Transcalvarial and Transdural Involvement of Skull Actinomycosis with Recurrence

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Abstract

Actinomycosis is caused by anaerobic gram-positive bacteria of Actinomyces genus, generally found as commensals in the human body and infection occurs when the immune system is compromised. We present the case of a 35-year-old diabetic woman, who presented with headache and sudden onset of left-sided weakness apart from scalp swelling, which on imaging showed transcalvarial transdural involvement, and subsequent surgical debridement and histopathologic analysis confirmed actinomycosis. Follow-up showed recurrence of the lesion at the same site following a period of remission. Calvarial involvement of actinomycosis is rare and can mimic even neoplastic or malignant lesions, hence requiring prompt diagnosis, treatment with surgery, and long-term antibiotic treatment to prevent complications.

Actinomycosis is a suppurative disease that involves chronic infection of the soft tissue and bone, caused by anaerobic gram-positive bacteria of Actinomyces genus, most commonly due to the organism Actinomyces israelii. These organisms are either anaerobic or microaerophilic, producing branching filaments resembling fungi. They were previously mistaken for fungi due to their capability to form filaments. The clinical infection can resemble several other chronic disorders such as tuberculosis or malignancy. Appropriate diagnosis and management are of paramount importance as the disease is treatable. Manifestations of the disease are generally in the form of abscess formation associated with complications such as fibrosis and discharging sinuses. The infection spreads diffusely throughout the soft tissue across the fascial barriers.

A 35-year-old woman presented with complaints of subacute onset of weakness of the left upper and lower limbs. The patient previously had history of headache for the previous few months with a diffuse swelling over the scalp. She was recently diagnosed with diabetes mellitus. There was no history of fever, and on examination, sensorium was normal. Power in the left upper limb and lower limb was 2/5. Reflexes were brisk, and plantar reflexes were extensor. The random blood sugar was elevated. Blood urea, serum creatinine, and electrolytes were normal. The patient’s hemoglobin was normal, and blood count showed mild eosinophilia (7%). She was referred for imaging evaluation of her symptoms. Routine noncontrast and contrast-enhanced computed tomography was done, which showed a hyperdense lesion over the right parietal lobe associated with dural thickening. There was sclerosis in the overlying skull bone with irregularities in the inner table of the skull (►Fig. 1A). There were hypodense areas within the brain adjacent to the lesion. Minimal postcontrast enhancement was noted.

Contrast-enhanced magnetic resonance imaging (MRI) was done for further characterization of the lesion. MRI showed a large T1 isointense, T2 hypointense lesion in the epidural space infiltrating into the brain parenchyma causing confluent FLAIR hyperintensities in the subcortical, deep, and periventricular white matter in the right cerebral hemisphere due to cerebritis or meningitis with vasogenic edema. It was associated with midline shift to the left side and apposition of the lateral
ventricles. Altered signal changes were noted in the diploic spaces with thickening of the parietal bone. An extracranial component was also noted, which was communicating with the intracranial lesion through the sagittal suture. Moderate postcontrast enhancement was noted in the lesion. Superior sagittal sinus was infiltrated by the lesion (►Fig. 1B–D).

The initial imaging differential diagnoses were en plaque meningioma, tuberculosis, metastasis, and lymphoma due to the presence of soft tissue lesion along with underlying bony sclerosis and erosion of the inner table. Biopsy was obtained from the lesion as was subjected to histopathologic and microbiologic examination. Fungal microscopy and cultures were negative. The histopathologic examination was suggestive of actinomycosis. Right parietal craniotomy was done after the first MRI, and the lesion was debrided. On opening the skull, there was a mass with grayish black color that was submitted for frozen section with differential of a primary melanocytic lesion of the meninges to categorize the type of lesion. The frozen section showed granulomatous inflammation with some organisms that could not be categorized and was reported as inflammatory lesion likely to be a mycetoma, without any evidence of any neoplastic lesion.

On paraffin processing the tissue, the sections showed numerous well-formed granulomas (►Fig. 2) with filamentous organisms in the center surrounded by a lymphohistiocytic collection and giant cell reaction. In addition, there was a distinct “Splendore-Hoeppli phenomenon” in the form of a nice annular eosinophilic to pinkish material around the bacterial organisms representing host response in the form of an antigen-antibody reaction. The higher-power images highlighted the bluish hematoxyphilic filamentous bacterial profile. No fungal organism was seen even on the special stains. With these findings, final diagnosis of *A. israelii* was made.

The patient was started on penicillin after the biopsy reports were available. Follow-up MRI done 5 months later revealed residual/recurrent lesion of smaller size than preoperative lesion. A second follow-up MRI performed 1.5 years after surgery showed persistence and mild increase in the size of the lesion compared with the second MRI. The patient was continued on antibiotics with plan for surgical decompression if symptomatically worsened.

The very first report of central nervous system (CNS) actinomycosis was described in 1889 by Delepine. Many isolated cases have been reported both in the preimaging era and after the advent of CT/MRI. The CNS infection is generally secondary to a source in the respiratory tract, abdomen, pelvis, or cervicofacial routes such as dental or paranasal sinus manipulation and posttraumatic events. The source of CNS infection spreads through hematogenous routes. However, direct spread of the infection from dental infections, or from paranasal sinuses or other sites of cervicofacial region is also possible.

The type of CNS involvement is predominantly either focal lesion or dural/epidural involvement with adjacent parenchymal involvement. Though dura is considered a relative barrier, indolent infections such as tuberculosis are known to cause dural involvement. Local spread and invasion across tissue planes is considered one of the characteristic features of actinomycosis as seen across pleura in thoracic...
involvement, deep fascial planes of the neck in cervicofacial type, and dura in the CNS.6

Few case reports have described mainly the second type of dural/epidural actinomycosis with adjacent parenchymal involvement.3,5,7,12 The first report of an epidural actinomycotic infection was described by Kirsch and Stears in 1970 and hypothesized it due to spread from an indolent dental/mandibular focus.3 In the two cases of cranial actinomycosis reported by Sundaram et al in 2004, one had an underlying penetrating injury,6 but other reports of calvarial/dural involvement did not show any obvious predisposition of either dental source of infection or trauma for the initiation of infection. However, actinomycosis has been ascribed to be the cause of mandibular osteoradionecrosis and this is considered to be due to preferential localization of actinomycosis to devitalized tissue as reported in thorax and in relatively less vascular mandible after radiotherapy.3,14 It is possible that actinomycosis may become indolent once it establishes in a relatively avascular ischemic milieu. Similar involvement of relatively less vascular mastoid and petrous temporal bone involvement have been described.9 The associated actinomycosis-induced arteritis might prolong the indolent course by the involvement of both scalp vessels and dural–side arteries giving rise to relatively avascular environment. This may explain the chronicity and recurrence that is encountered with actinomycosis.

The bacterial cultures can take long time to yield results and are positive in less than 50% of cases. Medical management is the primary mode of treatment for actinomycosis, and penicillin is the drug of choice. However, the duration of treatment can be as long as few months, and serial radiologic follow-up is necessary to assess resolution of the lesion. Inadequate treatment can result in relapses. Adequate surgical debridement will help control the infection and can shorten the duration of antibiotic treatment.

In our case, the patient was a recently diagnosed case of diabetes mellitus who presented with subacute onset of neurologic deficits on the left side. The patient had a history of chronic headache for the past 1 year with a scalp swelling. She had no other extracranial source of infection in the body. Uncontrolled diabetes with a state of chronic hyperglycemia causes changes in microenvironment of the tissue with altered wound healing and control of infection. Hence adequate control with oral hypoglycemic agents is necessary for the control of ongoing infections in the patients. Craniotomy and debridement with antibiotic therapy initially caused symptomatic improvement. However, the patient had a recurrence of the lesion on follow-up, which is known to occur in actinomycosis.

In conclusion, actinomycotic osteomyelitis of the calvarium is rare but a known complication of actinomycosis. The disease has a chronic course and generally presents with nonspecific symptoms. In imaging it can masquerade as either malignancy or other chronic diseases. A high clinical suspicion and adequate histopathologic examination of the lesion are important for diagnosis and follow-up of the patient. The patient will generally require debridement of the lesion with medical management for long durations for complete resolution. Even with adequate treatment, recurrences are known to occur.

Conflicts of Interest
None declared.

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