New emerging drug-resistant malaria

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Clinical question: What is the best treatment for artemisinin-resistant malaria?
Results: There is still no better treatment than the presently used artemisinin-based combination therapies. A new antimalarial drug for this problem needs to be found.
Implementation: Pitfalls to avoid when treating drug-resistant malaria:
Keywords: malaria, drug resistance

Drug-resistant malaria
Definition: A “drug-resistant malaria” is a malarial infection that is not responding to the standard basic antimalarial treatment. Artemisinin is a drug used to treat multi-drug resistant strains of falciparum malaria. The World Health Organization has recommended that a switch to artemisinin combination therapies be made in all countries where the malaria parasite has developed resistance to chloroquine. Their site of action within the parasite remains controversial. At the chemical level, one theory states that when the parasite that causes malaria infects a red blood cell, it consumes hemoglobin within its digestive vacuole, liberating free heme, an iron–porphyrin complex. The iron reduces the peroxide bond in artemisinin, generating high-valency iron-oxo species and resulting in a cascade of reactions that produce reactive oxygen radicals that damage the parasite, leading to its death. Recently, emergence of new artemisinin drug-resistant malaria has been reported. This problem for falciparum malaria, of which artemisinin-based combination therapies are the recommended standard treatments, is specifically focused on herein.

Etiology: Resistant strain of malarial parasite.

Incidence: The incidence of drug-resistant malaria is different in different settings. The highest incidence is reported in Indochina (Thai–Cambodia border and Thai–Myanmar border).¹

Economics: No published study has addressed the economic issues of drug-resistant malaria. However, Foster et al reports that about US$1800 million is spent annually.²

Level of evidence: Systematic reviews, meta-analyses, randomized controlled trials (RCTs).
**Search sources:** PubMed, Cochrane Library, NHS Evidence, DARE, clinical evidence.

**Outcomes:** From the patient perspective the main outcomes are:

**Consumer summary:** The standard classical antimalarial drugs are not fully effective against emerging strains of drug-resistant malaria. The period of taking drugs has to be extended from that normally given. The control of drug use and surveillance of drug-resistant strains are the present means to fight this problem. New antimalarial drugs to solve the forthcoming problem are required.

**The evidence**

**How does artemisinin-resistant malaria affect the present standard antimalarial treatment?**
- Systematic reviews: 1
- Meta-analyses: 3
- RCTs: 1

There is a systematic review on artemisinin-resistant falciparum malaria. Partial artemisinin resistance is the present problem. The emerging problem is proposed to be due to selection of the resistant phenotype which is the result of exposure of the parasite population to artemisinin monotherapies in subtherapeutic doses for over 30 years, and the availability of substandard artemisinins. Focusing on the meta-analyses, for non-artemisinin-resistant cases, it is noted that “The addition of 3 days of artesunate to standard antimalarial treatments substantially reduces treatment failure, recrudescence, and gametocyte carriage.” Similar notes are reported in the other two meta-analyses. However, there is no specific meta-analysis on artemisinin-resistant falciparum malaria. The randomized trial concluded “Resistance is characterized by slow parasite clearance in vivo without corresponding reductions on conventional in vitro susceptibility testing.”

**Which treatments are best for cases of drug-resistant malaria?**
- Systematic reviews: 0
- Meta-analyses: 0
- RCTs: 1

There are no systematic reviews or meta-analyses for artemisinin-resistant falciparum malaria. The randomized trial showed that the present artemisinin-based combination therapies are still successful treatments but the clearance of malaria took longer than for the non-resistant cases. Prolonged drug administration is required and there is no report of adverse drug reaction due to this extended drug administration.

**Conclusion**
There is evidence on emergence of artemisinin-resistant malaria from highly endemic areas in Southeast Asia. Longer administration of standard artemisinin-based combination therapies is required for treatment of drug-resistant malaria. If resistance is complete, failure of present artemisinin-based combination therapies can be expected, and development of a new antimalarial drug for artemisinin-resistant malaria is required.

**The practice**

**Potential pitfalls**
Entering the endemic areas without concern for the existence of new drug-resistant malaria might lead to a lack of prevention. A history of exposure is required for early diagnosis and adjustment for the longer artemisinin-based combination therapies.

**Management**
Artemisinin-resistant malaria should not be managed by non-specialists. Suspected cases should be referred to specialists in infectious diseases or tropical medicine.

**Assessment**
- Traveling or living in the endemic areas (Indochina, Southeast Asia) poses a high risk.
- If history suggests a possibility of exposure and the diagnosis confirms falciparum malaria, refer to a specialist.
- If the primary standard treatment does not clear malaria, reassessment of patient history is needed and referral to a specialist is suggested.

**Treatment**
- Treatment must be by a specialist.
• Prolonged artesmin treatment is the present standard recommendation and there is no suggested alternative treatment.
• Long artesinin-based combination treatment until clearance of malaria.
• Close monitoring of clinical appearance and parasite clearance during treatment.
• Reporting to the local disease control authority is required.

Indications for specialist referral
• A history of visiting endemic areas.
• Failure of primary standard artesinin-based combination treatments.

Further reading


References