Cerebral Toxoplasmosis in a Treatment Naive HIV Patient with High CD4 Count Responding to Treatment with a Regime of Cotrimoxazole and Pyrimethamine: Do We Need to Start Prophylaxis for Toxoplasmosis at a Higher CD4 Count?

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ABSTRACT

Cerebral toxoplasmosis is one of the commonest opportunistic infection of the nervous system in HIV patients. We present a case of gradual onset hemiparesis in an ART naïve HIV patient with high CD4 count who was subsequently diagnosed to be a case of cerebral toxoplasmosis based on radiological and serological investigations. The patient responded to a regime of Cotrimoxazole and Pyrimethamine. His CD4 count at diagnosis was 299/µl. HAART was started after completion of treatment of cerebral toxoplasmosis. Prophylaxis against toxoplasmosis is recommended with cotrimoxazole if the CD4 count is below 200/µl. However, in this case as the patient had developed toxoplasmosis at a CD4 count value above the cut-off value for prophylaxis for cerebral toxoplasmosis, it may be worth considering to starting prophylaxis at a higher CD4 count than 200/µl and continuing for a longer time than the current guidelines.

KEYWORDS: Cerebral toxoplasmosis; Opportunistic infections; HAART; Cotrimoxazole.

INTRODUCTION

Opportunistic infections are often the presenting diagnosis in patients with HIV/AIDS. Though most of the opportunistic infections occur at low CD4 counts, we present a case of cerebral toxoplasmosis presenting with progressive hemiparesis, who was subsequently diagnosed to have HIV infection with a CD4 count of 299/µl and made good recovery with a regime comprising of pyrimethamine, sulfadoxime and cotrimoxazole. The case is of interest as cerebral toxoplasmosis has been rarely reported with high CD4 count in treatment naïve HIV patients. The completeness of recovery with commonly available drugs before the initiation of Highly Active Antiretroviral Therapy (HAART), thereby makes a case for initiation of prophylaxis with cotrimoxazole for toxoplasmosis at higher CD4 count in HIV infected patients.

CASE HISTORY

A 47 year old trucker was admitted with history of progressive weakness of right half of the body with headache that started about four weeks back. The patient had become non-
ambulatory a week before the admission and was unable to feed or clothe himself on his own. His speech had become slurred. He had history of contact with multiple sexual partners. Examination revealed a male patient of BMI 23.5 kg/m² with normal blood pressure and pulse rate. He was conscious with Grade 3 upper motor neuron type of weakness of both upper and lower limbs on right side along with dysarthria. Early papilledema was seen on both eyes on examination.

CT scan of head - both plain and contrast enhanced, showed multiple ring enhancing lesions on both sides of brain with surrounding cerebral edema. He tested positive for HIV 1. Subsequent MRI of brain showed multiple target lesions on both sides of the brain. The CD4 count of the patient was 299/µl. Blood examination of the patient showed mild normocytic, normochromic anemia with normal biochemical parameters. He tested negative for Hepatitis B and C. Chest X-ray and ultrasound examination of abdomen did not reveal any pathology. He tested positive for IgG anti toxoplasma antibody (Figure 1).

A diagnosis of Stage IV HIV infection with toxoplasma encephalitis was made. The patient was initiated on a regime of pyrimethamine 200 mg and sulfadoxime 4 gm (Day 1) in two divided doses followed by pyrimethamine 50 mg and sulfadoxime 1 gm and cotrimoxazole double strength – 2 tabs thrice daily from Day 2. Inj. dexamethasone 8 mg IV thrice daily was started for cerebral edema. Folic acid supplementation was also given. The patient responded to treatment and by the end of 2nd week was able to walk with support. Follow-up CT scan of head showed resolution of the lesion by 3rd week. The dose of dexamethasone was gradually tapered and the patient was continued on cotrimoxazole double strength tablets with pyrimethamine and sulfadoxime for a total four weeks of therapy. HAART was started 2 weeks after completion of treatment of toxoplasma encephalitis.

DISCUSSION

Cerebral toxoplasmosis is a major cause of morbidity and mortality among HIV-infected patients, particularly from developing countries. Cerebral toxoplasmosis is an HIV-indicative event in 35% of patients and an AIDS-defining event in 75% of cases. Globally, T. gondii causes the most common focal brain lesion in HIV-infected patients. In most of the studies, incidence of toxoplasma encephalitis have been reported with CD4 count of less than 100 cells/µl. In a study involving 97 HIV patients with toxoplasma encephalitis, the median CD4 count was 68 cells/µl. Similar findings have been observed in studies from Puerto Rico and Brazil where toxoplasma encephalitis was observed in patients with low CD4 count. The incidence of toxoplasmosis varies by country and depends on the prevalence of T. gondii infection in the general population. T. gondii has an unusual clonal population structure consisting of three widespread lineages known as I, II, and III. Differences in genotypes of T. gondii isolates, races and ethnicities and the mode of transmission also seem to influence the occurrence of the infection.

Cerebral toxoplasmosis causes unifocal or, more commonly, multifocal lesions and, less frequently, diffuse encephalitis. Patients usually present with subacute symptoms. The clinical manifestations depend on the location and number of lesions. More frequent complaints include: headache (49-63%), fever (41-68%), focal deficits (22-80%), seizures (19-29%), mental confusion (15-52%), ataxia (15-25%), lethargy (12-44%), cranial nerve palsies (12-19%) and visual alterations (8-15%). Other manifestations include dysarthria, cognitive dysfunction, raised intracranial pressure and involuntary movements. The definitive diagnosis of cerebral toxoplasmosis requires the presence of the tachyzoite form of the parasite in cerebral tissue to be directly demonstrated by brain biopsy. In clinical practice, presumptive cerebral toxoplasmosis diagnosis depends on an association of serological, clinical and radiological findings. Detection of Immunoglobulin G (IgG) is possible within 2 weeks of infection using the Enzyme-linked immunosorbent assay (ELISA) test. Diagnosis is confirmed with a response to empiric anti-Toxoplasma therapy. A favorable clinical and radiological response is expected within 10-14 days of specific treatment.

The treatment of T.gondii infection is usually by a combination of pyrimethamine, sulfadiazine and leucovorin though
HIV Patients with CD4 count <100 cells/µl are given prophylaxis against *T. gondii* with a single double-strength-tablet daily dose of cotrimoxazole and continued till CD4 count >200/µl for more than 6 months. However, in a recent study, *T. gondii* cases have been reported in patients with high CD4 count >200/µl with 299 cases Per Year Follow up (PYFUP) with more than 50 cases reported at CD4 count >500/µl PYFU. Toxoplasma encephalitis has also been reported in patients with CD4 count >300 cells/µl but these patients had a history of CD4 count of <100 cells/µl at one stage and the rise of CD4 count was attributed to HAART. These cases of toxoplasma encephalitis occurring with high CD4 count have been attributed to persistence of immune dysfunction in spite of HAART or poor compliance. As toxoplasma encephalitis responds well to cotrimoxazole, it may be worth considering prophylaxis for toxoplasmosis at higher CD4 count (>200/µl) that may be continued till CD4 count remains well over 200/µl for a longer period.

CONCLUSION

Cerebral toxoplasmosis has been rarely been reported in treatment naïve HIV patients with CD4 count >200 µl. As cerebral toxoplasmosis respond well to a combination of cotrimoxazole and pyrimethamine in a HIV patient, it may be worth considering to start prophylaxis against toxoplasmosis with cotrimoxazole at a CD4 count higher than 200/µl as cases of toxoplasmosis have been reported with higher CD4 count.

CONFLICTS OF INTEREST: None.

CONSENT

The patient has provided written permission for this publication.

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