KASP: a genotyping method to rapid identification of resistance in Plasmodium falciparum

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Research

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Abstract

Background

The emergence and spread of antimalarial resistance continues to hinder malaria control. *Plasmodium falciparum*, the species that causes most human malaria cases and most deaths, has shown resistance to almost all known antimalarials. This antimalarial resistance arises from the development and subsequent expansion of Single Nucleotide Polymorphisms (SNPs) in specific parasite genes. A quick and cheap tool for the detection of drug resistance can be crucial and very useful for use in hospitals and in malaria control programs. It has been demonstrated in different contexts that genotyping by Kompetitive Allele Specific PCR (KASP), is a simple, fast and economical method that allows a high-precision biallelic characterization of SNPs, hence its possible utility in the study of resistance in *P. falciparum*.

Methods

Three SNPs involved in most cases of resistance to the most widespread antimalarial treatments have been analyzed by PCR plus sequencing and by KASP (C580Y of the Kelch13 gene, Y86N of the *Pfmdr1* gene and M133I of the *Pfcytb* gene). A total of 113 *P. falciparum* positive samples and 24 negative samples, previously analyzed by PCR and sequencing, were selected for this assay. Likewise, the samples were genotyped for the MSP-1 and MSP-2 genes, and the Multiplicity of Infection (MOI) and parasitaemia were measured to observe their possible influence on the KASP method.

Results

The KASP results showed the same expected mutations and wild type genotypes as the reference method, with few exceptions that correlated with very low parasitaemia samples. In addition, two cases of heterozygotes that had not been detected by sequencing were found. No correlation was found between the MOI or parasitaemia and the KASP values of the sample. The reproducibility of the technique shows no oscillations between repetitions in any of the three SNPs analyzed.

Conclusions

The KASP assays developed in our study were efficient and versatile for the determination of the *Plasmodium* genotypes related to resistance. The method is simple, fast, reproducible with low cost in personnel, material and equipment and scalable, being able to core KASP arrays, including numerous SNPs, to complete the main pattern of mutations associated to *P. falciparum* resistance.

Introduction
Malaria remains an important problem in global public health, being a major cause of morbidity and mortality that continues to claim more than 400,000 lives every year [1]. The continuing devastating impact of this disease is partly due to the emergence and spread of resistance to antimalarials [2].

*Plasmodium* parasites quickly develop resistance to antimalarials and evade the immune system through mutations in their genome. Since an effective vaccine has not yet been developed, control and surveillance of antimalarial resistance is crucial for saving lives [3]. *Plasmodium falciparum*, the species that causes most human malaria cases and most deaths, has shown resistance to almost all known antimalarials [4]. In fact, recently, cases of delayed parasite clearance following treatment with an artemisinin-based combination therapy (ACT) have been reported in the Greater Mekong sub-region. This represents a major threat to the ability to control and treat malaria, since this is the current first line treatment for uncomplicated *P. falciparum* infections [3, 5]. This antimalarial resistance arises from the development and subsequent expansion of Single Nucleotide Polymorphisms (SNPs) in specific parasite genes [6, 7]. To identify and monitor the propagation of these resistances, SNPs can be detected by molecular analysis. The usual method of resistance detection is the Polymerase Chain Reaction (PCR) and sequencing or digested with restriction enzymes [7]. Moreover, next generation sequencing genotyping is an emerging method of genotyping SNPs that is increasingly being adopted for both diagnosis and research [8]. Despite these advances, rapid, simple, affordable method that can be transferred to the daily clinic are necessary to detect possible resistance in the patient and to a fast population screening of resistance at low cost. It has been demonstrated that genotyping by Kompetitive Allele Specific PCR (KASP), is a simple, fast and economical method that allows a high-precision biallelic characterization of SNPs, as well as insertions and deletions in specific loci, hence its possible utility in the study of resistance in *P. falciparum*.

This method has been used, for example, for the detection of SNPs associated with efficacy of specific drugs [9]; for genotyping candidate genes associated with the development of genetic diseases, such as Huntington's disease [10]; or for the genotyping of SNPs associated with G6PD deficiency [11].

Given the importance of rapid response for guidance in the administration of antimalarial treatment in certain patients, a quick and cheap tool, as KASP, for the detection of drug resistance can be crucial and very helpful for use in hospitals and in malaria control programs. The objective of this study is to verify the possible utility of the KASP technique in the analysis of resistance to antimalarial drugs. To validate the method, three SNPs involved in most of the resistance of the most widespread treatments have been selected: i) SNP C580Y of the gene that codes for the Kelch13 protein (Gene PF3D7_1343700) that is present in 80% of cases of resistance to artemisinin treatments [12]; ii) SNP Y86N of the *Pfmdr1* gene (gene PF3D7_0523000) that is involved in resistance to different drugs such as quinine, chloroquine, mefloquine, halofantrine, amodiaquine and lumefantrine; the latter frequently used in combination with artemisinin, and where a natural selection of this SNP has been observed after treatment [13, 14, 15]; and iii) SNP M133I of the *Pfcytb* gene (gene MAL MYTH 3) that is related to resistance to atovaquone [16] and mefloquine [17], being the most frequent mutation in *in vitro* tests.
Methods

Sample collection

The *P. falciparum* resistance genotype of those patients who presented single malaria infection was analyzed as part of a prospective, observational, multicenter study, approved by the Medical and Health Research Ethics Committee (CEIC) of the Hospital Universitario 12 de Octubre (CEIm: 18/021).

Final selection was based on species-specific identification by nested-PCR [18, 19] and only samples with unique infection with *P. falciparum* were chosen for the study.

A total of 112 *P. falciparum* positive samples and 24 negative samples were selected for this assay. In addition, a strain, donated by the Parasitology Department of the Institute for Medical Research in Malaysia with the mutation C580Y of the gene that codes for the Kelch13 protein, was included.

Sample processing

DNA was extracted from 200 µl of whole blood, collected in ethylenediamine tetraacetic acid (EDTA) tubes, using the QIAamp DNA mini blood kit (QIAGEN®, Hilden, Germany) according to the manufacturer's instructions, resuspended in a final volume of 100 µl of distilled water. The DNA is stored at 4°C until use, using 5 µl per reaction.

Plasmodium falciparum resistance genotyping

Genotyping of *Pfmdr1*, *Pfcytb* and *PfK13* genes was carried by real time PCR modified from the original methods [20, 21, 22]. Reaction mixture consists of 1x QUANTIMIX HotSplit Probes kit (Biotools, Madrid, Spain) 0.25 µM of each primer (Table 1) and 1 µM EvaGreen® dye (Biotium, Hayward, CA, USA) in a final volume of 20 µl. The thermal cycle was performed in a Qiagen Rotor Gene Q 5 Plex HRM (QIAGEN®, Hilden, Germany). The amplification conditions for the *Pfmdr1* and *Pfcytb* genes were an initial cycle of denaturation of 10 minutes at 95 ºC, followed by 45 cycles of 15 seconds at 95 ºC, 15 seconds at 54 ºC and 30 seconds at 68 ºC, ending with a melting step (62 ºC to 92 ºC). In the case of the *PfK13* gene, the conditions were an initial denaturation cycle of 7 minutes at 94 ºC, followed by 40 cycles of 30 seconds at 94 ºC, 1 minute at 60 ºC and 1 minute at 72 ºC, ending with a melting step (62 ºC to 92 ºC).
Table 1
Primers name, gene target, sequence and method for the *P. falciparum* resistance genotyping.

| Target | Primer name/Reference | Method | SEq. 5’→3’ |
|--------|-----------------------|--------|------------|
| MDR    | MDR1-pfmdr1-1FK76T²¹  | RT-PCR MDR A & Sequencing | GTTGAACAAAAAGAGTACCAGCTG |
|        | MDR2-pfmdr1-1RK76T²¹  | Sequencing | TCGTACCAATTCTGAAGTCAC |
|        | MDR3-pfmdr1-2FK76T²¹  | Sequencing | TTTCCGTTTTAATGTTCACCTGC |
|        | MDR4-pfmdr1-2RK76T²¹  | Sequencing | CCATCTTGATAAAAAACTTTTCTT |
|        | MDR1-1246-pfmdr1-1FD1246Y²¹ | RT-PCR MDR B & Sequencing | ATGACAAATTTTCAGATTA |
|        | MDR2-1246-pfmdr1-1FD1246Y²¹ | Sequencing | ACTAACACGGTTAAACATTT |
|        | MDR3-1246-pfmdr1-1FD1246Y²¹ | Sequencing | AATGTAAATGAATTTCACAAC |
|        | MDR4-1246-pfmdr1-1FD1246Y²¹ | Sequencing | CATCTTCTTTCCAAATTGATA |
| Cyt B  | CytB1-1F²²           | RT-PCR / Sequencing | CTCATATATTAGTTAAAGACAC |
|        | CytB2-1R²²           | ACAGAATAATCTTAGCACC |
|        | CytB3-2F²²           | Sequencing | AGCAGTAAATTTGGATATGGAGG |
|        | CytB4-2R²²           | ATTTTTAATGCTGTATCATACCCT |
| K13    | Pfal-k13-2-2PCR²⁰     | RT-PCR / Sequencing | GCCAAGCTGCCATTTCATTG |
|        | Pfal-k13-3-2PCR²⁰     | GCCTTGTGGAAAGAAGCA |

Specificity of amplification was determined by post reaction analysis using the melting temperature (Tm) curve of the amplified fragments (76.8 °C and 76.5 °C for *Pfmdr1* fragment A and B respectively, 79.5 °C for *Pf cyt b*, and 82.0 °C for *PfK13*).

Amplicons were purified using Illustra DNA and Gel Band Purification Kit (General Electric Healthcare, England), then sequenced, with its specific primers (Table 1), using Big Dye Terminator v3.1 Kit in an ABI PRISM® DNA Analyzer 3700. All amplified products were sequenced in both directions, twice. Blast tool from NCBI was used to confirm correct target amplification. Multiple nucleotide sequence alignments and analysis were performed using BioEdit version 7.0.5.3 [23] using sequences from 3D7 strain as wild-type for comparison.

*Plasmodium falciparum* genotyping
*Plasmodium falciparum* genotyping was performed by characterizing two merozoite surface membrane genes (MSP-1 and MSP-2) by PCR described elsewhere [24, 25]. Multiplicity of infection (MOI), defined as the number of genetically distinct parasite strains co-infesting a single host, was determined as the largest number of different alleles present at one of the two loci studied.

**Kompetitive allele-specific PCR (KASP)**

Sequences flanking SNPs were submitted for KASP™ assay design to Biosearch Technologies (California, USA). KASP assay was carried out following the instructions of the manufacturer [6]. The reaction mix per reaction consists of 5.1 µl of the KASP Master Mix, which include the two allele specific primers and one reverse primer (Table 2), and 0.138 µl of the Assay Mix, containing universal fluorescent probes, Taq polymerase and dNTPs in an optimized buffer solution.

| ID   | Primer_AlleleX & Allele Y | Primer_Common       |
|------|--------------------------|---------------------|
| Cyt b M133I | AATTACAGTTGCACCCCAATAACTC | AACTGCTTTGTTGTTATGTCTTACCAT  |
|       | GTAATTACAGTTGCACCCCAATAACTT |                       |
| K13 C580Y    | ATACCCCTAGATCATCAGCTATGTG | CTCACCATTAGTTCCACCAATGACATAAA |
|       | AATACCCCTAGATCATCAGCTATGTA |                       |
| MDR N86Y     | GTGTTTGGTGTAAATATAAGAAACATGA | GTACTAAAACCTATAGATACTAATGATAATA |
|       | CTGTTTTGGTGTAAATATAAGAAACATGT |                       |

To five µl of this reaction mix, five µl of DNA from the sample to be analyzed, are added.

PCR and fluorescent readings were performed in a Qiagen Rotor Gene Q 5 Plex HRM (QIAGEN®, Hilden, Germany) following the recommended thermal cycling conditions (Table 3).

**Tabla 3.** Thermal cycle conditions for KASP genotyping reactions
To analyze and interpret genotypic data, an Excel sheet is used, although it can also be done using the Thermo Fisher Cloud Genotyping application [26]. The FAM and HEX data, normalized by the internal reference ROX dye, corresponding to the wild and mutant allele, are plotted on the X and Y axes in a scatter plot, grouping the different allele clusters, while the negative controls are located near the intersection of the axes in the zero.

### Results

Analysis for SNPs resistance genotyping showed that only 12 out of 112 samples show any mutations. In all cases this corresponded to amino acid position 86, replacing asparagine with tyrosine (N86Y) of the *Pfmdr1* gene. In the other two genes studied, *Pfcytb* and *PfKelch13*, no mutations were found. The *P. falciparum* strain from Malaysia was found to show the expected C580Y mutation in the *PfKelch13* gene.

The KASP showed that none of the 24 negative samples were amplified, as did NTCs (no template control) included in each reaction. On the contrary the KASP showed positive results in all the expected samples except one for the SNP of *Pfmdr1*, one more in the case of *Pfcytb* SNP and in 3 for *PfK13* SNP. In all cases this lack of results was related to low parasitaemia; in the first two cases, where the samples were not amplified, presented a parasitaemia, quantified by real-time PCR, lower that 0.05 parasites/µl, being less than 1 parasites/µl for the *PfK13*. Despite these results, no statistical correlation has been observed between the level of parasitaemia and KASP values which are between −0.20 and −0.67 for wild genotypes and 0.20 for mutated genotypes.

The KASP result for *Pfcytb* M133I was that all of the amplified samples were wild type, as expected. Likewise, in the case of *Pfk13* C580Y all the samples showed the wild genotype, and only the control strain showed the mutated genotype. In the case of *Pfmdr1* Y86N, 10 samples gave a mutated genotype and two were characterized with a heterozygous genotype; the rest of the samples showed the wild genotype.
A factor that may also influence is the MOI, which ranges between 1 and 7 in the samples analyzed, but no statistical correlation has been observed with the KASP values, obtaining values between −0.12 and 0.24, except in one case that reaches 0.75.

The reproducibility of the method was analyzed by repeating two samples between four and six times in independent tests in the three SNPs, observing that the obtained values did not vary substantially as demonstrated by the low values of the standard deviation (Table 4).

| Gen    | Sample | Type | Average FAM filter | St. Desviations | Average HEX filter | St. Desviations |
|--------|--------|------|--------------------|-----------------|-------------------|----------------|
| Pfmdr1 | 222    | M    | 0.08               | 0.00            | 0.59              | 0.04           |
|        | 223    | WT   | 0.55               | 0.07            | 0.14              | 0.02           |
| PfK13  | 222    | WT   | 0.76               | 0.04            | 0.10              | 0.04           |
|        | 223    | WT   | 0.76               | 0.07            | 0.07              | 0.008          |
| Pfcytb | 222    | WT   | 0.92               | 0.003           | 0.02              | 0.001          |
|        | 223    | WT   | 0.48               | 0.003           | 0.08              | 0.0005         |

The expected graphical KASP results should show three clusters, in addition to the negative controls, corresponding to wild type samples, to mutated samples and to heterozygous samples (in our case, populations with both alleles). In the three SNPs studied the grouping of the samples according to the expected values are observed. For the Y86N in the Pfmdr1 gene the three clusters are present (Fig. 1a); in the case of the C580Y in the PfK13, the cluster of the wild type genotype and that of the mutated strain are observed (Fig. 1b) and in the case of M133I in Pfcytb only wild samples are observed (Fig. 1c). By grouping all the SNPs in a single graph, the same result is obtained, with the four clusters well defined (Fig. 1d).

**Discussion**

The development and utilization of genetic markers play a pivotal role in the study of malaria pathology in general and in resistance analysis in particular. Among molecular markers, SNPs have become the most promising due to their wide distribution within genomes and suitability for high-throughput automated genotyping [27]. Different methodologies, such as TaqMan, KASP, and rhAmp, have been proposed for the analysis of SNPs using high-resolution automated systems [26]. The Kompetitive Allele Specific PCR genotyping system (KASP™) is a homogeneous, fluorescent, endpoint genotyping technology whose use is expanding in plant genotyping [26, 27]. In malaria, especially in the study of *P. falciparum*, the design of a set of core SNPs array based on this technique could be very interesting for the study of resistance and other SNPs associated with pathology [28]. Firstly, it is necessary to validate
this methodology for *Plasmodium* due to the special characteristics of this parasite and the multiple infections it produces.

The system is designed in principle to characterize the presence of bi-alleles in samples of diploid organisms, although its efficacy has also been seen in hexaploid organisms such as wheat [26]. *Plasmodium* in humans is haploid, the only phases where this organism is diploid occurs in the anopheline vector, but the infection, in general, is produced by a number of genetically distinct parasite strains co-infecting a single host (MOI), therefore the genotype of a sample for a given gene can cover all allelic variants, being able to present the homozygote of both types or the heterozygote.

The KASP values obtained could be influenced either by the parasitaemia or by the MOI of the samples. Parasitaemia, whose detection limits ranges from 1 parasites/µl in the case of the analysis of *Pfk13* to 0.05 parasites/µl in *Pfmdr1* and *Pfcytb*, does not influence the KASP values, as demonstrated by the low statistical correlation values (-0.67 to 0.20). Similarly, no correlation was found between the MOI and the KASP values of the sample (-0.12 to 0.75). In both cases, the values obtained are very far from the expected values around ±1 in the event of a correlation.

In the three SNPs studied, the values obtained correspond to the expected, except in two samples in the Y86N mutation in the *Pfmdr1* gene, which according to KASP we would have both alleles in the sample. This type of case cannot be solved by direct sequencing of the amplified product since in this case only the majority form is detected or in any case an indeterminacy would occur in the involved nucleotide. The cloning and sequencing of multiple clones or pyrosequencing could resolve this heterozygosis but at a higher cost and with more delay than the KASP.

The reproducibility of the technique is very good, showing no oscillations between repetitions in any of the three SNPs analyzed (Table 4).

The KASP assays developed in our study were efficient and versatile for the determination of the *Plasmodium* genotypes related to resistance, not being influenced by the parasitaemia of the infection nor by the number of populations involved on it (MOI), and showing high reproducibility.

In conclusion, the method is simple, fast, reproducible and scalable, being able to core KASP arrays developed including numerous SNPs, and, on the other hand, the cost in personnel, material and equipment is lower than with other methodologies.

**Abbreviations**

Amino acids C: Cysteine; Y: Tyrosine; N: Asparagine; M: Methionine; I: Isoleucine

KASP: Kompetitive allele-specific PCR; MOI: Multiplicity of infection; *Pfmdr1*: *P. falciparum* Multi Drug Resistance gene 1; *Pfcytb*: *P. falciparum* gene cytochrome b gene. *PfK13*: *P. falciparum* Kelch13 gene. SNP: Single nucleotide polymorphism
Declarations

Ethics approval and consent to participate

This work is part of a prospective, observational, multicenter study, approved by the Medical and Health Research Ethics Committee (CEIC) of the Hospital Universitario 12 de Octubre (CEIm: 18/021), Madrid, Spain.

Consent for publication

No aplicable.

Availability of data and materials

All data analyzed during the study are included in this published article. The full datasets of *P. falciparum* genotyping generated during the current study are not publicly available due to its future publication but are available from the corresponding author on reasonable request.

Competing interests

The authors declare that they have no competing interests.

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Authors' contributions

JMR, SH and APA conceptualized the study. JMR, AAF, MJB, AMR, NAY designed the experiments. JMR provided training and supervision for the procedures. IFV, APA recruited patients and maintained databases. AAF, MJB, AMR, NAY, MLS performed the experiments. All authors contributed to the writing of the final version of the manuscript. All authors read and approved the final manuscript.

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Figures

Figure 1

Graphics showing genotypes of the Pfmdr1 (A), Pfcytb (B) and Pfk13 (C) genes, as well as a grouping of the data set (D). Red ovals correspond to WT genotype, blue to mutated genotype, yellow to heterozygous genotypes and green negative controls samples and non template controls (NTC). Red dots correspond to positives samples that were not detected due to their low parasitaemia.