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## RESEARCH

# Risk factors for mortality from imported falciparum malaria in the United Kingdom over 20 years: an observational study

 OPEN ACCESS

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## Abstract

**Objectives** To determine which travellers with malaria are at greatest risk of dying, highlighting factors which can be used to target health messages to travellers.

**Design** Observational study based on 20 years of UK national data.

**Setting** National register of malaria cases.

**Participants** 25 054 patients notified with *Plasmodium falciparum* malaria, of whom 184 died, between 1987 and 2006.

**Main outcome measures** Comparison between those with falciparum malaria who died and non-fatal cases, including age, reason for travel, country of birth, time of year diagnosed, malaria prophylaxis used.

**Results** Mortality increased steadily with age, with a case fatality of 25/548 (4.6%) in people aged >65 years, adjusted odds ratio 10.68 (95% confidence interval 6.4 to 17.8),  $P<0.001$  compared with 18–35 year olds. There were no deaths in the  $\leq 5$  year age group. Case fatality was 3.0% (81/2740 cases) in tourists compared with 0.32% (26/8077) in travellers visiting friends and relatives (adjusted odds ratio 8.2 (5.1 to 13.3),  $P<0.001$ ). Those born in African countries with endemic malaria had a case fatality of 0.4% (36/8937) compared with 2.4% (142/5849) in others (adjusted odds ratio 4.6 (3.1 to 9.9),  $P<0.001$ ). Case fatality was particularly high from the Gambia. There was an inverse correlation in mortality between region of presentation and number of cases seen in the region ( $R^2=0.72$ ,  $P<0.001$ ). Most delay in fatal cases was in seeking care.

**Conclusions** Most travellers acquiring malaria are of African heritage visiting friends and relatives. In contrast the risks of dying from malaria once acquired are highest in the elderly, tourists, and those presenting in areas in which malaria is seldom seen. Doctors often do not think of these as high risk groups for malaria; for this reason they are important groups to target in pre-travel advice.

## Introduction

Malaria is a major global infection, causing around 250 million clinical cases a year.<sup>1</sup> It causes around 800 000 deaths a year, almost all in people living in poverty with limited access to healthcare, but it also causes avoidable deaths every year from imported malaria in non-endemic countries, mainly in people who are otherwise well. Travel to countries where malaria is endemic is increasing, and most general practitioners and acute care physicians are involved either in pre-travel advice or in diagnosing and treating cases on their return. The number of cases of imported falciparum malaria has increased over the past four years in the United Kingdom,<sup>2,3</sup> with 5774 reported cases in the period 2007–11. The UK, other European countries, and the United States also saw increases throughout the 1990s, which stabilised over the past decade.<sup>3–8</sup> Case fatality has remained stable at around 1%, both in the UK and in other high income countries.<sup>9–11</sup>

Migrants from malaria endemic countries and their descendants are at particular risk of acquiring malaria and make up over half of the UK's imported malaria cases.<sup>3</sup> Risk factors for acquiring malaria may, however, differ from risk factors for dying from malaria. Previously described risk factors associated with death from imported malaria include older age, European origin, travel to east Africa, failure to take chemoprophylaxis, and delays in seeking medical care and in the diagnosis and treatment of suspected malaria.<sup>9,10,12,13</sup> Preventing malaria associated deaths is the ultimate aim in malaria prevention strategies, as those diagnosed and treated promptly generally make a full recovery. We reviewed in detail all deaths reported to be caused by malaria infection in the UK over a 20 year period and compared them with all reported cases of malaria over the same period. The UK

has extensive links with malaria endemic countries and has one of the highest rates of imported malaria in the developed world.<sup>14</sup> We hypothesised that the groups most likely to acquire malaria may be different from the groups who die from malaria. We aimed, firstly, to identify factors associated with mortality to find high risk groups who could be targeted in prevention efforts by general practitioners and public health campaigns and, secondly, to learn clinical lessons relevant to the early identification of malaria in non-endemic countries.

## Methods

Malaria is a notifiable infectious disease in the UK, so clinicians have to report cases by law and all cases of malaria or suspected malaria are treated free by the National Health Service, including among non-residents, which reduces financial barriers to accessing treatment and to being recorded. The Malaria Reference Laboratory, the national reference laboratory and part of the UK Health Protection Agency, runs a passive case detection system; the data collection system has been described previously.<sup>3</sup> When a case is notified (via statutory notification through local authorities, clinicians sending standardised malaria reports to the Malaria Reference Laboratory, or laboratories sending blood films for diagnostic verification), the responsible clinician is contacted and asked to complete a data collection form, which covers demographic and clinical data. All cases included are parasitologically confirmed and speciated by microscopy or histology (and sometimes polymerase chain reaction) either at the referring hospital or at the Malaria Reference Laboratory. In most cases the diagnosis is confirmed by the Malaria Reference Laboratory. This system has been shown to identify 66% of cases of falciparum malaria in the UK.<sup>15</sup> Reporting methods did not change over the period of the study.

The notifying laboratory and clinician are requested to provide further information about patients: personal details (date of birth, sex, country of birth, country of usual residence), details of travel (date of arrival in UK, country or region visited, purpose of travel, duration of visit), prophylaxis taken, and details of illness (date of onset, date of treatment, and method of diagnosis). (The data collection sheet is available from the authors on request.) Where data are not clear they are followed up at the time by telephone with the referring clinician.

All malaria associated deaths in the UK are separately directly notified to the Malaria Reference Laboratory by the Office for National Statistics. Following this, death certificates and coroner's reports are obtained where available. In the case of a death in hospital, clinical and demographic details are requested from the responsible clinician as above. Cases are parasitologically confirmed and speciated by the above methods or by autopsy.

## Data analysis

Data on malaria cases over the 20 year study period were prospectively double-entered (by VS and MB) into the Malaria Reference Laboratory malaria database. The data were rechecked for cases (by AS) and deaths (AMC) to identify duplicate or uncertain data, and any variations manually checked. Data on cases were entered into dBase IV and subsequently cleaned and validated using Microsoft Access 10, and deaths were additionally entered in Microsoft Excel. Data were analysed using Stata 11 (Statacorp, College Station TX, USA).

Almost all deaths are from falciparum malaria, and our detailed analysis was therefore restricted to falciparum cases. Bivariate methods and multivariable logistic regression were conducted

to examine potential risk factors, comparing deaths from falciparum malaria with the denominator of all imported falciparum malaria cases to identify risk factors for case fatality. Unadjusted odds ratios and odds ratios adjusted for these potentially confounding factors on univariate analysis are quoted. There were no time series data.

Potential risk factors for death from imported falciparum malaria were studied by univariate analysis. These were age divided into groups ( $\leq 5$ ,  $>5-18$ ,  $>18-35$ ,  $>35-50$ ,  $>50-65$ , and  $>65$  years), sex, tourist versus traveller visiting friends and relatives, presentation by calendar month, reported use of effective malaria chemoprophylaxis versus reported use of no prophylaxis, birth in an African country with endemic malaria versus birth elsewhere, and country and region visited. Countries were classified as endemic or not endemic for malaria transmission according to the World Health Organization classification.<sup>1</sup> The biologically likely potential confounding factors age, born in a malaria endemic country, reason for travel, and reported use of prophylaxis were predetermined to be included as potential confounding factors in all logistic models. Age results are presented in clinically relevant groups (such as young adults  $>18-35$  years old, young children  $0-5$  years), but adjustment for confounding in all other multivariable models was performed using age as a continuous variable. Additionally factors found to be associated with mortality with a probability value  $P < 0.05$  in univariate analysis were introduced into the logistic regression model to adjust further for confounding. Where data were missing we used multiple imputation (STATA mi command) with five imputations. An additional analysis restricted to those cases with complete data was undertaken to confirm findings; this restricted analysis led to no change in interpreting these results (data not shown but available on request).

Cases were analysed by the UK government office region from where they were reported (North East, North West, Yorkshire and The Humber, East Midlands, West Midlands, East of England, London, South East, and South West), Northern Ireland, Scotland, Wales, and non-UK regions<sup>16</sup> to determine whether mortality was associated with number of cases of malaria seen locally, and a regression line fitted for log transformed data comparing mortality rate against number of cases seen.

This study used routine data collected within the Health Protection Agency official remit and did not require separate ethical board approval. Data were made anonymous for analysis.

## Results

Between 1987 and 2006, 191 malaria associated deaths were reported to the Malaria Reference Laboratory, from a total of 39 302 cases of confirmed malaria in which the outcome was known. Of the deaths, 184 were associated with *P falciparum* infection. There were 25 054 cases of *P falciparum* malaria, giving a case fatality of 0.73%. This rate did not change over the study period,<sup>3</sup> and adding year to models made no appreciable difference so it was excluded. Seven deaths were associated with non-falciparum malaria, (four from *P vivax*, and one each from *P ovale* and *P malariae*) among 13 804 cases, giving a case fatality for non-falciparum malaria of 0.05%. Among the non-falciparum cases, two patients had severe pre-existing comorbidities, and one died of splenic rupture.

The remainder of our analysis was performed on the 25 054 cases of falciparum malaria where the outcome was known, which included 483 cases of mixed infection. Completeness of data are summarised in table 1; 15 717 had full data, with use of prophylaxis the most likely to be missing.

Mortality from *P falciparum* malaria increased steadily with increasing age (fig 1 and table 2), with a case fatality of 4.6% (25/548) in those aged >65 years. The adjusted odds ratio of dying of malaria was 10.68 (95% confidence intervals 6.4 to 17.8,  $P<0.001$ ) in those >65 years old compared with the >18–35 year old group. Mortality among infants and children was low, with no deaths in the  $\leq 5$  age group, and a case fatality of 0.33% (11/3347) in those aged >5–18 years.

Table 3 summarises the risk of death from *P falciparum* malaria associated with different factors. Case fatality varied depending on purpose of visit to a country with endemic malaria, from 2.96% (81/2740) in tourists to 0.32% (26/8077) in travellers visiting friends and relatives (table 4). The adjusted odds ratio of death associated with tourism compared with visiting friends and relatives was 8.2 (95% confidence interval 5.1 to 13.3,  $P<0.001$ ). This remained true even when restricting the analysis to individuals born in African countries with endemic malaria; among these individuals, case fatality was 1.6% (3/189) in those visiting for tourism compared with 0.4% (19/5308) in those visiting friends and relatives ( $P=0.009$ ). Those born in an African country with endemic malaria had a case fatality of 0.4% (36/8937) compared with 2.4% (142/5849) in those born outside such countries ( $\chi^2$  test,  $P<0.001$ ; adjusted odds ratio of death 4.6 (3.1 to 9.9),  $P<0.001$ ). There were marked regional and national differences, with case fatality of 0.6% (86/14 174) after visiting west Africa, 1.5% (60/4064) for east Africa, and 3.3% (12/363) for southern Africa, although these reduced after adjusting for potential confounding factors such as reason for travel. Case fatality was particularly high in individuals visiting the Gambia (3.9%, (28/726)) compared with any other west African country (0.4% (58/13,448),  $\chi^2$  test,  $P<0.001$ ; adjusted odds ratio of death 4.7 (2.7 to 8.1),  $P<0.001$ ). Restricting the analysis to tourists increased this difference, with a case fatality of 6.0% (20/333) for cases from the Gambia compared with 1.4% (8/565) for tourists visiting the rest of west Africa ( $\chi^2$ ,  $P<0.001$ ).

There was a striking seasonal peak in deaths from imported *P falciparum* malaria, with a case fatality of 2.6% (49/1922) in those notified in December compared with 0.6% across all other months (135/23 132) ( $\chi^2$  test,  $P<0.001$ ) (fig 2), meaning that over a quarter of deaths occurred in cases notified in December. The peak for number of imported cases is September.<sup>3</sup> In the December cases 25.0% (11/44) of individuals died at home, compared with 17.4% (21/121) in other months (difference not significant). More tourists (9.3%) than those visiting friends and relatives (5.4%) were notified in December, although the greater mortality in December remained after correcting for purpose of visit as well as other factors (table 3).

There was a clear trend in mortality based on the UK government office region where the patient presented, which was inversely correlated with number of cases seen (fig 3) (linear regression  $R^2=0.72$ ,  $P<0.001$ ). The highest case fatality of 8.4% was in a region where only 119 patients were seen, with 10 deaths; the lowest case fatality (0.3%) was in the region where most cases were seen, with 15 993 patients seen and 50 deaths (adjusted odds ratio 18.2 (8.6 to 38.3),  $P<0.001$ ). Restricting the analysis only to tourists, case fatality was 6/20 (30%) among cases presenting in the region where fewest cases were seen, compared with 7/979 (0.7%) in the region where most cases were seen ( $\chi^2$  test,  $P<0.001$ ).

Data on reported chemoprophylaxis was inevitably limited and has to be treated with caution in fatal cases for reasons of reporting bias. Where the data were available, 35.8% (571/15 933) individuals reported taking prophylaxis against malaria infection, and, among these, 5.7% (903/15 933) reported taking

one of the three regimens effective in Africa (mefloquine, atovaquone-proguanil, or doxycycline). Of those taking either no prophylaxis or an ineffective regimen, 64.2% took no prophylaxis, 6.7% took chloroquine, 15.4% took chloroquine and proguanil, 4.1% proguanil, 0.7% pyrimethamine, and 3.5% another regimen. Among individuals not born in an African country with endemic malaria, case fatality was 0.6% (3/509) among those who reported taking effective prophylaxis, 1.4% (42/3046) among those who took any prophylaxis, and 3.5% (80/2304) among those who took no prophylaxis.

Median time intervals in fatal cases are summarised in fig 4. Among the fatal cases, 75.1% of deaths were in hospital, 18.1% at home, and 3.4% in an ambulance, and the median percentage of red blood cells parasitised was 8.5% (interquartile range 1–25%). These data were available only in the fatal cases.

## Discussion

This study shows that risk factors for dying from imported falciparum malaria in the UK are different from those for first acquiring the infection, and highlights groups of travellers who should be targeted for pre-travel health advice when they seek medical care. Those visiting friends and relatives in countries with endemic malaria make up the majority of cases of falciparum malaria in the UK, but the risks of this group dying from malaria are much smaller than for other travellers,<sup>3</sup> with most deaths occurring in tourists. Older age is strongly associated with mortality, and, in contrast to cases in Africa, mortality in children is minimal. There were both seasonal and geographical risks in the UK for dying from malaria, with a clear peak in December, and consistently higher risks of dying for patients presenting in UK regions where malaria is less regularly seen and treated. Travellers to the Gambia, a popular destination for winter sun, were at particular risk. Most delays occurred before presentation to medical services. These findings are biologically plausible (see discussion below), compatible with other data, and have implications for targeting pre-travel health messages and treatment when travellers return.

### The effect of previous exposure on mortality

Most cases of malaria in the UK are in individuals born in African countries with endemic malaria who are returning from visiting friends and relatives. Public health efforts have therefore concentrated on encouraging such people to take up chemoprophylaxis.<sup>3</sup> This remains an important message and is backed up by European data showing that rates of chemoprophylaxis use are low and of malaria are high in this group.<sup>17 18</sup> However, our study shows that, both in absolute numbers and in relative risk, tourists and those born outside African countries with endemic malaria are more likely to die once malaria is acquired. These data complement other studies from Europe and case series in the UK.<sup>4 12 19 20</sup>

There are several possible reasons for these findings, some biological and some behavioural. Those exposed to malaria in childhood develop immunity to severe malaria, with more protection the higher the degree of exposure; although they may still develop symptomatic infection, the risk of complications and death is greatly reduced.<sup>21</sup> This protection is incomplete, and wanes with time, but studies on imported malaria and in areas where malaria occurs in occasional epidemics suggest that some degree of protection continues for a prolonged period.<sup>4 12 19 22</sup> Other factors may also play a role; some genetic polymorphisms more prevalent in regions of malaria transmission are associated with protection from severe malaria.<sup>23 24</sup>



Immunity to malaria is however unlikely to be the full reason, as analyses restricted to those born in the UK (second or third generation migrants) or to tourists compared with those visiting friends and relatives when born in Africa still show significant differences. It is likely, although not possible to test with this study design, that differences in health seeking make up at least some of the difference. Those with an African heritage and staying with African families may be exposed to an understanding that malaria is common and dangerous, and may know the symptoms of early disease; malaria is, if anything, overdiagnosed as a cause of fever in medical practice in west and east Africa.<sup>25-27</sup> Conversely most tourists to Africa (and even their healthcare providers) may be unaware that several attacks of falciparum malaria a year are typical in residents living in highly endemic areas (considerably more common than influenza in the United Kingdom), and that around 20% of fevers in those returning from Africa to the UK are due to malaria.<sup>27</sup>

### The effect of age on mortality

Increasing age was strongly and independently associated with an increasing risk of death from malaria, and there was no evidence of the high case fatality seen in children in malaria endemic countries. These data are supported by previous reports of increased case fatality<sup>11 28 29</sup> and higher levels of parasitaemia in elderly people.<sup>30</sup> The mechanism is unclear, but it is unlikely that it is explained solely by increasing comorbidities in older age groups, as our study and others<sup>9 11</sup> show a continuous increase in case fatality from childhood upwards.

Our findings in children are also in keeping with a smaller prospective study of children in the UK, which found a substantial number of severe cases but no deaths.<sup>31</sup> A low case fatality in children has been documented in settings with low levels of endemic malaria, such as in the highlands of Kenya,<sup>32</sup> KwaZulu-Natal,<sup>33</sup> and Ceylon (Sri Lanka),<sup>34</sup> and also in non-immune individuals migrating to an Indonesian island hyper-endemic for malaria transmission.<sup>35</sup> These data are compatible with an explanation that the reason for high mortality from childhood malaria in Africa is limited access to health services.

### The effect of region within Africa, season and number of cases seen in regions on mortality

The higher case fatality in visitors to east Africa compared with west Africa was accounted for by a greater proportion of tourists visiting east Africa. The exception in west Africa is the Gambia, which had a strikingly high case fatality. Visitors to this country typically have other risk factors, including being tourists, older, and born in the UK, but, even after adjustment for these factors, the odds ratio remained high. This previously reported high case fatality<sup>36 37</sup> may reflect the predominance of Europeans travelling on package holidays, with low levels of use of chemoprophylaxis and little awareness of the risk of malaria on return.

Case fatality had a marked seasonal variation independent of peaks in the numbers of cases, which occurred in September and January (fig 2).<sup>3</sup> Higher case fatality occurs over the winter in several conditions; peaks in all cause and cardiac mortality occurred on Christmas Day and New Year's Day in a study of all deaths recorded in the United States between 1973 and 2001,<sup>38</sup> and all cause mortality peaked on New Year's Day in north east England.<sup>39</sup> Delayed presentation to hospital, lower levels of hospital staffing over the holiday period, and delays in communication of the malaria diagnosis from laboratory to physician may be contributory factors. Other possibilities include initial misdiagnosis of a febrile illness as influenza-like illness,

and the different demographics of travellers who travel in search of "winter sun." The non-significant increase in the proportion of deaths occurring at home during this month may reflect delayed presentation to medical services during this time.

There was an inverse correlation between the number of cases seen per UK region and case fatality. This is partly explained by the younger age and higher proportion of African nationals presenting in London. However, after adjusting for confounding factors, we found the odds of dying from malaria remained much higher in those presenting in regions where fewer cases of malaria were seen, and the increase in case fatality with decreasing number of cases in fig 3 is striking. Less awareness of malaria among friends and neighbours may lead to travellers being less likely to seek advice from doctors before travel, or to recognise that malaria is a possibility if they experience a fever on return. Equally, in regions where travel to malarious areas is less common, general practices advising on prophylaxis and doctors seeing febrile returned travellers may be less aware of the risk of malaria, and this is an area where public health advice and medical training should be targeted.

### Prophylaxis and delay to treatment

The use of effective prophylaxis was associated with lower risk of death from malaria in those not born in an African country with endemic malaria. There was no association between prophylaxis and case fatality in those born in African countries with endemic malaria. These data should, however, not be over-interpreted as a combination of reporting bias and incomplete data make it difficult to reach firm conclusions.

Delays in presentation, diagnosis, and administration of effective treatment are critical in contributing to deaths from malaria.<sup>7 10 40</sup> The median delay from the development of symptoms to the diagnosis of malaria in our study is similar to the reported time intervals of 4–8.5 days in other studies.<sup>12 19</sup> Reassuringly, our results suggest that treatment was usually initiated on the day of diagnosis, although we were unable to comment on individual regimens. These data also suggest that most patients who died presented late in the course of their illness, since the time interval between diagnosis and death was short. This suggestion is supported by the observation that 26% of individuals died either at home or in an ambulance. Reducing delays to presentation may be the single most important factor that could reduce mortality, although we are unable to test this from these data. There was no significant difference between time from onset of symptoms to diagnosis of malaria, from diagnosis to initiation of treatment, and from diagnosis to death, but numbers were small (data not shown). Data on delays were only available in fatal cases, so associations with mortality cannot be directly tested.

### Potential limitations of the study

There are limitations associated with the use of national data. Under-reporting is inevitable in a passive case detection system, although capture-recapture data suggest that levels of reporting are high.<sup>15</sup> The reporting of deaths known to be due to malaria is likely to be more complete than that of surviving cases, as deaths may also be reported via coroners as well as by clinical staff, although malaria may not have been suspected in all people who died at home without autopsy. Data on certain factors, such as those relating to delay in presentation, were available only for fatal cases, so their contribution to mortality cannot be directly assessed. Certain data, in particular the use of chemoprophylaxis, are likely to be associated with considerable recall bias in fatal cases. Lastly, although we adjusted for

important potential confounding factors we identified, we cannot exclude the presence of residual confounding. It seems unlikely, however, that the clear associations of case fatality with age, reason for travel, being born in a non-endemic country, the calendar month, and the number of cases seen per region could be largely explained by residual confounding.

## Implications for practice

These data describe two different groups of people—ethnic Africans (mainly of west African heritage), who travel mainly to west Africa to visit friends and relatives and bear the main burden of morbidity from falciparum malaria, and Europeans, who travel as tourists to east and southern Africa and to the Gambia in west Africa and have a higher case fatality.

Additionally, case fatality was higher in those presenting in December, and in those presenting in UK regions where malaria is less often seen.

Our results suggest that certain groups may particularly benefit from interventions to reduce the mortality from imported malaria, in particular pre-travel advice to encourage prompt presentation to healthcare facilities on the development of a febrile illness as well as stressing the need for antimalaria measures and chemoprophylaxis. These groups include the older traveller, travellers born in countries not endemic for malaria, and those travelling on “winter sun” holidays, in particular to the Gambia. It should not be forgotten, however, that individuals born in African countries with endemic malaria still bear the brunt of morbidity from imported malaria cases, so the importance of pre-travel advice and education in this group remains. Doctors in the UK advising patients who seldom travel but take a holiday to Africa must make them aware that malaria is very common, potentially fatal, may start initially with mild symptoms and needs early diagnosis.<sup>41</sup> Older visitors to malaria endemic countries, especially older tourists, are in particular need of targeting for pre-travel advice, both to take prophylaxis and to seek care early if unwell on return.

**Contributors:** The study was devised by AC, PLC, DB, CW. Data collection was by MB and VS. Data validation on deaths was by AC, and on cases by AS. Data analysis was by AC, AS and CW. All authors contributed to writing the paper. AC is the guarantor.

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**Competing interests:** All authors have completed the Unified Competing Interest form at [www.icmje.org/coi\\_disclosure.pdf](http://www.icmje.org/coi_disclosure.pdf) (available on request from the corresponding author) and declare: PLC had support from the Malaria Reference Laboratory for the submitted work; PLC has received an honorarium from GlaxoSmithKline for a lecture on malaria in the previous 3 years. No other financial relationships with any organisations that might have an interest in the submitted work in the previous 3 years; no other relationships or activities that could appear to have influenced the submitted work.

**Data sharing:** Data on cases imported by year available from the corresponding author.

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**What is already known on this topic**

Falciparum malaria is a common cause of fever in travellers returning to the UK from Africa

Individuals whose families originate from malaria endemic countries visiting friends and relatives are at greatest risk of importing malaria infection to the UK

Most deaths from malaria in endemic countries are in children, but deaths from malaria in children are rare in the UK

**What this study adds**

Holiday makers and tourists are at greatest risk of dying from malaria in the UK

Mortality from malaria increases steadily with age in the UK

Presentation in the month of December and to UK regions where malaria is rarely seen are associated with marked increase in mortality

Elderly travellers in particular need to be targeted for pre-travel advice

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## Tables

Table 1 | Completeness of data for notified cases of *Plasmodium falciparum* malaria in UK between 1987 and 2006

Factor studied	No (%) of cases with complete data
<b>All cases of <i>P falciparum</i> malaria (n=25 054)</b>	
Age	24 244 (97)
Sex	23 653 (94)
Country of birth	14 787 (59)
Country visited	21 856 (87)
Purpose of travel	17 252 (69)
Month reported	25 054 (100)
UK county where case reported	24 751 (99)
Use of malaria prophylaxis	15 933 (64)
<b>All deaths from <i>P falciparum</i> malaria (n=184)</b>	
Time from arrival in UK to onset of symptoms	135 (73)
Time from arrival in UK to death	142 (77)
Time from onset of symptoms to diagnosis	146 (79)
Time from diagnosis to starting specific treatment	92 (50)
Time from diagnosis to death	161 (88)
Location of death	177 (96)
Receipt of specific treatment before death	150 (82)
Percentage of red blood cells parasitised	61 (33)



**Table 2** Risk of death from imported *P falciparum* malaria in UK between 1987 and 2006 by age

Age group (years)	Odds ratio (95% CI) of death			
	Crude	P value	Adjusted*	P value
0–5	0 deaths	—	—	—
>5–18	0.73 (0.4 to 1.4)	0.36	0.81 (0.4 to 1.6)	0.54
>18–35	1.0 (reference group)	—	1.0 (reference group)	—
>35–50	1.49 (0.97 to 2.3)	0.07	1.63 (1.1 to 2.5)	0.02
>50–65	4.6 (3.1 to 6.8)	<0.001	4.6 (3.1 to 6.9)	<0.001
>65	10.66 (6.5 to 17.6)	<0.001	10.68 (6.4 to 17.8)	<0.001

\*Adjusted for risk factors listed in table 3.

Table 3| Factors other than age associated with risk of death from imported *P falciparum* malaria in UK between 1987 and 2006

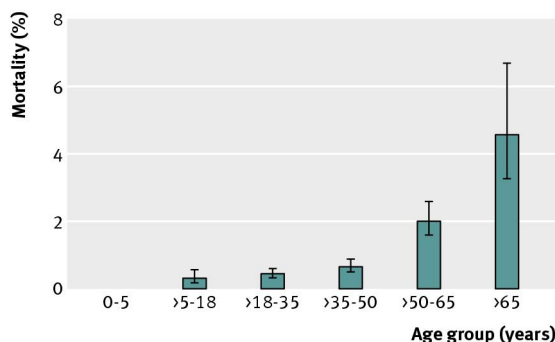
Risk factor	No (%) of fatal cases	Odds ratio (95% CI) of death			
		Crude	P value	Adjusted*	P value
Purpose of visit to country with endemic malaria:					
Tourism	81/2740 (2.96)	9.4 (6.1 to 14.7)	<0.001	8.2 (5.1 to 13.3)	<0.001
Visiting friends and relatives	26/8077 (0.32)				
UK region where disease presented:					
Where least malaria is seen	10/119 (8.40)	29.3 (14.5 to 59.2)	<0.001	18.2 (8.6 to 38.3)	<0.001
Where most malaria is seen	50/15 993 (0.31)				
Calendar month of presentation:					
December	49/1922 (2.55)	4.5 (3.2 to 6.2)	<0.001	3.7 (2.6 to 5.2)	<0.001
All other months	135/23 132 (0.58)				
Reported use of effective prophylaxis:					
No	99/10 222 (0.97)	1.8 (0.7 to 4.3)	0.220	2.0 (0.81 to 4.9)	0.13
Yes	5/903 (0.55)				
Birth in African country with endemic malaria:					
No	142/5849 (2.43)	6.2 (4.3 to 8.9)	<0.001	4.6 (3.1 to 9.9)	<0.001
Yes	36/8937 (0.40)				

\*Adjusted for age (continuous) and for other listed risk factors.

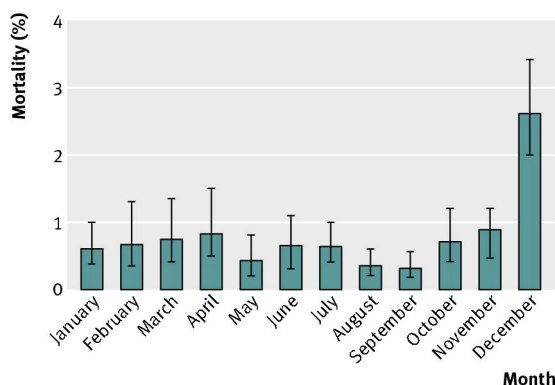
**Table 4| Purpose of visiting a country with endemic malaria among cases of imported *P falciparum* malaria in UK between 1987 and 2006 and associated case fatality**

<b>Purpose of travel</b>	<b>No (%) of fatal cases</b>
Tourist from UK visiting malaria endemic country	81/2740 (2.96)
Visiting UK from malaria endemic country	17/3841 (0.44)
Visiting friends and relatives in malaria endemic country	26/8077 (0.32)
Other or unstated	60/10 396 (0.58)
Total	184/25 054 (0.73)

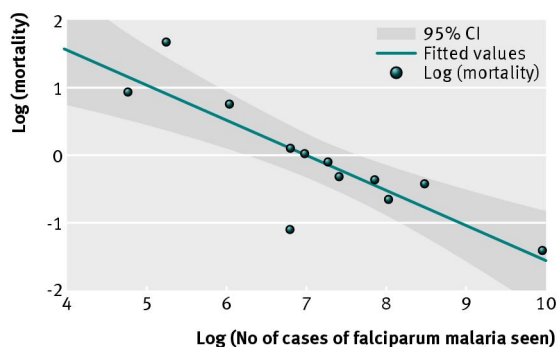
## Figures



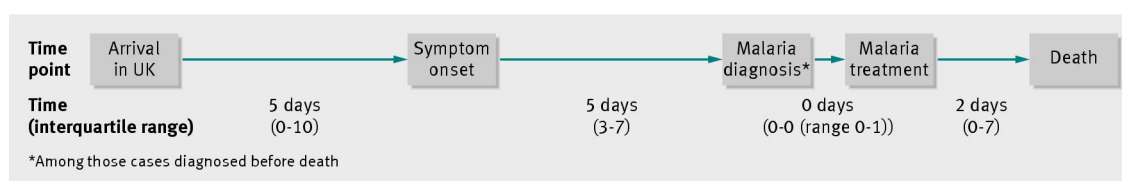
**Fig 1** Mortality from imported *P falciparum* malaria in UK between 1987 and 2006 by age



**Fig 2** Mortality from imported *P falciparum* malaria in UK between 1987 and 2006 by month of notification of illness



**Fig 3** Mortality from imported *P falciparum* malaria as a function of number of cases seen in a UK region between 1987 and 2006 ( $R^2=0.67$ ,  $P<0.001$ )



**Fig 4** Relevant time intervals in fatal cases of imported *P falciparum* malaria in the UK between 1987 and 2006