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RDTs as a source of DNA to study *Plasmodium falciparum* drug resistance in isolates from Senegal and the Comoros Islands

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**Abstract**

**Background:** The World Health Organization has recommended rapid diagnostic tests (RDTs) for use in the diagnosis of suspected malaria cases. In addition to providing quick and accurate detection of *Plasmodium* parasite proteins in the blood, these tests can be used as sources of DNA for further genetic studies. As sulfadoxine-pyrimethamine is used currently for intermittent presumptive treatment of pregnant women in both Senegal and in the Comoros Islands, resistance mutations in the *dhfr* and *dhps* genes were investigated using DNA extracted from RDTs.

**Methods:** The proximal portion of the nitrocellulose membrane of discarded RDTs was used for DNA extraction. This genomic DNA was amplified using HRM to genotype the molecular markers involved in resistance to sulfadoxine-pyrimethamine: *dhfr* (51, 59, 108, and 164) and *dhps* (436, 437, 540, 581, and 613). Additionally, the *msp1* and *msp2* genes were amplified to determine the average clonality between Grande-Comore (Comoros) and Thiès (Senegal).

**Results:** A total of 201 samples were successfully genotyped at all codons by HRM; whereas, in 200 *msp1* and *msp2* genes were successfully amplified and genotyped by nested PCR. A high prevalence of resistance mutations were observed in the *dhfr* gene at codons 51, 59, and 108 as well as in the *dhps* gene at codons 437 and 436. A novel mutant in *dhps* at codon positions 436Y/437A was observed. The *dhfr* I164L codon and *dhps* K540 and A581G codons had 100% wild type alleles in all samples.

**Conclusion:** The utility of field-collected RDTs was validated as a source of DNA for genetic studies interrogating frequencies of drug resistance mutations, using two different molecular methods (PCR and High Resolution Melting). RDTs should not be discarded after use as they can be a valuable source of DNA for genetic and epidemiological studies in sites where filter paper or venous blood collected samples are nonexistent.

**Keywords:** *Plasmodium falciparum*, RDT, DNA source, *dhfr*, *dhps*, Comoros, Senegal

**Background**

Malaria remains a major public health problem as it is responsible for 207 million cases and 627,000 deaths worldwide [1]. Currently, 90% of the deaths attributable to this infection occur in sub-Saharan Africa. To combat this disease, many programmes have implemented control measures such as the distribution of long-lasting insecticide-treated mosquito nets, the use of artemisinin-based combination therapy (ACT) for the treatment of uncomplicated *Plasmodium falciparum* malaria, and the introduction of rapid diagnostics tests (RDTs) in health facilities for malaria diagnosis. Such interventions have...
contributed to a dramatic drop in malaria-attributable deaths worldwide [1].

Widespread RDT use has greatly assisted in the accurate diagnosis of presumed malaria cases in regions where microscopy (or molecular methods) are non-existent. RDTs are based on antigen detection and these antigens, as well as parasite DNA, remain on the discarded RDT after it has been used for patient diagnosis. The widespread use of RDTs for diagnosis creates a valuable resource for population-based studies as RDTs have been shown to be suitable material for genetic studies, as DNA can be extracted from RDTs [2–5].

To validate the utility of field-collected RDTs for typing of drug resistance markers, two markers involved in sulfadoxine-pyrimethamine (SP, Fansidar®) resistance were selected: \( dhfr \) and \( dhps \) genes, as these both drugs are routinely used in the populations of interest (Senegal and Comoros). Until recently, SP was one of the main molecules for the treatment of uncomplicated malaria [6, 7]. However, resistance to SP has evolved quickly [8]. SP is a combination of two drugs that act at two successive stages of the parasite folate pathway. Sulfadoxine inhibits the dihydroyperoate synthetase gene (\( dhps \)), while pyrimethamine inhibits dihydrofolate reductase gene (\( dhfr \)). Point mutations in the \( dhfr \) and \( dhps \) genes confer resistance to pyrimethamine and sulfadoxine, respectively, with a decrease of in vitro susceptibility of \( P. falciparum \) in relation to the number of mutations of each gene [9–13]. Although this molecule has seen a relative decline of its therapeutic efficacy, it is now used in Senegal for the Intermittent Preventive Treatment (IPT) of pregnant women and children against malaria, and in Comoros, for IPT of pregnant women, a policy change that was implemented in 2003 and 2004, respectively.

While many studies measuring the prevalence of \( dhfr \) and \( dhps \) mutations have been conducted previously in Senegal [14–19], few have been performed in Comoros [20–22], largely due to a lack of clinical studies in which venous blood or filter paper are collected to study parasite molecular markers. The objectives of this study were (1) to assess the feasibility of DNA extraction from field-collected RDTs, (2) to study gene mutations \( dhfr \) (S108N, N51I, C59R and I164L) and \( dhps \) (A437G, S436F, K540E, A581G and A613T/S) from samples of Senegal and Comoros with high-resolution melting technology and compare the prevalence between these two countries. \( Msp \) typing was used as a tool to compare the average multiplicity of infection between the two countries.

Methods

Study sites and sample collection

The Comoros islands are located on the southeastern coast of Africa in the Mozambique canal, and represent a country where malaria transmission is high [23]. In Grand-Comore, the largest of the four islands of Comoros, the malaria transmission is meso- to hyperendemic (EIR between 10 and 200) [20]. In contrast, Senegal has Sahalien characteristics marked by a transmission that is seasonal and short. Samples collected for this study were from Thiès, an urban area 70 km east of Dakar, where malaria is hypoendemic [24], with an EIR between 1 and 5.

In Grande-Comore, RDT samples were collected from 2012 to 2013 by the National Malaria Control Programme in the city of Moroni, where malaria is hypoendemic; and in two hospitals Mitsamiouli and Mbeni where malaria is mesoendemic and meso to hyperendemic respectively [20]. In Senegal, RDT samples were collected in 2010 at the Service de Lutte Anti-Parasitaire (SLAP) clinic, in Thiès. Malaria positive RDTs were stored at room temperature with desiccant at both sites.

Rapid diagnostic tests and DNA extraction

In Senegal, SD BIOLINE Malaria rapid test for Pf alone (SD Bioline, 05FK50) was used while in Comoros two different tests were used: Malaria pLDH/HRP2 Combo (Access Bio, PBX-KM30003) and SD BIOLINE Malaria Ag Pf/Pan (SD Bioline, 05FK60). The proximal third of the nitrocellulose membrane was chosen for DNA extraction as previously described [2]. DNA was extracted with the QIAamp DNA Mini kit (Qiagen) according to the manufacturer’s recommendations for filter paper. In total, 204 samples (124 from Grande-Comore and 80 from Thiès) were extracted and genotyped.

Single nucleotide polymorphism analyses by high resolution melting

SNP analysis for mutations in \( dhfr \) (S108N, N51I, C59R and I164L) and \( dhps \) (A437G, S436F, K540E, A581G and A613T/S) genes were conducted by High Resolution Melting (HRM) analysis using a Light Scanner LS-32, according to previously published methodology [18]. Asymmetric PCR was performed with final primer concentrations of 0.25 µM reverse primer; 0.05 µM forward primer; and a final probe concentration of 0.02 µM. The cycling and melting conditions for \( dhps \) amplification were as follows: 95 °C denaturation for 1 min, followed by 55 cycles of (95 °C for 5 s and 66 °C for 30 s) a pre-melt cycle of 5 s each at 95 °C and 37 °C, followed by a melt from 45 to 90 °C at a 0.30 °C/s. For the 437 \( dhps \) assay, cycling and melting conditions were as follows: 95 °C denaturation for 1 min, followed by 55 cycles of (95 °C for 5 s, 66 °C for 30 s, 74 °C for 30 s) a pre-melt cycle of 5 s each at 95 and 37 °C, followed by a melt from 45 to 90 °C at a 0.30 °C/s. The cycling and melting conditions for \( dhfr \) amplification were as follows: 95 °C denaturation
for 1 min, followed by 55 cycles of (95 °C for 5 s, 56 °C for 30 s) a pre-melt cycle of 5 s each at 95 and 37 °C, followed by a melt from 45 to 90 °C at a 0.30 °C/s.

DNA sequencing

PCR sequencing (Sanger sequencing) of the dhps 436/437 amplicon was performed by using the same primers used in the HRM reaction (Forward: GAATGTGG AAATGATAATGAAGGTGCTA and Reverse: CAGG AACACGCTATGACGAAATAATTGTAATACAGG TACTACTAAATCCTCT). Sequencing was performed in both the forward and the reverse direction by Macrogen and contigs were assembled using Lasergene 10.

Allelic typing of Plasmodium falciparum msp1 and msp2

The polymorphic regions (block 2 of MSP1 and block 3 of MSP2) were amplified as previously described [25]. All PCR reactions were carried out in a total volume of 20 µl containing 6 µl Gotaq, 0.5 µM of each primer, and 11 µl reagent grade water. In the first round reaction (nest 1), 1 µl of genomic DNA was added as a template. In the second nested reaction (nest 2), 1 µl of the nest 1 PCR product was used as DNA template.

The cycling conditions for the nest 1 PCR were as follows: initial denaturation at 95 °C for 5 min, followed by 35 cycles of (94 °C for 1 min, 58 °C for 2 min, 72 °C for 2 min), with a final extension cycle of 72 °C for 3 min. The cycling conditions for the nest 2 PCR were as follows: initial denaturation at 95 °C for 5 min, followed by 35 cycles of (94 °C for 1 min, 61 °C for 2 min, 72 °C for 2 min), with a final extension cycle of 72 °C for 3 min. The polymerase chain reactions (PCR) with a final extension cycle of 72 °C for 3 min. Positive controls (3D7 for K1 and IC27 alleles, Dd2 for MAD20 and FC27 alleles, and 7G8 for the RO33 allele) and negative control (reagent grade water alone) were run in each PCR reaction. PCR products were analyzed on 2 % agarose gels stained with ethidium bromide and visualized by UV trans-illumination (Biorad Gel Doc™ XR + System with Image Lab). The size of resulting DNA bands was approximated using Gene Ruler 100 bp DNA ladder (papalinger, Quick Load).

Statistical analyses

Statistical analyses were performed using the Z-test for two population proportions to compare mutant allele prevalence between Comoros and Senegal samples. For all tests, the significance level was α = 0.05.

Results

The prevalence of mutations in dhfr and dhps

Genotyping of dhps and dhfr genes was performed for 201 samples (n = 124 for Grande-Comore and n = 77 for Thiès). For dhfr codons 51/59, only 62 samples from Thiès and 96 samples from Grande-Comore yielded a successful genotyping result; whereas at all other loci, genotyping was successful for all samples (Table 1).

In Thiès, for the dhfr gene, 90 % (54/60) of the samples had the mutant allele at codons dhfr N51C/C59R, and 95.8 % (68/71) had the mutant allele at codon dhfr S108 N. For the dhps gene I33 % (1/75) had the A613T mutant allele and 53.9 % (41/76) of the samples had the A437G mutant allele. At the S436 codon of dhps 12 % (9/77) of the samples had the mutant allele S436F, a recently characterized mutation described previously [18]. For all other codons (namely, dhfr I164, dhps K540, and dhps A581) 100 % of the samples had the wild-type allele. Previously unreported mutations were observed for dhps gene at codons 436/437 by HRM (Fig. 1a) and were confirmed to be S436Y/437A by Sanger sequencing (Fig. 2).

In Grande-Comore, for the dhfr gene, 93.4 % (115/123) had the S108 N mutation, 62.5 % (60/96) of the samples had the N511 mutant allele, and 69.8 % (67/96) of all samples had the C59R mutant allele. For the dhps gene, at position A437, 29.8 % (37/124) of samples had the wild-type allele A437G. For all other codons (namely, dhfr I164, dhps K540, and dhps A581) 100 % of the samples had the wild-type allele.

A statistically significant difference was observed in the prevalence of mutations between the Comoros and Senegal for the dhfr gene codons 51 (0.0001), and 59 (p = 0.003), and dhps gene codons 436 (p = 0.0001) and

### Table 1 Utility of RDTs as a source of DNA for genotyping loci involved in diversity and drug resistance

| RDT Brand             | Number RDT | HRM<sup>a</sup> Positive | HRM<sup>a</sup> Negative | PCR<sup>b</sup> Positive | PCR<sup>b</sup> Negative | Fragment used RDT |
|-----------------------|------------|---------------------------|--------------------------|--------------------------|--------------------------|-------------------|
| SD Bioline malaria Rapid test pf | 80         | 77                        | 3                        | 77                       | 3                        | 1/3 NC            |
| SD Bioline malaria Ag Pf/Pan      | 13         | 13                        | 0                        | 13                       | 0                        | 1/3 NC            |
| Malaria pLDH/HRP2 combo    | 111        | 111                       | 0                        | 110                      | 1                        | 1/3 NC            |
| Total                  | 204        | 201<sup>c</sup>          | 3                        | 200                      | 4                        | –                 |

<sup>a</sup> The HRM method was used for genotyping dhfr and dhps genes

<sup>b</sup> PCR was used to genotype msp1 and msp2

<sup>c</sup> For codons 51/59, only 62 samples from Thiès and 96 samples from Grande-Comore gave a comprehensive genotyping
Fig. 1 HRM peak profiles for wild-type and mutant dhps 436/437 alleles. 

a. The 3D7 peak represents the profile for the wild-type at codon 436 and mutant profile for the 437 codon. The Dd2 peak presents the profile for the two mutant Codons 436/437. The HB3 peak presents the wild-type profile of the two codons. The 436A/437A peaks, represented the mutations found by Daniels et al. in Senegal [18]. The blue peak from sample Th50 RDT represents a new mutant allele.

b. The peaks C03 and C05 represent mixed samples (mutant + wild). The Dd2 peak presents the mutant profile for both codons 436 and 437. The HB3 peak represents the wild-type profile for both codons 436 and 437. The 3D7 peak represents the profile for the wild-type at codon 436 and mutant profile for the 437 codon.
The prevalence of mutations on the \textit{dhfr} gene at codon 51 and 59 and \textit{dhps} gene at codon 436 and at codon 437 were higher in Thiès compared to Grande-Comore.

The single mutation (S108N alone, in the absence of other \textit{dhfr} mutations) was not found in Thiès; whereas, in Grande-Comore it was present at 1.6% (2/122) (Fig. 3). The prevalence of the double mutation (\textit{dhfr} C59R and S108N) was 9% (6/67) in Thiès and 25.4% (31/122) in Grande-Comore (Fig. 3). However, the triple mutation (\textit{dhfr} S108N, N51I, and C59R) was present at similar prevalences in both countries with 44.3% (28/67) in Grande-Comore and 42% (28/67) in Thiès; whereas the quadruple mutation (\textit{dhfr} S108N, N51I, C59R, and \textit{dhps} A437G) was observed at 28.7% (35/122) in Grande-Comore and 49% (33/67) in Thiès (Fig. 3). No quintuple mutations (\textit{dhfr} S108N, N51I, C59R, \textit{dhps} A437G, and K540E) were observed in either country. Statistically, there was no significant difference in the prevalence of single (p = 0.29) and triple mutation (p = 0.74); however there was significance difference in the prevalence of double mutation (p = 0.0063) and the quadruple mutation (p = 0.0048) between the two countries using Z-test for 2 population proportions (Fig. 3).

When comparing the number of mixed alleles (mutant + wild) detected by HRM, overall mixed alleles were more prevalent in Grande-Comore samples than in samples from Thiès. An example of the HRM output for mixed alleles is shown in Fig. 1b. The difference in mixed alleles at \textit{dhfr} position 108 and \textit{dhps} position 437 was not significantly different between the two sites (\textit{dhfr} 108: Grande-Comore 4 mixed samples (3%); Thiès 3 mixed...
(4 %) samples ($p = 0.77$); *dhps* 437: Grande-Comore 5 mixed samples (4 %); Thiès 2 mixed samples (2.6 %) ($p = 0.60$). In contrast, in Grande-Comore a significant difference in the prevalence of mixed alleles at *dhfr* 51/59 was observed. In Grande-Comore, 27 mixed samples (28 %) at *dhfr* codons 51/59 compared to six mixed samples (9 %) in Thiès ($p = 0.003$) were observed.

**Genotyping using msp1 and msp2**

Having validated the RDT-extracted DNA to study markers of drug resistance, the average clonality in both populations was determined by msp typing. With samples from Thiès, a total of 77 samples were positive for *msp1* and *msp2* genes, with 3 samples failing to amplify (Table 1). In total, 36 samples (64.2 %) contained polyclonal infection at least with 2 clones. The average multiplicity of infection (MOI) for all isolates from Thiès was estimated to be 1.57.

With samples from Grande-Comore, a total of 123 samples were positive for *msp1* and *msp2* genotyping, with only a single sample failing amplification (Table 1). In total, 42 samples (43.7 %) had polyclonal infections at least with 2 clones. The average multiplicity of infection (MOI) was 1.47.

**Discussion**

The WHO recommends that all suspected malaria cases be confirmed by RDTs. RDTs facilitate the diagnosis of malaria by providing evidence of the presence of specific *Plasmodium* antigens in human blood without the need for electricity or advanced microscopy training. Previous studies have shown that it is possible to extract *Plasmodium* DNA from RDTs [2, 3, 5]. This study sought to address whether the use of stored, discarded RDTs from field sites can serve as source of DNA for genomic studies in populations in which whole blood or filter paper-based DNA samples are unavailable.

There are many factors that could affect the efficacy of extracting DNA from field-based RDTs. Some of these factors include variable preparation of RDTs (variable blood volumes), and storage conditions. Storage conditions in field sites can differ, introducing the possibility of bacterial or fungal contamination as well as DNA degradation. Another variable to consider is the proportion of human DNA to *Plasmodium* DNA on each RDT as the ability to detect *Plasmodium* genes will depend on the specific concentration of *Plasmodium* DNA. Taking all these field-based realities into consideration, the methodology outlined by Cnops et al. [2] was applied to the field as overall, RDTs represent a cost-efficient method for the conservation, transport, and storage of DNA samples. This study addresses the reality of using RDTs from malaria endemic sites as sources of genetic material for performing molecular studies.

To test the performance of field-RDT extracted DNA for genetic experiments, two genetic methodologies were employed: high resolution melting (HRM) for drug resistance markers (*dhfr* and *dhps*) and *msp* typing by PCR to determine clonality. Differences in malaria transmission reported between Grande-Comore (meso-hyperendemic) and Thiès (hypoendemic) were compared by *msp* typing, and a small, but insignificant difference was observed in the multiplicity of infection for *msp* genes (*msp1* and *msp2*) (MOI = 1.57 in Thiès and MOI = 1.47 in Grande-Comore). This small difference is interesting given the predicted higher endemicity of Grande-Comore compared to Thiès.

HRM is a method that uses post-PCR melting analysis used to identify SNPs in nucleic acid sequences, and has a limit of detection of 100 fg of DNA [18]. In this study, successful genotyping by HRM was performed from DNA extracted from RDTs. An additional methodological advantage of HRM is that known mutations can be classified and novel mutations can be identified, such as the 436Y/437A mutations in *dhps*, which were identified in a sample from Thiès (Fig. 1a). When results are compared from RDT-extracted DNA to previous results obtained with the filter-paper extracted DNA using the same HRM method from the same population (Thiès, Senegal), similar frequencies for *dhfr* mutations at codons 51, 59 and 108 and *dhps* mutations at codon 437 (2012–2013) compared to what was shown by Rebaudet *et al.* [20] (Additional file 1: Table S1), specifically at *dhfr* codon 51 (38.5 % in 2010 compared to 62.5 % in 2012–2013; $p = 0.02$), *dhfr* position 108 (50 % in 2010 compared to 93 % in 2012–2013; $p < 0.05$) and *dhps* codon 437 (4 % in 2010 compared to 29.8 % in 2012–2013;
p = 0.006). A significant increase in the number of mixed alleles at \(dhfr\) C59 and C59R (7.7 % in 2010 compared to 28 % in 2012–2013; p = 0.03) was observed and a trend increasing in the number of mixed alleles at \(dhps\) A437 and A437G (Fig. 1b); albeit non-significant (0 % in 2010 compared to 4 % in 2012–2013, p = 0.3) (Additional file 1: Table S1). However; a decrease in mixed alleles \(dhfr\) S108 and S108 N (15.4 % in 2010 compared to 3.25 % in 2012–2013; p = 0.01) was observed. Of concern, a statistically significant increase in the triple mutation \(dhfr\) N51I/C59R/S108 N (50 % in 2010 compared to 89.6 % in 2012–2013; p < 0.05) was observed in this study. Other studies performed in Comoros in 2004–2006 [26] and 2006 [27] showed similar triple mutation frequencies (53 and 45 %, respectively) to the 50 % reported by Rebaut out this study. Other studies performed in Comoros in 2004–2006 [26] and 2006 [27] showed similar triple mutation frequencies (53 and 45 %, respectively) to the 50 % reported by Rebaut et al. [20], indicating a very recent increase in triple mutation prevalence.

This high prevalence of mutations could potentially be explained by the use of SP for IPT but also by the fact that SP has been used as treatment for uncomplicated malaria [28]. On the other hand, the widespread use of trimethoprimsulfamethoxazole (Cotrimoxazole® or Bactrim®) in Comoros, a drug similar to SP that has the ability to select for resistant mutants [29, 30] could also potentially explain the higher frequencies of mutant alleles between 2010 and 2012–2013.

Drug resistance mutations highly associated with drug failure in both countries were compared, and encouragingly, the quintuple mutant \(dhfr\) 51I/59R/108N + \(dhps\) 437G/540E which is a key mutation associated with clinical failure to SP [31, 32] was not observed, in either Grande-Comore or in Thiès, Senegal. This lack of the quintuple mutation has been reported in previous studies [14–19], with the exception of one study by Andriantsoainina et al. where a 0.4 % prevalence of the mutation A437G/K540E was observed in Comoros [27]. While, SP can be still used for ITP as recommended by World Health Organization [33], vigilance is needed as the WHO recommends replacing SP monotherapy if the prevalence of the K540E mutation exceeds 50 % in the population [33, 34]. These results taken together suggest that yearly monitoring of SP drug resistance in Comoros needs to be a policy priority as it will be important to assess the future efficacy of SP as a drug for IPT.

**Conclusion**

In conclusion, drug resistance monitoring in Comoros is a critical area of future ongoing research. DNA extracted from RDTs can be a useful source of DNA for genetic epidemiological studies. DNA extracted from RDTs can be an alternative to filter paper or venous blood when such samples are not available.
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