Prevalence of the molecular marker of \textit{Plasmodium falciparum} resistance to chloroquine and sulphadoxine/pyrimethamine in Benin seven years after the change of malaria treatment policy

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Abstract

\textbf{Background:} In Benin, the National Malaria Control Programme (NMCP) changed the policy of malaria treatment in 2004 following increasing of failure rate of treatment with chloroquine (CQ) and sulphadoxine-pyrimethamine (SP). The objective of this study was to determinate the prevalence of \textit{Plasmodium falciparum} molecular markers that are associated with resistance to CQ and SP in Benin seven years after the new policy was instituted.

\textbf{Methods:} The study was conducted in southern Benin, a region characterized by a perennial malaria transmission. Blood samples were collected in 2011 from children presenting with symptomatic and asymptomatic \textit{P. falciparum} infections and living in the same area. The prevalence of critical point mutations in the genes of \textit{pfcrt} (codon 76), \textit{pfmdr1} (codon 86), \textit{pfdhfr} (codons, 51, 59 and 108) and \textit{pfdhps} (codons 437, 540) was examined in parasite isolates by mutation-specific restriction enzyme digestion of nested PCR products.

\textbf{Results:} A high prevalence of parasites carrying point mutations in all studied targets was found: T\(_{76}\): 93.9% [89.8; 96.7], I\(_{51}\): 96.2% [92.7; 98.4], R59: 93% [89.7; 96.7], N108: 97.6% [94.6; 99.2] and G437: 71.4% [64.8; 77.4]. No mutation was found at codon 540 of the \textit{pfdhps} gene. The proportion of parasite isolates carrying triple mutation in the \textit{pfdhfr} gene \textit{IRN} (I\(_{51}\), R59 and N108) and quadruple mutation on the combination of \textit{pfdhfr/pfdhps} IRNG (I\(_{51}\), R59, N108 and G437) was 91.5% [86.9; 94.9] and 65.7% [58.9; 72.1], respectively. Analysis of mutation in relation to the clinical status (symptomatic or asymptomatic) and according to age (younger or older than 10 years) showed similar very high frequencies in each category without significant difference between two groups.

\textbf{Conclusions:} These results suggest a persistence level of resistance of \textit{P. falciparum} to CQ and SP, seven years after the recommendation of the change of malaria treatment policy in Benin. The distribution of mutations studied was neither related to age nor to clinical status.

\textbf{Keywords:} Prevalence-mutation, \textit{P. falciparum}, Chloroquine, Sulphadoxine-pyrimethamine
Background
In Benin, the anti-malarial treatment policy has long been based on the use of chloroquine (CQ) and sulphadoxine-pyrimethamine (SP) as first- and second-line treatments, respectively. In vivo efficacy studies, conducted in 2002 by the National Malaria Control Programme (NMCP) according to the WHO protocol, had revealed treatment failures rates by region ranging from 15.0–61.3% with CQ and 3.3–45.9% with SP in under-fives followed up to 14 days [1]. These studies were performed in five regions of the country located in the south (Lokossa), centre (Dassa Zoume, Abomey) and north (Kouandé, Malanville), with an overall failure rate of 35.2 for CQ (11.5% and 23.7% early and late treatment failures respectively) and 22.8% for SP (8.3% and 24.5% early and late treatment failures respectively) (unpubl. data from NMCP, Ministry of Health). Based on these observations, the national anti-malarial drug policy was changed in 2004 by the official withdrawal of CQ and SP in the treatment of uncomplicated malaria. However, SP is still used for the prevention of malaria in pregnancy through intermittent preventive treatment (IPTp), as recommended by WHO. The current policy is based on the use of artemether-lumefantrine (AL)(Coartem®) and artesunate amodiaquine (ASAQ)(Arsucam®) as first-line treatment for uncomplicated malaria. It should be noted that between 2004 and 2005 there was an overlap of treatments with CQ, SP and AL, ASAQ, due to the unavailability of the new treatment in some health facilities. In 2005, Aubouy [2] conducted an in vivo study according to the WHO protocol over 28 days [3], which showed very high failure rates (85.7% and 50%), to CQ and SP, respectively. Since the first reports pointing the decreased efficacy of anti-malarial drugs on malaria parasites in endemic regions, important investigations led to the discovery of the involvement of genetic mutations in these parasites conferring resistance [4,5]. Thus, mutations occurring on the gene pfCRT (T76), and secondarily pfmdr1 (Y86), pfdhps (G437, E 540, A436), pf dhfr (I51, R59, N108) have respectively been associated with resistance of P. falciparum to CQ, and SP [6-9]. Moreover, it has been increasingly reported in some African settings, that P. falciparum parasites again become susceptible to CQ several years after the withdrawal of the molecule [10,11]. In Benin, no data on the prevalence of parasite molecular markers of resistance was available before the treatment policy change and only a limited number of studies of molecular markers of anti-malarial drug resistance in P. falciparum have been carried out [12,13]. These studies have focused on genes pf dhfr and pf dhps and reported high proportion of quadruple mutant parasites (above 80%). Thus, it seemed appropriate to assess the prevalence of different markers of resistance to CQ and SP in the population of Benin seven years after the official withdrawal of those drugs in the treatment of malaria in Benin. The main objective of this study was to determine the prevalence of P. falciparum molecular markers that are associated with resistance to CQ and SP by analysing the point mutations in pfCRT, pfmdr1, pf dhfr and pf dhps gene using samples from asymptomatic children and those with uncomplicated malaria in southern Benin.

Methods
Study sites and population
The study was conducted in two highly endemic regions of southern Benin, including the departments of Littoral and Ouémé. Southern Benin is characterized by a sub-equatorial climate and a perennial malaria transmission with two peaks corresponding to the rainy seasons (April–July and mid-September–November) [14]. Children aged between six months and 15 years, who resided in the study sites for more than a period of six months were enrolled from May through August 2011. Asymptomatic children were recruited among the nursery and primary school pupils in the study area when they showed a positive thick smear regardless of parasite density. A census of all nurseries and primary schools in the study area was initially conducted, from which a draw of five nurseries and five primary schools was made. In each school included, all attending students in all open classes on the day of the survey were involved in the study. The age ranged from 3 to 5 years in nurseries and 6 to 18 years in primary schools (children over 15 years were screened to evaluate the carriage rate of P. falciparum in the study area, but were removed from the sample for molecular analysis). Written informed consent from the principal and parents were obtained. For the selection of symptomatic children, WHO protocol for in vivo studies was applied to avoid severe malaria or uncomplicated malaria associated with other diseases. Thus, children visiting the health facilities in the study area and who met the criteria below were enrolled in the study: (i) fever (axillary temperature ≥ 37.5°C) or a history of fever within the past 48 hours, (ii) P. falciparum mono-infection with parasite density ≥ 1,000 asexual forms per microlitre, identified by microscopy on blood smears; (iii) no evidence of a concomitant febrile illness; iv) no sign/symptoms of severe malaria as defined by WHO [15] and (v) written informed consent from parents.

Sample collection and laboratory procedure
Venous blood from symptomatic children fulfilling the above criteria was collected systematically on the filter paper. Samples of asymptomatic children containing parasites were also stored as spots on filter paper. Thick and thin blood smears were prepared and were stained
with 10% Giemsa for rapid diagnostic. All thick blood smears were examined against 500 leucocytes. Parasite densities were recorded as the number of parasites/μl of blood, assuming an average leucocyte count of 8,000/μl of blood. All slides were read in the laboratories of the health centres, with external quality control performed on 10% of the negative slides and all positives in the reference Parasitology Laboratory of the Centre National Hospitalo-Universitaire in Cotonou. All malaria-infected patients, based on microscopy results, were treated according to malaria treatment policy based on ACT: artemether/lumefantrine.

DNA extraction, PCR- RFLP assay
Parasite DNA was extracted using the Chelex methods [16] and stored at −20°C until use. The regions of the pfcrt, pfmdr1, pfdhfr and pfdhps genes containing the polymorphisms of interest were amplified by nested PCR described previously [17-19]. The RFLP was performed by digesting 6 μl (pfcrt and pfmdr1) or 10 μl (pfdhfr and pfdhps) of PCR products with specific restriction enzymes (that varied according to the studied codons) and buffers obtained from New England Biolabs (Medinova, Glostrup, Denmark) according to supplier’s specifications. The pfcrt mutation at codon 76, pfmdr1 mutation at codon 86, pfdhfr mutations at codons 51, 59, 108, and pfdhps mutations at codons 437 and 540, were characterized. Five laboratory clones (3D7, FCR3, HB3, D10 and Dd2) used as a standard positive controls and negative controls (without DNA) were amplified and also digested with the samples. The digested products were visualized by electrophoresis on 1.5% agarose gels containing ethidium bromide. Gels were recorded by digital photography. Four samples of each series were randomly selected and tested repeatedly to check the reproducibility of the technique. Enzymes and control used, the size of the fragments obtained after cleavage of mutant or wild-type strains are shown in Table 1. Control strains were used for each analysis and mutations were determined to be present or absent on the basis of the presence or absence of the expected bands on the gel. Markers of 100 bp (Pharmacia Biotech) were used to size the bands. Each codon was characterized as wild-type (no mutation present), pure mutant (only mutant genotypes detected). Cases of mixed infection (wild type and mutant) were categorized as mutant throughout the analysis. All molecular analyses were performed in the Molecular Biology Laboratory of Center for integrated malaria control (CLIP).

Data analysis
The data were entered in the software Reversion 2.12.0 (R Foundation for Statistical Computing, Vienna, Austria). The frequency of a particular mutant was calculated as the proportion of the specific mutant samples among the total number of samples successfully analysed for this mutation. Similarly, the frequencies of double, triple and quadruple mutants were determined as the proportion of subjects with two, three and four mutations among the total numbers of samples tested for the each. To investigate the relationship between the mutation and age, children were segregated into two categories: children below and above 10 years of age. The reason for this division is that recent intensification of malaria control activities in the country, such as the widespread distribution and use of insecticide-treated nets, large-scale indoor residual spraying is likely to impact the acquisition of immunity, usually achieved at about five years of age in endemic areas [20]. Wilcoxon test and Student test were used to compare the distribution of age according to the clinical status and distribution of the parasite density respectively. The chi-square test or Fisher’s exact test was used for proportion comparisons. The significance level (P < 0.05) was used

| Restriction enzymes | Wild type alleles/control used | Number and size of bands obtained (bp) | Mutant alleles/control used | Number and size of bands obtained (bp) |
|---------------------|---------------------------------|----------------------------------------|-----------------------------|----------------------------------------|
| **pfdhfr** : Nested PCR product size : 594 | | | | |
| Alu I | S108/3D7 | 2bands : 280+ 314 | N108 or T108 HB3, Dd2 | 1 band : 594 |
| Tsp 509 I | N51/3D7 FRC3 | 2 bands : 125+ 150 | IS1/Dd2 | 2bands : 125+ 250 |
| Xmn I | C59/3D7, FCR3 | 2bands : 250+ 344 | RS9/Dd2 | 2bands : 250+ 320 |
| **pfdhps** : Nested PCR product size : 711 | | | | |
| Ava II | A437/FCR3 | 1band : 711 | G437/3D7, Dd2 | 1band : 650 |
| Fok I | K540/3D7, Dd2, FCR3 | 1band : 711 | E540/D10 | 2bands : 311 + 400pb |
| **pfcrt** : Nested PCR product size : 145 | | | | |
| Apo I | K76/3D7, HB3 | 2 bands : 99 + 46 | T76/Dd2 | 1 band : 145 |
| **pfmdr1** : Nested PCR product size : 521 | | | | |
| AFI III | N86/3D7, HB3 | 1 band : 521 | Y86/FCR3 | 2 bands : 346 + 175 |
to make the link between mutations, clinical status and age.

Ethical approval
This study was approved by Ethical Committee of the Faculté des Sciences de la Santé, Cotonou, Benin.

Results
Characteristics of the study population
A total of 2,249 asymptomatic children were screened for malaria parasitaemia; 208 samples were found positive for \( P. falciparum \) and were collected and stored as spots on filter paper. The carriage rate of \( P. falciparum \) was 9.2%. From these positive cases, 143 samples were selected for molecular analysis (after the withdrawal of children above 15 years age). A total of 73 samples of symptomatic subjects were collected and analysed. Thus a total of 216 samples were analysed for point mutations. The characteristics of subjects enrolled are shown in Table 2. For each target studied, the number of isolates actually analysed depended on the success of PCR amplification and were 213 for \( pfcrt \), 212 for \( pfmdr1 \) and \( pfdhfr \), and 210 for \( pfdhps \). Retested samples to check the reproducibility results were consistent with those found initially.

Prevalence of \( pfcrt \) and \( pfmdr1 \), alleles and mutations
The mutant allele T76 of \( pfcrt \) gene was present in 92% (196/213) of samples from patients that were analysed for this mutation. In contrast, only 28.3% (60/212) of samples analysed carried mutation type allele Y86 of \( pfmdr1 \). 42.9% carried the wild-type allele (N86), and 28.8% carried a mixture of the wild type and the mutant allele (N86Y86) (Figure 1).

The T76 mutation was present in 93.9% (200/213) [89.8; 96.7], which was higher in that population than Y86: 57.1% (121/212) [92.7; 98.4], R59: 93.9% (199/212) [89.7; 96.7], N108: 97.6% (207/212) [94.6; 99.2], \( Pfldhfr \) triple mutant IRN (I51, R59 and N108): 91.5% (194/212) [86.9; 94.9], \( Pfldhfr/Pfdhps \) quadruple mutant IRNG (\( Pfldhfr \) I51, R59, N108, and \( Pfdhps \) G437): 65.7% (138/210) [58.9; 72.1] (Table 3).

Mutation and clinical status and age
According clinical group, Y86 and double mutation T76Y86 were more represented in asymptomatic group, \( P = 0.03 \). Otherwise, there was no significant difference between prevalence of single, double, triple or quadruple mutation and clinical status (Table 3). When the data were analysed by age categories similar frequencies of single, triple and quadruple mutant parasite were found in children younger than 10 years and in older children, regardless of the clinical status. The purpose here was to see if the distribution of mutant parasites could be affected by age and therefore probably by the gradual acquisition of immunity. However, a tendency towards elevation of Y86 and double mutation T76Y86 rate among symptomatic children less than 10 years was noted but no statistically significant difference (Table 4).

Table 2 Characteristics of the study population

|                      | Symptomatic children | Asymptomatic children | p-value |
|----------------------|----------------------|-----------------------|---------|
| Number (%)           | 73 (33.8%)           | 143 (66.2%)           |         |
| Age (years)          | Mean (SD)            | 7.8 (3.0)             | 8.3 (2.5) | 0.09<sup>w</sup> |
|                     | Median (IQR)         | 8 (7–9)               | 9 (8–10) | 0.008<sup>b</sup> |
| Children below 10 years n (%) | 56 (76.7%)          | 82 (57.3%)            | 0.85<sup>b</sup> |
| M/F ratio            | 39/34; 1.15          | 73/70: 1.04           |         |
| Parasitaemia         | Geometric mean 95% CI| 6822.58               | 656.69  | <0.001<sup>f</sup> |
|                      | [5072.29; 9176.86]   | [474.74; 908.36]      |         |
|                     | Median (IQR)         | 7000 (2400–15000)     | 1000 (150–2520) | <0.001<sup>xx</sup> |

<sup>w</sup>: Wilcoxon’s test; <sup>b</sup>: Chi 2 test; <sup>t</sup>: Student test.


Discussion

The main objective of this study was to evaluate the prevalence in Benin of *P. falciparum* resistance markers to CQ and SP, two conventional anti-malarials that have been used for a long time. This study was justified by the need of surveillance of *P. falciparum* resistance to anti-malarials drugs in Benin through well-characterized molecular markers of resistance to contribute to the monitoring in the subregion. This monitoring is intended on the one hand to watch for a possible return of sensitivity of *P. falciparum* to CQ and secondly, to describe the current patterns of molecular markers associated with parasite resistance to SP, the latter still used as a preventive treatment of malaria in pregnant women. This work was carried out on parasites obtained from two distinct clinical groups of children, to investigate a possible relationship between molecular markers and clinical status. Mutation at codon 164 of *pfdhfr* gene has not been studied because several studies have noted the absence or rarity of this mutation in African isolates [13,21-24].

High rates of single, double, triple or quadruple mutation observed in this study reflect the current level of sensitivity of *P. falciparum* to CQ and SP in Benin, which is still expected to be low although the two drugs were officially withdrawn from curative treatment of uncomplicated malaria in 2004. These high rates of mutant parasites had been previously found by other authors [23,25-28]. The fact that the frequency of mutations at codon site 437 of *pfdhps* is lower than those observed on the *pfdhfr* gene confirms that the mutations associated with parasite resistance to SP appeared earlier on the *pfdhfr* than those affecting the *pfdhps* [29,30].

Similar to other reports from the sub-region [31,32], the parasites harbouring the mutation at codon 540E were not found in Benin. Regarding the *pfcr* gene, analysis of the T76 mutation (93.9%) in isolates from Benin showed a high prevalence of parasites carrying this

![Figure 1 Prevalence of pfcr and pfmdr alleles.](http://www.malariajournal.com/content/12/1/147)

Table 3 Prevalence of molecular markers associated with *P. falciparum* resistance to CQ and SP in symptomatic and asymptomatic children

| Molecular marker | Population n (%) | Symptomatic n (%) | Asymptomatic n (%) | p-value |
|------------------|------------------|-------------------|--------------------|---------|
| T76 (n = 213)    | 200 (93.9%)*     | 68/72 (94.4%)     | 132/141 (93.6%)    | 1       |
| Y86 (n = 212)    | 121 (57.1%)      | 29/71 (40.8%)     | 92/141 (65.2%)     | 0.03⁵   |
| T76+Y86 (n = 212)| 117 (55.2%)      | 28/71 (39.4%)     | 89/140 (63.6%)     | 0.03⁵   |
| G437 (n = 210)   | 150 (71.4%)      | 48/73 (65.8%)     | 102/137 (74.5%)    | 0.42²   |
| E540 (n = 210)   | 0%               | 0%                | 0%                 |         |
| I51 (n = 212)    | 204 (96.2%)      | 69/73 (94.5%)     | 135/139 (97.1%)    | 0.45⁴   |
| R59 (n = 212)    | 199 (93.9%)      | 69/73 (94.5%)     | 130/139 (93.5%)    | 1       |
| N108 (n = 212)   | 207 (97.6%)      | 71/73 (97.3%)     | 136/139 (97.8%)    | 0.72⁵   |
| IRN¹ (n = 212)   | 194 (91.5%)      | 67/73 (91.8%)     | 127/139 (91.4%)    | 0.61²   |
| IRNG² (n = 210)  | 138 (65.7%)      | 44/73 (60.3%)     | 94/137 (68.6%)     | 0.15⁵   |

¹ *pfdhfr* triple mutants (I51, R59, and N108).
² *pfdhfr/pfdhps* quadruple mutants (*pfdhfr* I51, R59, N108 and *pfdhps* G437).
³ *p* < 0.001 of Chi 2 test (comparison between T76 and Y86, T76 and T76Y86).
⁴ *p*-value of fisher test.
⁵ *p*-value of Chi 2 test.
mutation. Unlike Malawi, where there has been a reversal of the prevalence of mutant T76 a few years after the withdrawal of CQ [10,11], it is clear that this has not been the case in Benin. Self-medication with fake medicines especially with regard to CQ (despite its formal withdrawal from the treatment policy of malaria) could be a leading cause for maintaining this high prevalence of T76 mutant parasites as a result of continued and frequent use of CQ [33] involving either insufficient doses or too short a duration of administration. It is a common practice in the south of the country characterized by large markets (with neighbouring Nigeria) where illegitimate distribution of fake drugs is common [2,34]. This could also be marginally explained by a possible cross-resistance shared between CQ and AQ [35,36], since AQ is present in the ASAQ combination currently used for malaria management in Benin. WHO in vivo drug efficacy studies conducted in 2008 with this combination in two localities of the country noted an adequate clinical and parasitological response (ACPR) of 100% and 76.6% respectively without PCR correction. The second locality (Dassa Zoume) with a decrease of ACPR (23.44% of late parasitological failure) had posted 38.8% of late parasitological failure in 2002 during in vivo drug efficacy studies with CQ (unpublished data from Ministry of Health, Benin).

Thus, the re-emergence of sensitive parasite strains after the withdrawal of CQ depends not only on the time elapsed since the withdrawal of the treatment, but also on the speed of replacement and implementation of the new drugs, to allow the cessation of use and consumption of drugs replaced. It is important that the health authorities of the country ensure the effective withdrawal of CQ by emphasizing the education about the risks of self-medication and implementing control methods against the entry of fake medicines. The results obtained

![Figure 2 Prevalence of *pfdhps* alleles.](image2)

![Figure 3 Prevalence of *pfdhfr* alleles.](image3)
with the gene pfcrt are consistent with those reported in China by Zang et al. [37], but is in contrast to those reported by Kamugisha et al. in Tanzania [38] and Raman et al. in Mozambique [39].

Regarding SP, it should be noted that the use of other sulpha drugs, such as trimethoprim-sulphametoxazole (co-trimoxazole) often prescribed for bacterial infections in children but also in prevention and treatment of opportunistic infections in persons living with HIV, contribute in maintaining drug pressure on malaria parasites. The use of SP in intermittent preventive treatment in pregnant women (IPTp) since its withdrawal from the treatment of uncomplicated malaria may also explain the maintain of the selective pressure on SP targets as shown in the study conducted by Bertin et al. in Benin and Pearce et al. in Tanzania [13,40]. Similarly to what has been reported by other authors [17,27], this study shows that regardless of the gene, mutations are not associated with the clinical status of children. On the other hand, to determine whether immunity contributed to the ability to clear infections by parasites carrying resistance, the proportion of infections by parasites carrying different mutations compared between children younger than 10 years and older children suggests that age does not influence the distribution and carriage of resistant parasites whatever the clinical status and type of mutation.

**Conclusions**

This study showed high prevalence of pfcrt76 and quadruple pfdhfr/pfdhps mutants parasites (triple mutant pfdhfr + single 437 pfdhps mutant) confirming the persistence of parasite resistance to CQ and SP in Benin several years after officially withdrawal of these two drugs in the treatment of uncomplicated malaria. Clinical status and age do not seem to influence the frequency of individual mutants or the distribution of the triple and quadruple mutant parasites. This study has generated data which will be useful in monitoring the dynamics of drug resistant malaria parasites in Benin and in the sub-region.

**Competing interests**

The authors declare that they have no competing interests.

**Authors’ contributions**

DKZ designed the study protocol, supervised the study and corrected the manuscript; AQH participated in the design of study, supervised the study and laboratory tests, drafted the manuscript; NTN supervised laboratory tests, monitored laboratory quality and participated in the manuscript writing; SA, LR, EO, CA, MB performed patients enrollment, molecular analysis and participated in the manuscript writing. JPC has coordinated and helped to draft the manuscript. AM participated in the design of the study, coordination, and helped to draft the manuscript. All authors read and approved the final manuscript.

**Acknowledgements**

We are grateful to the children who participated in the study, as well as to their mothers and head of their schools. We are pleased to thank caregivers from St Luc and Bethesda hospitals, Semé Kpodji health centres, IRD laboratory workers. This work was financed by French Development Agency and the Ministry of Health of Benin. This funding has contributed to data collection, laboratory testing and statistical analysis.

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**Published: 3 January 2013 Accepted: 28 April 2013**

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**Table 4**  
Prevalence of mutations conferring resistance to CQ and SP in *Plasmodium falciparum* according age

| Molecular marker | Symptomatic | Asymptomatic |
|-----------------|------------|--------------|
|                 | < 10 ans   | ≥ 10 ans     | < 10 ans   | ≥ 10 ans     |
| T 76            | 53 (94.6%) | 15 (93.8%)   | 74 (91.4%) | 58 (96.7%)   |
| Y86             | 25 (45.5%) | 4 (25%)*     | 49 (60.5%) | 43 (71.7%)   |
| T76 + Y86       | 24 (43.6%) | 4 (25%)**    | 47 (58.8%) | 42 (70%)     |
| G437            | 36 (64.3%) | 12 (70.6%)   | 54 (68.4%) | 48 (82.8%)   |
| E540            | 0%         | 0%           | 0%         | 0%           |
| IS1             | 52 (92.9%) | 17 (100)     | 77 (96.2%) | 58 (98.3%)   |
| R59             | 53 (94.6%) | 16 (94.1%)   | 77 (96.2%) | 53 (89.8%)   |
| N108            | 54 (96.4%) | 17 (100)     | 78 (97.5%) | 58 (98.3%)   |
| IRN1            | 51 (91.1%) | 16 (94.1%)   | 76 (95%)   | 51 (86.4%)   |
| IRNG2           | 33 (58.9%) | 11 (64.7%)   | 52 (65.8%) | 42 (72.4%)   |

*pfdhfr* triple mutants: (I51, R59, and N108).

*pfdhfr/pfdhps quadruple mutants: (pfdhfr I51, R59, N108 and pfdhps G437).

**p = 0.24 (fisher test).**
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Cite this article as: Ogouyèmi-Hounto et al.: Prevalence of the molecular marker of Plasmodium falciparum resistance to chloroquine and sulphadoxine/pyrimethamine in Benin seven years after the change of malaria treatment policy. Malar J 2013 12:147.