to eliminate the parasite reservoir and assess if and to which degree transmission is still occurring.31 As prevalence decreases, additional measures to reduce transmission, such as infection control in the animal reservoir, improved sanitary conditions, and health education, will be of increasing importance.32

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Authors’ addresses: Tore Lier, Gunnar S. Simonsen, Hanne H. Haukland, and Joachim Hegstad, Department of Microbiology and Infection Control, University Hospital of North Norway, PO Box 56, N-9038 Tromsø, Norway. E-mails: tore liar@unn.no, gunnar.skov.simonsen@unn.no, hanne.husom.haukland@unn.no, and joachim.hegstad@unn.no. Tiaping Wang, Anhui Institute of Parasitic Diseases, 207 Dongjiao Road, Wuha 241000, Anhui, Peoples Republic of China, E-mail: wangtiaping@hotmail.com, Dabing Lu, Department of Infectious Diseases, Faculty of Medicine, Imperial College, London SW7 2AZ, United Kingdom, E-mail: dabling2001wyz@yahoo.com.cn. Birgitte J. Vennervald and Maria V. Johansen, DBL–Centre for Health Research and Development, Faculty of Life Sciences, Thorvaldsensvej 57, University of Copenhagen, DK-1871 Frederiksberg C, Denmark, E-mails: bjv@life.ku.dk and mvj@life.ku.dk.

REFERENCES


Appendix to Paper 3:

Figure 2 in the printed version of the paper can be difficult to read due to the small size. The figure and figure text is reproduced in a larger size below.

Figure 2. Agreement between the indirect hemagglutination assay (IHA) and stool-based PCR, Kato-Katz thick smear (KK) and hatching test (H) for Schistosoma japonicum egg detection. The size of each column corresponds to the number of positive stool-based samples in each group. The percentage in each column represents the proportion positive by IHA in the group.