Misconcepts in Management of Severe Malaria

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Abstract

Early diagnosis and proper management is very important in management of severe malaria. Although there were many update information’s regarding to management of severe malaria, some misconceptions of the clinicians in management remained and should be reconsidered and corrected to avoid harmful management to the patients.

Keywords: Severe malaria; Misconcepts

Malaria is the most important parasitic infection in the world since it causes high mortality. Nearly one million people died due to malaria each year and most of them were children in African countries. No other parasite can cause high fatality patients like malaria. In management of severe malaria, early diagnosis and properly appropriate management of patients is most important and essential. Although World Health Organization (WHO) has worldwide disseminated informative books and handbook regarding management of malaria including pathophysiology, common errors in diagnosis and management, management of complications [1-4], some misconceptions in management of severe malaria have been found in many hospitals in the world. Mortality rates of severe malaria in many hospitals varied due to many factors and one factor is misconception of clinicians. Although many hospitals have well-equipments for supportive care of malaria complications such as dialysis facility for acute renal failure and volume ventilators for adult respiratory distress syndrome (ARDS), misconceptions in management of severe malaria may contribute to higher mortality rate of the patients. We would like to comment some misconceptions in management of severe malaria found in one western country. The misconceptions were published in one journal regarding to intensive care [5] as an example as followings:

Neurological Manifestations [5]: The authors mentioned “Computed tomography (CT) scan of the brain may show cerebral edema”. There were earlier discreet studies. Some studies mentioned CT [6] and even magnetic resonance imaging (MRI) [7] of non-fatal cerebral malaria showed no cerebral edema. Another studies of CT [8] and MRI [9] showed cerebral edema. In fatal cerebral malaria patients, mildly swollen brain is commonly found [10]. Cerebral edema may be also found in post-mortem of other non-malarial diseases too. WHO (2010) mentioned no significant difference in mortality between dexamethasone used in cerebral malaria, but gastrointestinal bleeding and seizure were more common with dexamethasone [2,11] and dexamethasone is not recommended as adjunctive treatment in cerebral malaria at presence. However the trails were small and there was no reported effect on disability [2].

Antimalarial Therapy: Quinine Dihydrochloride and Quinidine Gluconate [5]: The authors mentioned “Intravenous therapy with quinine or quinidine should be switched to oral therapy as soon as the patient is able to take oral medications. It is prudent to administer at least 5 to 7 days of quinine”. The present update of severe malaria by WHO [2] in 2010 recommended that parenteral quinine should be given to treat severe malaria patients for a minimum of 24 hours, once started (irrespective of the patient’s ability to tolerate oral medication earlier), and thereafter, complete treatment by given an oral course of quinine plus clindamycin or doxycycline or artemisin combination therapy. It is correct that total quinine should be given for 7 days. Shorter courses of quinine (e.g., 5 day-courses) are less effective [10]. In areas of multidrug resistant malaria, quinine should be combined with clindamycin or doxycycline if there is no parental artesunate available. Intravenous arsunate was proved to be better than intravenous quinine in treatment of severe malaria in both adults and children [12,13] since artesunate reduced mortality rates more than quinine. Quinidine is recommended outside malaria endemic area where there may be delays in obtaining other antimalarial drugs for severe malaria (e.g. artesunate or quinine). However quinidine should be given by careful rate-controlled intravenous infusion with continuous electrocardiographic monitoring [10]. The authors (using reference number 71 published in 2001) mentioned that “adverse outcome of either death or neurological sequelae was significantly less common in artemether group and treatment with artetherem was associated with significantly faster parasite clearance. In subgroup analyses, artemether was associated with a significantly lower mortality than quinine in adult with multisystem failure”. However, WHO in 2010 [2] showed that “two systematic reviews and 3 subsequent randomized controlled (RCT) trials found no significant difference in death rates between the groups receiving artetherem and quinine for severe malaria”.

The authors [5] also mentioned the large RCT comparison of intravenous artesunate and quinine in 1461 patients “in Thailand” or SEAQUAMAT study. The study was actually multicenter study conducted in many countries “outside” Thailand (e.g. Bangladesh, India, Indonesia, and Myanmar) [12].

The authors mentioned [5] that maintenance dose of artesunate should be given for 3 days which was incorrect. Three days of intravenous artesunate administration alone is not enough for treatment of severe malaria. In 2010, WHO [2] recommended that parenteral antimalarials (including artesunate) should be given to treat severe malaria patients for a minimum of 24 hours, once started (irrespective of the patient’s ability to tolerate oral medication earlier), and thereafter, complete the

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In conclusion, clinicians who manage severe malaria should get updated information’s otherwise misconcepted management may cause improper management and unfavorable outcome to the patients.

References