Case report

What about the treatment of asymptomatic forms of congenital malaria: case report and review of the literature

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Abstract

We report in this manuscript a case of newborn baby with asymptomatic form of congenital malaria; the screening of the peripheral blood smear of the baby after a positive result in the mother allowed the diagnosis. The authors were permitted through this case to discuss the therapeutic possibility in these cases.

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Introduction

In congenital malaria, infants are infected by the passage of parasites from mother to child before or during birth, it is a fatal condition that occurs at low rates in endemic countries [1]. Malaria is defined as an important public health problem in sub-Saharan Africa endemic countries to Plasmodium falciparum, recent data suggest that this is a more frequent situation than expected [2]. In these areas, even if the risk of contamination from the mother is high, newborns are protected against congenital malaria by maternal antibodies.

We report a case of asymptomatic congenital malaria; to our knowledge the articles available in the literature talk about symptomatic cases of the disease. On the other hand, the authors through this case discuss the therapeutic possibilities of asymptomatic congenital malaria. Congenital malaria is the presence of malarial parasites in the peripheral blood smear of the newborn from the first day until the seventh day of life [3]. It is due to transplacental transfer of parasites which infect the infant in utero or during delivery. In endemic regions congenital malaria is principally caused by Plasmodium falciparum [4]. In these countries mortality is over 86% in children under 5 years of age [5, 6]. The most common clinical symptoms are fever, anaemia, and splenomegaly in 80% of cases [7, 8].

Patient and observation

A 32 years old woman was admitted during labor, to the Department of Gynecology of the Moroccan Field hospital deployed in South Sudan. The patient was gravida 3, para 2, abortus 1 (2 living children) at the 39th week of gestation, prenatal screening tests were not performed. The clinical examination showed normal fundal height, with perception of uterine contractions and normal fetal heart activity, the cervix was effaced and dilated to 4cm. Trans-abdominal ultrasound was performed; the placenta was localized in the posterior side of the fundus; the amniotic fluid and the fetal growth were normal. A healthy male infant weighing 2600g was successfully delivered via spontaneous vaginal delivery, the apgar score was 8/9/10 with no blood loss during delivery, and no congenital abnormality was detected. 24 hours after delivery, the mother presented acute fever associated to chills and excessive sweating; she had anaemia with hemoglobin 8.2g/dl, hematocrit 25.8%, MCV 86.1fl, MCH 28.5pg, MCHC 31.5g/dl, total leucocyte count 5.5x10^3/ul and platelet count was low 56,000/ul. C-reactive protein (CRP) was normal (5,2mg/l), with normal range of serum electrolytes. Blood and urine cultures were sterile and peripheral blood smear showed malarial parasites of Plasmodium falciparum parasitemia at 4%.

The mother reported after evaluation a possible malaria infection in the 32th week of pregnancy, managed with unspecific treatment. The newborn baby was healthy and vigorous, exclusively breastfed, the clinical examination was normal; biological analysis showed hemoglobin 16.3g/dl, total leucocyte count 7.5x10^3/ul and platelet count 72,000/ul. C-reactive protein (CRP) was normal (< 5mg/l), with normal range of serum electrolytes. Peripheral blood smear was negative for malaria parasites; unfortunately, the other methods of diagnosis (Detection of PfHRP2 antigen, DNA detection and quantification) were not available for further investigations, the newborn was placed under close observation and a second examination of the peripheral blood, was done within 3 days; a thick and thin blood film revealed P. falciparum trophozoites, with a parasitaemia of 1,5%. The mother was treated according to local guidelines with artemether + lumefantrine and the infant was treated by oral chloroquine 10mg/kg for two days and 5mg/kg in the third day. Four days after treatment haemoparasites were cleared, with normal follow-up.

Discussion

Congenital malaria was described the first time in 1876; it is defined by the presence of asexual forms of malaria parasites in peripheral blood within the first week of life [9]. The occurrence of congenital malaria remains rare. The asymptomatic form is the most frequently reported [10], only 7 to 10% of newborns develop the disease form [11, 12]. The physiopathology still misunderstood, possible theories include transfusion of the fetal circulation by maternal blood during pregnancy, direct penetration in the chorionic villi due to premature separation of the placenta [13]. Clinical signs in order of frequency are anemia, fever, hepatosplenomegaly, hypotonia (poor feeding, lethargy), irritability and jaundice [14, 15]. Severe thrombocytopenia is also frequently reported [16]. The diagnosis is simple, based on microscopic examination of peripheric blood films, other diagnostic methods are possible if the blood film examination is negative: RDT (rapid diagnostic test) or PCR (polymerase chain reaction) [17]. In our case, the first thick blood smear examination was negative and could be explained by several factors such as fetal hemoglobin composition [18], protection by maternal antibodies specific to malaria after passive transplacental transfer, exclusive
breastfeeding [19] and by the reduction of red blood cells in the circulation due to their sequestration [20]. The second examination was positive at the age of 5 days [21].

The genomic amplification and DNA quantification increases the prevalence of diagnosis but they were not possible in our conditions. There are no clearly guidelines established to treat congenital malaria. Primaquine is not indicated because the infection is acquired congenitally (it doesn't include the persistent hepatic phase); chloroquine has been the drug of choice in case of congenital falciparum malaria [22]. However, a high incidence of chloroquine resistance has been reported by a recent study; in Nigeria 89.1% of patients were resistant to oral chloroquine, it was replaced by the association of sulfadoxine-pyrimethamine with good result [23]. Quinine plus clindamycin was also used in another study and have been reported as an effective treatment [24]. The efficacy of artesunate over quinine have been demonstrated [25]. The authors noted that it can be used as drug of first choice to treat congenital malaria. Preventive therapy using the association sulfadoxine-pyrimethamine to pregnant women during the second and third trimester, according to the world health was beneficial [26]. For patients who develop the symptomatic form of congenital malaria, the morbidity and the mortality associated is very high; although, in different studies, the majority of the infected neonates were asymptomatic during the observation period.

However, in various studies, the cases of congenital falciparum malaria, whether symptomatic or not, should be treated [11, 27, 28], but unfortunately none of the patients was followed until the 15th week corresponding to the necessary time for the elimination of the passive immunity transmitted by the mother [29], therefore the clinical evaluation of the asymptomatic patients was incomplete and they didn't receive any treatment. In our case, the patient received his treatment because the access to hospitals and healthcare facilities in such an emerging country was very difficult which impeded regular clinical monitoring and was impossible for him.

**Conclusion**

The repetition of thick drop examination has helped us to assert the diagnosis of congenital malaria in asymptomatic and healthy patient. When close clinical monitoring of patients over a 15-week period is feasible, antimalarial treatment should not be initiated unless a clinical signs of infection occurs and after diagnostic confirmation.

**Competing interests**

The authors declare no competing interests.

**Authors’ contributions**

All authors contributed to this work. They read and approved the final version of this manuscript.

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