How Not to Miss a Case of Malaria in Emergency Department in Malaria Non-Endemic Areas? Practical Approach & Experiences in Hong Kong

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Abstract

Human malaria is a life-threatening mosquito-borne protozoan parasitic infection in human involving female anopheline mosquitoes as vector for transmission. It is caused by *Plasmodium* species, most commonly, *P. vivax*, *P. ovale*, *P. malariae* and *P. falciparum*, and rarely *P. knowlesi*. Malaria remains a significant global health issue and is a medical emergency. It is also an important cause of morbidity and mortality in endemic areas, particularly in at-risk groups. In Hong Kong, where malaria is non-endemic, more than 20 cases of malaria per year have been notified in recent years. We still have chances encountering patients with malaria presented to public or private emergency departments. High index of clinical suspicious is utmost important for not missing a case of malaria. A practical approach for prompt identification of patients with severe malaria is essential, followed by appropriate initiation of appropriate effective antimalarial treatment within 24 to 48 hours of symptoms onset after blood taken for thick and thin smears for diagnosis. Vigilance with increased awareness of not falling into common diagnostic traps has to be alerted. The risk of missing any case of malaria presenting to emergency department could be largely minimized.

Keywords

Malaria, *Plasmodium falciparum*, *Plasmodium* species, Uncomplicated Malaria, Severe Malaria
1. What Is Malaria?

Human malaria is a life-threatening mosquito-borne protozoan parasitic infection in human caused by *Plasmodium* species, most commonly, *P. vivax*, *P. ovale*, *P. malariae* and *P. falciparum* [1] [2] [3]. *P. knowlesi* which causes primate malaria has been found to cause human infections [2] [4].

The route of transmission is mainly through infected female Anopheline mosquitoes taking blood meal from and inoculating parasites to human and infecting their erythrocytes [1] [2] [5]; other routes of transmission in human include blood transfusion, organ transplantation, shared use of contaminated needles or syringes, and maternal-to-child transplacental or perinatal transmission [6] [7] [8].

After incubation period of 6 days to up to 1 year, depending on different species of *Plasmodium*, malaria manifest clinically in a spectrum; from non-specific uncomplicated symptoms of acute febrile illness, intermittent fever, chills, sweating and headache, with or without anaemia and jaundice, to severe complication with seizure, confusion, coma, renal failure, shock, acute respiratory distress syndrome and death [1] [2] [3] [5] [9].

In human malaria, the life cycle of *Plasmodium* species ([Figure 1](#)) involved hepatic and erythrocytic infection [1] [2] [5]. And due to the cyclical parasitaemia, patients often report experiencing symptoms every 48 hours (in *Plasmodium falciparum*, *Plasmodium vivax* and *Plasmodium ovale* infection) to 72 hours (in *Plasmodium malariae* infection) [1] [5].

Globally, despite the incidence and death rate of malaria have been decreasing in recent decades, malaria remains an important cause of morbidity and mortality in endemic areas [6]. It is estimated that about 3.2 billion people, that is almost half of the world’s population are at risk of malaria ([Figure 2](#)) [1] [6]. The most at risk groups are young children, pregnant women, non-immune travelers from malaria-free areas and patients co-infected with HIV [1] [5] [6].

*Plasmodium vivax* and *P. falciparum* malaria are the most important cause of malaria. Falciparum malaria is the most severe form of malaria leading to deaths [5] [10], vivax malaria is considered with increasingly importance as a cause of severe malaria and sometimes mortality [11]. The mechanism of severe malaria in *Plasmodium falciparum* infection is mainly due to cytoadherence and then sequestration of *P. falciparum*-infected erythrocytes leading to occlusion of blood flow resulting in decrease oxygen delivery to tissues and end organ damage ([Figure 3](#)) [5] [10] [12].

2. Why Do We Need to Be Aware of Malaria? Experiences in Hong Kong

Hong Kong is a malaria non-endemic area; while Sub-Saharan Africa carries a disproportionately high share of the global malaria burden, and *P. falciparum* is the most prevalent malaria parasite [5] [6]. We still have to be vigilant to maintain a high index of suspicion, conduct a thorough history including travel history, and interpret the constellations of symptoms and signs, and preliminary investigation findings accordingly.
Figure 1. Life cycle of malaria infection in human and proposed mechanism of fever induction. Adapted from [5] [12] [13]. In human malaria, the life cycle of malaria involved hepatic and erythrocytic infection. When an infected female anopheline mosquito bite and take a blood meal from human, the sporozoites released from the mosquito are carried to human liver cells where multiplication take place and then exit into the bloodstream as merozoites, which infect other erythrocytes in the blood stream to start the asexual blood-stage cycle. This cycle completes in 48 to 72 hours, depending on the infecting Plasmodium species. The process of Plasmodium species infecting erythrocytes, and its associated inflammatory response elicited, correlates with the development of clinical symptoms. These infected erythrocytes are then phagocytosed by the spleen, which intended to clear the infection, but through cytoadherence and sequestration that also lead to profound anaemia and folic acid deficiency.

Figure 2. Countries with ongoing transmission of malaria, 2000 and 2015 [6].
Figure 3. Pathophysiology of severe malaria, particularly in *Plasmodium falciparum* infection—sequestration of infected erythrocytes due to cytoadherence and rosetting. Adapted from [5] [10] [13]. In *Plasmodium falciparum* infection, the infected erythrocytes are capable to roll on and cytoadhere themselves via respective receptors to vascular endothelial wall. This leads to sequestration of infected cells in small blood vessels, and subsequent hemorrhage or infarct resulting in end organ damage. The spontaneous binding of uninfected erythrocytes to the infected erythrocytes also contributes to the vessel occlusions. The parasite antigens stimulate the release of interferon-γ and tumor necrosis factor-α in addition to immunoglobulin complexes production.

to initiate prompt and appropriate management [12]. It is because malaria is a life-threatening condition, which is preventable by appropriate prophylaxis and measures and treatable with prompt appropriate effective treatment within 24 to 48 hours of symptom onset [4] [5].

Case presentation—Have you encountered similar situations?

A 36-year-old lady presents to an emergency department of Hong Kong with on and off fever for 10 days with chills without other symptoms. She is a Nigerian lawyer living in Hong Kong for 3 years. She travels frequently between Hong Kong and Nigeria. She has just visited her family and returned from Nigeria, and her symptoms were actually developed during her stay in Nigeria. Her past medical history include malaria diagnosed few months prior symptom onset and has been treated with 3 days of medication; however she cannot recall any names of infecting parasite species and medica-
tions. She is currently pregnant at 26 weeks gestation. Her drug allergy history includes augmentin, gabapentin and chloroquine. What would you do and how would you manage this patient?…

3. Malaria in Hong Kong

Malaria is a notifiable infectious disease in Hong Kong and has been well controlled over decades that around 20 to 30 cases of malaria per year have been notified for the last few years (Figure 4) [13] [14] [15]. Therefore, we still have chances encountering patients with malaria presented to private or public emergency departments. Majority of these patients were Hong Kong residents followed by tourists/transients, migrants and immigrants; almost half of them had travelled to and considered as imported cases from India, Pakistan and Nepal [3]. Other imported cases by country were from Africa, Southeast Asia & Pacific Islands, Latin & South America, and the Mainland China [3]. Majority of these malaria-infected Hong Kong residents acquired the infection during travel and amongst more than 80% of them did not have pre-travel chemoprophylaxis taken [3].

In Hong Kong, the clinical presentation of patients with malaria presented to emergency department usually non-specific which overlaps with symptoms of other diseases associated with other mosquito-borne illnesses or other conditions that commonly found in returned travelers [3] [12].

Similar to the global situation, about 1/3 of malaria cases notified in Hong Kong were due to *P. vivax*, followed by *P. falciparum*, the rest were by *P. malariae* and *P. ovale*. Mixed infections of two or more parasites have also been observed [3].

Moreover, appropriate local malaria vectors are present in Hong Kong, the female

![Number of Malaria Cases Notification to CHP, Hong Kong, Year 1997 to as of October 2016](image)

**Figure 4.** Number of malaria cases notification to Centre for Health Protection (CHP), Department of Health, the Government of the Hong Kong Special Administrative Region (Hong Kong) [15].
anopheles mosquitoes, including *Anopheles jeyporiensis* which breed in paddy field, waterlogged field and stream, and *Anopheles minimus* which breed along slow running, unpolluted stream with diffused sunlight and marginal vegetation [3] [16]. The presence of these vectors theoretically allows malaria transmission in Hong Kong when they take blood meals from patients with malaria and parasitaemia.

4. What Can We Do in Emergency Departments for Diagnosis of Malaria?

Malaria is a medical emergency and making diagnosis of malaria in emergency department is challenging [12]. In general, with background clinical knowledge on malaria, a high index of clinical suspicion is the utmost importance in assessment of anyone with fever or history of fever, and has returned from or previously visited a malaria-endemic area. And do not assume malaria can be excluded from diagnosis just simply by history of anti-malaria chemoprophylaxis use or history of a negative test for malaria.

Be aware of the non-specific presenting symptoms of malaria, most commonly fever, headache and generalized malaise, followed by gastrointestinal disturbances, jaundice or respiratory symptoms [3] [4] [12]. These are often misdiagnosed as non-specific viral infection, especially in children whose presentation is even particularly vague [4].

For those with uncomplicated malaria, the unremarkable findings in physical examination, and the temperature measured at single or a few time points only during the short period stay in emergency department, are not only unable to completely reveal any specific fever pattern, but also would have perpetuated the overlooking of malaria as diagnosis. Patients with severe malaria present late may present with confusion, seizures or jaundice, and in children, hepatomegaly or splenomegaly may be palpable [4] [12].

The basic history taking including travel history, physical examination and preliminary findings are our tools to minimize the chance of missing a case of malaria [9] [12]. At the same time, it is important to go through the respective lists of differential diagnosis of common causes of fever in returned travelers, and other vector-borne infection presented with fever and various clinical syndromes accordingly. These include typhoid, rickettsiosis, leptospirosis, hepatitis, influenza, HIV, dengue, agents of viral hemorrhagic fever, meningitis and encephalitis [12].

5. Assessment of Patients with Suspected Malaria—A Practical Approach

Upon approach to patients, general examination of the patient and assessment of vital signs including blood pressure, pulse, oxygen saturation, rapid measurement of glucose and haemoglobin levels by glucometer and haemocue respectively, are important to ascertain the haemodynamic stability of patient and to determine whether immediate resuscitation and management is the prime priority. Rapid assessment of airway, breathing and circulation are essential, in addition to mental state examination coupled with immediate establishment of intravenous access, monitoring and management of the
abnormal vital signs accordingly whenever clinical indicated.

6. History and Physical Examination

It is important to identify those at-risk of having malarial infection and secondary complications. These include patients at age of extremes, pregnancy, history of blood transfusion, organ transplantation, shared use of contaminated needles or syringes, HIV infection, and travel history [6] [7] [8]. It is particularly important to diagnose malaria in pregnancy because higher rates of poor maternal outcomes including profound maternal anaemia, cerebral malaria, severe malaria and maternal mortality, in addition to poor fetal or neonatal outcomes including abortion, stillbirths, preterm delivery and low birth weight, congenital malaria and neonatal mortality have been reported [5] [11]. Moreover, expertise of critical care medicine has to be urgently consulted for admission of these patients for subsequent monitoring and management in collaboration with obstetricians [5].

7. Why Do We Need to Take Travel History?

Travel history including country and area of travel, stopovers and date of return, is particularly informative especially when travel history to malaria endemic areas could be elicited [11]. Not only has it raised our clinical alertness in considering malaria as possible diagnosis, it gives a clue to the likelihood of infecting Plasmodium species. It is because P. falciparum is considered to be the most prevalent malaria parasite on the African continent, and is most likely to occur within 3 months or no later than 6 months of return from an endemic area, while P. vivax in many countries outside of Africa, P. vivax and P. ovale commonly present later than 6 months after return [1] [6].

Moreover, travel history also gives clues to the likelihood of the antimalarial resistance of the infecting Plasmodium species, the effectiveness of antimalarial chemoprophylaxis if any have been taken pre- and or during travel; both of which have clinical implications on subsequent selection of appropriate antimalarial treatment for prompt management [4] [11] [16]. Chloroquine-resistant Plasmodium falciparum malaria have been documented in Caribbean (Haiti and Dominican Republic) and Central America (Mexico, Belize, Guatemala, Honduras, El Salvador, Nicaragua, Costa Rica, northwest of Panama); while mefloquine-resistant Plasmodium falciparum malaria in South America (southwest Panama, Columbia, Venezuela, Ecuador, Peru, Brazil and Bolivia), and Southeast Asia (eastern Myanmar, Thailand, Cambodia and southern Vietnam) & Oceania [16]. For those who have travelled to border areas of Thailand, Cambodia, and Myanmar are particularly at the highest risk for multiple drug resistant infections [4] [17].

With further emergence of antimalarial drug resistance, extreme cautious is needed to those with suspected malaria and with travel history to the 5 countries of the Greater Mekong subregion: Cambodia, Lao People’s Democratic Republic, Myanmar, Thailand and Viet Nam, because artemisinin resistance has been found in these areas [4] [16]. And multi-drug resistant P. falciparum have been found in parts of Cambodia and
In addition, travel history gives clues for risk assessment regarding the effectiveness of preventive measures. Insecticide resistances have been reported in malaria-endemic areas of sub-Saharan Africa and India [4].

8. Investigation

Without any delay, whenever malaria is suspected, blood taking for 1 - 5 ml blood sample in EDTA tubes prior to initiation of anti-parasitic therapy, and urgently sending for thick and thin smear for microscopic examination by experienced microscopists is the most important investigation to be performed [4] [12]. The result is expected to be available within 2 hours upon laboratory receipt of specimens [4] [12]. Clinicians must not hesitate to contact and communicate with laboratory colleagues for urgent test and whenever results not readily available after a reasonable time frame [12]. Remember to note the result of percentage of red blood cells that are parasitized in falciparum malaria [4] [9].

In situations where and when there is lack of expertise in the laboratories, particularly out of hours, commercially available rapid diagnostic tests (RDT) play the role for prompt diagnosis (Figure 5). These are based on immunochromatography, which depends on different kits of RDTs, to detect *Plasmodium* species antigen, histidine-rich protein II (HRP-2) or enzymes such as *Plasmodium* lactate dehydrogenase (pLDH) and adolase, are commonly used in some laboratories in Hong Kong and other malaria non-endemic countries such as the United States and the United Kingdom [4] [18] [19] [20]. These tests are easy to perform requiring low skill levels with results readily available within 20 minutes [16]. It has been shown that RDT for malaria is superior to a single set of blood smears performed in laboratories, and its use has been advocated for rapid diagnosis or exclusion of *P. falciparum* malaria, especially for evaluation of patients with compatible clinical contexts in outpatient settings [18]. RDTs for detecting *Plasmodium falciparum* HRP-2 can also be useful for patients with negative blood smears resulted from recent exposure but inadequate dose of artemisinin derivatives [4]. However, mis-identification of *Plasmodium falciparum* malaria as non-falciparum malaria by RDT has been occurred in non-endemic area during evaluation of returned traveler from endemic area, and revealed by subsequent blood smears examination [18]. Moreover, false negative result may occur in patients with very high parasitaemia due to prozone effect [19]. Nevertheless, the use of RDT does not exclude the must tests of thick and thin smear as investigation for suspected malaria [12] [18] [20].

Moreover, molecular diagnosis with qualitative and/or quantitative analysis by species-specific polymerase chain reaction (PCR) is considered as adjunct in initial diagnosis of malaria when species determination cannot be made from blood smears [20]. Likewise, it cannot replace the must tests of thick and thin smear because the turnover time of PCR is currently still not within the clinically comparable and relevancy for acute diagnosis of malaria, despite of its high sensitivity at detection limit 5 parasites/µl and high specificity [20].
Figure 5. Different commercially available rapid diagnostic tests (RDTs) for diagnosis of malaria and the antigen tested. Adapted from [19] [20] [21]. BinaxNOW is the only brand that the U.S. Food and Drug Administration (FDA)-approved RDT for diagnosis of malaria in the United States. All results from RDTs must be confirmed by microscopic examination of thick and thin smears. The BinaxNOW RDTs (A, left) showed a faint line in the aldolase (T2) and control window, suggestive of infection due to *Plasmodium vivax*, *P. malariae* or *P. ovale*; and lines indicating HRP-2 (T1), aldolase (T2), and control window are present (A, right), suggestive of *P. falciparum* or mixed infection; Wright-Giemsa-stained thin smear showing classic thin, delicate ring trophozoite, supporting *P. falciparum* infection.

The microscopic examination of thick smear is highly sensitive with detection limit 50 parasites/µl which is equivalent to 0.001% of red blood cell infected, and a large volume of blood could be screened, as compared to the detection limit of $\geq 100$ parasites/µl in RDT kits [20]. Thin smear takes the advantages of preservation of morphology of parasites in erythrocytes that allow species identification under microscopic examination [20].

The initial negative blood smears does not exclude the diagnosis of malaria. Whenever clinical suspicious of malaria, repeat blood smears whenever the temperatures spike up, and for the second sets of thick and thin smears at 6 to 12 hours after the initial tests, the third sets at 12 to 24 hours after the second sets [4] [9]. Negative smears...
must be interpreted with cautions for pregnant patients because of placental sequestration of parasites, and in other high-risk groups with suspicious of malaria [9]. Negative or delay of parasitological diagnosis should not delay an immediate start of antimalarial treatment [4].

Other blood tests including i-stat point-of-care blood gas analyses, laboratory investigation of complete blood picture, clotting profiles, liver and renal function tests, random glucose, lactate and pH should also be taken and monitored [12]. Blood for glucose-6-phosphate dehydrogenase (G6PD) status is needed in selection of antimalarial treatment, because primaquine which used for elimination of hypnozoites from liver can cause haemolysis in G6PD deficient patients [9]. Baseline electrocardiography and chest X-ray are needed as well. Blood culture and other investigations should be undertaken accordingly for diagnosis or rule out other differential diagnosis high on the list [12].

9. Clinical Pathway for Management of Patients with Malaria in Emergency Department

After clinical assessment of symptoms and signs, with the presence of preliminary laboratory parameters, these allow identification and differentiation of patient with severe from uncomplicated malaria; these in turns guide the initial antimalarial treatment [12]. Severe malaria is characterized by organ damage or hematological abnormalities, and those with poor prognosis and require admission to intensive care unit can also be identified and stratified [12].

Major features of severe falciparum malaria in adults include shock with blood pressure < 90/60 mmHg, hypoglycaemia with blood glucose < 2.2 mmol/L, haemoglobinuria, haemoglobin ≤ 8 g/dL, impaired consciousness or seizures, pulmonary edema or acute respiratory distress syndrome, spontaneous bleeding or disseminated intravascular coagulation, renal impairment with oliguria < 0.4ml/kg body weight or creatinine > 265 µmol/L, acidosis with pH < 7.3 [4] [9] [12].

Severe malaria in children are characterized by hypoglycaemia with blood glucose < 2.2 mmol/L, severe anaemia with haemoglobin < 8 g/dL, impaired consciousness or seizures, respiratory distress or acidosis with pH < 7.3, prostration, presence of parasitaemia > 2% red blood cells parasitized [4] [9].

The distinction between falciparum and non-falciparum malaria is also important, as life-threatening infection is commonly seen in the former, and the urgent initial antimalarial treatment regimen differs (Table 1) [4] [9] [17].

10. Supportive Management

Supportive management in emergency department settings is crucial. Antipyretics should be considered if the core temperature >38.5°C [4] [12]. Paracetamol at a dose of 15 mg/kg body weight every 4 hourly given orally or as rectal suppository for young children is safe and well tolerated, and a more rapid fever clearance time had been observed when used in combination with quinine as treatment for uncomplicated
Table 1. Antimalarial regimens for severe and uncomplicated malaria [4] [17] [23].

<table>
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<tr>
<th>Severe malaria</th>
<th>Uncomplicated/ P. falciparum (or species not identified)</th>
<th>Chloroquine-sensitive regions:</th>
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<tbody>
<tr>
<td>For adults—</td>
<td>For Adults—</td>
<td>For Paediatrics—</td>
</tr>
<tr>
<td>Doxycycline 100 mg intravenous (IV)/oral (po) every 12 hourly for 7 days, AND</td>
<td>Chloroquine phosphate 1 gm salt (600 mg base) po, then 0.5 gm in 6 hours, then 0.5 gm daily for 2 days. Total 2500 mg salt;</td>
<td>Chloroquine phosphate 10 mg/kg of base po; then 5 mg/kg of base at 6, 24 &amp; 48 hours. Total 25 mg/kg base (never exceed adult dose).</td>
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<td>OR</td>
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<td>Quinidine gluconate in normal saline [10 mg/kg (salt) IV over 1 - 2 hours then 0.02 mg/kg/min by constant infusion] or (24 mg/kg IV over 4 hours, then 12 mg/kg over 4 hours every 8 hourly).</td>
<td>Hydroxychloroquine 800 mg salt (620 mg base) po, followed by 400 mg salt (310 mg base) po at 6, 24, and 48 hours.</td>
<td>Hydroxychloroquine 10 mg base/kg po immediately, followed by 5 mg base/kg at 6, 24, and 48 hours. Total 25 mg base/kg.</td>
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<td>OR</td>
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<td>then quinine sulfate 650 mg po three a day for completion of 3 days of quinidine/quinine (7 days if Southeast Asia)</td>
<td>Artemether-lumefantrine 4 tablets (80 mg/480 mg) as a single dose, then 4 tablets again after 8 hours, then 4 tablets every 12 hours for 2 days</td>
<td>Artemether-lumefantrine (20/120 mg tablets) • &lt;5 kg: not recommended; • 5 kg to &lt;15 kg: 1 tablet (20 mg/120 mg) as a single dose, then 1 tablet again</td>
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<th>Alternatives</th>
<th>Alternatives</th>
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<tr>
<td>Doxycycline 100 mg IV loading dose, then 5 mg/kg IV or po every 8 hourly for 7 days instead of doxycycline in pregnant women</td>
<td>Substitute clindamycin 10 mg/kg IV loading dose, then 5 mg/kg IV or po every 8 hourly for 7 days instead of doxycycline in pregnant women</td>
<td>IV or intramuscular artesunate for at least 24 hours and until they can tolerate oral medication, then complete artemisinin-based combination therapies (ACT, see below) for 3 days (add single dose primaquine in areas of low transmission).</td>
</tr>
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</table>

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**Parenteral artesunate:**

Children with BW <20 kg: 3 mg/kg BW per dose;

Children with >20 kg BW and adults: 2.4 mg/kg BW per dose.

If parenteral artesunate is not available, use artemether in preference to quinine for treating children and adults with severe malaria.

**For adults and pediatrics except pregnant women in 1st trimester:**

Any of the (ACT) for 3 days:

- Artemether + lumefantrine;
- Artesunate + amodiaquine;
- Artesunate + mefloquine;
- Dihydroartemisinin + sulfadoxine-pyrimethamine (SP)

**For 1st trimester pregnancy:** (quinine sulfate + clindamycin) for 7 days;
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Pyronaridine/artesunate 4 tablets (180 mg/60 mg) as a single dose each day for 3 days for >65 kg or 3 tablets if 45 kg to <65 kg after 8 hours, then 1 tablet every 12 hours for 2 days;
- 15 kg to <25 kg: 2 tablets (40 mg/240 mg) as a single dose, then 2 tablets again after 8 hours, then 2 tablets every 12 hours for 2 days;
- 25 kg to <35 kg: 3 tablets (60 mg/360 mg) as a single dose, then 3 tablets after 8 hours, then 3 tablets every 12 hours for 2 days;
- ≥35 kg: as per adult dose;

Pyronaridine/artesunate 2 tablets (180 mg/60 mg) as a single dose each day for 3 days for 24 kg to >45 kg or 1 tablet if 20 kg to <24 kg.

Infant <5 kg BW: ACT at same mg/kg BW target dose as for children with 5 kg BW;

For patients co-infected with HIV: avoid artesunate + SP if treated with co-trimoxazole, and avoid (artesunate + amodiaquine) if treated with efavirenz or zidovudine.

Non-immune travelers: ACT;

Uncomplicated hyperparasitaemia: Chloroquine-resistant or unknown resistance regions:

For Adults—
Atovaquone-proguanil (1 gm/400 mg) (4 adults tabs) po in single dose daily for 3 days with food;

OR
Atovaquone-proguanil (pediatric tablet 62.5/25 mg)
- <5 kg: not recommended;
- 5 kg - 8 kg: 2 pediatric tablets in a single dose daily for 3 days;
- 9 kg - 10 kg: 3 pediatric tablets in a single dose daily for 3 days;
- 11 kg - 20 kg: 1 adult tablet in a single dose daily for 3 days;
- 21 kg - 30 kg: 2 adult tablets in a single dose daily for 3 days;
- >40 kg: as per adult dose

Pyronaridine/artesunate 4 tablets (180 mg/60 mg) as a single dose each day for 3 days for >65 kg or 3 tablets if 45 kg to <65 kg.

For Paediatrics—
Artemether-lumefantrine (20/120 mg tablets): weight-based dosing as above;

Artemether-lumefantrine 4 tablets (80 mg/480 mg) as a single dose, then 4 tablets again after 8 hours, then 4 tablets every 12 hours for 2 days (take with food);

OR
[(quinine sulfate 650 mg po three times a day for 3 days (7 days if Southeast Asia)] + [(doxycycline 100 mg po two times a day) or (tetracycline 250 mg po four times a day)]

Pyronaridine/artesunate 4 tablets (180 mg/60 mg) as a single dose each day for 3 days for >65 kg or 3 tablets if 45 kg to <65 kg.

[(quinine sulfate 10 mg/kg po three times a day for 3 days); (7 days if Southeast Asia)] + (clindamycin 20 mg/kg per day divided in three times a day)] both for 7 days under age of 8;

[(quinine sulfate 10 mg/kg po three times a day for 3 days); (7 days if Southeast Asia)] + (doxycycline 2.2 mg/kg/two times a day) or (tetracycline 25 mg/kg/day divided four times a day) for 7 days.

Pyronaridine/artesunate 2 tablets (180 mg/60 mg) as a single dose each day for 3 days for 24 kg to >45 kg or 1 tablet if 20 kg to <24 kg.
| Uncomplicated/| Chloroquine-sensitive regions: | For blood stage infection: (also include P. vivax & P. ovale) |
| P. ovale, P. malariae or P. knowlesi | | For unknown species, treat as for uncomplicated P. falciparum malaria. |

For adults—
Chloroquine phosphate 1 gm salt (600 mg base) po, then 0.5 gm in 6 hours, then 0.5 gm daily for 2 days. Total 2500 mg salt;
OR
Hydroxychloroquine 800 mg salt (620 mg base) po, followed by 400 mg salt (310 mg base) po at 6, 24, and 48 hours.
OR
Artemether-lumefantrine 4 tablets (80 mg/480 mg) as a single dose, then 4 tablets again after 8 hours, then 4 tablets every 12 hours for 2 days (take with food).
PLUS (P. ovale only)
+ primaquine phosphate 52.6 mg (30 mg base = 2 tablets) po once daily for 14 days.

For paediatrics—
Chloroquine phosphate 10 mg/kg of base po; then 5 mg/kg of base at 6, 24 & 48 hours. Total 25 mg/kg base (never exceed adult dose).
OR
Hydroxychloroquine 10 mg base/kg po immediately, followed by 5 mg base/kg at 6, 24, and 48 hours. Total 25 mg base/kg.
OR
Artemether-lumefantrine (20/120 mg tablets)
- 5 kg to <15 kg: 1 tablet (20 mg/120 mg) as a single dose, then 1 tablet again after 8 hours, then 1 tablet every 12 hours for 2 days;
- 15 kg to <25 kg: 2 tablets as a single dose, then 2 tablets again after 8 hours, then 2 tablets every 12 hours for 2 days;
- 25 kg to <35 kg: 3 tablets as a single dose, then 3 tablets again after 8 hours, then 3 tablets every 12 hourly for 2 days;
- ≥35 kg: as per adult dose;
PLUS (P. ovale only)
+ Primaquine phosphate 0.5 mg/kg po base once daily for 14 days.

OR
Hydroxychloroquine 800 mg salt (620 mg base) po, followed by 400 mg salt (310 mg base) po at 6, 24, and 48 hours + primaquine phosphate 52.6 mg (30 mg base = 2 tablets) po once daily for 14 days.
OR
Artemether-lumefantrine 4 tablets

| Uncomplicated/ P. vivax | Chloroquine-sensitive regions (everywhere but Papua New Guinea, Indonesia): |
| | For G6PD deficiency: consider primaquine base 0.75 mg/kg BW once a week for 8 weeks, with close medical supervision for potential primaquine-induced haemolysis. |

For Adults—
Chloroquine phosphate 1 gm salt (600 mg base) po, then 0.5 gm in 6 hours, then 0.5 gm daily for 2 days. Total 2500 mg salt;
OR
Hydroxychloroquine 800 mg salt (620 mg base) po, followed by 400 mg salt (310 mg base) po at 6, 24, and 48 hours + primaquine phosphate 52.6 mg (30 mg base = 2 tablets) po once daily for 14 days.
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OR
Hydroxychloroquine 10 mg base/kg po immediately, followed by 5 mg base/kg at 6, 24, and 48 hours. Total 25 mg base/kg + primaquine phosphate 0.5 mg po base once daily for 14 days;
OR
Artemether-lumefantrine (20/120 mg tablets)
(80 mg/480 mg) as a single dose, then
4 tablets again after 8 hours, then
4 tablets every 12 hours for 2 days
(take with food) + primaquine phosphate
52.6 mg (30 mg base = 2 tablets) po once
daily for 14 days.

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• 15 kg to <25 kg: 2 tablets (40 mg/240
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again after 8 hours, then 2 tablets
every 12 hours for 2 days;
• 25 kg to <35 kg: 3 tablets as a single
dose, then 3 tablets again after
8 hours, then 3 tablets every
12 hourly for 2 days;
• ≥35 kg: as per adult dose;

+ Primaquine phosphate 0.5 mg/kg po
base once daily for 14 days

Chloroquine-resistant regions: same as chloroquine-resistant as below

### Uncomplicated/ P. vivax

<table>
<thead>
<tr>
<th>For adults—</th>
<th>For paediatrics—</th>
</tr>
</thead>
<tbody>
<tr>
<td>Atovaquone-proguanil 4 adult tablets (1000/400mg) po in a single dose daily for 3 days (take with food) + primaquine phosphate 52.6 mg (30 mg base = 2 tablets) po once daily for 14 days.</td>
<td>Artemether-lumefantrine (20/120 mg tablets) weight-based dosing as above + primaquine phosphate*</td>
</tr>
<tr>
<td>OR</td>
<td>OR</td>
</tr>
<tr>
<td>Artemether-lumefantrine 4 tablets (80 mg/480mg) as a single dose, then 4 tablets again after 8 hours, then 4 tablets every 12 hours for 2 days (take with food) + primaquine phosphate 52.6 mg (30 mg base = 2 tablets) po once daily for 14 days.</td>
<td>Artemether-proguanil (pediatric tablets 62.5/25 mg) (weight-based dosing as above) + primaquine phosphate*</td>
</tr>
<tr>
<td>OR</td>
<td>OR</td>
</tr>
<tr>
<td>[quinine sulfate 650mg po three times a day for 3 days (7 days if Southeast Asia)] + [(doxycycline 100 mg po two times a day) or (tetracycline 250 mg po four times a day)] + primaquine phosphate 52.6 mg (30 mg base=2 tablets) po once daily for 14 days.</td>
<td>[(quinine sulfate 10 mg/kg po three times a day for 3 days); (7 days if Southeast Asia)] + (clindamycin 20 mg/kg per day divided in three times a day)] for 7 days under age of 8 + primaquine phosphate*</td>
</tr>
<tr>
<td>OR</td>
<td>OR</td>
</tr>
<tr>
<td>[quinine sulfate 10mg/kg po three times a day for 3 days (7 days if Southeast Asia)] + (doxycycline 2.2 mg/kg/two times a day) or (tetracycline 25 mg/kg/day divided four times a day) for 7 days if aged ≥ 8 + primaquine phosphate*</td>
<td></td>
</tr>
</tbody>
</table>

Remarks: * denotes primaquine phosphate 0.5 mg/kg po base once daily for 14 days

### Chloroquine-resistant regions

*Plasmodium falciparum* malaria as compared to quinine alone [4] [22]. Aspirin and non-steroidal anti-inflammatory drugs are not recommended as antipyretics for patients with suspected or diagnosed malaria because of the risk of gastrointestinal bleeding, renal impairment and Reye’s syndrome [12].
Anti-emetic is another commonly prescribed medications in emergency departments. Cautious use of anti-emetics in patients with vomiting is needed because these potentially sedative medications, with the occurrence of their potential neuropsychiatric adverse effects, can mask the diagnosis of cerebral malaria [4] [6] [9].

Suspected patients with more than two seizures within 24 hours should be treated as severe malaria. When seizure continues, airway should be maintained and anticonvulsants, such as parenteral or rectal benzodiazepines or intramuscular paraldehyde, are needed [4].

11. Subsequent Treatment

The subsequent treatment regimens for severe and uncomplicated malaria due to different infecting *Plasmodium* species and according to the potential antimalarial resistance are also shown in Table 1 [4] [9] [17] [23].

12. Summary

Malaria is a medical emergency and more than 20 cases per year have been notified in recent years in Hong Kong where malaria is non-endemic. High index of clinical suspicion is utmost important for not missing such a case, followed by prompt identification of patients with severe malaria to start appropriate effective antimalarial treatment within 24 to 48 hours of symptom onset after blood have been taken for thick and thin smears for diagnosis, and admission to intensive care unit for further management.

In addition, we have to be vigilant with increased awareness of not falling into common diagnostic traps. Be aware not to rule out malaria simply based on history of chemoprophylaxis use, not to assume that the chemoprophylaxis is the correct regimen, not to assume the compliance of chemoprophylaxis, not to initiate inappropriate treatment, not to delay treatment for cases of severe malaria and not to delay treatment beyond 6 hours after *Plasmodium falciparum* is confirmed by laboratory test [9] [12].

Whenever in doubt, do not hesitate to contact laboratories for urgent microscopic examination of thick and thin smears for diagnosis, consult infectious diseases physicians for assessment and management, in particular when encountering at-risk groups or patients with G6PD deficiency.

When we practice in this approach, the risk of missing any case of malaria presenting to emergency departments could be largely minimized.

Conflict of Interest

All authors have no conflict of interest.

References


Diseases, 49, 908-913. https://doi.org/10.1086/605436


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