Hemozoin Pigment: An Important Tool for Low Parasitemic Malarial Diagnosis

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Abstract: Low parasitic condition in malaria remains a diagnostic challenge; as the available diagnostic methods failed to detect. Currently, hemozoin (Hz) pigment is gaining attention in the diagnosis of malaria. The major drawback is ease of detection of Hz in routine practice. A pilot study was conducted to evaluate the role of Hz pigment and to compare the performance of quantitative buffy coat assay (QBC) and PCR in such conditions. Clinically suspected cases of malaria were examined by both Giemsa stain and immunochromatographic test (ICT). Samples positive by ICT and negative by Giemsa stain were further examined by nested PCR targeting 18S rRNA and QBC for the presence of malaria parasites and pigments. Thirty blood samples fulfilled the inclusion criteria out of which 23 were Plasmodium vivax (Pv), 4 Plasmodium falciparum (Pf), and 3 mixed (Pv and Pf) by immunochromatographic test. Twenty-one out of 30 (70%) were positive by nested PCR in comparison to 25/30 (83%) by QBC. Samples containing both malaria parasites and Hz pigment by QBC completely showed concordance with the PCR result. However, 61% of total samples containing only Hz pigment were observed positive by PCR. Hz pigment remains an important tool for malaria diagnosis. Identification of leukocytes containing pigments by QBC not only indicates recent malarial infections but also puts light on severity of the disease. QBC assay is a rapid, highly sensitive, and cost-effective method to detect malaria parasites and Hz pigment especially in low parasitic conditions.

Key words: Plasmodium vivax, Plasmodium falciparum, low parasitemia, hemozoin pigment, QBC, nested PCR

INTRODUCTION

In malaria, identification of low parasitemic conditions is always a diagnostic challenge. The malaria parasites are maintained as a source of reservoir in these conditions and help in transmission in the community [1]. Correct identification with appropriate treatment of such conditions is highly essential for malaria elimination. It is not only helpful in decreasing the morbidity and mortality but also prevents the transmission. Microscopy of peripheral blood smear and rapid diagnostic tests (RDT) detecting malarial antigen remain the main stay for malaria diagnosis [2]. Although these diagnostic modalities are quite popular in endemic countries, they have their own limitations. Microscopic examination of peripheral blood smears stained with Giemsa needs expertise. In addition, parasitemia below levels of microscopic detection will be falsely diagnosed as negative and continue as potential sources of disease transmission. Similarly, performance level of an RDT can be affected due to varying antigen concentrations in blood and parasitemia levels due to multiple host and parasite factors [3]. Detection of Plasmodium nucleic acid by PCR offers the advantage of high analytical sensitivity in terms of detecting 1 Plasmodium falciparum or 3 Plasmodium vivax parasites per ml of blood when conventional methods will be negative [4]. However, the usage of PCR requires higher infrastructural support and economic restraints can be a hindrance in its implementation in routine usage in resource-limited settings.

Currently, the role of hemozoin (Hz) pigment in malaria diagnosis is gaining attention [5]. Hz pigment is an important biomarker of malarial infections [6]. These pigments are released during rupture of infected erythrocytes and malaria parasites and phagocytized by the leukocytes during their life cycle. Hz pigments can be easily detected under a microscope because of their typical morphology. Despite the presence of various sensitive methods like quantitative buffy coat assay (QBC) and PCR for malaria diagnosis, the available literatures...
on Hz pigment mostly use flow cytometer for its detection on routine basis [7,8]. Here, in this study, it was aimed to look for the role of Hz pigment in the diagnosis of low parasitemic malarial conditions. We also tried to compare the performance of diagnostic methods like QBC and PCR for the detection of Hz pigment in such conditions.

MATERIALS AND METHODS

A pilot study was conducted over a period of 3 months. Thirty patients with clinical diagnosis of malaria were included in this study. Ethical clearance was not required for this study as all the samples were routinely sent to the parasitology laboratory of Microbiology Department, and all the tests were performed for all the samples. All the samples were positive by immunochromatographic test (ICT) detecting the LDH antigen and negative by microscopic examination of the peripheral (thin) smear by Giemsa stain. These cases were defined as low parasitemic conditions as no malaria parasites were detected by microscopic examinations after 3 independent observations and positive by ICT (Advantage Mal Card, J Mitra & Co Pvt. Ltd, New Delhi, India). Blood samples fulfilling the above conditions were further tested by QBC and nested PCR. QBC capillary tubes were examined for presence of malaria parasites and Hz pigment by all the examiners. All the blood samples were also tested for nested PCR targeting 18S rRNA.

DNA extraction

DNA was extracted from blood (50 µl) using Qiagen DNA isolation kit following manufacturer’s instructions [9,10].

Nested PCR based on 18S rRNA amplification

For nested PCR, the species-specific nucleotide sequences of 18S rRNA genes of P. falciparum, and P. vivax were amplified as described previously, with slight modifications. For each 20-µl reaction mixture for Nest 1 amplification, 5 µl of DNA template (corresponding to approximately 0.25 to 0.5 µl of blood) was added to reaction mixture containing 10 pmoles of each primer (rPLU1 TCAAAGAATAAGCCATGCAAGTGA and rPLU2 TAACCCCGTGTTGTTGCCCTTAAACTCC), 4 mM MgCl₂, PCR buffer (50 mM, KCl, 10 mM Tris-HCl), 200 mM of each deoxynucleoside triphosphate, and 1.25 units of Taq DNA polymerase. For Nest 2 assay, 1 µl of Nest 1 PCR product was used with 10 pmoles of respective primers in given PCR conditions. P. vivax VIV F 5’CGCTTCTAGTTCATCCACATATGATA3’; VIVR 5’ACTTCCAAGGAGGAAAGTCTCA-3’ and P. falciparum FAL F 5’TAAACTGGGTITGGGAAAACAAATATATT-3’; FAL R ACACAATGAATCAATGACTACCCGT. The amplified products were visualized in 2% agarose gels stained with ethidium bromide.

RESULTS

All the blood samples were positive by ICT. Twenty-three out of 30 samples were detected positive for P. vivax, 4 were P. falciparum, and 3 were mixed infection showing test bands for both P. vivax and P. falciparum. All the samples were negative by Giemsa stain examined by 3 independent observers. Twenty-five (83%) out of 30 samples were found positive by QBC either for malaria parasite or Hz pigment or both in comparison to 21/30 (70%) by nested PCR. Considering PCR as the gold standard for malaria diagnosis, 70% (21 out of 30 samples) were considered as true positive (Table 1).

QBC result analysis

QBC assay was performed using fluorescent dye (acridine orange)-coated microcentrifuge capillary tube, where the blood cells are layered as per their specific gravity. The lighter cells remained towards the periphery of the capillary tube whereas the heavier cells were positioned towards the cap (Fig. 1). In the buffy coat area, the malarial parasites (with or without Hz pigments) were found. Leucocytes (e.g. neutrophils, monocytes, and macrophages) containing Hz pigments were also found in this zone. According to the QBC observation, our study group was categorized into 3 different subgroups (blood samples containing both malaria parasites and Hz pig-

| Groups                        | PCR positive | PCR negative | Hz in monocytes | Hz in macrophages | Hz in neutrophils |
|-------------------------------|--------------|--------------|-----------------|------------------|-------------------|
| Malaria positive and Hz pigment (n= 7) | 7            | 0            | 7               | 7                | 1                 |
| Hz pigment (n= 18)            | 11           | 7            | 18              | 18               | 5                 |
| Negative for malaria and Hz pigment (n= 5) | 3            | 2            | 0               | 0                | 0                 |
| Total (n= 30)                 | 21 (70%)     | 9            | 25              | 25               | 6                 |
ment, samples showing only Hz pigment, and samples without malaria parasites and Hz pigment). QBC was positive in 83% (25/30) of the total samples. The majority of the samples (18/25) positive by QBC showed only Hz pigment. Seven samples were positive for both malaria parasites and Hz pigment, and 5 samples were negative for both. All the samples positive for both malaria parasites and Hz pigment by QBC were found positive by nested PCR. Monocytes and macrophages were the commonest leukocytes containing pigments followed by neutrophils (Table 1).

**Table 2.** Correlation of hemozoin (Hz) pigment detection with PCR

|                         | PCR positive | PCR negative |
|-------------------------|--------------|--------------|
| Hz positive             | 18           | 7            |
| Hz negative             | 3            | 2            |

PCR sensitivity 18/21 (85%), PCR specificity 2/9 (22%).

**Fig. 1.** Quantitative buffy coat (QBC) photograph showing leucocytes containing hemozoin pigment.

**Fig. 2.** Gel picture of nested PCR (Pf, 220 bp; Pv, 120 bp).

**DISCUSSION**

Low parasitemic condition in malaria remains a diagnostic challenge. Correct diagnosis of such conditions is highly essential as they are the mainstay of malaria transmission in community. Highly sensitive diagnostic tools are required for the detection of such conditions. Hz pigment remains to be the pathognomonic biomarker of malaria pathogenesis [5]. During the intraerythrocytic cycle, *Plasmodium* synthesizes Hz pigment by detoxifying the hemoglobin of RBCs. These pigments are formed in all parasitic stages after the early trophozoite stage and persist till the end. During the pathogenesis, the parasites (alone/within the RBC) rupture to release Hz pig-
ment into the peripheral circulation. Various phagocytic cells such as monocytes, macrophages, and lastly neutrophils engulf the malaria parasites and Hz pigments. The phagocytic cells are unable to digest these pigments, as they are acid resistant. They can be easily visible under a microscope. They persist within the phagocytic cell till the cells circulate in the blood (life span ranges from 72-296 hr) and finally get deposited within reticuloendothelial cells and various organs [12].

Detection of hemozoin pigment within the phagocytic cell in blood always indicates a recent malarial infection [11]. As a pilot project, we undertook the study to test our hypothesis that detection of hemozoin pigment by QBC of peripheral blood is an alternative diagnostic modality for malaria with low parasitemia by including cases which were diagnosed as positive by RDT but were negative by microscopy of thin smear of peripheral blood. In our study, the majority of samples (defined as low parasitic) contained only Hz pigments without presence of any malarial parasite. All the above samples were negative by Giemsa stain and only 70% of total samples were detected by PCR. Considering PCR as a gold standard for malarial diagnosis, Hz pigment detection shows a sensitivity of 85% (18/21) and a specificity of 22% (2/9). (Table 2) Hence, it proves that Hz pigment has a certain role in the diagnosis of low parasitic conditions. The decreased specificity could be as a result of remaining Hz particle after the treatment.

QBC is a better tool for detection of Hz pigment in comparison to PCR. In the present study, the samples were negative for malarial parasites by QBC. Hence, these samples were unlikely to be positive for malaria by a thick smear since the sensitivity of QBC and thick smears for parasite detection are almost similar. Moreover, the focus of our study was detection of intracellular Hz pigment in QBC particularly in the lymphocyte-monocyte layer of the buffy coat. Since Hz pigment is concentrated in the lympho-monolayer in QBC it is a more sensitive method of pigment detection than the thick smear. Hence, thick smear examination was not done in our study. We also found that the overall positivity of QBC was 83% in comparison to 70% by nested PCR. Some of the patients had received antimalarials outside the hospital before the start of investigations. The antimalarials were proved to be inhibitory to PCR. This may be the reason of decreased sensitivity by PCR.

Hz pigment is a crystalline product having a very typical morphology [12]. It may be linear or non-linear, coarse or fine, and does not require any chemical stain for its detection. Hz-based sensitive tools are highly desirable for the detection of low parasitic conditions. QBC is a rapid, cost-effective, and highly sensitive tool for Hz pigment detection [8]. QBC is a much sensitive and specific technique in comparison to others because of 2 reasons. Firstly, the sample is centrifuged in a very high revolution speed so that all the parasites, infected RBCs, and WBCs are concentrated in the buffy coat region. Hence, a very low number of parasites can also be detected and increases its sensitivity. Secondly, the tube is coated with fluorescent dye (acridine orange), which increases the level of detection in comparison to conventional microscopy. It takes less time for sample preparation and interpretation (approximately 30 min). In addition to the detection of malaria parasites and Hz pigment, we can characterize the pigment morphology. Neutrophils containing Hz pigment have already been proved as a marker for severe malaria and can be easily detected by the QBC assay as shown in Fig. 1. In the present study, we found that 6 out of 30 samples were positive for neutrophils containing Hz pigment, and 3 out of 5 were having various complications. Samples with neutrophils containing pigment can be alerted from the beginning, and adequate care can be taken to reduce the morbidity and mortality. Other advantages using QBC assay include semi-quantitative detection of packed cell volume or hemoglobin level. This assay can also detect other hemoparasites (such as Babesia, microfilaria, trypanosome, and Donovan bodies of Leishmania donovani).

Molecular methods are proven to have highest sensitivity and specificity among all the diagnostic modalities available for malaria. However, in the current study, few cases were found positive by ICT and QBC but negative by PCR. In a retrospective data analysis, many of these patients were found to receive 1 or 2 doses of antimalarials prior to the sampling. Moreover, few samples exhibited only Hz pigments in QBC, and positive by both ICT and PCR. Hence, the detection of Hz pigments in low parasitic conditions can be an adjunct for malarial diagnosis.

The limitation of this study was that we did not examine the role of Hz pigment detection by QBC in samples which were both negative by microscopy and RDTs. This was a pilot study by which we wanted to validate the utility of QBC for Hz detection for which we needed a substantial number of samples, which were negative for parasitic forms by microscopy. The majority of such samples received during the study period were RDT positive, and hence we used such samples to see
whether Hz detection by QBC serves any purpose.

Low parasitemic conditions in malaria need to be evaluated carefully. Hz pigment is a very good marker for malarial infection. Its use in the detection of low parasitemic condition will be highly useful. It not only signifies the recent malarial infection but also puts light on severity of the disease. In the present study, we found that QBC is a rapid, highly sensitive, and cost-effective method to detect Hz pigment in comparison to PCR in low parasitemic conditions.

CONFLICT OF INTEREST

The authors declare that they have no competing interest.

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