Brain Studded with Innumerable Larger Stones: Starry Sky at Night Appearance: Multiple Calcified Giant Tuberculoma with 15 Years of Follow-up

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Abstract

Keywords

- multiple intracranial tuberculoma
- stones
- starry night appearances
- hydrocephalus
- histopathological diagnosis

Tuberculosis is rapidly reemerging as a major health problem due to rising incidence of HIV cases across the globe. Central nervous system (CNS) involvement due to tuberculous affection is rare and commonly presents as meningitis, ventriculitis, or, rarely, as intracerebral granulomas, which can occur as multiple smaller lesion or occasional solitary tuberculoma.

A 24-year-old woman presented in our OPD with headache and occasional vomiting for past 8 months, but it increased over past 2 weeks. She was a diagnosed case of pulmonary tuberculosis and kept on antituberculous drugs for past 10 months. On examination, she was consciously alert, her vital was stable, fundi showed bilateral papilledema, and the rest of the neurologic examination was essentially within normal limit. Noncontrast computed tomographic (CT) scan of the head showed larger-sized multiple calcified lesions, scattered throughout the supratentorial compartment, which were noticed in each axial section images, with absent perilesional edema in the bilateral cerebral hemispheres, bilateral cerebellar hemisphere, right basal ganglia, and thalamus, as well as in left cerebral peduncle mimicking as brilliantly shining stars in the sky at night appearance (►Figs. 1, 2). In view to ascertain the definite diagnosis, a plan was made for right frontal craniotomy and excision of tuberculoma. She was taken up for surgery under general anesthesia. Right frontal osteoplastic flap craniotomy was made. After dural opening, a small cortical incision was made, just beneath the corticectomy incision. A well-encapsulated mass was seen and excised, and wound was closed in layers after achieving hemostasis. Ventriculoperitoneal (VP) shunt was also performed to relieve persistent hydrocephalus.

Histopathologic examination of specimen was suggestive of tuberculoma granuloma, and the patient was advised addition antituberculous medication for further 8 months as she had already received 10 months medication. At last follow-up at 15 years following surgery, she was doing well.
The patient underwent a right temporal tuberculoma excision surgery, and she had no focal neurologic deficit. The cranial CT scan at the present follow-up revealed the presence of multiple intracranial tuberculomas. Although the sizes of the multiple lesions were slightly reduced, still considerable sizes are persisting (\textbf{Figs. 3, 4}).

Intracranial tuberculoma is not a common entity, and multifocal tuberculosis incidence is increasing due to the rise in immunocompromised conditions such as HIV.

CNS involvement in tuberculosis is usually caused by hematogenous spread from either the lungs or gastrointestinal tract, which in the beginning causes formation of small subpial or subependymal infective focus also called \textit{Rich focus} and acts as a reservoir from which intracranial spread can subsequently occur either during the primary infection or later as reactivation causing a postprimary infection.\textsuperscript{1–3} CNS tuberculosis is usually secondary to hematogenous dissemination of bacilli from an infection focus elsewhere in the body. All intracranial compartments can be affected. Intracranial involvement broadly can be divided into intra- and extra-axial affection. Extra-axial involvement includes tuberculous meningitis that is more common compared to pachymeningitis that is extremely uncommon. Extra-axial tuberculous involvement can manifest as tuberculous granuloma formation also called \textit{tuberculoma}, with others being focal cerebrites, tuberculous abscess, and encephalopathy.

These CNS involvements such as meningitis, solitary tuberculoma, abscess, infarct, or miliary parenchymal lesion...

\textbf{Fig. 1} Noncontrast computed tomography scan of head showing presence of multiple calcified lesions in the bilateral temporal lobes and cerebellar hemisphere, and right basifrontal regions mimicking starry sky at night appearance (preoperative scan).

\textbf{Fig. 2} Cranial computed tomography scan of head showing presence of multiple calcified lesions in the bilateral frontal and parietal region in the supratentorial compartment (preoperative scan).

\textbf{Fig. 3} Postoperative computed tomography scan of head at follow-up period after 15 years following craniotomy and resection of right frontal calcified tuberculoma in brainstem, and supratentorial compartment.
are seen in approximately 5 to 10% of tuberculosis cases. The intracranial tuberculoma account for 5 to 30% of all intracranial space-occupying lesions in developing countries.1

The tuberculosis leads to intracranial calcifications in 10 to 20% cases. It is usually located supratentorial in adults and infratentorial in children.2 Intracranial tuberculomas are usually seen in the basal section of the brain. Tuberculoma may show a spectrum of clinical manifestations depending on location, size, associated edema, number of tuberculoma, and obstruction to cerebrospinal fluid (CSF) pathway. The most frequent symptoms include headache, epileptic attacks, focal neurologic deficits, and features of raised intracranial pressure.3,4

It is difficult to diagnose when the patient has no history of tuberculosis. Radiologic diagnosis is also difficult due to varied and nonspecific imaging presentation.

Our patient was a known case of pulmonary tuberculosis, who was receiving antituberculous therapy, and she presented with headache. Noncontrast CT of the head showed multiple calcified lesions in the brain with single larger lesion in the right frontal region.

With history of tuberculosis, first differential of tuberculosis was made and further histopathologic confirmation was made after excision of lesion in the right frontal region. In the developing countries, granulomatous infections such as tuberculosis and parasitic infections such as neurocysticercosis and hydatid cyst can also result in the development of characteristic intracranial calcifications. Rising number of HIV cases may lead to increased incidence of tuberculosis, and other associated infections including toxoplasmosis, rubella, cytomegalovirus, and herpes simplex may promote occurrence of intracranial calcifications.2 The other differentials can be multiple calcified meningioma, craniopharyngioma, oligodendroglioma, dystrophic calcification in chronic infarction, chronic vasculitis, and exaggerated physiologic calcifications.

CT can identify and clearly delineate and characterize brain stones defining their locations, size, number, involvement of supra- and infratentorial compartment, extent, associated perilesional edema, and associated hydrocephalus. However, magnetic resonance imaging (MRI) of the brain remains the gold standard imaging modality, and advent of newer sequences such as gradient echo T2-weighted images and susceptibility-weighted image is adjunctive and clearly categorizes the tuberculoma. The calcified tuberculoma can appear as “broken shell” or dense centrally located lobulated calcification.5,6

Neurocysticercosis is a major cause of adult-onset seizures in developing countries; here calcifications represent dead larvae.7 The hydatid disease is considered a rare cause of intracranial calcification, and lesion is usually solitary, septated, or multilocular representing the dead parasite.8

In countries where suspicion for tuberculoma is high and clinical and imaging findings suggests high probability of tuberculous mass lesion, a trial of anti-tuberculous medication course should be tried. However, the surgical excision or biopsy is usually reserved for the cases, in which the diagnosis of the intracranial calcification or mass lesion is in doubt or lack of appropriate response to empirical medical therapy after receiving for at least 2 months or paradoxical response to antituberculous medications. However, in cases associated with severe obstructive hydrocephalus or raised intracranial pressure, CSF diversion surgery such as VP shunt surgery is required.9,10

Our patient was already on ATT for past 10 months, and she developed the lesions so the diagnosis was doubtful. Therefore, to confirm that, single lesion was excised and additionally VP shunt was also placed. Dawar et al observed that the calvarial tuberculosis is rare and can present with a localized focal scalp swelling with overlying skin discoloration and solitary or multiple discharging sinus. They reported a case of calvarial tuberculosis in 17-year-old girl, who had a large painless scalp swelling spreading over both the frontal areas from supraorbital margin to coronal suture without discoloration or sinus formation, and neuroimaging showed absence of extradural granulation tissue deposits with very subtle bony architecture changes of adjacent calvarium. The swelling completely subsided with medical therapy alone.11

To conclude, early diagnosis and appropriate management including medical or surgical intervention should be tailor-made, and should constitute an important step in terms of reducing mortality and morbidity. In our country tuberculosis is an endemic disease and the physician should be aware that size of tuberculoma can be variable and sometimes can be of giant sizes, which may persist up to many years as current case persisted for at least 15 years despite receiving full course of antituberculous therapy.
References