Pseudotumor Cerebri Presenting Unilateral Papilledema Associated with Iron-deficiency Anemia

Pseudotumor cerebral apresentando papiledema unilateral associado a anemia ferropriva

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Abstract
Pseudotumor cerebri is a relatively common pathology that is characterized by intracranial hypertension in the absence of mass lesions. It commonly affects young and obese women, and its presentation with visual loss and bilateral papilledema is well-described in the literature. We present a case of a 44-year-old, non-obese, female patient presenting with unilateral papilledema and iron-deficiency anemia. This case emphasizes this unusual presentation and the rare association with iron deficiency.

Keywords
► pseudotumor cerebri
► idiopathic intracranial hypertension
► unilateral papilledema
► iron-deficiency anemia

Palavras-chave
► pseudotumor cerebral
► hipertensão intracraniana idiopática
► papiledema unilateral
► anemia ferropriva

Introduction
Pseudotumor Cerebri (PTC), also known as idiopathic intracranial hypertension (IIH) or benign intracranial hypertension (BIH), is a medical condition characterized by increased intracranial pressure (ICP) without evidence of intracranial mass, hydrocephalus, infection or hypertensive encephalopathy. It usually affects overweight women of childbearing age, and its pathogenesis is uncertain.¹⁻⁴ Some studies indicate that this is an absorbing problem at the arachnoid level,⁴ but there is much controversy in the literature. The diagnosis of PTC is based on clinical symptoms and signs, normal findings in imaging studies, increased cerebrospinal fluid (CSF) opening pressure, and exclusion of other probable pathologies.⁵
Case Report

A 44-year-old female patient, who was previously healthy, underwent an abdominoplasty under epidural anesthesia. The surgery was uneventful, and the patient was discharged in good general condition 24 hours after the procedure. About 48 hours after discharge, the patient sought care with an ophthalmologist due to loss of vision in the right eye. In addition, she had a sudden headache of strong intensity and holocranial in nature, which was associated with an emetic episode, diplopia and dizziness. Papilledema was observed in the ophthalmologist’s office, and the patient was referred to our clinic.

Upon admission, the general physical examination revealed a heart rate of 120 bpm, and the carotid auscultation revealed bilateral bruits. The body mass index (BMI) was 28 kg/m². A neurological examination showed a reduction in visual acuity in the right eye, maintaining only the perception of movement. In addition, an edema was found in the ipsilateral papilla. The other aspects of the neurological examination did not suggest other abnormalities.

Due to the rapid onset of symptoms, the patient underwent magnetic resonance imaging (MRI) with and without contrast, which revealed only relatively small ventricles (Fig. 1). In addition, arterial and venous angio-MRIs were performed, and the results were considered normal (Fig. 2). After the radiological investigation, the patient was admitted for a clinical investigation.

The laboratory investigation revealed levels of serum hemoglobin of 5.2 g/dL. In addition, the hemogram showed microcytosis and hypochromia. The cardiologic evaluation revealed no abnormalities. A lumbar puncture was performed, and the opening pressure was of 34 cm H₂O. The diagnosis of BIH was made, and the treatment for anemia associated with blood transfusion and iron supplements was initiated. Due to severe vision loss, intravenous (IV) methylprednisolone 250 mg was prescribed every 6 hours associated with acetazolamide 1,000 mg twice a day.

The patient presented a dramatic response to the treatment, with complete improvement of the symptoms, except for the visual deficit, which remained unchanged. She was discharged 72 hours after admission, with no complaints, maintaining only the visual deficit. The ophthalmologic evaluation after discharge suggested a probably irreversible lesion in the right optic nerve.

Discussion

Pseudotumor cerebri is a disorder that affects overweight/obese women of childbearing age, and it has an annual incidence of 1–2/100,000 inhabitants among the general population. However, it is a disease that can affect men, children, the elderly, and patients who are not obese. When it affects younger people, gender and weight appear to have no significance in the incidence.

A considerable number of systemic conditions, drugs and vitamin deficiency have been associated with PTC. One peculiar condition mentioned in the literature is iron-deficiency anemia, whose prevalence in patients diagnosed with PTC is ~ 10%. However, the statistically significant association between PTC and the other conditions is unclear in many cases.

Typically, PTC is a self-limited but recurrent pathology, with a recurrence rate ranging from 9–43%. Severe visual deficits develop in 4–12% of the patients, although there is no relation to the duration of the symptoms, the degree of the papilledema, headache or the number of recurrences.

Due to the intrinsic mechanisms of the body, it is conclusive that the increase in intracranial pressure is originated in at least one of the following body components: CSF, blood, interstitial fluid or cerebral cells. Cerebrospinal fluid...
production is relatively constant and independent of its internal pressure, at least to some extent. Hence, the increase in pressure is not quickly normalized with the reduction in CSF production, facilitating the development of the condition, once the process is initiated.21–23 An increase in CSF production has been suggested in PTC,24 but there is no evidence to support such a theory, or any structural changes in the choroid plexus, such as hypertrophy. There is some evidence to suggest that CSF production is normal in PTC.25

Most of the focus on PTC pathogenesis studies has been on resistance to the CSF absorption mechanism. Several studies involving CSF infusion have been published, and all showed a greater resistance to CSF absorption in PTC.26–28 Gjerris et al found a lower CSF flow in 12 out of 14 patients with PTC, with 2 normal values measured after the beginning of the treatment.27 Janny et al demonstrated resistance to the flow in all 16 patients with PTC studied.28 Although most studies have focused on the intracranial CSF compartment, new imaging techniques using dynamic contrast phases suggest that there is an increase in extraventricular CSF volume and a decrease in jugular venous output in PTC compared with control individuals.29 Alterations in spinal canal compliance may be a contributory factor.30 It seems likely that the decrease in absorption is the core of the pathophysiology, possibly due to a local effect involving the epithelial membrane of the CSF output routes.

Magnetic resonance venography demonstrates that the majority of PTC patients have venous sinus stenosis.31 It is unclear if such stenosis is the cause or consequence of PTC, although cases of demonstrated resolution of collapsible venous sinus segments after CSF drainage and pressure normalization have suggested that the latter is true.32,33 Cerebral venous sinus narrowing usually develops at the transverse sinuses or transverse/sigmoid junction, and may be bilateral or unilateral.34

For the cases of venous sinus stenosis, the placement of a stent seems to be a good therapeutic option. In a study conducted with a large cohort composed of 52 patients who underwent unilateral cerebral venous sinus stenting, all patients had resolution of the papilledema; 23 out of 30 patients had resolution of the visual field loss, and headache only persisted in 3 patients.34

**Etiology – Association with Iron-deficiency Anemia**

Even though the condition is also known as idiopathic intracranial hypertension (IIH), the nomenclature of pseudotumor cerebri (PTC) syndrome is more inclusive, because it covers both the idiopathic cause and the secondary etiologies.

Giuseffi et al17 reported a prospective controlled study adjusted for age and sex that investigated the associations of diseases and drugs in PTC. They found that iron-deficiency anemia was not more common in patients with intracranial hypertension than in controls. This result was similar to the ones published by Ireland et al,16 who reported that iron-deficiency anemia was more frequently found in the control group. However, the association between the two conditions seems to be strong, since PTC symptoms resolve when the patient is treated for iron-deficiency anemia, as it happened in the case presented here.17,35

In a study by Mollan et al,35 8 patients with PTC and iron-deficiency anemia were reported, and 7 of them had remission of the symptoms with the treatment for anemia. The patients were treated with blood transfusion and oral iron replacement, depending on the severity of the symptoms.

**Clinical Features**

The most prevalent symptoms in PTC are headache (88%), transient visual obscurations (70%), pulsatile tinnitus (56%), nausea (32%), diplopia (28%), and visual loss (14%). The presentation can be acute with recurrences or chronic (26–32%).3,19

The characteristics of the headache can be variable. Unilateral or bilateral headaches that are pulsatile and with a retro-ocular location have been reported.36 Transient visual obscurations occur in two thirds of the patients with papilledema, and they may be unilateral or bilateral. The frequency ranges from rare episodes to multiple cases per day. However, the presence of this symptom does not seem to be correlated with the severity of the disease.17,37 It is believed that pulsatile tinnitus represents vascular pulsations transmitted by the CSF to the venous sinuses at high pressure.38 When it occurs along with headache, it is highly suggestive of PTC.17,39,40

The principal signs of PTC are papilledema, visual field loss, and abducens nerve palsy.

Papilledema is generally symmetrical and bilateral, and is one of the hallmark features of PTC.41–43 The presence of unilateral papilledema is rare, and the degree of visual field loss is greater in the eye with the higher grade of papilledema.42 This uncommon unilateral commitment was present in our patient, and it was associated with the permanent loss of the visual field.

Abducens nerve palsy may present unilaterally or bilaterally, and it reflects the effect of the elevated intracranial pressure on this nerve, which has the largest intracranial pathway.19,44

In the presence of signs of intracranial hypertension, neuroradiology should be performed to discard a secondary cause, such as tumor, intracranial hemorrhage or venous sinus thrombosis. In the absence of altered imaging, lumbar puncture and CSF analysis are indicated to discard other causes. It is also necessary to request a blood count to verify the presence of anemia, a condition that may be associated with PTC.45

In general, the patients present spontaneous resolution of the symptoms within a year. Therefore, the interventional treatment is commonly suggested whether the patients are symptomatic or did not have resolution of the set.46–48 However, recurrence may occur in those patients who received the medical treatment, as well as in those who did not need it. In obese patients, weight loss has been associated with complete resolution of the papilledema.49,50 Weight loss has shown such good results that bariatric surgery has been studied as an option of treatment for obese patients. Nevertheless, this improvement may be too slow for cases of highly-threatened vision; thus, in these cases, pharmacological treatment would be helpful.50 The interventional treatment is based on fluid and
salt restriction combined with diuretics. Acetazolamide, a carbonic anhydrase inhibitor, has been widely used to reduce the intracranial pressure with good results; therefore it seems to be a good drug to be administered as a first choice, even though no randomized clinical trial data have confirmed its effectiveness. A long-term follow-up study demonstrated that 60% of the patients had recurrent episodes during a period of 6.2 years; however, none of these episodes occurred during the time the patients were taking acetazolamide. Other drugs, such as furosemide, methazolamide and topiramate, have also been used to manage the symptoms, but they do not seem to be as effective as acetazolamide when used alone. Corticosteroids do not seem to be recommended as a routine choice of treatment, especially because of their relationship to weight gain, but in a critical situation, they may be useful to relieve the symptoms.

Additionally, as PTC appears to be related to different causes, it is plausible that the underlying cause is corrected. In our case, we prescribed acetazolamide and corticosteroid, since the patient presented a high risk of visual loss. We treated the patient for iron-deficiency anemia with ferrous sulfate. In the cases that cannot be managed with drugs, ventricule or lumbar-peritoneal shunts have been shown to be effective choices. However, some patients may develop chronic daily headaches, and this may be understood with drugs, ventricle or lumbar-peritoneal shunts have been used to manage the symptoms, but they do not seem to be effective as acetazolamide when used alone. Corticosteroids do not seem to be recommended as a routine choice of treatment, especially because of their relationship to weight gain, but in a critical situation, they may be useful to relieve the symptoms.

**Conclusion**

Our case highlights an uncommon presentation of PTC with unilateral papilledema and the still poorly-understood association between iron deficiency and BIH. As reported in the literature, our case presented a dramatic response to PTC symptoms with the treatment for iron-deficiency anemia. This response points to a possible association between these two conditions, but more studies should be conducted to elucidate this correlation.

**Conflicts of Interest**

The authors have no conflicts of interest to disclose.

**References**