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Nsobya, Sam L Walakira, Andrew Namirembe, Elizabeth [et al.](https://escholarship.org/uc/item/90j8x90x#author)

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Deletions of *pfhrp2* and *pfhrp3* genes were uncommon in rapid diagnostic test-negative *Plasmodium falciparum* isolates from Uganda

Sam L. Nsobya^{1,2*}, Andrew Walakira¹, Elizabeth Namirembe¹, Moses Kiggundu¹, Joaniter I. Nankabirwa^{1,3}, Emmanuel Ruhamyankaka¹, Emmanuel Arinaitwe¹, Melissa D. Conrad⁴, Moses R. Kamya^{1,3}, Grant Dorsey⁴ and Philip J. Rosenthal⁴

Abstract

Background: Rapid diagnostic tests (RDTs) play a key role in malaria case management. The most widely used RDT identifes *Plasmodium falciparum* based on immunochromatographic recognition of *P. falciparum* histidine-rich protein 2 (PfHRP2). Deletion of the paralogous *pfhrp2* and *pfhrp3* genes leads to false-negative PfHRP2-based RDTs, and has been reported in *P. falciparum* infections from South America and Africa. However, identifcation of *pfhrp2/pfhrp3* deletions has usually been based only on failure to amplify these genes using PCR, without confrmation based on PfHRP2 protein expression, and understanding of the true prevalence of deletions is incomplete.

Methods: Deletions of *pfhrp2*/*pfhrp3* in blood samples were investigated from cross-sectional surveys in 2012-13 in three regions of varied malaria transmission intensity in Uganda. Samples with positive Giemsa-stained thick blood smears, but negative PfHRP2-based RDTs were evaluated by PCR amplifcation of conserved subunit ribosomal DNA for *Plasmodium* species, PCR amplifcation of *pfhrp2* and *pfhrp3* genes to identify deletions, and bead-based immunoassays for expression of PfHRP2.

Results: Of 3516 samples collected in cross-sectional surveys, 1493 (42.5%) had positive blood smears, of which 96 (6.4%) were RDT-negative. Of these 96 RDT-negative samples, *P. falciparum* DNA was identifed by PCR in 56 (58%) and only non-falciparum plasmodial DNA in 40 (42%). In all 56 *P. falciparum*-positive samples there was a failure to amplify *pfhrp2* or *pfhrp3*: in 25 (45%) *pfhrp2* was not amplifed, in 39 (70%) *pfhrp3* was not amplifed, and in 19 (34%) neither gene was amplifed. For the 39 *P. falciparum*-positive, RDT-negative samples available for analysis of protein expression, PfHRP2 was not identifed by immunoassay in only four samples (10.3%); these four samples all had failure to amplify both *pfhrp2* and *pfhrp3* by PCR. Thus, only four of 96 (4.2%) smear-positive, RDT-negative samples had *P. falciparum* infections with deletion of *pfhrp2* and *pfhrp3* confrmed by failure to amplify the genes by PCR and lack of expression of PfHRP2 demonstrated by immunoassay.

Conclusion: False negative RDTs were uncommon. Deletions in *pfhrp2* and *pfhrp3* explained some of these false negatives, but most false negatives were not due to deletion of the *pfhrp2* and *pfhrp3* genes.

Keywords: *pfhrp2*, *pfhrp3*, *Plasmodium falciparum*, HRP2, Rapid diagnostic test

Background

*Correspondence: samnsobya@yahoo.co.uk

¹ Infectious Diseases Research Collaboration, Kampala, Uganda Full list of author information is available at the end of the article

Malaria is among the leading health threats in Africa. Sub-Saharan Africa carries the largest malaria burden in the world, with an estimated 213 million cases and 381,000 deaths, primarily from *Plasmodium falciparum*,

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in 2018 [[1\]](#page-6-0). The World Health Organization (WHO) recommends that all cases of suspected malaria should have the diagnosis confrmed by either microscopy or malaria rapid diagnostic test (RDT) before treatment. The WHO gold standard for malaria diagnosis remains microscopic examination of Giemsa-stained thick and thin blood films $[1]$ $[1]$. This method requires an experienced reader to provide accurate diagnosis. RDTs offer a number of benefts over microscopy, as they are less labour-intensive, do not require electricity, require less sophisticated laboratory personnel, and specifcally detect *P. falciparum*.

Over 259 million malaria RDTs have been used in sub-Saharan Africa annually since 2018 [[1\]](#page-6-0). Malaria RDTs can target a number of antigens, including histidine-rich protein 2 (PfHRP2), lactate dehydrogenase, and aldolase [\[2](#page-6-1)]. PfHRP2 is abundantly expressed by erythrocytic stages of *P. falciparum*, and it is the antigen most commonly used in malaria RDTs. RDTs that detect lactate dehydrogenase or aldolase have the advantage of also detecting nonfalciparum species, but these tests are less sensitive and more susceptible to degradation from heat and humidity than are PfHRP2-based RDTs [[3\]](#page-6-2).

Some PfHRP2-based RDTs test positive also in the presence of the paralogous PfHRP3 protein, as the proteins share antigenic epitopes [[4,](#page-6-3) [5\]](#page-6-4). PfHRP2-based RDTs are more sensitive than those directed against other antigens because of higher levels of circulating PfHRP2 and PfHRP3, superior antigen–antibody binding kinetics, and amplifcation of secondary antibody binding due to repeated epitopes in the HRPs [\[6](#page-6-5), [7](#page-6-6)]. PfHRP2-based RDTs have sensitivity and specifcity for *P. falciparum* diagnosis similar to those for expert examination of Giemsa-stained thick blood smears [[8\]](#page-6-7).

An important limitation of PfHRP2-based RDTs is potential false-negative results due to deletion of the *pfhrp2* and *pfhrp3* genes [\[9](#page-6-8)[–11\]](#page-6-9). Existence of *P. falciparum* lacking the *pfhrp2/pfhrp3* genes poses a major risk to malaria control programmes because, if these parasites circulate widely, infected individuals may not be diagnosed and treated, and thus may serve as parasite reservoirs enabling continued transmission. Parasites with *pfhrp2/3* deletions were frst reported in Peru about 10 years ago, after investigation of microscopy-positive/ RDT-negative infections, with subsequent reports from other countries in the America. [\[5](#page-6-4), [12](#page-6-10)]. More recently, *pfhrp2/3* deletions have been described in *P. falciparum* from parts of Africa, most notably Eritrea, where prevalence is reported to be much higher than in other areas for unknown reasons [\[13](#page-6-11)]. Data for *pfhrp2/3* deletion prevalence from other parts of Africa are limited, but deletions have been reported in *P. falciparum* from a number of countries in west (Senegal, Mali) [\[9](#page-6-8), [14](#page-6-12)], central (Rwanda, Democratic Republic of Congo) [\[15,](#page-6-13) [16](#page-6-14)],

and east (Kenya, Tanzania, Uganda) Africa [[17,](#page-6-15) [18\]](#page-6-16). A previous study of samples collected in Uganda in 2014-15 noted that seven of 116 (6.0%) microscopy-positive/RDTnegative isolates had deletion of *pfhrp2*; two of these also had deletion of *pfhrp3* [[18](#page-6-16)]. Another study of 300 isolates collected from 48 Ugandan districts in 2017-19 reported that 3.3% had deletion of only *pfhrp2*, 3.0% had deletion of only *pfhrp3*, and 3.3% had deletions of both genes [\[19](#page-6-17)]. Importantly, in many studies reports of *pfhrp2/3* deletions have been based only on failure to amplify one or both of these genes by PCR, and it is not clear if amplifcation failure reliably identifes true deletions. Of note, in samples from Peru [[12](#page-6-10)] and Eritrea [\[13](#page-6-11)] *pfhrp2/3* deletions identifed by PCR were validated by demonstration of lack of expression of PfHRP2 protein $[18]$ $[18]$ $[18]$. The aim of this study was to determine whether deletions in *pfhrp2/ pfhrp3* explain occasional observations of negative PfHRP2-based RDTs in individuals with positive blood smears for malaria parasites. The studied samples were from cross-sectional surveys at three sites in Uganda with positive blood smears but negative malaria RDTs, considering both amplifcation of *pfhrp2* and expression of PfHRP2.

Methods

Source of samples

Cross-sectional surveys, including blood collection, were conducted in 2012 and 2013 in 200 randomly selected households at each of three districts located in diferent epidemiological settings: Kihihi, Kanungu District, a rural area with relatively low transmission intensity in southwestern Uganda; Walukuba, Jinja District, a periurban area with moderate transmission intensity in central Uganda; and Nagongera, Tororo District, a rural area with high transmission intensity in eastern Uganda. Of note, since the time of this study transmission has decreased considerably in Walukuba, likely due its periurban characteristics, and in Nagongera, associated with regular rounds of indoor residual spraying of insecticide since 2014 [[20\]](#page-6-18). Participants for this study were children ages 6 months to 15 years who were full-time residents of the recruited households, selected as previously described $[21]$ $[21]$. This study was approved by the Makerere University Research and Ethics Committee, the Uganda National Council of Science and Technology, and the University of California, San Francisco Committee on Human Research.

Sample collection and malaria diagnosis

Blood was obtained by fnger prick for thick blood smears, malaria RDTs and drying on flter paper for molecular studies. Thick blood smears were stained with 2% Giemsa for 30 min [[22\]](#page-6-20) and read by laboratory

technologists at the feld sites. Parasite densities were calculated by counting the number of asexual parasites per 200 leukocytes (or per 500 leukocytes, if the count was<10 asexual parasites/200 leukocytes), assuming a leukocyte count of 8000/μL. For quality control, all slides were read by a second microscopist, and discrepancies resolved by a third microscopist at the feld sites. In addition, all positive blood smears with parasite densities \leq 20,000/µL based on the field readings were re-read by an expert microscopist in Kampala; confrmation of parasitaemia was required for inclusion in the fnal analyses. RDTs (SD BIOLINE Malaria Ag Pf, a PfHRP2-based test from Standard Diagnostics Inc; Suwon City, Republic of Korea) were performed immediately after blood collection following manufacturer's instructions. Samples for study were all those that were positive for malaria parasites by microscopy but negative by RDT.

Plasmodium species identifcation

DNA was extracted using Chelex100, as previously described [\[23](#page-6-21)]. Species identifcation was performed by nested species-specifc PCR with primers specifc for the 18S small subunit ribosomal DNA gene of all human plasmodial species, as previously described [[24](#page-6-22)]. PCR reactions were performed in 25 μ l containing 1 \times standard *Taq* buffer (New England Biolabs), 200 µM deoxynucleoside triphosphates, 200 µM of each primer, 2 µl of template DNA (from Chelex extraction or the prior cycle of PCR), and 1 unit of *Taq* polymerase (New England Biolabs). All reactions included negative controls (water) and positive controls, obtained from the Biodefense and Emerging Infections Research Resources Repository (BEI U.S).

PCR products were resolved by electrophoresis on 2% agarose gels stained with ethidium bromide and visualized by UV illumination. Sizes of amplicons were identifed based on comparison with standard fragments of known size.

Detection of deletions in *pfhrp2* **and** *pfhrp3* **genes**

To identify deletions, we PCR-amplifed fragments spanning exon 1, the intron, and exon 2 of the *pfhrp2* and *pfhrp3* genes, as previously described [\[5](#page-6-4)]. *P. falciparum* Dd2 strain DNA was a negative control for *pfhrp2* and a positive control for *pfhrp3. P. falciparum* HB3 strain DNA was a negative control for *pfhrp3* and a positive control for *pfhrp2*. All PCR reactions were performed in duplicate. PCR products were separated and visualized on 2% agarose gels. In the event of discordant replicates, reactions were repeated, and the result recorded was that seen in multiple assays. Deletions were identifed by the absence of amplifcation of *pfhrp2/3* in the setting of successful amplifcation of ribosomal DNA in the sample and amplifcation of *pfhrp2/3* in positive control DNA.

Multiplicity of infection (MOI)

MOI, the number of diferent parasite genotypes coexisting within a host, is a metric of transmission dynamics [\[23](#page-6-21), [24\]](#page-6-22). To determine MOI, the *3D7* and *FC27* alleles of the merozoite surface protein-2 (*msp2)* gene, which each have extensive size polymorphism, were amplifed as previously described [\[25](#page-6-23)]. Amplicons were identifed on 2% agarose gels, the size of products was compared to standards on densitometric digitized gel images analysed by GelCompar II software (Applied Maths NV Belgium), and the number of diferently sized amplicons in each sample was determined.

Bead‑based immunoassay for detection of PfHRP2 protein

As the absence of gene amplifcation does not defnitively prove the presence of a gene deletion, we also assessed expression of PfHRP2 in study samples. Unfortunately, adequate material was available for this analysis for only 39 of the 56 RDT-negative *P. falciparum* samples. For these assays recombinant PfHRP2 (Microcoat Biotechnologie GmbH, Bernried am Starnberger See, Germany) was used as a positive control and blood from persons not infected with malaria as a negative control. PfHRP2 levels was quantifed using a bead-based immunoassay with a MAGPIX instrument (Luminex Corp., Austin, TX), as previously described [[26–](#page-6-24)[28](#page-6-25)]. Briefly, the beadbased HRP2 immunoassay, which relies on antigen capture, is capable of detecting PfHRP2 at sub-picogram levels, allowing fast processing and screening of large numbers of samples. As in prior studies, the cut-off for a positive PfHRP2 antigen result was the mean plus three standard deviations based on a panel of 92 antigen negative blood samples [[28\]](#page-6-25).

Results

Study samples

Of 3516 samples collected in cross-sectional surveys at three sites in Uganda, 1493 (42%) were positive for malaria parasites by Giemsa-stained thick smear (Fig. 1). Of the 1493 smear-positive samples, 96 (6.4%) were negative by PfHRP2-based RDT. These 96 samples were further investigated.

Amplifcation of plasmodial DNA in RDT‑negative samples

Of the 96 microscopy-positive/RDT-negative samples, 56 (58%) had *P. falciparum* ribosomal DNA amplifed, and in 40 (42%) only non-falciparum plasmodial DNA was amplifed. In these samples the species identifed was *Plasmodium vivax* in 12 (30%), *Plasmodium ovale* in 10 (25%), and *Plasmodium malariae* in 18 (45%). The range and SD for parasite densities for *P. falciparum* positive/ RDT-negative samples were $48 - 3400 \ (\pm 660)$ parasites/ µl. The multiplicity of infection for *P. falciparum* positive/ RDT-negative samples was low at all three sites (mean 1.5; Table [1\)](#page-4-0).

Amplifcation of *pfhrp2* **and** *pfhrp3* **in** *P. falciparum* **positive/RDT‑negative samples**

To analyse the 56 *P. falciparum* positive/RDT-negative samples for potential deletions in the *pfhrp2* and *pfhrp3* genes, these genes were amplifed. For all 56 samples there was a failure to amplify *pfhrp2* or *pfhrp3*. Of these samples, in 25 (45%) *pfhrp2* was not amplifed, in 39 (70%) *pfhrp3* was not amplifed, and in 19 (34%) neither gene was amplifed.

Immunoassay for PfHRP2 protein expression

Of the 56 *P. falciparum* positive/RDT-negative samples, 39 had adequate remaining material for additional study. These 39 samples were evaluated for expression of PfHRP2 by bead-based immunoassay. With this assay, four of the 39 samples (10.3%) had no detectable PfHRP2 antigen (Table [2](#page-4-1)). These four samples all had failure to amplify both *pfhrp2* and *pfhrp3* by PCR. Overall, in 35/39 (89.7%) samples with failure to detect *pfhrp2* and/ or *pfhrp3* by PCR, PfHRP2 was detected by immunoassay. The mean parasite density for four samples with no PfHRP2 protein expression was 279 parasites/ul, with MOI of 1.8 (Table [2\)](#page-4-1).

Discussion

To explore the basis of false-negative malaria RDTs, the presence of deletions in *pfhrp2/pfhrp3* genes were investigated in Ugandan blood samples that were positive for malaria parasites by blood smear, but negative by PfHRP2-based RDT. To evaluate for potential deletions in RDT-negative samples amplifcation of the *pfhrp2* and *pfhrp3* genes, amplifcation of sub-unit ribosomal DNA (for species identifcation), amplifcation of *msp2* (for MOI determination), and expression of PfHRP2 by immunoassay, were assessed following an established protocol [\[28](#page-6-25)]. Importantly, no samples characterized as *P. falciparum*-negative by PCR demonstrated expression of *pfhrp2* by immunoassay. In the small subset of samples (6.4%) that were microscopy-positive, but RDTnegative, the false negative RDT results were explained by non-falciparum malaria infection in 42%. The identifcation of non-falciparum malaria infections was consistent with our recent identifcation of non-falciparum infections in 8.2% of subjects diagnosed with malaria at 10 sites in Uganda, although most of these were mixed falciparum/non-falciparum infections [[29\]](#page-6-26). For *P. falciparum* samples that were false negative by RDT, only 10.3% had both failure to amplify *pfhrp2/pfhrp3* by PCR and no detection of PfHRP2 by immunoassay, consistent with false negative RDT results caused by absent expression of PfHRP proteins. Thus, the data suggest that in Uganda, *pfhrp2/pfhrp3* gene deletion is present, as described in other recent studies [\[18](#page-6-16), [19\]](#page-6-17), but that absence of PfHRP2 expression was uncommon, and that this absence did not explain most false-negative RDT results.

Table 1 Characteristics of study districts and results of *P. falciparum* **surveillance**

n (%), number of samples; MOI, multiplicity of infection; PD, geometric mean parasite density/µL

Deletions of the *pfhrp2/pfhrp3* genes, leading to lack of expression of PfHRP proteins, has been well documented in *P. falciparum* from a number of regions, most notably parts of South America [\[12](#page-6-10)] and, in Africa, Eritrea [\[13](#page-6-11)]. Lower prevalence of deletions has been reported in *P. falciparum* from many African countries [\[9](#page-6-8), [14–](#page-6-12)[18](#page-6-16)], including Uganda [\[18](#page-6-16), [19\]](#page-6-17). However, methods used in these studies have not been consistent. In some studies, only *pfhrp2*, but not *pfhrp3* has been studied, despite the fact that expression of PfHRP3 may yield a positive PfHRP2 based RDT. In many studies, *pfhrp2/pfhrp3* gene deletion has been documented based only on failure to PCRamplify the genes. However, even with controls demonstrating amplifcation of other *P. falciparum* genes, there is concern that failure to amplify *pfhrp2/pfhrp3* might be due to technical difficulties rather than true deletions. To address this concern, efficient methods are now available to assess expression of PfHRP2 using a bead based immunoassay $[27]$ $[27]$. This technology was utilized to further characterize samples with *pfhrp2/pfhrp3* gene deletions based on PCR results [[5\]](#page-6-4).

Results from the PfHRP2 immunoassay were revealing. Most samples that had apparent deletions of *pfhrp2* or *pfhrp3* based on PCR actually showed expression of PfHRP2 by immunoassay. This result suggests that failure to amplify *pfhrp2/pfhrp3* was, in some cases, due to technical challenges (with identifcation of the false negatives facilitated by the high sensitivity of the immunoassay), rather than true deletions. A less likely possibility is that the *pfhrp2*/3 deletions suggested by PCR were real, and that the immunoassay yielded false positive results. PCR might have failed to amplify *pfhrp2/pfhrp3* due to the presence of enzyme inhibitors in samples, PCR primer mismatch due to mutations that did not afect PfHRP2/3 expression, inadequate quantities of DNA for successful amplifcation, or other technical factors. Consistent with these explanations, our samples were extracted without full purifcation, potentially allowing some PCR inhibitors in reactions. In addition, parasite densities and MOI were low, and samples were extracted after long-term storage on flter paper, all consistent with limited quantities of DNA for reactions. Results suggest that defnitive detection of *pfhrp2/pfhrp3* deletion should incorporate assays for both DNA and protein.

The study had important limitations. First, the study was completed well after collection of samples in 2012- 13, and so ofers limited insight into the current prevalence of the deletion of *pfhrp2/pfhrp3* may have been identified. Third, this was a cross-sectional study of households, with the large majority of subjects asymptomatic, so it offers relatively little insight into the prevalence of *pfhrp2/3* deletions in those presenting with symptomatic malaria. Fourth, the full *pfhrp2/pfhrp3* genes in study samples was not sequenced, thus the mechanisms behind failure to amplify *pfhrp2/pfhrp3* is not known. Additional studies are needed in Africa to determine the cause of false-negative RDTs, and specifcally whether, as is the case in parts of South America and in Eritrea, true deletions of *pfhrp2/3* are responsible for a substantial number of false negative diagnostic assays.

Conclusion

In summary, false negative PfHRP2-based RDTs were uncommon in Uganda, and that false negatives were explained by non-falciparum infections, PCR amplifcation failures, and, only in a small subset of samples, true absence of expression of PfHRP2. Nonetheless, the prevalence of *pfhrp2/pfhrp3* deletions may be increasing in various regions $[13]$ $[13]$. As this phenomenon threatens a primary method of malaria diagnosis in Africa, continued surveillance for *pfhrp2/pfhrp3* deletions across Africa, ideally using multiple experimental methods, is a high priority. Results suggest that defnitive detection of *pfhrp2/pfhrp3* deletion should incorporate assays for both DNA and protein; a multiplex format will facilitate high throughput screening.

Abbreviations

PfHRP2: *P. falciparum* histidine rich protein 2; PfHRP3: *P. falciparum* histidine rich protein 3; *pfhrp2*: *Plasmodium falciparum histidine rich protein 2 gene*; *pfhrp3*: *Plasmodium falciparum histidine rich protein3 gene*; RDT: Rapid diagnostic test; MSP1: *Merozoite surface antigen 1*; PCR: Polymerase chain reaction; WHO: World Health Organization; DNA: Deoxyribonucleic acid; DBS: Dried blood spots; MOI: Multiplicity of infection; PBS: Phosphate buffered saline.

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Authors' contributions

SLN, PJR, and GD designed the study. SLN, AW, EN, ER, EA, JN, MDC, and MRK did the sample analysis. SLN, AW, PJR, GD, and MRK supported the data analysis. SLN, AW, and PJR drafted the manuscript. SLN, AW, MK, EN, JN, ER, EA, MRK, MDC, GD, and PJR reviewed the manuscript. All authors read and approved the fnal manuscript.

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Competing interests

All authors declare that they have no competing interests.

Author details

¹ Infectious Diseases Research Collaboration, Kampala, Uganda. ² Department of Pathology, College of Health Science, Makerere University Kampala,

Kampala, Uganda. 3 Department of Medicine, Makerere University College of Health Sciences, Kampala, Uganda. 4 Department of Medicine, University of California, San Francisco, CA, USA.

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