Performance of OptiMAL® in the Diagnosis of *Plasmodium vivax* and *Plasmodium falciparum* Infections in a Malaria Referral Center in Colombia

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Alternative, non-microscopic methods for the diagnosis of malaria have recently become available. Among these, rapid dipstick methods stand out. One such test, OptiMAL®, is based on the immunochromatographic detection of *Plasmodium* lactate dehydrogenase (pLDH) and has the capacity to detect and distinguish infections caused by *P. falciparum* and *Plasmodium* sp. This capacity is particularly important in countries where different species of *Plasmodium* co-exist. In this study we evaluated the performance of OptiMAL® in an urban referral center for malaria diagnosis. Two sets of patients were included: one (n = 112) having predetermined infections with *P. falciparum* or *P. vivax* and individuals with negative blood smears; and another consisting of all eligible consecutive patients (n = 80) consulting for diagnosis at the referral center during one month. The overall diagnostic efficiency of OptiMAL® for both sets of patients was 96.9%. Efficiency was higher for *P. vivax* (98.1%) than for *P. falciparum* (94.9%). These results corroborate the diagnostic utility of OptiMAL® in settings where *P. vivax* and *P. falciparum* co-exist and support its implementation where microscopic diagnosis is unavailable and in circumstances that exceed the capacity of the local microscopic diagnosis facility.

Key words: rapid diagnosis - malaria - OptiMAL® - dipstick - Colombia

Rapid and accurate diagnosis is critical to the effective management of malaria and is one of the principal interventions in the control of this disease (WHO 1993). Better utilization of microscopy, the standard method of diagnosis, and the development of alternative diagnostic methods, could substantially improve the control of malaria (WHO 2000).

Microscopic diagnosis is highly sensitive and specific. This test allows discrimination of asexual and sexual parasite stages, differentiation of the four species that cause human malaria, and quantification of parasitemia. However, requirements such as electricity, a functioning microscope, highly trained technicians, and strict quality control, result in this method being unavailable in remote endemic regions.

In places where microscopic diagnosis is not available, the diagnosis of malaria is based on the clinical presentation of illness. This practice is unreliable because of the non-specific nature of malaria symptoms, which can readily be confused with those of other tropical febrile illnesses (WHO 2000). Reliance on clinical diagnosis results in inappropriate treatment of patients not having malaria with antimalarial drugs, self-medication and incomplete treatment upon improvement of symptoms. All of these practices could favor the development of drug resistance.

Notwithstanding the obvious virtues of microscopy, alternative methods of malaria diagnosis that overcome its limitations are needed. In response to this need, several new diagnostic methods have been developed including serological tests, PCR (polymerase chain reaction), and fluorescent microscopy like QBC® (quantitative buffy coat) (Makler et al. 1998). Although these technologies have provided alternatives for some diagnostic circumstances, the requirement of sophisticated equipment and training has prevented the displacement of microscopy as the routine diagnostic method.

More recently, immunochromatographic tests, also known as rapid diagnostic tests of malaria, have been developed. The histidine rich protein II (HRPII) (Shiff et al. 1993) and *Plasmodium* lactate dehydrogenase (pLDH) (Makler & Hinrichs 1993), are the most common antigens used. OptiMAL® dipstick test (Flow Inc., Portland, OR) is able to differentiate between *P. falciparum* and non *P. falciparum* malaria based on the ability of monoclonal antibodies to react with *P. falciparum* and non *P. falciparum* specific epitopes of pLDH. In countries where different species of *Plasmodium* co-exist, the ability to detect *Plasmodium* sp. infections is an advantage of OptiMAL® over other currently available rapid diagnostic tests that only detect *P. falciparum*.

In Colombia during the past five years, approximately 55% of malaria cases were caused by *P. vivax*, 45% by *P. falciparum* and a low proportion (less than 0.1% overall), by *P. malariae* (Ministerio de Salud 1999, Vergara et al. 2001). Treatment regimens for *P. falciparum* differ from
those for other species of Plasmodium because of the high frequency of resistance to chloroquine (Osorio et al. 1999) the absence of hypnozoite liver stages and the risk of lethal complications of infection. Therefore, diagnostic tests that both detect and distinguish P. falciparum and Plasmodium sp. infection are essential to the appropriate treatment and management of malaria in Colombia.

In this study we have evaluated the rapid diagnostic test OptiMAL® in a referral center under the conditions of a public malaria diagnostic service. Sensitivity, specificity, predictive values and efficiency in comparison with conventional microscopy as “gold standard” were determined. Additionally, PCR was used as a second gold standard based on its greater sensitivity in cases of low parasitemia (Snounou et al. 1993).

MATERIALS AND METHODS

Study site - This study was conducted in Cali, the second largest city in Colombia, within the Malaria Diagnostic Service of the Universidad del Valle and the Secretaria de Salud Publica Municipal. Although malaria is not transmitted in the city of Cali, this urban center is located near endemic areas and serves as referral center for malaria diagnosis in South Western Colombia.

Patients and samples - Patients consulting the Malaria Diagnostic Service were invited to participate in the study if they had fever and a history of traveling to an endemic area. Informed written consent was obtained from all patients included. A first study group, designated Group I, was composed to conduct an exploration of the test performance in malaria patients with P. falciparum and P. vivax, and in controls having negative thick smears. The second group, Group II, consisted of consecutive patients consulting the Malaria Diagnostic Service during a period of one month. The latter group was enrolled in order to evaluate the performance of the diagnostic for the case mix (including a high proportion of non-malarial febrile illness) typically consulting this service. Thick smear and OptiMAL® were performed for both groups. In addition, PCR was performed for Group II. A fingerprick blood sample was obtained from each patient and was used to prepare thick and thin smears and to fill a heparinized capillary tube used for the OptiMAL® test. Additionally, for Group II blood samples were collected on Whatman filter paper #3 for DNA extraction.

Diagnosis by thick/thin smear - Thick smears were treated with methylene blue to remove hemoglobin and thereby facilitate reading of the slides, then stained with 1% Giemsa in phosphate buffer (pH 7.2). Thin smears were fixed in methanol and stained with 2% Giemsa and were evaluated only if parasite species confirmation was needed. All the smears were evaluated using light microscopy at 1000X magnification by two trained medical technologists with more than a year of experience in reading malaria slides. A thick smear was considered negative if no parasites were seen in at least 200 fields. In positive smears parasitemia was calculated as follows:

\[
\frac{\text{Number of parasites}}{100 \text{ leukocytes}} \times 8000 \text{ leukocytes/µl} = \text{parasites/µl}
\]

Diagnosis using OptiMAL® - The OptiMAL® test (Flow Inc, Portland, OR) employs immunochromatography in a dipstick that contains monoclonal antibodies (Mabs) directed against the enzyme pLDH. There are two Mabs: one is specific for P. falciparum and is located at the bottom of the dipstick and the other is specific for Plasmodium sp. and is located in the middle of the dipstick. A polyclonal goat anti-mouse antibody located at the top of the dipstick functions as a reaction control by capturing the mouse anti-pLDH monoclonal colloidal gold conjugate. The whole process was conducted following the manufacturer’s instructions. In accordance with the species prevalent in Colombia, positive results to Plasmodium sp. were considered to be P. vivax in our analysis.

PCR DNA extraction - DNA was extracted from the blood spots impregnated on filter paper using a modified Saponin-Chelex protocol (Giraldo et al. 1998). The blood spots were cut up into small pieces, suspended in 1% Saponin in distilled water (Sigma, St. Louis, MO) and incubated on ice to produce lysis of the cells, followed by a second incubation at 56°C and 100°C with 5% Chelex-100 resin (BioRad, Hercules, CA) that captures and releases the DNA in response to the change of temperature.

Nested diagnostic PCR - A nested diagnostic PCR was used as described previously (Snounou et al. 1993). This PCR is based on the amplification of the sequences of the gene that code for the 18S subunit of the four malarial parasites. Ten microliters of the extracted DNA were used as template for the first amplification reaction. Five microliters of the amplification product of the first reaction were used for the second amplification. All the amplifications were performed in a PTC-100 thermocycler (MJ Research Inc.) and the DNA amplified by PCR was electrophoresed in 2% agarose gels and visualized on a UV transilluminator after staining with ethidium bromide.

Data analysis - The sensitivity, specificity and predictive values of the OptiMAL® test were calculated using the thick smear as the gold standard. For Group II both the thick smear and PCR were independently used as gold standards. Overall, diagnostic efficiency of OptiMAL® for malaria and for each species of Plasmodium was determined based on the combined results of Group I and II, as described by Weigle et al. (1993) using the formula:

\[
\frac{1 - \text{False negatives + False positives}}{\text{Total samples}} \times 100
\]

The statistical analyses were conducted with SPSS 7.5 for Windows® and Epi-Info 6.04.

RESULTS

Diagnosis with thick smear and OptiMAL® in Group I - A total of 112 patients were included in Group I. Eighty-seven patients presented a positive thick smear: 55 with P. vivax, 28 with P. falciparum, 3 with mixed malaria (P. vivax and P. falciparum), 1 with P. malariae; while 25 patients were negative for malaria. Sensitivity was higher for P. vivax than P. falciparum, whereas specificity was comparable for both species (Table I).
Among cases of *P. falciparum* infection, there were three false negative tests in which OptiMAL® was positive but failed to correctly diagnose the species. In two of these cases the level of parasitemia was low (80 and 480 parasites/µl) and in one the parasitemia consisted only of sexual forms. Likewise, for *P. vivax* cases the only false negative case presented scarce parasitemia (80 parasites/µl). A false positive result for *P. vivax* occurred in a patient with a history of travel to an endemic area but negative thick smear on two occasions.

In the three cases of mixed *P. falciparum*/*P. vivax* malaria, OptiMAL® diagnosed only *P. falciparum* in two and *P. vivax* in one. In the single case of *P. malariae*, reactivity occurred with the Mab for *Plasmodium* sp., and therefore it was considered a *P. vivax* infection in our analysis.

**Diagnosis using thick smear, OptiMAL® and PCR in Group II** - During the sampling period (one month), 80 patients were included in this group. Fifteen samples were positive for *P. vivax*, four for *P. falciparum*, one had mixed infection and the remaining 60 samples were negative for malaria by thick smear. All the samples that were positive by thick smear in this group were also positive by OptiMAL® for the corresponding species. Sensitivity and specificity was high for both species but the low prevalence of malaria infection resulted in wide confidence intervals and false positive results for *P. falciparum* affected the corresponding positive predictive value (Table II).

In this group the performance of OptiMAL® was also analyzed using PCR as the reference standard. In this analysis the sensitivity of OptiMAL® for both species was lower; 83.3% for *P. falciparum* and 84.2% for *P. vivax* versus 100% using the conventional gold standard. The specificity was unaffected (data not shown). For *P. falciparum* the single false negative sample came from a patient in post-treatment control. For *P. vivax* there were three false negative cases and two were from patients with history of previous episodes of malaria.

The predetermined diagnosis of patients included in Group I and the small number of patients with malaria in Group II (n = 20) limited the generalizability of the results that were obtained. Therefore, we conducted an analysis of Group I and Group II combined to obtain a more representative assessment of the diagnostic performance of OptiMAL® compared with thick smear (Table III). The results were similar to those obtained for Group I, but confidence intervals were narrower. Overall, diagnostic efficiency for malaria was 96.9%. The diagnostic efficiency for *P. vivax* was higher (98.1%) than for *P. falciparum* (94.9%).

### TABLE I

Sensitivity, specificity and positive (PPV) and negative (NPV) predictive values of OptiMAL® for the diagnosis of *Plasmodium falciparum* or *P. vivax* infections in Group I

<table>
<thead>
<tr>
<th></th>
<th><em>P. falciparum</em></th>
<th><em>P. vivax</em></th>
</tr>
</thead>
<tbody>
<tr>
<td>% sensitivity (95%IC)</td>
<td>89.3 (70.6-97.2)</td>
<td>98.2 (89.0-99.9)</td>
</tr>
<tr>
<td>% specificity (95%IC)</td>
<td>100 (84.3-100)</td>
<td>96.0 (77.7-99.8)</td>
</tr>
<tr>
<td>% PPV (95%IC)</td>
<td>100 (83.4-100)</td>
<td>98.2 (89.0-99.9)</td>
</tr>
<tr>
<td>% NPV (95%IC)</td>
<td>89.3 (70.6-97.2)</td>
<td>96.0 (77.7-99.8)</td>
</tr>
</tbody>
</table>

*a:* composed to conduct an exploration, a total of 112 patients were included.

### TABLE II

Sensitivity, specificity and positive (PPV) and negative (NPV) predictive values of OptiMAL® for the diagnosis of *Plasmodium falciparum* or *P. vivax* infections in Group II

<table>
<thead>
<tr>
<th></th>
<th><em>P. falciparum</em></th>
<th><em>P. vivax</em></th>
</tr>
</thead>
<tbody>
<tr>
<td>% sensitivity (95%IC)</td>
<td>100 (39.6-100)</td>
<td>100 (75.9-100)</td>
</tr>
<tr>
<td>% specificity (95%IC)</td>
<td>95.0 (85.2-98.7)</td>
<td>98.0 (89.9-99.9)</td>
</tr>
<tr>
<td>% PPV (95%IC)</td>
<td>57.1 (20.0-88.2)</td>
<td>94.1 (69.2-99.7)</td>
</tr>
<tr>
<td>% NPV (95%IC)</td>
<td>100 (92.1-100)</td>
<td>100 (92.4-100)</td>
</tr>
</tbody>
</table>

*a:* consecutive patients consulting the Malaria Diagnostic Service during a period of one month, a total of 80 patients were included.

### TABLE III

Sensitivity, specificity and positive (PPV) and negative (NPV) predictive values and efficiency of OptiMAL® for the diagnosis of *Plasmodium falciparum* or *P. vivax* infections in Groups I and II combined

<table>
<thead>
<tr>
<th></th>
<th><em>P. falciparum</em></th>
<th><em>P. vivax</em></th>
<th>Malaria overall</th>
</tr>
</thead>
<tbody>
<tr>
<td>% sensitivity (95%IC)</td>
<td>90.6 (73.8-97.5)</td>
<td>98.6 (91.3-99.9)</td>
<td>99.1 (94.2-100)</td>
</tr>
<tr>
<td>% specificity (95%IC)</td>
<td>96.5 (89.3-99.1)</td>
<td>97.6 (91.0-99.6)</td>
<td>94.1 (86.2-97.8)</td>
</tr>
<tr>
<td>% PPV (95%IC)</td>
<td>90.6 (73.8-97.5)</td>
<td>97.2 (89.4-99.5)</td>
<td>95.5 (89.5-98.3)</td>
</tr>
<tr>
<td>% NPV (95%IC)</td>
<td>96.5 (89.3-99.1)</td>
<td>98.8 (92.6-99.9)</td>
<td>98.8 (92.4-99.9)</td>
</tr>
<tr>
<td>% efficiency (95%IC)</td>
<td>94.9 (90.9-98.9)</td>
<td>98.1 (96.0-100)</td>
<td>96.9 (94.4-99.3)</td>
</tr>
</tbody>
</table>
DISCUSSION

In this study the OptiMAL® test efficiently diagnosed malaria caused by either *P. vivax* or *P. falciparum*. However, the sensitivity and efficiency of the test was lower for *P. falciparum* infections. This lower efficiency of OptiMAL® for *P. falciparum* detection was due to false positive tests in a small number of cases. Studies conducted in settings having a high prevalence of *P. falciparum* have shown OptiMAL® to be highly sensitive and specific for infections having parasitemias > 500 parasites/µl (Hunt-Cooke et al. 1999). On the other hand, disparity in the sensitivity for *P. vivax* (94%) and *P. falciparum* (88%) infections has also been reported by Palmer et al. (1998) in a study conducted in Honduras. Characteristics of transmission such as intensity, endemicity of multiple species of *Plasmodium*, acquired resistance in the exposed population and treatment, can individually and collectively influence the results of this and other diagnostic tests for malaria. Hence, these variables need to be considered in the interpretation of diagnostic tests and management of the corresponding patients.

PCR was also used as reference standard to evaluate the OptiMAL® yielding a lower sensitivity than that obtained with thick smear. PCR detected more positive cases than thick smear or OptiMAL® because it detects the parasite DNA, which could be present in either dead or live parasites, whereas OptiMAL® detects an enzyme produced by live parasites. The fact that pLDH is present only when there are living parasites in the sample is especially relevant for post-treatment evaluations (Makler et al. 1998).

Although there were only four false negative cases in this study, these presented some characteristics that could influence OptiMAL® sensitivity including low parasitemia, gametocytoma without asexual forms and the apparent effects of treatment on parasite metabolism including pLDH. The presence of low parasitemias, especially below the detection level that has been established by the manufacturer as 100-200 parasites/µl, was associated with false negative results in this study. Gametocytes produce pLDH, but since gametocyte parasitemias are usually low, they are not detected efficiently by OptiMAL® (Moody et al. 2000). Two of the false negative cases in this study were from patients that had received previous treatment. OptiMAL® sensitivity has been shown to be influenced by the decrease of parasitemia and pLDH levels after treatment (Moody et al. 2000).

In all false negative cases of *P. falciparum*, OptiMAL® yielded a positive result with the monoclonal antibody for *Plasmodium* sp., which was interpreted as *P. vivax* infection. In cases of low or gametocyte parasitemias, *P. falciparum* pLDH may react only with the genus-specific antibody as has been described (Hunt-Cooke et al. 1998). This fact should be taken into consideration in regions where *P. falciparum* and *P. vivax* coexist since treatment regimens are different for these species.

During this study only one patient presented *P. malariae* infection, which by OptiMAL® was positive with the Mab for *Plasmodium* sp. In previous studies it has been observed that OptiMAL® is less efficient in detecting *P. malariae* and *P. ovale* (Srinivasan et al. 2000). In Colombia *P. malariae* is becoming an increasingly important species, especially in some regions of the Pacific Coast where up to 10% of infections may be caused by this species (Vergara et al. 2001). Overall 0.1% malaria infections in Colombia are due to *P. malariae* (Ministerio de Salud 1999).

Mixed infections account for 1% of malaria cases in Colombia (Ministerio de Salud 1999). In cases of mixed *P. falciparum*/*P. vivax* malaria, OptiMAL® presents reactions with both the *P. falciparum* and *Plasmodium* sp. Mabs, a pattern which corresponds to diagnosis of *P. falciparum* infection. Therefore, the actual format of OptiMAL® favors the diagnosis of *P. falciparum* but masks co-infection by *P. vivax* or other species of *Plasmodium* and hence constitutes a limitation of the current configuration of the test in settings of co-endemicity (Srinivasan et al. 2000).

Our results, which are similar to those obtained in previous studies in other endemic and non-endemic regions, indicate that OptiMAL® would be useful and should be implemented in areas of transmission and circumstances not covered by the traditional microscopic diagnosis of malaria. These situations include epidemic outbreaks, occupationally exposed groups, and displaced populations moving from or toward endemic areas and therefore represent a risk of spreading malaria. All of these circumstances are common in Colombia and in other developing countries. However, some issues of rapid diagnosis of malaria by OptiMAL® and other tests are still unresolved. The applicability of such tests in assessing treatment efficacy, and the cost-effectiveness of the implementation of these alternative methods have neither been extensively nor definitively investigated. Furthermore, efforts to develop a more sensitive test that can detect all the four species of *Plasmodium* and quantify the parasite burden should continue.

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This article is dedicated to the memory of Luis Ernesto Giraldo, MD PhD, who was the mentor of this research.

REFERENCES


