Post-Malaria Neurological Syndrome - A Case of Bell’s Palsy After Plasmodium Vivax Malaria

Singh AK¹, Chakraborti S², Subhranag S³

¹Dr. Arvind Kumar Singh, MBBS, MD, Senior Resident, Kalawati Saran Children’s Hospital, Lady Hardinge Medical College, New Delhi, India. ²Dr. Snehansu Chakraborti, MBBS, MD, Professor, Department of Paediatrics, BSMC, Bankura, West Bengal, India. ³Dr. Shankha Subhra Nag, MBBS, MD, Senior Resident. Dept of Paediatrics, BC Roy Post Graduate Institute of Paediatric Sciences, Kolkata, India.

Abstract

Post-malaria neurological syndrome (PMNS) is defined as the acute onset of neurological or neuropsychiatric syndrome in a patient who had recently recovered from malaria and have negative blood film at the time of onset of neurological symptoms. It is relatively rare, with various clinical symptoms. We report first case of Bell’s palsy developing on 10th day of afebrile period after successful treatment of Plasmodium vivax (P.vivax) malaria and which completely recovered in next two weeks.

Introduction

PMNS is usually a short-lived, self-limiting condition with no long-term neurological sequelae. The time from eradication of systemic parasitaemia to the development of this syndrome can be up to two months¹. Clinical spectrum include generalised convulsions, acute confusional state, psychosis, tremors, cerebellar ataxia, motor aphasia, generalised myoclonus, bilateral facial nerve palsy²-⁴. It generally follows after recovery from Plasmodium falciparum (P.falciparum) malaria.

The Case

A six year old male was admitted in Paediatrics department with complaints of fever for 6 days, with headache. During physical examination, he was conscious, oriented and there was mild pallor but no icterus, lymphadenopathy or cyanosis. Mild splenomegaly was present. Other systems including respiratory, cardiovascular and central nervous system were normal. His blood examination revealed: Hb: 9.2 gm%, total leukocyte count: 7,800/cumm, neutrophile: 56%, lymphocyte: 40%, eosinophils: 03%, monocyte: 01%. Peripheral smear showed presence of ring form of P. vivax. Rapid diagnostic test kit was used to confirm the diagnosis of P. vivax malaria and rule out P. falciparum. Patient was put on chloroquine for three days. Fever subsided after three days and patient was discharged with primaquine on 5th day of admission. After 10 days of discharge patient presented with complaints of sudden onset of weakness in right side of face, difficulty in closing his right eyelid, deviation of angle of mouth towards left, and drooling of saliva from right side of mouth since morning (Fig. 1). He was fully conscious and well-oriented. Examination of facial nerve revealed it to be of right sided lower motor neuron type palsy. Other cranial nerves were within normal limit. There were no cutaneous lesions of herpes zoster in the external ear canal. There was no history of fever after discharge from hospital or any known pre-existing disease like diabetes mellitus. Investigation like complete blood count with peripheral blood smear, blood sugar, chest X-ray and MRI brain were normal. Patient was readmitted for observation and discharged after two days as symptoms were not progressing. No treatment was given and patient fully recovered in two weeks.

Fig 1: The patient showing features of right sided Bell’s palsy
Discussion

PMNS is defined as the acute onset of confusion, epileptic seizures, or any other neurological or psychiatric sign occurring with a latency of several days to weeks (generally within 2 months) after an episode of successfully treated *P. falciparum* malaria. Although it generally follows after infection with *P. falciparum* but few case reports of PMNS after *P. vivax* malaria are available in literature, one case presented with acute disseminated encephalomyelitis like picture, second one with acute inflammatory demyelinating polyneuropathy and third with bilateral facial nerve palsy. The prevalence of PMNS is 300 times more common in patients with severe malaria in comparison to uncomplicated malaria.

The first description of a post-infectious neurological complication of malaria was delayed cerebellar ataxia (DCA), identified in Sri Lanka in 1984. The prospective study, conducted in Vietnam and Thailand, first defining PMNS, described 22 patients, 19 adults, and 3 children, having neurological or psychiatric symptoms occurring within 2 months after an acute and cured malaria, among 18,124 patients with a treated *P. falciparum* malaria: overall incidence was 0.7 to 1.8 per 1000 in four regions were study was conducted. The syndrome was self-limiting, median duration 60 hours (range 24—240). A correlation between mefloquine therapy and PMNS was also noted.

Schnorf and others in 1998 proposed a classification of PMNS into 3 subtypes: a mild or localized form, characterized by isolated cerebellar ataxia or postural tremor; a diffuse, but relatively mild encephalopathic form, characterized by acute confusion or epileptic seizures; and a severe, corticosteroid-responsive encephalopathy that is characterized by motor aphasia, generalized myoclonus, postural tremor, and cerebellar ataxia.

The treatment of PMNS is mainly supportive. As reported previously, corticosteroids may be useful for patients with corticosteroid responsive encephalopathy. However no randomized trials of corticosteroid treatment have been conducted. Response to corticosteroids provides evidence that PMNS is probably an immunologically mediated phenomenon.

In our case, we found no evidence of cerebral Malaria. The patient developed Bell’s palsy after 10 days of full recovery and repeat blood test and other relevant investigations ruled out malaria parasite in blood or any other disease process. It fully recovered after 2 weeks without any medication. These findings are consistent with PMNS. It is unlikely that the patient’s symptoms were due to the toxic effect of antimalarial treatment. As the patient was treated with chloroquine and primaquine and Bell’s palsy is not described as side effects of either drug.

Conclusion

As the PMNS symptoms ranges from minor symptoms like bilateral facial nerve palsy and Bell’s palsy as in this case to severe generalized encephalopathy resembling an acute disseminated encephalomyelitis (ADEM), it is important to follow up the cases and distinguish PMNS from relapse of malaria and its complications. Clinicians should be aware of PMNS so that further studies could be done to explore its pathophysiology, range of clinical spectrum, and its relationship other post-infectious neurological syndromes.

References