Emergence of artemisinin-resistant malaria on the western border of Thailand: a longitudinal study

Aung Pyae Phyo, Standwell Nkhoma, Kasia Stepniewska, Elizabeth A Ashley, Shalini Nair, Rose McGready, Cant Ier Moo, Salma Al-Saai, Arjen M Dondorp, Khin Maung Lwin, Pratap Singhasivanon, Nicholas P J Day, Nicholas J White, Tim J C Anderson, François Nosten

Summary

Background Artemisinin-resistant falciparum malaria has arisen in western Cambodia. A concerted international effort is underway to contain artemisinin-resistant *Plasmodium falciparum*, but containment strategies are dependent on whether resistance has emerged elsewhere. We aimed to establish whether artemisinin resistance has spread or emerged on the Thailand–Myanmar (Burma) border.

Methods In malaria clinics located along the northwestern border of Thailand, we measured six hourly parasite counts in patients with uncomplicated hyperparasitaemic falciparum malaria (≥4% infected red blood cells) who had been given various oral artesunate-containing regimens since 2001. Parasite clearance half-lives were estimated and parasites were genotyped for 93 single nucleotide polymorphisms.

Findings 3202 patients were studied between 2001 and 2010. Parasite clearance half-lives lengthened from a geometric mean of 2·6 h (95% CI 2·5–2·7) in 2001, to 3·7 h (3·6–3·8) in 2010, compared with a mean of 5·5 h (5·2–5·9) in 119 patients in western Cambodia measured between 2007 and 2010. The proportion of slow-clearing infections (half-life ≥6·2 h) increased from 0·6% in 2001, to 20% in 2010, compared with 42% in western Cambodia between 2007 and 2010. Of 1583 infections genotyped, 148 multilocus parasite genotypes were identified, each of which infected between two and 13 patients. The proportion of variation in parasite clearance attributable to parasite genetics increased from 30% between 2001 and 2004, to 66% between 2007 and 2010.

Interpretation Genetically determined artemisinin resistance in *P falciparum* emerged along the Thailand–Myanmar border at least 8 years ago and has since increased substantially. At this rate of increase, resistance will reach rates reported in western Cambodia in 2–6 years.

Funding The Wellcome Trust and National Institutes of Health.

Introduction Artemisinin combination treatments are the recommended first-line therapy for falciparum malaria. *Plasmodium falciparum* parasites with reduced in-vivo susceptibility to artemisinin derivatives (eg, artesunate) have emerged in western Cambodia.1 This finding threatens worldwide initiatives to control and eliminate malaria.4 Resistance to the previous mainstays of anti-malarial treatment—namely, chloroquine and sulfadoxine–pyrimethamine—also arose in western Cambodia5–7 and spread across southeast Asia into Africa, resulting in the deaths of millions of children.8 If resistance to artemisinin is confined to the Cambodia–Thailand border, regional elimination of falciparum malaria will probably be necessary for containment.8 However, if resistant parasites have already spread or emerged elsewhere, then the containment zone will need to be extended and the strategy reconsidered.9

Artemisinin resistance is characterised by slow parasite clearance.10,11 Clearance (assessed by microscopy) of sensitive *P falciparum* is achieved within 2 days in 95% of patients,1 whereas artemisinin-resistant infections remain slide-positive for 3 or more days; treatment failure is more common in such infections after artemisinin combination treatment. High-grade artemisinin resistance has not been reported. Laboratory studies and mathematical modelling suggest that slow clearance of resistant parasites mainly results from decreased susceptibility of ring-stage parasites to artemisinin and its derivatives.4 No laboratory assays reliably identify artemisinin-resistant parasites.12 Patients from western Cambodia with similar parasite clearance rates are infected with genetically indistinguishable parasite clones, suggesting that parasite genetics have a central role in determination of this trait,15 although the genes responsible are unknown.

On the northwestern border of Thailand, 800 km from western Cambodia, treatment failure rates with artesunate and mefloquine combinations have increased.6 In this area, patients without signs of severity but with more than 4% parasitaemia (ie, parasites present in 4% or more of red blood cells) are given oral artesunate regimens, which have proven superior to intravenous quinine.24 High admission parasitaemias allow for accurate assessment of parasite clearance rates. To assess whether artemisinin resistance has emerged in this strategically important area, we analysed over 10 years the longitudinal changes in parasite clearance rates in a large prospective series of hyperparasitaemic patients given artesunate. Parasites were genotyped to establish the contribution of parasite genetics to parasite clearance rates.
region of the northwestern border of Thailand. Maela (refugee camp) and Wang Pha village are north of Mae Sot town (where the research unit is based), whereas Mae Kon Khen and Mawker Thai villages are south (appendix). Most patients came from adjacent Myanmar (Burma). We compared the parasite clearance data with results from 119 patients studied in Pailin, western Cambodia (2007–10), where artemisinin resistance has been confirmed. A subset of 30 parasite genotypes from Pailin was part of the genetic analysis. These studies were approved by the ethics review boards of the Faculty of Tropical Medicine, Mahidol University.

Falciparum malaria was diagnosed by microscopy of thick and thin peripheral blood smears stained with Giemsa. Parasite counts were read per 1000 red cells (thin film) or 500 white cells (thick film). Uncomplicated hyperparasitaemia was defined as 4% or more of red cells infected with malaria parasites without clinical evidence of severe malaria, and was the only criterion for inclusion in the analysis. Routine care was hospitalisation with a six-hourly blood smear to monitor parasite clearance until smears were slide-negative. Treatment was with a 7-day regimen of oral artesunate (4 mg/kg initially, then 2 mg/kg once daily for 7 days), usually combined with either mefloquine (25 mg/kg in two divided doses), or doxycycline (4 mg/kg per day for 7 days) or clindamycin (5 mg/kg three times daily for 7 days) if mefloquine was contraindicated. In cases of clinical deterioration parenteral treatment was substituted. We monitored treatment until smears became slide-negative.

### Procedures

DNA was extracted from admission blood spots by use of a two-step protocol to maximise DNA yield. Blood was eluted from six 3 mm diameter punches with the GenSolve kit (GenVault Corporation, Pleasanton, CA, USA), and DNA was then extracted with QIAamp 96 DNA blood kits (Qiagen, Valencia, CA, USA). Genotyping of 96 single nucleotide polymorphisms distributed across the *P falciparum* genome was done with the Illumina Goldengate platform (Illumina Inc, San Diego, CA, USA) (appendix). We judged samples to be multiple-clone infections if more than five single nucleotide polymorphisms showed heterozygous base calls.

We measured the heritability of parasite clearance half-life to assess the part that parasite genetics plays in determination of this trait. Multilocus parasite genotypes infecting two or more patients were identified. Such parasite genotypes are equivalent to identical twins reared apart and can be used to measure heritability of parasite clearance (panel 1).

The geometric mean parasite clearance half-life in patients with artemisinin-resistant falciparum malaria in western Cambodia was 6.2 h (n=36, 95% CI 5.7–6.6), corresponding to a log, normalised distribution with a mean of 1.8 (SD 0.22). Hyperparasitaemic patients on the northwestern border of Thailand in 2001 were used as a...
Reference artemisinin-sensitive population, with a mean log, parasite clearance half-life of 0·95 (0·33). The hypothesis that the changes in parasite clearance half-life in western Thailand resulted from emergence versus importation of parasites with the same resistance phenotype prevalent in western Cambodia was tested.

**Statistical analysis**

We assessed parasite clearance using a standardised fitting method that separates the variable initial lag phase, during which parasite counts level off or rise, from the subsequent log-linear decline. The slope of this phase was calculated and expressed as the parasite clearance half-life, which is the time taken for parasitaemia to fall by half during log-linear decline. Patients with parasite clearance curves showing a poor fit (R²<0·8) to the log-linear model were excluded from the analysis.

We used Stata (version 11) to examine the association between parasite clearance half-life and the following covariates: age, sex, history of malaria, treatment regimen (artesunate with mefloquine vs artesunate with other partner drugs or artesunate alone), date of admission, duration of fever before presentation (<3 days vs ≥3 days), admission parasitaemia, haematocrit (<30% vs ≥30%), gameto cyt- 

Univariate regression analysis of all listed covariates was done on log-transformed parasite clearance half-lives, followed by stepwise multivariate analysis (excluding parenteral rescue therapy). Variables that only significantly improved the final model (likelihood ratio test p<0·05) were included (tables 1, 2).

To assess heritability, variance of parasite clearance was compared within and between clonally identical parasites recovered from more than two patients. Heritability (HR) was estimated from the mean squares terms in ANOVA. HR was estimated for normalised parasite clearance and for residuals remaining after regression of clearance rates against patient and environmental factors.

We examined the hypothesis that changes in parasite clearance resulted from importation or emergence of parasites with the western Cambodian phenotype. The clearance rate distribution of a mixture of the western Cambodia and western Thailand populations was calculated as: π×normal([log(x)−1·8] / 0·22) + (1−π)×normal([log(x)−0·95] / 0·33), where π is the proportion of infections derived from the resistant population. For each year, we used the Kolmogorov–Smirnov test (50 tests for each year) to compare recorded distributions of log-transformed half-lives with these theoretical distributions (for π between 0·01 and 0·50).

**Role of the funding source**

The sponsor of the study had no role in study design, data collection, data analysis, data interpretation, or writing of the report. The corresponding author had full access to all the data in the study and final responsibility for the decision to submit for publication.
Results

Between July, 2001, and December, 2010, 3202 hyperparasitaemic patients were treated. Most patients were younger than 15 years (appendix). Most patients (2403) were given oral artesunate and divided doses of mefloquine. The remaining patients were given artesunate monotherapy (321 patients), artesunate in combination with either doxycycline (382 patients) or clindamycin (71 patients), or other combinations (25 patients). Patients who got blood transfusions within 24 hours of admission, or an incomplete course of or more than one parenteral dose of artesunate were excluded from the analysis. 2855 patients (1268 in Wang Pha, 975 in Mawker Thai, 467 in Maela, and 145 in Mae Khon Ken) remained eligible, of whom 1778 (62%) were men or boys. In 1759 of 2855 patients, artesunate monotherapy was given, or else the partner drug (mefloquine, doxycycline, or clindamycin) was given more than 48 h from the start of artesunate treatment. All patients survived, confirming the effectiveness of oral artesunate in uncomplicated hyperparasitaemia.18

Between 2001 and 2010, the median age of hyperparasitaemic patients increased slightly but significantly from 11 years to 15 years (p<0·0001). The proportion of patients developing severe malaria did not increase over time, nor did the ratio of severe to uncomplicated falciparum malaria cases, malaria mortality, nor the proportion of patients developing anaemia. We noted a significant increase in the proportion of patients with gametocytaemia on admission (p<0·0001; 7·5×10⁻⁶ test for trend).

Parasite clearance half-life lengthened from a geometric mean of 2·6 h (95% CI 2·5–2·7) in 2001, to 3·7 h (3·6–3·8) in 2010 (p<0·0001), compared with 5·5 h (5·2–5·9) in the 116 patients from western Cambodia between 2007 and 2010 (clearance data from three patients fitted poorly to the linear model [R²<0·8] and were thus excluded) (figures 1, 2A, table 1). We used a stringent threshold of a parasite clearance half-life of more than 6·2 h to categorise infections with slow or fast parasite clearance; the proportion of patients at our study centres with slow clearance rose from 0·6% in 2001, to 20% in 2010, compared with 42% of patients in western Cambodia in 2010 (figure 2B). Parasite clearance rates in patients given artesunate monotherapy or given artesunate alone for more than 48 h (n=1759) before receiving partner drugs were similar to those in patients who were given combination treatments immediately, suggesting that increasing resistance to the partner drug cannot explain the temporal changes reported.

No relation existed between parasite clearance half-life and age (p=0·439) or starting parasitaemia (p=0·598) in patients given artesunate monotherapy or given artesunate alone for more than 48 h (n=1759) before receiving partner drugs, nor in all patients (p=0·124 and p=0·569, respectively). The variable most strongly associated with increases in parasite clearance half-life was the date of patient admission (appendix). This trend was strongly significant in all clinics sampled except Mae Khon Ken, Thailand (table 2; appendix). Only 7·4% of the variation in half-life could be explained by other admission variables. Location had a significant effect on parasite clearance; infections in northern locations (ie, Maela and Wang Pha) had slower clearance than those sampled in the south (ie, Mawker Thai and Mae Khon Ken) (appendix; table 2).
Of 96 single nucleotide polymorphisms assessed, 93 gave robust genotype data. Of the 1583 infections genotyped, 1029 were single-clone infections (appendix). From these, 148 unique 93-locus parasite genotypes were identified, each infecting between two and 13 patients (29 from 2001 to 2004, and 119 from 2007 to 2010) (appendix). Patients harbouring parasites with the same 93-locus genotype had similar parasite clearance half-lives (figure 3), showing the important part played by parasite genotype. Between 2001 and 2004, parasite genotype had a significant effect on parasite clearance half-life (p=0·0166), even after adjusting for significant covariates (p=0·01). Two of the three patients with the longest parasite clearance half-lives recorded before 2004 had the same 93-locus genotype (figure 3). From 2007 to 2010, parasite genotype had a much stronger effect on parasite clearance half-life (p<0.0001) than it had between 2001 and 2004, and remained highly significant after adjusting for other covariates (F=5·219, df=118, p<0·0001). Mean \( H^2 \) increased from 0·30 (SD 0·14) in 2001–04, to 0·66 (0·04) in 2007–10. After adjustment for the effects of covariates, \( H^2 \) rose from 0·32 (0·14) between 2001–04, to 0·58 (0·05) in 2007–10 (appendix). The increase in \( H^2 \) was tested by permutation and was significant both before (p<0·01) and after (p=0·04) adjustment for covariates (appendix).

To investigate whether the increase in \( H^2 \) was driven by alleles conferring slow parasite clearance, we removed 30% of the slowest clearing infections from the 2007–10 dataset, which caused \( H^2 \) to decrease from 0·66 (SD 0·04) to 0·17 (0·07). Removal of 30% of the fastest clearing infections increased \( H^2 \) to 0·70 (0·05), whereas random removal of 30% of infections did not alter \( H^2 \) (mean \( H^2 \) from 100 randomisations 0·67 (0·03); range 0·56–0·74). These in-silico experiments show that high \( H^2 \) is driven by parasite genes that determine slow parasite clearance.

Admixture of parasites with the Cambodian phenotype could explain the parasite clearance distributions in 2001–02, and 2009–10, but not those between 2003 and 2008 (p<0·01). This finding suggests that the gradual increases in parasite clearance half-life are unlikely to be explained by a single-step process of importation or de novo selection of a parasite population with the same characteristics as the resistant parasites prevalent in western Cambodia.

None of the 93-locus genotypes was common to patients from the Thailand–Myanmar border and western Cambodia. Pairwise allele sharing (p9) between the 23 slowest and 25 fastest clearing parasites from the Thailand–Myanmar border was compared with 30 slow-clearing parasites from western Cambodia. Cambodian parasites were more closely related (p9=0·453) to slow-clearing parasites than to fast-clearing parasites (p9=0·450) from the Thailand–Myanmar border. However, permutation tests show that this difference is not significant (p9=0·137, 10 000 permutations).

Discussion

This large prospective series of patients with hyperparasitaemia during 10 years provides clear evidence that parasite clearance responses after artesunate treatment are slowing on the northwestern border of Thailand (panel 2). The rate of decline is consistent with theoretical expectations and empirical data for the spread of resistance loci through a parasite population under drug selection.\(^27\text{–}29\) If the rate of decline continues then parasite clearance half-lives on the Thailand–Myanmar border will be similar to those in western Cambodia within 2–6 years.

Parasite clearance after treatment with artesinin derivatives is determined mainly by drug effectiveness, although host immunity and the partner drug can also contribute.\(^21\) The frequency of malaria and the proportion of multiple-clone infections (appendix) in the study area in northwestern Thailand have fallen over the past 10 years; thus immunity to malaria has probably also declined. Because patients with hyperparasitaemia are unable to control their infection, changes in immunity are unlikely to have contributed substantially to the slowing of parasite clearance. The absence of any relation with age in this study provides strong support for this contention.

Similarly, although previous malaria exposure has a weak effect on parasite clearance half-life (table 1), this finding cannot explain the temporal trends noted (appendix). The artesinin combination treatment partner drug mefloquine has been used in this region for more than 25 years, and resistance developed rapidly after initial deployment. The proportion of \( P.\ falciparum \) isolates with increased \( pfmdr1 \) copy numbers (the main determinant of
continuing to drive down the incidence of malaria is of the highest priority.

Artemisinin resistance has been suspected because of diminishing treatment responses to artemisinin combination therapies on the western border of Thailand4,5 and changes in parasite clearance rates on the Kenyan coast.16

Interpretation

Although artemisinin resistance has been suspected outside western Cambodia, this study provides the first unequivocal proof that resistance has emerged or spread westward. It confirms (with a large sample size) findings showing that this phenotype is explained mainly by a heritable genetic trait in the parasite population. The research suggests a potential transmission advantage associated with resistance to artesunate through increased gametocyte carriage on admission. These findings suggest that regional artemisinin resistance-containment strategies should be reviewed. Increasing the availability of artemisinin combination therapies in southeast Asia will reduce the incidence of malaria but provides a selection pressure driving artemisinin resistance. The development of strategies to reduce the selection and spread of artemisinin-resistant Plasmodium falciparum while continuing to drive down the incidence of malaria is of the highest priority.

mefloquine resistance) has been increasing over the past decade.16 However, the temporal changes in parasite clearance half-lives for patients given artesunate monotherapy for more than 48 h before administration of partner drugs were closely related to those reported in the full dataset, showing that resistance to the partner drug does not contribute substantially to the changes recorded (figure 2).

The genotyping data provide compelling evidence that parasite genetic factors explain the fall in parasite clearance rates. In this low transmission setting, different patients are often infected with genetically very similar parasites, allowing assessment of the contribution of genetic factors to in-vivo phenotypes.14-16 After adjustment for covariates, the heritability of parasite clearance increased during the study period, from 32% between 2001 and 2004, to 58% between 2007 and 2010, which is consistent with an increasing prevalence of alleles determining slow parasite clearance.25 The increased contribution of genetics was confined largely to the parasites with the slowest clearance rates. The high heritability of slow parasite clearance suggests that this trait will spread rapidly under continued selection.

This study shows the importance of longitudinal detailed monitoring to detect the early emergence of antimalarial drug resistance. Hyperparasitaemic patients provide important information on temporal trends in artemisinin susceptibility because confounding by immunity and partner drug effects is minimised and clearance rates can be assessed accurately (>10 sequential parasite counts for individual patients in this analysis). The slope, and thus the derived parasite clearance half-life of the exponential decline (ignoring the variable lag phase) is judged the key pharmacodynamic determinant of the artemisinins.22 No evidence suggests saturation or density dependence in parasite clearance rates. Thus parasite clearances in patients with hyperparasitaemia can be compared with data from patients with lower parasitaemias, notably the artemisinin-resistant P. falciparum infections in western Cambodia.

This study cannot establish whether resistance alleles present in western Cambodia and northwestern Thailand have a common origin. Parasites with identical 93-locus genotypes were not recorded at the two sites, nor were slow clearing parasites from northwestern Thailand more closely related to Cambodian parasites than were fast clearing parasites from western Thailand. The evolution of artemisinin resistance will possibly be understood when the genetic determinants of parasite clearance rates have been identified.2,7

Parasite clearance rates in northwestern Thailand are continuously distributed. Modelling of the changes in parasite clearance over time did not support selection of a single artemisinin-resistant subpopulation with the same clearance phenotype as in Cambodia—ie, a one-step process. Together with the finding that clonally identical parasites with slow parasite clearance were reported in different patients in northwestern Thailand 8 years ago, this result argues against recent importation of Cambodian parasites as the explanation for the reduction in parasite clearance.

The artesunate–mefloquine combination has been the first-line treatment for falciparum malaria on the western border of Thailand since 1994. Artesunate was added to a failing mefloquine regimen, and the high effectiveness of the combination (>90% day 42 cure rates) has been sustained since then, although evidence suggests that effectiveness is decreasing.31 With a day 42 cure rate in 86 patients in 2010 of 88·9% (95% CI 77·8–94·7). Declining artemisinin effectiveness will have an adverse effect on the treatment of uncomplicated falciparum malaria by slowing therapeutic responses and increasing treatment failure rates, and will reduce the remarkable life-saving effectiveness of artesunate in the treatment of severe malaria and hyperparasitaemic patients. The degree to which current rates of resistance compromise these benefits has not been established, although the proportion of patients developing severe malaria did not increase concomitantly with the decline in artesunate effectiveness in this location. No patient died in this series, whereas in the past the mortality rate in patients given quinine for uncomplicated hyperparasitaemia was 3%.

A potential radical approach to containment of artemisinin resistance is to try to eliminate P. falciparum malaria from western Cambodia, but would this strategy be justified if resistance has already spread or emerged elsewhere? Genetically determined resistance to artemisinins is now prevalent on the Thailand–Myanmar border, contiguous with a malaria endemic area in which a large
burden of uncontrolled disease exists, so containment efforts will need to be expanded and surveillance and control strategies re-examined. Whether the fitness and transmissibility of the resistant parasites in the two locations are similar cannot be established yet, and thus elimination of falciparum malaria in western Cambodia might still be beneficial through prevention of the spread of higher degrees of resistance.

Identification of a molecular marker will be crucial to monitor the distribution and spread of resistance and to understand the evolution of this trait and the mechanism of action of artemisinin. The large numbers of patients infected with malaria, high heritability, and the broad range of parasite clearance half-lives make the Thailand–Myanmar border region ideal for powerful association studies. Clinical studies are needed urgently to map further spread and to establish the effect of different degrees of artemisinin resistance on treatment effectiveness and transmissibility.

Contributors
TJCA, SN, EAA, FN, and NJW contributed to study design. EAA, RM, AD, APP, CIM, KMI, and FN collected clinical data. SN, ShN, and SA-AS did molecular work. SN, KS, EAA, TJCA, and FN analysed the data. SN, TJCA, EAA, FN, PS, NPJD, and NJW wrote the manuscript. APP and SN contributed equally to this work. TJCA and FN also contributed equally.

Conflicts of interest
NJW is co-chairman of the WHO antimalarial treatment guidelines committee. All other authors declare that they have no conflicts of interest.

Acknowledgments
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References
Supplementary appendix

This appendix formed part of the original submission and has been peer reviewed. We post it as supplied by the authors.

APPENDIX MATERIAL

Emergence of artemisinin-resistant malaria on the western border of Thailand: a longitudinal study

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Appendix 1. Geographical variation in $t_{1/2}$ P. The section of the Thai-Myanmar border containing the four study clinics is shown, with bar charts showing changes in the proportions of slow clearing parasites ($t_{1/2}$ P $\geq$ 6.2 hrs) from 2001-10. Years in which no patients were treated are marked “nd”. There is a sharp rise in proportions of slow clearing parasites in clinics north of Maesot between 2008-10.
### Appendix 2: SNPs genotyping.

SNPs are named by chromosome and position in genome version 6.2. Alternate bases at the SNP are shown in square brackets, with Minor allele.”

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### Appendix 3. Age structure of patient population

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Appendix 4. Multiple regression analysis of t1/2P, stratified by location. Locations (from North to South) are Maela (MLA), Wang Pha (WPA), Mae Khon Ken (M KK) and Mawker Thai (MKT). The influence of patient age, prior malaria exposure, gender and sampling date on log t1/2P were examined using least squares regression in (A) all samples and (B) only patients with clearance data showing a good fit to a linear model ($r^2\geq0.8$). In both analyses sampling date strongly influences t1/2P in three of the four locations (black shading, white text). Significance of associations with patient age, gender and malaria exposure do not fall below p=0.025 and fail to reach Bonferroni corrected table-wide significance levels (0.05/16=0.003) in both analyses.

(A) t1/2P measures calculated from slopes with $r^2>0.8$ only

<table>
<thead>
<tr>
<th></th>
<th>MLA n=433</th>
<th>WPA n=1244</th>
<th>MKK n=141</th>
<th>MKT n=931</th>
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</thead>
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<tr>
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<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>F</td>
<td>p</td>
<td>F</td>
<td>p</td>
</tr>
<tr>
<td>82.834</td>
<td>8.9E-20</td>
<td>12.678</td>
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<tr>
<td>Age (yrs)</td>
<td>1.440</td>
<td>0.230</td>
<td>2.686</td>
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<tr>
<td>Malaria exposure$^1$</td>
<td>0.016</td>
<td>0.899</td>
<td>2.457</td>
<td>0.117</td>
</tr>
<tr>
<td>Gender</td>
<td>3.301</td>
<td>0.069</td>
<td>0.639</td>
<td>0.424</td>
</tr>
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(B) All data

<table>
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<th>MLA n=444</th>
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<th>MKK n=150</th>
<th>MKT n=1038</th>
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<td>F</td>
<td>p</td>
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<tr>
<td>79.154</td>
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<td>Age (yrs)</td>
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<tr>
<td>Malaria exposure$^1$</td>
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<td>0.956</td>
<td>4.086</td>
<td>0.043</td>
</tr>
<tr>
<td>Gender</td>
<td>5.006</td>
<td>0.025</td>
<td>0.661</td>
<td>0.416</td>
</tr>
</tbody>
</table>

$^1$Malaria exposure was defined by a documented malaria episode prior to the current episode.
Appendix 5. Genotyping data for one SNP. A typical SNP graph for the A/T SNP on chromosome 13, position 2158308, showing clustering of samples based on their genotypes. Genotype calls are made based on the normalized ratio of fluorescence intensity signals for the “A” and the “B” allele in the sample. The X-axis shows Norm Theta values, which correspond to the composition of the “B” allele in each sample. Multiple-clone infections are easily identified and were excluded from CR heritability analysis.
Appendix 6. Sampling breakdown for epidemiology and genetic analyses. Exclusion criteria are detailed in the text. The flow chart details how parasites with identical 93-locus genotypes infecting multiple patients were identified in the data set.
Appendix 7. Genetic relatedness among parasites. The tree shows the relatedness among 148 93-locus parasite genotypes infecting 441 patients. Blue genotype labels indicate infections collected between 2001 and 2004, black labels those collected from 2007-2010, while red labels show 30 infections from Cambodia. The numbers of patients infected with each genotype are shown in square brackets.
Appendix 8. Change in heritability of clearance rates on the Thai-Myanmar border. The proportion of variation in $t_{1/2}$ P that is due to parasite genetic factors (heritability) is shown for 2001-4 and 2007-10 time windows. Grey bars show heritability of $t_{1/2}$ P, while white bars show heritability analysis of residuals after removing effects of significant covariates. The error bars are 1 SD. The increase in heritability observed between 2001-4 and 2007-10 periods were tested by permutation (Appendix 9) and were significant for both $t_{1/2}$ P ($p<0.01$) and residuals ($p=0.04$).
Appendix 9. Permutation-based comparison of heritability in 2001-4 and 2007-10. In 2001-4, 29 groups of patients are infected with indistinguishable parasite genotypes, while in 2007-10, there were 119 groups. We randomly resampled 250 patients from 2007-10 generating samples containing on average 29 parasite genotypes infecting >1 patient. This resampling excersise provides a direct comparison with the 2001-4 period. Heritability estimates from 100 resampled 2007-10 datasets (range=0.48-0.85 exceeded that observed in 2001-4, demonstrating that the observed increase in $H^2$ is significant (p<0.01). For residuals (following removal of effects of covariates) all but 4 resampled datasets exceeded $H^2$ values for 2001-4 (p=0.04). These analyses are shown below. The frequency distribution shows $H^2$ values from 100 resampled 2007-10 datasets for A. t1/2P measures and B. residuals.