Eosinophilic Granuloma of the Occipital Bone Presenting as Intracranial Venous Hypertension

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ABSTRACT: Intracranial venous sinus thrombosis is an uncommon cause of pseudotumor cerebri. The diagnosis is often not confirmed on the rationale that treatment will not be altered. We report a case presenting a pseudotumor cerebri where the underlying pathology disclosed dural sinus thrombosis resulting from compression by an eosinophilic granuloma of the occipital bone. Routine CT of the head and Technetium-99m brain scan initially demonstrated neither tumor nor thrombosis. Plain skull x-rays subsequently revealed a lytic lesion of the occiput. When reinvestigated with CT using bone density windows the tumor was revealed. Excision of the tumor and a short course of cobalt therapy was curative. Special techniques in nuclear scanning, CT and MRI designed to improve the sensitivity for diagnosing venous sinus thrombosis are described. This case illustrates the importance of establishing a definitive diagnosis and shows the importance of pre-test consultation between clinicians and radiologists to ensure that specific investigative techniques are properly utilized.


Intracranial venous sinus thrombosis is a well recognized but uncommon condition in children and adults. Historically it has been associated with infective intracranial processes such as mastoiditis and periorbital cellulitis. As a result of widespread use of antibiotics it is now more frequently aseptic in origin associated with states predisposing to thrombosis. Venous sinus thrombosis has been suspected (but less frequently confirmed) as a cause for pseudotumor cerebri. In spite of major advances in diagnostic imaging with increased efficacy and safety, not all clinicians have pursued this specific diagnosis, accepting instead the less definitive diagnosis pseudotumor cerebri. This is especially true when no change in treatment is anticipated. We describe a case initially diagnosed as pseudotumor cerebri where pursuit of a definitive diagnosis resulted in significant modification in treatment and outcome for the patient.

CASE REPORT

A ten-year-old Caucasian male was well until one month prior to admission when he developed symptoms of an upper respiratory infection. He was treated with two courses of antibiotics before his symptoms completely resolved. Approximately one week later he developed a severe headache which became persistent, associated with recurrent vomiting, photophobia and decreased energy. His headache was worse in the mornings and he would awaken at night with the pain. After six days he saw a pediatrician. Past history included a congenital partial ptosis of his right eye and tonsillectomy at age five. On exam he was afebrile with normal vital signs. His general medical examination was unremarkable except for the fundoscopy which revealed bilateral disc edema.

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He was admitted to hospital where cranial CT with enhancement was done and reported as normal; it was noted that he had small ventricles. Lumbar puncture was performed which revealed an elevated opening pressure of 51 cm water. His cerebrospinal fluid (CSF), cell count, and chemistry were normal. Following the LP he had significant improvement in his headache and vomiting.

One week later he had recurrence of severe headache and vomiting associated with worsening of his papilloedema. Repeat LP revealed an opening pressure of 52 cm water. CSF was removed until the pressure was reduced to 25 cm water. Analysis of the fluid was again normal. His symptoms improved transiently but then recurred and he was transferred to our centre.

On arrival he was agitated and crying. His vital signs were normal. There was mild meningismus. Papilloedema was present bilaterally. He had a partial right abducens nerve palsy. There was one centimeter, mildly tender, mid-occipital nodule interpreted as lymphadenopathy. Cranial CT was unchanged from previous studies. Mastoid x-rays were normal. Technetium brain scan with static and dynamic imaging suggested normal distribution in the dural sinuses.

A plain occipital skull x-ray was done assessing the tender occipital lump initially suspected as lymphadenopathy. This revealed a large lytic lesion of the occiput at the torcular, extending slightly to the right of the midline (Figure 1). Review of the initial CT scan done with brain density windows only, had failed to demonstrate this abnormality. A repeat CT scan of the occipital region with bone density windows revealed a diploe/inner table tumor extending through the outer table corresponding with the tenderness of his right occiput and the lesion seen on plain skull x-ray (Figure 2a, b).

A craniotomy was performed and at surgery an occipital bone tumor was found infiltrating and compressing the superior sagittal sinus and dura adjacent to it on the right. The superior sagittal sinus was distended and the right lateral sinus had no demonstrable filling, presumably a result of the compressive effects of the tumor. Pathology of the tumor revealed an eosinophilic granuloma. Post-operative cobalt therapy consisting of 300cGy in three doses was delivered to the affected area. Following radiotherapy a cranioplasty was performed. At this second surgery there was still some distension of the superior sagittal sinus but filling of the right lateral sinus was demonstrable. Bone scan of his whole body revealed no other evidence of increased uptake suggesting his eosinophilic granuloma was solitary. He has done well in follow up.

**DISCUSSION**

Pseudotumor cerebri, first described by Quinke in 1893, is now considered a disorder of intracranial pressure regulation in which CSF pressure is elevated without clinical, laboratory or radiologic evidence of focal lesions, meningeal inflammation or hydrocephalus. It is an uncommon diagnosis in children and adults and recent reviews have summarized various conditions associated with the syndrome. The diagnosis is based on four criteria (Table 1).

Dural sinus thrombosis can present as pseudotumor cerebri and has frequently been cited as an association or cause.
Ahlskog and O’Neill’s recommendation that these patients be considered a separate entity bears clinical importance as treatment may be modified by demonstration of sinus thrombosis.10,11

Intracranial venous sinus thrombosis has been associated with a variety of conditions (Table 2). Early reviews noted up to 39% of cases in children resulting as a complication of middle ear infection.8 In Couch et al.’s more recent series, only 8% resulted from otitis media or its complications.7

We believe this is the first reported case of eosinophilic granuloma causing dural sinus thrombosis and presenting as pseudotumor cerebri. Routine CT imaging failed to demonstrate the bony defect later clearly demonstrated with CT, utilizing bone density windows. The large lytic lesion seen on plain skull x-ray prompted the repeat CT. Several authors have published cases where contrast enhanced CT scanning has demonstrated both the presence and resolution of thrombus in the posterior superior sagittal sinus (Empty Triangle Sign or Delta Sign).13,14,15 Zilkha and Diaz offer advice on how to increase the sensitivity of this method and its major pitfalls.12

Dynamic radionuclide scanning with technetium-99m DTPA has also been reported to be useful in demonstrating venous sinus thrombosis. Barnes and Winestock describe increased sensitivity of this procedure using an 80-lens optical camera with the patient’s head in the posterior oblique position.14 Front et al. have reported their five year experience with camera with the patient’s head in the posterior oblique position.15 Static imaging was found to be 100% sensitive and 86% specific for SST when compared to angiography. In our patient, the diagnostic sensitivity of both CT and nuclear scans could have been increased had these techniques specific to investigation of venous sinus thrombosis been employed.

Recently there have been reports describing the superiority of MRI in demonstrating both superficial and deep cerebral venous thrombosis16,17 and it is now the preferred method for demonstrating cerebral venous thrombosis. Where MRI is not available, cerebral angiography may be helpful in demonstrating dilated corkscrew veins, shunting of blood and filling defects in the involved sinus.

This case confirms the importance of ruling out venous sinus thrombosis as a cause of pseudotumor cerebri. The diagnosis in our patient led to successful management of the tumor and underlying intracranial hypertension. Plain skull films, frequently ignored in favor of more advanced technology, were helpful on this occasion. Specialized CT and radionuclide techniques, and now MRI imaging are the procedures of choice in diagnosing venous sinus thrombosis. Consultation with the radiologist may enhance their diagnostic sensitivity.

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REFERENCES