Detection of Malaria Parasites in Mosquitoes from the Malaria-Endemic Area of Chakaria, Bangladesh

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Malaria is one of the major public health problems of Bangladesh. We investigated the mosquito populations infected with malaria parasites in a malaria-endemic area Chakaria, Bangladesh, where Anopheles dirus and Anopheles minimus are the principal vectors. Anopheles mosquitoes were collected with a CDC miniature light trap from inside households in June 2007. A total of 868 mosquitoes were collected, among which females numbered 669 (77.1%). The species of female Anopheles mosquitoes were identified morphologically, and 651 were A. minimus and the remaining 18 were other Anopheles species. Malaria parasite DNA from individual female mosquitoes was extracted and distinguished using the microtiter plate hybridization (MPH) technique targeting the 18S rRNA of human malaria parasites. Nineteen mosquitoes were malaria parasite positive: 12 for Plasmodium falciparum, 1 for Plasmodium vivax, and 6 for both P. falciparum and P. vivax. This is the first time that the MPH technique was used for distinguishing malaria parasites in mosquitoes and the first report from Chakaria. Our results may contribute to planning and assessing malaria control strategies in Chakaria.

Key words  malaria; Anopheles; polymerase chain reaction; microtiter plate hybridization

Malaria is caused by the protozoan parasite of the genus Plasmodium which is transmitted to humans when female Anopheles mosquitoes feed on an infected host.1) It has become one of the biggest health concerns in Bangladesh.2) Malaria disappeared from the plains after an eradication campaign with dichloro-diphenyl-trichloroethane (DDT) from 1961—19763,4) but slowly spread again with migration. Thirteen of the 64 districts on the northern and eastern borders are seriously affected by malaria. Reported clinical cases numbered 378000, laboratory-confirmed cases reached 60000, and deaths totaled 528 in 1998.5) The most cases of malaria are found in the three districts of the Chittagong Hill Tracts bordering India and Myanmar, and Plasmodium falciparum is the predominant species.6) In Chakaria, which is under the division of Chittagong, there was community-based surveillance in 2002 which detected 1543 clinical cases.7)

Malaria distribution is usually determined by the presence of its mosquito vector. Climatic factors such as humidity, rainfall, and temperature control the development of both malaria parasites and vectors.8) Four hundred anopheline species are distributed worldwide; only 40 are important malaria vectors.9) In Bangladesh, 34 Anopheles species have been recorded and among them the four species Anopheles dirus, Anopheles philippinensis, Anopheles sundaicus, and Anopheles minimus are considered important malaria vectors.9,10) Among these species, A. dirus and A. minimus are the most widespread, are known to be forest-related vectors, and can expand their range under favorable conditions such as the monsoon period.9)

Malaria transmission ultimately depends upon the sporozoite rate of mosquitoes,11) and determining the presence of malaria sporozoites in Anopheles mosquitoes caught in the wild is an important factor in epidemiologic studies of malaria-endemic areas.12) The standard method for detection of malaria sporozoites in mosquito salivary glands uses light microscopy. Specimens must be dissected when fresh, and examination of individual mosquitoes is time-consuming and labor intensive.13,14) In recent years, the polymerase chain reaction (PCR) has been used to amplify the specific DNA sequences of P. falciparum for highly sensitive detection of parasites in the mosquitoes.14,15) In most cases, the target sequence used for detecting malaria parasites in mosquitoes using PCR is the Plasmodium species-specific small subunit ribosomal RNA (SSUrRNA).13,16,17) On the other hand, it is sometimes difficult to detect parasites in blood-engorged mosquitoes using the PCR method,18) because mosquito extracts contain strong PCR inhibitors that prohibit amplification of parasitic DNA in mosquito vectors.19)

Previously, we developed the microtiter plate hybridization (MPH) technique, which is a combination of PCR amplification targeting the 18S rRNA of human malaria parasites and DNA hybridization. The MPH technique not only detects the malaria parasites but also can distinguish the different human malaria parasite species. Our technique is simple, highly sensitive, and specific for human malaria parasites and has already been used for human malaria parasite diagnosis in the Solomon Islands,20) Vietnam,21) Thailand,22) and Korea.23) Few data are available on the natural transmission of malaria, in particular mosquito bionomics and the infectivity of the malaria vectors occurring in the Chittagong division, Bangladesh. The aim of our study was to determine the infection rate and distinguish different Plasmodium species in the mosquitoes in a malaria-endemic area in Bangladesh using the MPH technique. To the best of our knowledge, this was the first time that the PCR-based MPH technique was used to detect and distinguish malaria parasite species infection in Anopheles mosquitoes.

MATERIALS AND METHODS

Study Area The study was conducted in Chakaria,
Bangladesh, and the protocol was approved by the International Centre for Diarrhoeal Disease Research, Bangladesh (ICDDR, B). Chakaria is a subdivision in the district of Cox’s Bazaar under the division of Chittagong. It has a population of 550,000 with an area of 643.46 km², and is surrounded by many rivers such as the Matamuhuri, Maheshkhali, and Kutubdia Channel (Fig. 1). The east side of the Chakaria area is hilly with forests, and the west side is low lands bound by the Bay of Bengal. Two unions (Kakara and Harvang) of Chakaria were selected for mosquito collection. Traditionally, the main economic activities in the area have been agriculture, forestry, and fishing. Most of the people are very poor, wear very scanty clothes, and are exposed to the vector as they live in houses that sometimes have one or more side open. Household walls are usually made of bamboo and mud and very few are made of tin, wood, or cement; floors are earthen; roofs are made of leaves and straw and very few are of tin. The domesticated animals are cows, goats, and chickens that live in the areas surrounding houses. Cattle sheds are open without walls and located adjacent to households. The climate of Chakaria from May to September is characterized by heat and heavy rainfall, while the remainder of the year is mostly dry. The annual maximum temperature is 28—29 °C and the minimum temperature is 20—21 °C. Most rain occurs during the monsoon (June—September) and little in winter (November—February); annual rainfall is 3000—3500 mm. During the study period in June 2007, there was heavy rainfall of 425 mm/d in Chittagong division, and flooding made two-thirds of the area unreachable.

Mosquito Collection Adult mosquito collection was carried out June 12—30, 2007, just after the heavy rainfall. The mosquitoes were collected using six standard miniature CDC light traps (Model 512; John W. Hock Company, Gainesville, Florida, U.S.A.). In Chakaria, mosquitoes were collected from the three villages Goalmar, Pahartoli, and Manikpur in two unions upon receiving consent from the household heads. People living in these houses were protected with nonimpregnated bed nets. The trap was hung inside the room of sleeping persons as described previously. The light traps were switched on each day at 18:00 and were collected at 06:00 of the next morning.

The captured mosquitoes were transported to the laboratory in plastic containers inside desiccators over silica gel. They were then separated into Culicinae and Anophelinae and counted. The individual female *Anopheles* mosquitoes were identified based on the taxonomic keys and stored in tubes inside the desiccators over silica gel until completely dry. The mosquitoes were stored under the same conditions until transported to the main laboratory, where they were stored at −80 °C until processed for DNA extraction.

Extraction of Malaria Parasite DNA from Mosquitoes DNA was extracted from the individual mosquitoes with MX-16 DNA purification kits (Promega, U.S.A.) that were used with the MX-16 instrument to provide an easy method for efficient automated purification of genomic DNA from solid biomaterials. This system is based on magnetic bead technology and composed of a magnetic bar, bore, and purified cartridge. In this system, the movement of the bore is controlled by motors, which allows easy DNA extraction from crude lysate magnetic beads. All the necessary reagents for purification of nucleic acid from a single mosquito are supplied in one reagent cartridge. Each well of the cartridge is prefilled with magnetic beads, wash buffer (I, GuSCN 5.25 M; II, 70% EtOH), and lysis buffer (GuSCN 4.7 M). The DNA isolation was performed by adding sample into the lysis chamber. The procedures for DNA purification was performed automatically, and after DNA extraction it was transferred to a fresh tube and stored at −80 °C until use.

Fig. 1. Map of the Study Area Chakaria, Bangladesh, Where Mosquitoes Were Collected
Target Sequences of Malaria Parasites  The oligonucleotide primer set, biotin-labeled MPH1 of 5′-CAGATACGGTCGTAATCTTA-3′ and MPH2 of 5′-CCAAAGACTTTGATTTCTCAG-3′, which are specific to the 18Sr RNA genes of the human malaria parasites of P. falciparum and Plasmodium vivax, Plasmodium ovale, Plasmodium ovale-variant, and Plasmodium malariae were described previously.20,23,26,28) These two primers were used for the DNA amplification of the malaria parasites in mosquitoes with PCR. After PCR amplification, the PCR products were used to distinguish the malaria parasite species using the MPH technique. The specific probes of P. falciparum 5′-GTCACCTCGAAAGATGACTT-3′, P. vivax 5′-TAAACTCCGAGGAAATATC-3′, P. ovale 5′-AATTCCCCGAAAGGAATTTTC-3′, P. ovale-variant 5′-GAAATTTCCAAAGGAATTTC-3′, and P. malariae 5′-ACTCATATATAAGAATGTCACCT-3′ were captured in the microtiter plate as described previously.21,27)

Preparation of DNA for PCR  Five microliters of extracted DNA from Anopheles mosquitoes was added to the 0.2 ml microcentrifuge tube containing 40 µl of lysis solution (Tris–HCl 1% w/v, pH 8.9, MgCl2 1.5 mM, KCl 80 mM, BSA 500 µg/ml, 0.1% sodium cholate, 0.1% Triton X-100, Proteinase K 200 µg/ml, 0.45% Tween 20, 0.45% Nonidet p-40). Then the sample was incubated at 60 °C for 20 min for destruction and proteolysis, at 95 °C for 13 min to inactivate the Proteinase K, and then cooled at 50 °C for 5 min.

PCR and Electrophoresis  PCR was carried out to detect the presence of malaria parasites in mosquitoes. The sample was spun down and mixed with 10 µl of PCR reagent mixture (Tris–HCl 10 mM, pH 8.9, MgCl2 1.5 mM, KCl 80 mM, BSA 500 µg/ml, 0.1% sodium cholate, 0.1% Triton X-100, 5 µg/ml primer, dNTPs 1 mM). The mixture was spun down, vortexed, and subjected to 30 cycles. The conditions were as follows: denaturation at 92 °C for 60 s, annealing at 52 °C for 90 s, and extension at 72 °C for 90 s. The amplified DNA was denatured by heating at 95 °C for 10 min and then rapid cooling on ice for 10 min. Positive controls were DNA of P. falciparum, P. vivax, P. ovale, P. ovale-variant, and P. malariae from clinical patients. Sterilized water was used as a negative control. The PCR products (10 µl) were electrophoresed at 100 V for 40 min using 1.2% agarose gel, stained by ethidium bromide, and the bands were visualized with a UV transilluminator.

Hybridization and Colorization of the MPH  The procedure for hybridization and colorization in the MPH technique have been described previously.23,26,28) Briefly, microtiter plate wells are coated with probes specific for P. falciparum, P. vivax, P. ovale, and P. ovale-variant, and P. malariae was added at 100 µl/well of 5×SSC (1×SSC is NaCl 0.15 M plus sodium citrate 0.015 M) and 5 µl of PCR product was added to the wells. Then the plate was incubated at 58 °C for 1 h, then the solution was removed, and the wells were washed three times with 250 µl/well of 1×solution (Tris–HCl 0.1 M, pH 7.5, NaCl 0.1 M, MgCl2 2 mM, 0.05% Triton X-100). Then 100 µl of alkaline phosphatase-labeled streptavidin was added and incubated at 28 °C for 15 min. After that the solution was removed and the wells were washed three times with 250 µl/well of 1×solution. Then 100 µl/well of coloring substrate, PNPP solution (diethanolamine 1 M, pH 9.8, MgCl2 0.5 mM, and para-nitrophenyl phosphate 10 mM) were added and incubated at 28 °C for 30 min. The absorbance of each well was read at 405 nm using a microtiter plate reader (MPR-A4, Tosoh, Tokyo, Japan). The data were obtained by subtraction of the background corresponding to the PNPP solution.

RESULTS

Proportion of Anopheline Mosquitoes  Mosquitoes were captured only one time from each household of two unions in Chakaria during June 2007. In this survey, a total of 868 mosquitoes were collected from the three villages in two unions. Among the total collected mosquitoes, 669 (77.1%) mosquitoes were found to be female anopheline mosquitoes. The species of anopheline mosquitoes were morphologically identified as A. minimus, 651 (97.3%); 18 (2.7%) mosquitoes were other Anopheles species. The remaining 199 (22.9%) of the total collected mosquitoes were Culex species. Among the 868 mosquitoes, 838 were collected from the village of Pahartoli (Table 1).

<table>
<thead>
<tr>
<th>Union</th>
<th>Village</th>
<th>Mosquito species</th>
<th>Anopheles minimus</th>
<th>Other Anopheles species</th>
<th>Culex species</th>
<th>Total No.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Kakhara Manikpur</td>
<td>2</td>
<td>70</td>
<td>9</td>
<td>617</td>
<td>21</td>
<td></td>
</tr>
<tr>
<td>Harvang Goalmara</td>
<td>4</td>
<td>17</td>
<td>2</td>
<td>199</td>
<td>9</td>
<td></td>
</tr>
<tr>
<td>Pahartoli</td>
<td>649</td>
<td>182</td>
<td>7</td>
<td>838</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total</td>
<td>651 (75.0%)</td>
<td>18 (2.1%)</td>
<td>199 (22.9%)</td>
<td>868</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Fig. 2. Agarose Gel Electrophoresis Pattern Showing the Malaria Parasite-Positive PCR Product from A. minimus (Lanes 1—19), Negative Control (Lane 20), and Positive Control with DNA of P. falciparum, P. vivax, P. ovale, P. ovale-variant, and P. malariae (Lanes 21—25) M, DNA ladder marker of 100bp. PCR products represent the results of only malaria parasite-positive mosquitoes; malaria parasite-negative mosquitoes are not included.
Electrophoresis Results of PCR Product in Mosquitoes

The extracted DNA of 669 mosquitoes was amplified using the primer pair set of MPH1 and MPH2 for detecting the malaria parasite. The amplified PCR product was subjected to agarose gel electrophoresis and this revealed a prominent band at 135 basepairs of malaria parasites in *Anopheles* mosquitoes. A total of 19 *Anopheles* mosquitoes were found to be malaria parasite positive (Fig. 2).

### Results of MPH

After PCR amplification, the PCR products were analyzed using the MPH technique. The total malaria parasite-positive rate was 2.8% (19/669) and it showed the same results as PCR. Among the 19 positive samples, 12 (63.2%) were positive only against the *P. falciparum* probe and 1 (5.3%) against the *P. vivax* probe. Six samples (31.6%) showed strong positive reactions against both the *P. falciparum* and *P. vivax* probe. Malaria parasite-positive mosquitoes were all *A. minimus* species and all were collected from the village of Pahartoli. Eleven malaria parasite-positive mosquitoes were collected on June 23, five on June 25, one on June 22, one on June 28, and one on June 30 (Table 2). The absorbance values of 19 samples of malaria-positive mosquitoes using the MPH technique are listed in Table 3.

### DISCUSSION

Malaria vector surveillance is a prerequisite for malaria vector control. When vector surveillance is carried out on large numbers of mosquitoes, they must be tested for the presence of parasites. Salivary gland dissection is impractical when processing large numbers of mosquitoes. Moreover, microscopic examination is time-consuming and labor intensive and cannot distinguish between different species of malaria parasites. On the other hand, MPH is very sensitive.
in distinguishing malaria parasite species. In Bangladesh, malaria is restricted to the hilly districts in the eastern and southern border areas. Very few studies have been carried out in Bangladesh on the population densities of Plasmodium sporozoite-infected mosquitoes, especially in the endemic area of Chakaria, and the malaria infection rate in mosquitoes has not been determined. Therefore we tried to determine the malaria vector population infected with malaria parasites responsible for malaria transmission in the highly endemic area of Chakaria, Bangladesh. In the area, mosquito collection was conducted in June 2007, when there was unusually heavy rainfall and most of the study area was under water. Therefore it was difficult to collect Anopheles mosquitoes from all three villages, and most mosquitoes were collected from the village of Pahratoli.

In our study using CDC light traps, 77.1% of mosquitoes collected were anopheline, of which 97.3% were A. minimus mosquitoes. In another study, the mosquitoes were collected using the human landing aspiration technique, the anopheline mosquito collection rate was 37.0%, and among the Anopheles mosquitoes, A. minimus comprised 50% in the same area. Other researchers reported that the A. minimus mosquito collection rate was 57.3% in the Chittagong Hill Tracts, which were also collected using human bait. Data from our study suggest that rainfall may increase the Anopheles mosquito population. During the study, the Matamuhuri River surrounding the study area overflowed and may have increased the breeding area for A. minimus. The increase in population is related to the increase in breeding sites in which Anopheles eggs are deposited and develop into adults. Usually rainfall is associated with increased humidity that stimulates the resting gravid mosquitoes to oviposit and seek new hosts. In our study, the maximum number of mosquitoes was collected 2 weeks after the rainfall as it takes 10—14 days for mosquitoes to oviposit. The same result was found when numerous mosquitoes were concentrated in 2—4 weeks after rainfall.

The sporozoite rate in mosquitoes can be detected using different methods such as the enzyme-linked immunosorbent assay (ELISA) and rapid dipstick immunochromatographic assay (Vec-Test Malaria). But ELISA requires training and is not easy, and not time-consuming for malaria parasite detection. Further research may be required to detect malaria parasites-positive mosquitoes that can transmit malaria parasites. But our investigations revealed that the MPH technique is simple, sensitive, easy, and not time-consuming for malaria parasite detection in numerous mosquitoes. Our results also indicate that there may be a relation between rainfall and the A. minimus mosquito population. As one of the principal malaria vectors in Chakaria, A. minimus mosquitoes may play a greater role in malaria transmission in the area after heavy rainfall. Therefore special attention should be given to reducing the population of this mosquito for malaria control. This was the first report of distinguishing malaria parasites in individual female Anopheles mosquitoes using the MPH technique and also the first report of the malaria-positive rate in Anopheles in Chakaria, Bangladesh. The MPH technique is recommended for use in identifying malaria transmitters and in epidemiologic studies for promoting effective control measures not only for Bangladesh but also for other malaria-endemic areas. Our data will play an important role in initiating malaria control in Bangladesh.

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