Host candidate gene polymorphisms and clearance of drug-resistant *Plasmodium falciparum* parasites

Mahamadou Diakite1*, Eric A Achidi2, Olivia Achonduh3, Rachel Craik4, Abdoulaye A Djimde1, Marie-Solange B Evehe3, Angie Green4, Christina Hubbart4, Muntasir Ibrahim5, Anna Jeffreys4, Baldip K Khan6, Francis Kimani7, Dominic P Kwiatkowski4,8, Wilfred F Mbacham3, Sabah Omar Jezan7, Jean Bosco Ouedraogo9, Kirk Rockett4, Kate Rowlands4, Nawal Tagelsir10, Mamadou M Tekete1, Issaka Zongo9 and Lisa C Ranford-Cartwright11

**Abstract**

**Background:** Resistance to anti-malarial drugs is a widespread problem for control programmes for this devastating disease. Molecular tests are available for many anti-malarial drugs and are useful tools for the surveillance of drug resistance. However, the correlation of treatment outcome and molecular tests with particular parasite markers is not perfect, due in part to individuals who are able to clear genotypically drug-resistant parasites. This study aimed to identify molecular markers in the human genome that correlate with the clearance of malaria parasites after drug treatment, despite the drug resistance profile of the protozoan as predicted by molecular approaches.

**Methods:** 3721 samples from five African countries, which were known to contain genotypically drug resistant parasites, were analysed. These parasites were collected from patients who subsequently failed to clear their infection following drug treatment, as expected, but also from patients who successfully cleared their infections with drug-resistant parasites. 67 human polymorphisms (SNPs) on 17 chromosomes were analysed using Sequenom’s mass spectrometry iPLEX gold platform, to identify regions of the human genome, which contribute to enhanced clearance of drug resistant parasites.

**Results:** An analysis of all data from the five countries revealed significant associations between the phenotype of ability to clear drug-resistant *Plasmodium falciparum* infection and human immune response loci common to all populations. Overall, three SNPs showed a significant association with clearance of drug-resistant parasites with odds ratios of 0.76 for SNP rs2706384 (95% CI 0.71-0.92, P = 0.005), 0.66 for SNP rs1805015 (95% CI 0.45-0.97, P = 0.03), and 0.67 for SNP rs1128127 (95% CI 0.45-0.99, P = 0.05), after adjustment for possible confounding factors. The first two SNPs (rs2706384 and rs1805015) are within loci involved in pro-inflammatory (interferon-gamma) and anti-inflammatory (IL-4) cytokine responses. The third locus encodes a protein involved in the degradation of misfolded proteins within the endoplasmic reticulum, and its role, if any, in the clearance phenotype is unclear.

**Conclusions:** The study showed significant association of three loci in the human genome with the ability of parasite to clear drug-resistant *P. falciparum* in samples taken from five countries distributed across sub-Saharan Africa. Both SNP rs2706384 and SNP1805015 have previously been reported to be associated with risk of malaria infection in African populations. The loci are involved in the Th1/Th2 balance, and the association of SNPs within these genes suggests a key role for antibody in the clearance of drug-resistant parasites. It is possible that patients able to clear drug-resistant infections have an enhanced ability to control parasite growth.
Background

*Plasmodium falciparum* malaria remains a major cause of morbidity and mortality among children and pregnant women in sub-Saharan Africa. The most recent global figures show that malaria was responsible for over 863,000 deaths in 2008 and one fifth of the world’s population is at risk [1]. 85% of cases and 89% of deaths due to malaria are found in sub-Saharan Africa [1]. Over the last decade some African countries have seen a reduction in malaria cases and deaths, probably through increased funding for disease control measures such as the use of insecticide-treated mosquito nets. However parasite resistance to anti-malarial drugs, and mosquito vector resistance to insecticides, remain a major threat to the control of malaria.

Development of acquired immunity to malaria, which is only partially protective, requires persistent, sub-clinical infection over a period of several years (reviewed in [2]). The partial protection is strain-, stage- and species-specific. This may account for the observed higher malaria infection in children than in adults, and indicates that the immune status of the host influences the severity of malaria disease and the outcome of the treatment [3].

It is known that host genetic factors play a significant role in determining an individual’s susceptibility to many infectious diseases, including malaria [4-6]. Factors such as ethnic background [7], immunity [8,9], age [10], drug availability [11], co-infecting pathogens [12], socio-economic status [13], and parasite population structure [14] may impact on the outcome of infection, and the development of an effective immune response.

Advances in molecular biology have led to the discovery of genes involved in resistance to commonly used anti-malarial drugs such as chloroquine and sulphadoxine-pyrimethamine [15,16]. However the prevalence of parasites carrying the “resistant” alleles of these genes consistently exceeds *in vivo* treatment failure rates in malaria endemic settings [17], implying that some human hosts in malaria endemic-areas are able to clear genuinely drug-resistant malaria parasites. The ability to clear resistant parasites is associated with age [10,18], suggesting that host acquired immunity has a critical role in the clearance of drug-resistant *P. falciparum* infections in endemic regions. Several studies have supported the role of antiparasite immune responses in the therapeutic response to anti-malarial drugs during acute malaria ([19,20], reviewed in [3]). Host genetic factors such as sickle cell trait (HbAs), alpha-thalassaemia and haemoglobin E, as well as host pharmacogenetic differences, can also have an impact on the outcome of treatment with anti-malarial drugs [21-24]. The outcome of anti-malarial chemotherapy is, therefore, dependent on host genetic and immunological factors, as well as the level of drug resistance shown by the parasites.

In this study, known host genetic factors (other than haemoglobinopathies) that might account for individual differences in the clearance of drug-resistant parasites have been analysed in samples taken from subjects aged from 5 months old. The study included data from five African countries from both West and East Africa. The human gene variants investigated included cytokines and other immune mediators, thought to be involved in malarial pathogenesis, together with their receptors, and promoters. The overall objective of this study was to identify host immune factors that may be responsible for *in-vivo* clearance of drug-resistant *P. falciparum* by comparing allele frequencies of known SNPs in patients who clear genotypically resistant parasites with those patients who do not.

Methods

Study location and participant recruitment

Individuals were recruited to the study from five African countries: Burkina Faso, Cameroon, Kenya, Mali and Sudan (Figure 1). These countries were members of the International Atomic Energy’s Co-ordinated Research Project E15019 on “Improved accuracy and immunological markers for prediction of efficacy of anti-malarial drugs”. In all study sites, *P. falciparum* is responsible for > 95% of the clinical cases.

![Figure 1 Study sites in Africa](http://www.malariajournal.com/content/10/1/250)
Individuals aged from 5 months old, with uncomplicated *P. falciparum* malaria, who were treated with anti-malarial drugs including chloroquine, amodiaquine, sulphadoxine-pyrimethamine (SP) and artemisinins according to the policy within each country, were recruited to standard *in vivo* drug efficacy studies carried out in accordance with WHO protocols [25]. Details of these studies and their outcomes have been previously reported: Burkina Faso [26], Sudan [27], Cameroon [28,29], Kenya [30] and Mali [31], and a summary is provided in Table 1.

Fingerprick blood samples were collected onto filter paper from each individual at the time of recruitment to the study, for genotyping of the parasites present and for characterisation of the human SNP markers used in the study.

**Ethical considerations**

The study protocol was reviewed and approved by the Institutional Review Boards of the respective participant countries. Individuals were recruited to the study with the consent of their parents or guardians (for children), or with their own consent.

**Definition of *in vivo* drug resistance and sensitivity**

The clinical outcomes of treatment were defined according to WHO recommendations [25]. Samples were analysed for markers of parasite drug resistance from those patients who successfully cleared their infection ("sensitive" or "adequate clinical and parasitological response (ACPR)" as well as from those meeting the criteria for treatment failure. Briefly, "sensitivity" is defined as the clearance of parasites following drug treatment, without subsequent recrudescence within a defined period (28 days). An adequate clinical and parasitological response (ACPR) is defined as the absence of parasitaemia on day 28 irrespective of axillary temperature, without previously meeting any of the criteria for early and late treatment failure [25].

**Molecular characterisation of drug resistance**

Molecular analysis of parasite DNA from patients was performed according to standard IAEA protocols [32,33]. In all studies, parasites appearing during the follow-up period were characterised to distinguish possible reinfections from genuine recrudescence of resistant parasites, according to standard methodology [32].

DNA was extracted from the filter paper samples taken at admission to the study (i.e. before treatment), and amplified with primers to the genes in *P. falciparum* previously reported to be involved in resistance to chloroquine (*PfCRT, Pfmdr1*) and to SP (*dhfr, dhps*). The PCR product for each gene was then analysed using dot blot or RFLP to characterize the mutations present that have been linked to resistance [33]. The set of polymorphisms within drug resistance genes which were used to define drug resistance was defined for each country based on previous studies (Table 2).

**Table 1 Molecular Summary of *in vivo* drug efficacy trials carried out by the participant countries**

| Country               | Drugs studied in efficacy trials *in vivo* | Follow-up period (days) | Age range of study participants       |
|-----------------------|------------------------------------------|-------------------------|---------------------------------------|
| Burkina Faso          | Dihydroartemisin + piperazine            | 42                      | 6 months - 53 years                   |
|                       | Artemether + lumefantrine                | 28/42                   | 6 months - 39 years                   |
|                       | Amodiaquine                              | 28                      | 6 months - 18 years                   |
|                       | Amodiaquine + artemunate                | 28                      | 6 months - 30 years                   |
|                       | Amodiaquine + sulphadoxine-pyrimethamine| 42                      | 6 months - 55 years                   |
| Cameroon (Yaoundé)    | Sulphadoxine-pyrimethamine              | 28                      | 5 - 59 months                         |
|                       | Amodiaquine                              |                         |                                       |
|                       | Amodiaquine + sulphadoxine pyrimethamine|                         |                                       |
| Cameroon (Buea)       | Artesunate + sulphadoxine-pyrimethamine  | 28                      | 6 - 60 months                         |
|                       | Amodiaquine + artemunate                |                         |                                       |
| Kenya                 | Chloroquine                              | 14                      | 5 months-18 years                     |
|                       | Sulphadoxine-pyrimethamine              | 28                      |                                       |
|                       | Sulphadoxine pyrimethamine + Cotrimoxazole|                       |                                       |
| Mali                  | Chloroquine                              | 14                      | 6 - 60 months                         |
|                       | Amodiaquine                              | 28                      |                                       |
|                       | Sulphadoxine-pyrimethamine              |                         |                                       |
| Sudan                 | Chloroquine                              | 28                      | 6 months - 7 years                    |
|                       | Sulphadoxine-pyrimethamine              |                         |                                       |
|                       | Artesunate + sulphadoxine-pyrimethamine  |                         |                                       |
Cases of mixed infection, i.e. infections with both the wild-type and the resistance (mutant) allele, were considered as resistant. Only those samples that carried resistant alleles were included in the analysis and were divided into two groups (i) the cases: drug-resistant parasite genotype but infection was cleared following drug treatment, and (ii) the controls: drug-resistant parasite genotype and infection not cleared following treatment.

SNP genotyping of human DNA
Human DNA was extracted from filter paper blood samples (1 ml), drawn at the time of enrolment, using the Nucleon BACC2 DNA extraction Kit (Amersham Pharmacia Biotech, Buckinghamshire, UK), according to the manufacturer’s protocol. The concentration of DNA was determined using the PicoGreen® double strand (dsDNA) DNA Quantification Kit (Molecular Probes, Inc.). In order to increase the amount of human DNA required for high-throughput genotyping, all samples were subjected to whole genome amplification by primer extension pre-amplification PCR, using 15N base primers http://www.genetix.com[34]. The thermal cycling parameters were: 1 cycle at 94°C for 3 minutes for an initial denaturation, followed by 50 cycles of denaturation for 1 min at 94°C, primer annealing for 2 min at 37°C, 0.1°C/sec to 55°C, primer extension for 4 min at 55°C; and a final extension for 5 minutes at 72°C as described [35]. Amplified DNA samples were used at 1:10 dilution for genotyping on the SEQUENOM® iPLEX® platform according to the manufacturer’s instructions.

Selection of human immune response gene variants and genotyping
Known candidate gene variants were selected from the growing list of cytokines and other immune mediators that are thought to be involved in malarial pathogenesis, together with their receptors and promoters. In addition, lymphokines that regulate their expression and the adhesion molecules and inflammatory mediators that mediate their pathological effects were included. SNPs were selected using information from the literature and dbSNP [36], and reflected a compromise between SNP function, marker spacing and minor allele frequency (MAF). The initial SNP selection consisted of validated markers with minor allele frequency (MAF) ≥ 5%. This was narrowed down to an economic 67 known SNPs (Table 3) for which genotyping assays could be designed into two multiplex reactions for the Sequenom® iPLEX® mass spectrometry platform http://www.sequenom.com [37-39]. Genotyping accuracy was assessed by testing the conformation of the observed genotype distributions in the controls to the expected distributions under Hardy-Weinberg equilibrium (HWE). Assays which deviated from HWE at the 0.1% significance threshold were excluded from further analysis.

Statistical analyses
Comparisons of age and gender of participants, and parasitaemia at recruitment, for each country, and for all countries pooled, between those who did and did not clear genotypically resistant parasites, were compared using chi-squared tests (for frequency data), Kolmogorov-Smirnov (K-S) tests (for non-normally distributed values) or ANOVA (for normally distributed values). Each SNP was tested for association with the clearance phenotype using Odds Ratio (Univariate allele-based association tests). The data were then adjusted for confounding factors of age, ethnicity, gender and study location.

The P-values were not corrected for multiple testing. The Bonferroni correction, a commonly used correction which assumes independence between markers, was considered too stringent in this study as several SNPs may exhibit high degrees of dependence with one another, as measured by LD (D’) [40].

The large overall sample size resulting from combining studies at different sites increases the power to detect true positive associations and reject false-positives. Inter-study heterogeneity in association was assessed using Cochran’s chi-square test (Q-test) under the null hypothesis of homogeneity (significant heterogeneity P < 0.05). Individual SNPs were investigated using allele- and genotype-based models.

### Table 2 Molecular definition of drug-resistance according to participating countries

| Country       | Chloroquine | SP                 | Other                  |
|---------------|-------------|--------------------|------------------------|
| Burkina Faso  | Pfcrt76T    | Dhfr51V/S/59R/108N | AQ: Pfcr76T + Pfmdr1-86Y |
| Cameroon (Yaoundé) | r/a        | Dhfr51V/S/59R/108N + Dhps437G |                     |
| Kenya         | Pfcrt76T    | Dhfr108N + one or more of Dhfr51, Dhps59, Dhps436A, Dhps540E |                     |
| Mali          | Pfcrt76T    | Dhfr51V/S/59R/108N | AQ: Pfcr76T + Pfmdr1-86Y |
| Sudan         | Pfcrt76T + Pfmdr1-86Y | Dhfr51V/S/59R/108N + Dhps437G |                     |

Mutations in Pfcrt and Pfmdr1 were considered for resistance to chloroquine, and in dhfr and dhps for resistance to SP. SP = sulfadoxine-pyrimethamine. AQ = amodiaquine. r/a = not applicable.

http://www.malariajournal.com/content/10/1/250
## Table 3 Polymorphisms genotyped using SEQUENOM® iPLEX®

| Alternative Name* | rsnumber | gene | chr | coord | ancestral/reference allele | derived allele |
|-------------------|----------|------|-----|-------|---------------------------|---------------|
| Duffy - FyA/FyB   | rs2814778| DARC | 1   | 159174683 | T                         | C             |
|                   | rs2179652| RGS2 | 1   | 192769826 | T                         | C             |
|                   | rs3024500| IL10 | 1   | 206940831 | G                         | A             |
|                   | rs1800890| IL10 | 1   | 206946897 | T                         | C             |
|                   | rs1800890| IL10 | 1   | 206949365 | A                         | T             |
| McC (McCoy)       | rs17047660| CR1 | 1   | 207782856 | A                         | G             |
| SI (Swain-Lagley) | rs17047661| CR1 | 1   | 207782889 | A                         | G             |
| IL1A G485T        | rs17561 | IL1A | 2   | 113537223 | C                         | A             |
| IL1B A2           | rs1143634| IL1B | 2   | 113590390 | G                         | A             |
|                   | rs708567 | IL17RE | 3 | 9660070 | C                         | T             |
|                   | rs352140 | TLR9 | 3   | 52231737 | T                         | G             |
|                   | rs187084 | TLR9 | 3   | 52261031 | G                         | A             |
|                   | rs6780995| IL17RD | 3 | 57138419 | G                         | A             |
| IL-4-589          | rs2243250| IL4  | 5   | 132009154 | C                         | T             |
| ILTC +77          | rs2239704| LTA  | 6   | 31540141 | C                         | A             |
| ILTNCO1           | rs909253 | LTA  | 6   | 31540313 | A                         | G             |
| TNFa -1031        | rs1799964| TNF  | 6   | 31542308 | T                         | C             |
| TNF -376          | rs1800750| TNF  | 6   | 31542963 | G                         | A             |
| TNF -308          | rs1800629| TNF  | 6   | 31543051 | G                         | A             |
| TNF -238          | rs361525 | TNF  | 6   | 31543101 | G                         | A             |
| TNF +4851         | rs3093662| TNF  | 6   | 31544189 | A                         | G             |
|                   | rs2242665| CTL4 | 6   | 31839309 | C                         | T             |
|                   | rs1555498| IL20RA | 6 | 137325847 | C                         | T             |
|                   | rs2075820| NOD1 | 7   | 30492237 | C                         | T             |
| CD36 T1264G       | rs3211938| CD36 | 7   | 80300449 | G                         | T             |
| CD36 G1439C       | None assigned | CD36 | 7 | 80302110 | G                         | C             |
|                   | rs17140229| CFTR | 7   | 117230283 | T                         | C             |
|                   | rs4986790| TLR4 | 9   | 120475302 | A                         | G             |
|                   | rs4986791| TLR4 | 9   | 120475602 | C                         | T             |
|                   | rs8176746| ABO  | 9   | 136131322 | G                         | T             |
| HbE               | rs33950507| HBB | 11  | 5248173 | C                         | T             |
| Hbs               | rs334 | HBB | 11  | 5248232 | T                         | A             |
|                   | rs7935564| TRIM5 | 11 | 5718517 | G                         | A             |
|                   | rs542998 | RTN3 | 11  | 63487386 | T                         | C             |
|                   | rs2227507| IL22 | 12  | 68642647 | T                         | C             |
|                   | rs1012356| IL22 | 12  | 68644618 | A                         | T             |
|                   | rs2227491| IL22 | 12  | 68646521 | T                         | C             |
|                   | rs2227485| IL22 | 12  | 68647713 | G                         | A             |
|                   | rs2227478| IL22 | 12  | 68648622 | G                         | A             |
|                   | rs229587 | SPTB | 14  | 65263300 | T                         | C             |
A total of 3,721 samples from five countries (Table 4) were found to contain "genotypically resistant" parasites according to the criteria in Table 2. Of these patients, 2,057 (55%) were able to successfully clear their infection. With the exception of Kenya (47.7% cleared resistant infections), more than 50% of individuals in the study were able to clear genotypically resistant parasites (47.7%-76.4%).

In contrast to previous studies, there was no significant difference in age overall between those patients who successfully cleared their infection (median 5 years) compared to those who did not (median 5.3 years; K-S test = 47.0; P = 0.16; Table 4). This could be because the data were pooled from five countries with different levels of acquired immunity, and involving different age groups according to the study design chosen. Individuals from highly malaria-endemic areas would be expected to have a higher potential to clear parasites than much older individuals from less endemic areas, so the influence of age is masked by pooling.

There was no difference in the gender of patients who successfully cleared their infection and those who did not in Cameroon ($\chi^2$ test, P = 0.69) and Kenya ($\chi^2$ test, P = 0.16). However, in Burkina Faso and Sudan, significantly fewer males and more females than expected successfully cleared a drug resistant infection ($\chi^2$ tests: BF: P = $2.5 \times 10^{-48}$, RR = 0.2; Sudan P = $2.2 \times 10^{-7}$, RR = 0.74). By contrast, in Mali significantly more males and fewer females than expected were able to cure a drug resistant infection ($\chi^2$ test, P = $3.6 \times 10^{-6}$, RR = 1.87). These apparent gender effects could however be the result of significant differences in the age of male and female participants in some countries in the study. In Burkina Faso and Sudan, the median age of females was significantly higher than that of males (K-S test, P = 0.04 (BF); P = 0.02 (Sudan), whereas in Mali, male participants were older, although this did not quite reach statistical significance (K-S test, P = 0.06). Previous studies suggest that overall, older children are more likely to clear drug-resistant infections than younger children [10].

The parasitaemia at admission to the study in those who cleared and did not clear their infections was not significantly different for any of the five countries (K-S tests: Burkina Faso (P = 0.18), Cameroon (P = 0.84), Kenya (P = 0.65), Mali (P = 0.37), and Sudan (P = 0.29)).

### Table 3 Polymorphisms genotyped using SEQUENOM® iPLEX® (Continued)

| rs  | Polymorphism | Gene | Chromosome | Position | Allele 1 | Allele 2 |
|-----|--------------|------|------------|----------|----------|----------|
| rs2230739 | ADCY9 | 16 | 4033436 | T | C |
| rs10775349 | ADCY9 | 16 | 4079823 | C | G |
| rs1805015 | ILAR | 16 | 27374180 | T | C |
| rs2535611 | ADORA2B | 17 | 15861332 | C | T |
| rs2297518 | NOS2 | 17 | 26096597 | G | C |
| rs1800482 | NOS2 | 17 | 26128509 | C | G |
| rs9282799 | NOS2 | 17 | 26128728 | A | G |
| rs2195909 | NOS2 | 17 | 26129212 | A | G |
| rs373533 | EMR1 | 19 | 6919624 | C | A |
| rs461645 | EMR1 | 19 | 6919753 | A | G |
| rs1799969 | ICAM1 | 19 | 10394792 | G | A |
| rs5498 | ICAM1 | 19 | 10395683 | A | G |
| rs2057291 | GNAS | 20 | 57472043 | C | T |
| rs8386 | GNAS | 20 | 57485812 | C | T |
| rs1128127 | DERL3 | 22 | 24179132 | G | A |
| Amelogenin_SNP1 | None assigned | AMELX | X | 11313735 | G** | A*** |
| Amelogenin_SNP2 | None assigned | AMELX | X | 11316106 | T** | C*** |
| Amelogenin_SNP6 | None assigned | AMELX | X | 11316650 | A** | A*** |
| CD40LG -727 | CD40LG | X | 135729609 | T | C |
| CD40LG +220 | CD40LG | X | 135730555 | T | C |
| G6PD +376 | G6PD | X | 154110298 | T | C |
| G6PD +202 | G6PD | X | 154111023 | C | T |

All SNPs are referenced to dbSNP130 and Ensembl build 56.*Alternative name from the literature or from laboratory usage. ‡Ancestral alleles are taken from dbSNP130 and where not identified a reference allele is given based on the human reference sequence on Ensembl. All alleles are with respect to the positive strand. ** Allele represented on the x chromosome and *** Allele represented on the Y chromosome.
| Country   | Burkina Faso | Cameroon | Kenya | Mali | Sudan | Total |
|-----------|--------------|----------|-------|------|-------|-------|
|           | Cleared | Not cleared | Total | Cleared | Not cleared | Total | Cleared | Not cleared | Total | Cleared | Not cleared | Total | Cleared | Not cleared | Total |
| Number of samples | 264 | 235 | 499 | 517 | 1247 | 656 | 718 | 1374 | 115 | 104 | 219 | 292 | 90 | 382 | 2057 | 1664 | 3721 |
| Median age in years | 5.5 | 7.3 | - | 4 | 6 | - | 11 | 12 | - | 3 | 3 | - | 14 | 10 | 5 | 5.3 | - |
| Number male gender | 37 | 186 | 223 | 371 | 257 | 628 | 265 | 317 | 582 | 78 | 371 | 123 | 66 | 189 | 874 | 867 | 1741 |
| Number female gender | 227 | 49 | 276 | 359 | 260 | 619 | 391 | 401 | 792 | 37 | 66 | 103 | 169 | 24 | 193 | 1183 | 797 | 1980 |
| Parasitaemia: median parasite density (parasites per μl) | 19 960 | 21 070 | - | 27015 | 22075 | - | 22 160 | 21360 | - | 19 630 | 17215 | - | 23 360 | 24380 | - | 22 425 | 21 220 | - |
| Parasitaemia: range (parasites per μl) | 25 - 44 870 | 25 - 38 990 | - | 25 - 26 870 | 25 - 31 990 | - | 25 - 23 870 | 25 - 48 190 | - | 25 - 190 | 310 | - | 25 - 27 870 | 28 090 | - | 25 870 | 40 213 | - |
the application of meta-analysis (P < 0.05). An initial analysis of association of each of the 70 SNPs with clearance of drug resistant parasites revealed 17 SNPs those were significantly associated with the phenotype (Table 5 P ≤ 0.05). Three SNPs (rs1799969, rs1126535 and rs2814778) showed a strong association with the clearance phenotype with p-values less than 10⁻⁵ (Table 5).

Further analysis using genotype-based tests (Table 6) indicated a highly significant association (P < 0.01) of 9 SNPS with the clearance of resistant parasites. 19 of the 25 SNPs showed a significant association (P < 0.05) with clearance (Table 6). After adjusting the data for age, ethnicity, gender and study location, three SNPs remained significantly associated with the clearance phenotype: SNP rs2706384 (OR = 0.76 [95%, CI: 0.64 - 0.92]; p = 0.005), SNP rs1128127 (OR = 0.77 [95%, CI: 0.59 - 0.99], p = 0.05), and SNP rs2057291 (OR = 1.27 [95%, CI: 1.02 - 1.57], p = 0.03). No other SNPs were statistically associated with the clearance phenotype (Table 6).

Multiple SNP analysis
A multiple SNP analysis was performed in order to correct for covariate effects (multiple SNPs adjusted for ethnic group, age, site, and gender) (Table 6). The main predictive factor of clearance was age as has been reported previously by others [10,41-43]. Two of the three SNPs identified in the single SNP analysis remained significantly associated with the clearance phenotype: SNP rs2706384 (OR = 0.76 [95%, CI: 0.51 - 0.99]; p = 0.005) and SNP rs1128127 (OR = 0.67 [95%, CI: 0.45 - 0.99], p = 0.05). One additional SNP (rs1805015) was now found to be associated with the clearance phenotype (OR = 0.66 [95% CI: 0.45 - 0.97], p = 0.03).

Table 5 Univariate allele-based association tests

| SNP* | Allele 1/2 | Clearance | Non-clearance | Chi-squared | p-value |
|------|-----------|-----------|---------------|-------------|---------|
|      |           | Allele 1 | Allele 2 | Allele 1 | Allele 2 |       |         |
| rs1012356 | A/T       | 0.53     | 0.47   | 0.49     | 0.51     | 11.38 | 0.007   |
| rs2227491 | C/T       | 0.57     | 0.43   | 0.60     | 0.40     | 7.02  | 0.008   |
| rs2227485 | A/G       | 0.44     | 0.56   | 0.47     | 0.53     | 10.9  | 0.0009  |
| rs2227478 | A/G       | 0.64     | 0.36   | 0.66     | 0.34     | 5.89  | 0.02    |
| rs2706384 | A/C       | 0.41     | 0.59   | 0.46     | 0.54     | 18.9  | 0.00001 |
| rs2057291 | A/G       | 0.19     | 0.81   | 0.19     | 0.81     | 0.17  | 0.68    |
| CD36 G1439C | C/G     | 0.01     | 0.99   | 0.01     | 0.99     | 1.49  | 0.22    |
| rs1799969 | A/G       | 0.04     | 0.96   | 0.06     | 0.94     | 15.83 | 7.10*   |
| rs20541  | C/T       | 0.76     | 0.24   | 0.74     | 0.26     | 6.86  | 0.009   |
| rs1800750 | A/G       | 0.05     | 0.95   | 0.05     | 0.95     | 0.70  | 0.40    |
| rs3024500 | A/G       | 0.62     | 0.38   | 0.63     | 0.37     | 2.10  | 0.15    |
| rs1805015 | C/T       | 0.37     | 0.63   | 0.36     | 0.64     | 1.39  | 0.24    |
| rs17047660 | A/G      | 0.72     | 0.28   | 0.74     | 0.26     | 8.13  | 0.004   |
| rs17047661 | A/G    | 0.42     | 0.58   | 0.45     | 0.55     | 7.20  | 0.007   |
| rs1714022 | C/T       | 0.29     | 0.71   | 0.27     | 0.73     | 2.80  | 0.09    |
| rs1126535 | C/T       | 0.22     | 0.78   | 0.28     | 0.72     | 31.13 | 10*     |
| rs2230739 | A/G       | 0.86     | 0.14   | 0.86     | 0.14     | 0.54  | 0.46    |
| rs229587  | C/T       | 0.39     | 0.61   | 0.42     | 0.58     | 4.37  | 0.04    |
| rs2814778 | A/G       | 0.14     | 0.86   | 0.20     | 0.80     | 55.17 | 10*     |
| rs3092945 | C/T       | 0.32     | 0.68   | 0.29     | 0.71     | 10.69 | 0.001   |
| rs1128127 | A/G       | 0.44     | 0.56   | 0.47     | 0.53     | 8.19  | 0.004   |
| rs1803632 | C/G       | 0.51     | 0.49   | 0.53     | 0.47     | 4.84  | 0.03    |
| rs7935564 | A/G       | 0.46     | 0.54   | 0.45     | 0.55     | 2.39  | 0.12    |
| rs4833095 | C/T       | 0.84     | 0.16   | 0.82     | 0.18     | 5.23  | 0.02    |
| rs5743809 | C/T       | 0.06     | 0.94   | 0.05     | 0.95     | 4.92  | 0.03    |

For each SNP, chi-squared comparisons were made of between the allele frequencies found in patients who cleared and did not clear drug resistant parasites. Highly significant associations are highlighted in bold. *Of the 67 SNPs genotyped for association, 42 SNPs were not included in the analysis because they were either monomorphic in one or more countries (n = 19) or for deviation from HWE in one or more of the participant countries (n = 23).
## Table 6 Genotype association analysis

| SNP (genotype model) | Genotype-based tests* | Adjusted analysis** | Multiple SNP analysis*** |
|----------------------|------------------------|---------------------|-------------------------|
|                      | Chi-squared | p-value | OR [95%, CI] | p-value | OR [95%, CI] | p-value |
| rs1012356 (TT vs AT/AA) | 17.20       | 0.001   | 1.15 [0.90 - 1.46] | 0.27 | 1.23 [0.95 - 1.58] | 0.09 |
| rs2227491 (TT vs CT/CC) | 6.90       | 0.07    | 1.04 [0.80 - 1.34] | 0.78 | 0.97 [0.68 - 1.39] | 0.88 |
| rs2227485 (AA vs GG/AG) | 11.30      | 0.01    | 1.04 [0.82 - 1.32] | 0.74 | 0.89 [0.64 - 1.25] | 0.50 |
| rs2227478 (GG vs AA/AG) | 6.60       | 0.09    | 1.14 [0.86 - 1.51] | 0.36 | 1.21 [0.82 - 1.80] | 0.34 |
| rs2706384 (AA vs CC/AC) | 21.80      | 0.0001  | 0.76 [0.64 - 0.92] | 0.005 | 0.76 [0.71 - 0.92] | 0.005 |
| rs2057291 (AA vs GG/AG) | 11.90      | 0.02    | 1.27 [1.02 - 1.57] | 0.03 | 0.91 [0.71 - 1.17] | 0.47 |
| CD36 G1439C (CC vs GG/CG) | 3.30 | 0.03  | - | - | - | - |
| ICAM1 CODON241 (AA vs GG/AG) | 30.03 | 0.0001 | 0.98 [0.71 - 1.36] | 0.92 | 1.04 [0.74 - 1.44] | 0.84 |
| rs20541 (TT vs CC/CT) | 13.40      | 0.004   | 1.17 [0.89 - 1.51] | 0.56 | 0.95 [0.70 - 1.31] | 0.87 |
| TNF -376 (AA vs GG/AG) | 11.30      | 0.01    | 1.04 [0.86 - 1.25] | 0.79 | 1.10 [0.87 - 1.38] | 0.60 |
| RS1805015 (CC vs TT/CT) | 8.70       | 0.03    | 0.84 [0.65 - 1.08] | 0.21 | 0.66 [0.45 - 0.97] | 0.03 |
| rs17047660 (GG vs AA/AG) | 7.20       | 0.07    | 0.95 [0.72 - 1.23] | 0.68 | 1.32 [0.84 - 2.07] | 0.23 |
| rs17047661 (AA vs GG/AG) | 7.40       | 0.06    | 0.88 [0.66 - 1.16] | 0.36 | 0.72 [0.49 - 1.05] | 0.09 |
| rs17140229 (CC vs TT/CT) | 8.60       | 0.04    | 1.08 [0.79 - 1.46] | 0.62 | 1.41 [0.94 - 2.10] | 0.09 |
| rs1126535 (CC vs TT/CT) | 21.60      | 0.0001  | 0.91 [0.67 - 1.22] | 0.52 | 0.85 [0.51 - 1.40] | 0.51 |
| rs229587 (CC vs TT/CT) | 10.40      | 0.02    | 1.06 [0.80 - 1.39] | 0.69 | 1.26 [0.82 - 1.92] | 0.29 |
| rs2814778 (AA vs GG/AG) | 8.50       | 0.04    | 0.97 [0.76 - 1.22] | 0.76 | 1.02 [0.73 - 1.43] | 0.92 |
| rs3092945 (CC vs TT/CT) | 33.03      | 0.0001  | 0.82 [0.53 - 1.28] | 0.38 | 0.85 [0.53 - 1.36] | 0.50 |
| rs1128127 (AA vs GG/AG) | 12.04      | 0.007   | 0.77 [0.59 - 0.99] | 0.05 | 0.67 [0.45 - 0.99] | 0.05 |
| rs1803632 (GG vs CC/CG) | 7.04       | 0.071   | 0.78 [0.58 - 1.05] | 0.10 | 0.76 [0.49 - 1.17] | 0.21 |
| rs7935564 (AA vs GG/AG) | 21.10      | 0.0001  | 1.02 [0.84 - 1.25] | 0.81 | 0.93 [0.70 - 1.22] | 0.59 |
| rs4833095 (TT vs CC/CT) | 6.90       | 0.08    | 0.75 [0.52 - 1.07] | 0.12 | 0.86 [0.57 - 1.33] | 0.50 |
| rs5743809 (CC vs TT/CT) | 11.40      | 0.01    | 1.25 [0.57 - 2.74] | 0.59 | 1.23 [0.53 - 2.84] | 0.63 |

Highly significant associations (P < 0.01) are highlighted in bold; significant associations (0.01 < P < 0.05) are underlined. OR = odds ratio; *Single locus based tests; **Each SNP adjusted for age, ethnic group, gender, and study location. ***Multiple SNPs adjusted for ethnic group, age, gender, and study location.
Discussion
The malaria parasite has had a substantial evolutionary influence upon the genetic constitution of its human host (recently reviewed in [44]). Individuals living in malaria-endemic regions seem to develop an ability to clear drug-resistant parasites (following treatment) as they get older [10], which is presumably the result of increasing acquired immunity. The influence of human host polymorphisms in immune response-type genes on the likelihood of clearance of drug-resistant parasites has so far received little attention.

In the present study, a single SNP locus analysis was carried out to investigate the contribution of the host genetic factors in the clearance of drug-resistant parasites following treatment, by examining the classic, previously published SNPs, which may play a critical role in individuals’ ability to clear drug-resistant malaria parasites. The effects of polymorphisms in a number of genes, including β-globin, G6PD, TNF-α, IFN-γ, CD36, ICAM-1, IL10, IL4R, and LTA (Table 3), upon the clearance of malaria parasites in African individuals was investigated across five large association studies from Burkina Faso, Cameroon, Kenya, Mali, and Sudan.

Amongst the 70 SNPs investigated in this study, seventeen were found at significantly different frequencies (P < 0.05) in people who cleared drug resistant infections than those who did not (Table 5). Further analysis using genotype-based tests indicated that nine SNPs were strongly associated (P < 0.01) with parasite clearance (Table 6). Following adjustments for the possible confounding factors of age, ethnicity, gender and study location, and analysing multiple SNPs to correct for covariate effects, three SNPs remained significantly associated with the clearance phenotype, across Africa and with three different drugs. It is, however, important to note that the demonstration of association with clearance phenotype of these three SNPs does not necessarily imply that any of the SNPs are functional in the clearance of drug-resistant parasites. The SNPs may simply be reflecting the signal of a different functional variant(s) in moderate-to-high LD with them [45].

SNP rs2706384 is in the 5’ upstream region of the interferon regulatory factor IRFI gene, 1710 bp upstream of the ATG start codon and -415bp from the transcriptional start site. Individuals with a homozygous AA genotype at this locus were significantly less likely to clear drug resistant infections than those homozygous CC or heterozygous (OR = 0.76 [95% CI: 0.71 - 0.92]; P = 0.005). The A allele was found more frequently in individuals who did not clear their drug-resistant infection than in those who did (P = 0.00001).

IRF-1 is a transcription factor that has been shown to regulate expression of a number of genes involved in both innate and adaptive immunity, notably TLR9, MHC Class I and II genes, IL-15, iNOS in macrophages, IL-4 and IL-12/p40 [46]. Interferon-γ, the strongest inducer of IRF-1, is thought to be a key player in the control of pre-erythrocytic and blood stage infection, both in rodent malaria infections [47] and in human malaria infections [48,49]. Healthy individuals homozygous AA at rs2706384 were found to have significantly higher IRF-1 mRNA expression than CC homozygotes [50]. Thus individuals of the AA genotype may produce higher levels of IRF-1 in response to the same IFN-γ stimulus, which may shift the balance more towards a Th1 response and away from a Th2 response through repression of IL-4 transcription and increased IL-12/p40 expression. This suggests that antibody may play a key role in the control of drug-resistant parasites. In addition, the binding of NF-kappa B to the C allele was significantly higher than to the A allele [50]; this transcription factor may have a negative regulatory role in IFN-induced gene expression [51,52].

Previous work has shown an association of the same SNP, rs2706384, with protection against P. falciparum infection in two West African ethnic groups [53]. However, in that study the C allele was associated with a higher risk of having a P. falciparum infection for Mossi but not for Fulani, and in Fulani CC and AA individuals were more frequently parasitized than heterozygous individuals.

Rs1805015 is a missense mutation (Ser503Pro) within the insulin-IL4 receptor motif (I4R) of the alpha subunit of the interleukin4 receptor gene IL4R. Individuals with a homozygous CC genotype (encoding 503Pro) at this locus were significantly less likely to clear drug resistant infections than those who were homozygous TT or heterozygous (OR = 0.66 [95%, CI: 0.45 - 0.97], p = 0.03), but there was no significant difference in the frequency of the C allele in individuals who cleared or did not clear their drug-resistant infection (P = 0.24), suggesting that the failure to clear infections was associated with the CC homozygote.

The interaction of IL-4 with its receptor results in binding of JAK to the I4R motif of IL4R-α; however, this binding is unaffected by the Ser503Pro substitution [54]. The Ser503Pro substitution appears to reduce the subsequent binding and phosphorylation of STAT6 [54]. Since phosphorylated STAT6 controls cell differentiation and gene transcription [55], the Ser503Pro substitution could therefore lead to a reduction in response to IL-4, such as reduced B-cell proliferation and antibody production, further supporting the role of Th2 responses in the clearance of drug-resistant parasites.

However, a previous malaria case-control study in Sudan found the CC genotype to be at a significantly
lower frequency in malaria cases compared to non-malaria controls in Sudan [56], whereas in this study CC genotypes were less able to clear drug resistant infections than individuals of genotype AA or AC. Individuals with the Ser503Pro IL4R mutation were found to have lower IgE levels [54] and a separate study found a significant association with atopy and asthma-related phenotypes [57].

SNP rs1128127 is a missense mutation (Ala211Val) within the Der1-like domain family gene Derl3. Individuals with a homozygous AA genotype (encoding 211Val) at this locus were significantly less likely to clear drug resistant infections than those who were homozygous GG or heterozygous (OR = 0.77 [95%, CI: 0.45 - 0.99], P = 0.05). The A allele was also found more frequently in individuals who did not clear their drug-resistant infection than in those who did (P = 0.004). This suggests that GG or AG individuals have an advantage over AA genotypes in their ability to control drug-resistant infections.

The derlin family of proteins are found in the endoplasmic reticulum (ER) and are thought to be involved in the degradation of misfolded glycoproteins within the ER [58-61]. Derl3 is expressed at high levels in specific tissues such as the placenta, pancreas, small intestine and spleen, whereas other members of the family have more widespread expression [61]. There does not appear to be any previous study linking mutations in Derl3 to the control of infectious disease. The frequency of heterozygous AG individuals is much higher in sub-Saharan Africans (0.65) than in Europeans (0.183) [62], which could be explained by positive selection of heterozygous AG individuals in populations exposed to malaria, because of their enhanced ability to clear (drug-resistant) parasites.

Host genetic factors such as cytokines may be the key determinants of malaria severity and outcome. Several studies suggest that the balance between pro- (TNF-α, IFN-γ, IL-8) and anti-inflammatory (IL4, IL-10, TGF-β) cytokines determines the degree of malaria parasitaemia, the level of anaemia, the clinical severity, the presentation, and/or the outcome of infection [63-65]. IFN-γ has been suggested to be a key molecule in human anti-parasite host defence, and appears to be essential for the control of parasitaemia. The role of IL4 is less clear; some studies have not supported direct involvement of IL-4 (or IL-13) in the clearance of P. falciparum parasites [63], and IL-4 has been shown to suppress macrophage-mediated killing of P. falciparum in vitro [66].

This study is the first to assess the role of specific human genetic variants (SNPs) in the clearance of drug-resistant parasites after anti-malarial treatment. Three SNPs were found to be strong predictors of the clearance of drug-resistant parasites, even after correction for age, ethnicity, gender and study location. Two of the three SNPs identified are in loci associated with pro-inflammatory (interferon-γ) and anti-inflammatory (IL-4) cytokine responses.

The assessment of the role of human genetic determinants may improve understanding of the interface between host immunity and anti-malarial drug resistance. The relationship between host polymorphisms and malaria parasite clearance is complex, and larger studies in other settings will be required, both to confirm these associations, to investigate further the weak associations, and also to investigate the contribution of the host immunological factors and the parasite per se in the clearance of drug-resistant parasites.

Conclusions

The study has identified a significant association of three loci in the human genome with the ability of parasite to clear drug-resistant P. falciparum. One locus, a SNP in the promoter region of the IRF-1 gene, has previously been linked to the control of malaria parasite density, and it is possible that patients able to clear drug-resistant infections have an enhanced ability to control parasite growth, perhaps through a more Th2-biased T cell response. The association of clearance with a SNP within the IL-4R gene, that possibly reduces the response to IL-4, supports the hypothesis that a stronger Th2 response assists clearance of drug-resistant parasites. The third locus encodes a protein involved in the degradation of misfolded proteins within the endoplasmic reticulum, and its role, if any, in the clearance phenotype needs to be further investigated.

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Author details

1Malaria Research and Training Centre, Faculty of Medicine, Pharmacy and Odontostomatologyp, University of Bamako, Mali. 2University of Buea, Faculty of Health Sciences, Department of Medical Laboratory Science, B.P. 63 Buea, S.W.P. Province, Cameroon. 3Université de Yaounde I, Centre de Biotechnologie, B.P. 8094, Yaounde, Cameroon. 4Wellcome Trust Centre for Human Genetics, University of Oxford, Oxford, UK. 5Department of Molecular Biology, Institute of Endemic Diseases, University of Khartoum, Khartoum, Sudan. 6International Atomic Energy Agency (IAEA), Vienna, Austria. 7Kenya Medical Research Development, Malaria Unit, Centre for Biotechnology Research & Development, Mb bunch Road, P.O. Box 54840, Nairobi, Kenya. 8Wellcome Trust Sanger Institute, Hinxton, Cambridgeshire, CB10 1SA, UK. 9Direction régionale de l’Odontostomatologie, University of Bamako, Mali.

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Burkina Faso. 7 National Ministry of Health, National Health Laboratory, Department of Parasitology, P.O. Box 287, Khartoum, Sudan. 8 Institute of Infection, Immunity and Inflammation, College of Medical, Veterinary and Life Sciences, University of Glasgow, 120 University Place, Glasgow G12 8QQ, UK.

Authors’ contributions
Data were collected, and molecular genotyping for drug resistance markers performed, by individual teams from Burkina Faso (UBO, IZ), Cameroon (WFM, OA, MSBE, EAA), Kenya (FK, SOJ), Mali (MKW, AD) and Sudan (NT, IM). Human polymorphisms were genotyped by MD, AG, CH, AJ, KR, KR, DPK. LRC supervised molecular drug resistance genotyping and provided positive controls and SOPs. MD provided SOPs for human polymorphisms genotyping and data analysis. MD, LRC, DPK, BK, KR conceived of the study and participated in its design and coordination. MD and LRC drafted the manuscript, with additional comments and input from DPK, BK and KR. All authors read and approved the final manuscript.

Competing interests
The authors declare that they have no competing interests.

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