Unusual Presentation of Focal Tubercular Meningoencephalitis in an Elderly Female

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INTRODUCTION

Central nervous system tuberculosis (CNS TB) can manifest as meningitis, tuberculoma, abscess, and encephalitis. Overall approximately 10% of all cases of tuberculosis have CNS involvement. We present a case of a 64-year-old female who presented to us with fever, abnormal behaviour in form of rage reactions and convulsions and was later diagnosed to be a case of focal tubercular meningoencephalitis. Tuberculosis (TB) is a global health problem. According to the World Health Organization (WHO) analysis in 2007, 9.27 million new cases of TB and 1.3 million TB-related deaths occur in immunocompetent subjects.[1] Central nervous system (CNS TB) is the most catastrophic extrapulmonary form of TB which is associated with a mortality rate of up to 60%. In the survivors of CNS TB neurologic sequel may occur in up to 25% despite new neuroimaging diagnostic modalities and effective antituberculous therapy.[2] The incidence of tuberculosis varies from 9 cases per 100,000 population per year in the US to 110–165 cases per 100,000 population in the developing countries in Asia and Africa.[3,4] The estimated mortality due to tuberculous meningitis in India is 1.5 per 100,000 population.[5]

PRESENTATION OF CASE

A 64-year-old female presented to us with chief complaints of fever and headache since 15 days, abnormal behaviour in form of rage reactions with abusive language, and 2 episodes of GTCS on the day of admission. The fever was intermittent and low grade. There was no history of diplopia, vomiting, unconsciousness, cough, expectoration, haemoptysis. The relatives being of village background presumed the rage reactions to be of paranormal origin and opted to go to some village priest, but when she developed seizures, they brought her to this hospital.

On examination GCS was E3-M5-V4. Patient was irritable, Pulse-120 bpm, RR 20 cycles per minute, BP-140/76 in right arm. CVS, RS and per abdomen examination was normal. CNS examination: Cranial nerves normal, she was moving all 4 limbs, Bilateral plantars showed withdrawal response. She did not cooperate for sensory examination but, she was withdrawing limbs from pinprick. Signs of meningeal irritation in form of neck stiffness, Kernig’s sign and jolt accentuation test were positive.
On investigation CBS, KFT, LFT, electrolytes, X-ray chest, fundus examination were all normal. MRI brain revealed abnormal leptomeningeal enhancement in the right high frontoparietal concavity with associated infarction and oedema, suggestive of focal meningoencephalitis (Figure 1). CSF examination was done which revealed: TLC-220/mm³ with 80% lymphocytes. CSF glucose-22 mg/dl, proteins 250 mg/dl. CSF ADA – 48.6 IU (Strongly in favour of TB). CSF interferon gamma released assay (QuantiFERON-TB Gold test) was positive. CSF-PCR was negative for HSV.

A diagnosis of focal tuberculous meningoencephalitis was made and the patient was started with intensive phase anti TB therapy (Isoniazid 300 mg OD, Rifampicin 600 mg OD, Pyrazinamide 750 mg BID, Ethambutol 800 mg OD), Inj Dexamethasone 4 mg IV 8 hourly and Inj levetiracetam 500 mg IV 12 hourly for seizures.

After 1 week of hospital stay the patient’s general condition was improved fever subsided and the patient was discharged on oral Prednisolone 40 mg along with ATT and Tab. Levetiracetam 500 mg Bid. In the follow up after one month a repeat MRI of brain showed significant resolution of the lesion. (Figure 2)

**DISCUSSION**

Amongst the different variants of CNS TB, meningoencephalitis is the most severe, life-threatening form. It can occur in an acute or chronic variant. The predominant clinical presentations of TBM are headache, vomiting, fever and neck rigidity.[8] Our patient had these findings, along with that the patient’s abnormal rage reactions, abusive aggression and seizures which pointed towards encephalitis which was later confirmed by neuroimaging and CSF analysis. Factors that are associated with poor outcome in TBM are extreme ages, stage of illness and development of hydrocephalus. A study conducted by Hosoglu et al;[7] showed that drowsiness on admission is an important predictor for residual neurologic sequel.

The basic pathophysiology of CNS TB is the entry of the bacilli to CNS by hematogenous route from another focus. Rich and McCordock,[9] in their study suggested that CNS tuberculosis develops in two stages. The first stage heralds with formation of small tuberculous lesions known as “Rich’s foci” in the CNS, either during the stage of bacteraemia of the primary tuberculous infection. The most favored sites in the brain are meninges, the subial or subependymal surface of the brain or the spinal cord. These lesions may remain dormant for years after initial infection. In the second phase rupture or growth of one or more of these small tuberculous lesions produces various types of CNS tuberculosis.[9,10] Rupture of Rich foci into the subarachnoid space or into the ventricular system results in meningitis. Tuberculous encephalopathy, a syndrome is thought to be exclusively present in children. Diffuse cerebral oedema, convulsions, stupor or coma are characteristic of encephalitis. CSF is largely normal in pure encephalitis, but abnormal CSF picture suggests meningoencephalitis.[11]

As far as management of CNS TB is concerned; the first-line anti-TB agents recommended for the treatment include rifampicin (RIF), isoniazid (H), pyrazinamide (PZA), ethambutol (EMP) and/or streptomycin (S). Among the first-line drugs used in TBM, RIF, EMP and streptomycin have poor penetration across the blood-CSF barrier.[12] Currently, WHO recommends a 2-month treatment with 4 first-line drugs in the intensive phase, followed by a continuation phase with at least rifampicin and isoniazid for 4-10 months.[13] There is evidence that addition of corticosteroids improves both survival rate and neurological outcome in patients with tuberculous meningitis.[14-15]

**CONCLUSIONS**

Tuberculous meningoencephalitis is the most catastrophic form of CNS TB. Classical clinical features, neuroimaging and recent diagnostic laboratory studies (CSF ADA, Quantiferon TB Gold assay) has improved the detection rate, so that early treatment can be initiated to limit the mortality and improve clinical outcomes in such cases.

**REFERENCES**
