Transient, unilateral, complete, oculomotor palsy in an adult patient with idiopathic intracranial hypertension

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ABSTRACT

Idiopathic intracranial hypertension (IIH) is a well recognized condition of elevated intracranial pressure of unknown cause. Many etiologies have been proposed, but few—besides a high body mass index, hypervitaminosis A, steroid withdrawal, exposure to tetracyclines and female gender—have been proven. The main morbidity in this condition is visual loss and is often reversed if recognized early and treated promptly with weight reduction, a low-sodium diet and acetazolamide.

Key words: idiopathic intracranial hypertension, oculomotor palsy, headache, adult

Introduction

Idiopathic intracranial hypertension (IIH), also known as pseudotumor cerebri, is a clinical condition of elevated intracranial pressure of unknown etiology. Typical presentation consists of daily headache, pulse-synchronous tinnitus, transient blurred vision, papilledema with associated visual loss, and diplopia from abducens nerve paresis. (1)

Several conditions have been suspected, but only a few, including obesity and weight gain during the 12 months before diagnosis, steroid withdrawal, female gender, hypervitaminosis A and related compounds, have been proven. (2-4)

The treatment is focused on the correction of the presumed cause, including a low-sodium diet, diuretics and a weight-reduction program. (5)

To our knowledge, we describe, possibly, the first adult patient with complete oculomotor palsy.

Case report

A 53-year old, right-handed woman presented with an acute onset of sharp pain involving the top of her head associated with nausea, vomiting, photophobia and phonophobia two weeks prior to her neurological visit. At that time she was hospitalized and treated symptomatically for headache. During the patient’s hospitalization, left ptosis was noticed by her primary doctor. Initial metabolic work-up was unremarkable and an MRI (magnetic resonance imaging) was performed, remarkable for a partially empty sella, not reported by the radiologist (figure 1). No other abnormalities were seen. Cavernous sinus was normal and no abnormal enhancement with gadolinium was seen.

The patient had been headache free for an entire week prior to her neurological exam, and the only complaint she had was not being able to elevate her left eyelid. She denied any fever, visual obscuration, diplopia or tinnitus. Past medical history was remarkable for migrainous headaches, (she had been headache free for years prior to this event), hypertension and depression.

While being on the antidepressant, venlafaxine, she has gained 45 pounds during the previous 12 months. Physical exam revealed 5’8” height, 228 pounds weight, blood pressure of 150/90 mm Hg and regular heart rate of 70.

Fundoscopic examination revealed grade III papilledema OS (left eye), grade II OD (right eye). There was no ocular or carotid bruit. The cardiac examination was normal, without murmur. Neurological examination was remarkable for complete left ptosis, mydriasis, light stimulation revealed OS 5mm down to 4mm, OD 3mm down to 2mm, normal abduction and complete palsy of oculomotor innervated muscles on the left.

Right extraocular motor activity was entirely normal. No visual field defect was detected.

An emergent magnetic resonance angiogram (MRA) and venogram (MRV) were obtained without any abnormalities. In particular, there was no evidence of intracranial aneurysm or venous sinus thrombosis. Subsequently, the same day a spinal tap was performed and revealed ele-
vated intracranial pressure of 420 mm water. Closing pressure was 100 mm water after 30 ml of clear cerebrospinal fluid (CSF) was removed. Analysis of CSF was normal, cytology was also normal, no malignant cells were found. The repeated metabolic work-up, including sederate, C - reactive protein and antinuclear antibodies was normal.

The patient was immediately started on acetazolamide 500 mg twice daily, a weight reduction program and instructed to contact her psychiatrist regarding stopping venlafaxine and starting another antidepressant. The patient’s symptoms resolved two weeks later, with no residual left oculomotor deficit and normal pupillary response.

**Discussion.**

The annual incidence of IIH is 0.9/100 000 persons and 3.5/100 000 in females 15 to 44 years old. Among obese women the incidence is 19/100 000. (6) More than 90% of IIH patients are obese and more than 90% are women of childbearing age. Female preponderance and the association of obesity is true only for postpubertal patients. (7) The mean age at diagnosis is 30 years. (6) There have been no reports of venlafaxine induced IIH, but it can produce weight gain. (8) The most accepted hypothesis is that IIH is a syndrome of reduced CSF absorption and current analysis has concluded that histological features of the brain are normal. (9)

The presenting symptoms are headaches in 94% of patients, transient visual obscuration in 68%, pulse synchronous tinnitus in 58%, photopsia in 54% and retrobulbar pain in 44%. (10-14) Diplopia, which occurs in 38% of patients due to abducens nerve paresis, (15) and loss of vision, occurring in 30% of patients, (16) are less common. The major signs of IIH are papilledema and paresis of the abducens nerve. (16,17)

We are aware of just a few clinical reports of IIH and oculomotor nerve involvement in the pediatric population, (18,19) and two additional patients have been reported, but age was unknown at the onset (20,21) with sparing of pupillary fibers in all described cases. There has been no single report of complete oculomotor palsy to our knowledge.

**Figure 1.** Sagittal MRI of the brain with gadolinium shows a partially empty sella.

This case report raises several important issues. First, even a relatively late onset of headaches (after age 50) and signs of oculomotor nerve paresis must raise the suspicion of IIH if other etiologies have been previously excluded, such as an intracranial aneurysm or any other compressive lesion. Second, despite the “empty sella” abnormality on MRI, very often underreported by an interpreting radiologist, and if reported usually interpreted of no clinical significance, it might be, in a particular setting, very important information. (22,23) Third, even though there have been no previous reports of venlafaxine induced IIH, we still believe the underlying mechanism is weight gain; this possibility cannot be completely excluded. Fourth, prompt diagnosis of IIH can reverse a patient’s symptoms and signs.
REFERENCES