Recurrent Pseudotumoral Demyelinating Disease in an Adolescent Patient

Osman Farooq1*, David Hojnacki1, Lucia Balos3 and Bianca Weinstock-Guttman2

1Women & Children Hospital of Buffalo, SUNY University of Buffalo, USA
2Jacobs MS Center, SUNY University at Buffalo, USA
3Department of Pathology, SUNY University at Buffalo, USA

*Corresponding author: Osman Farooq MD, Women & Children Hospital of Buffalo, Department of Neurology, 219 Bryant Street, Buffalo, NY 14222, USA, Tel: 716-878-7840; E-mail: ofarooq3@buffalo.edu

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Introduction

Multiple sclerosis (MS) is a chronic progressive inflammatory demyelinating and degenerative condition affecting the nervous system. Typical magnetic resonance imaging (MRI) findings consist of multiple asymmetric white matter lesions, with a predilection for periventricular and subcortical white matter [1]. The number and extent of lesion accumulation can be predictive of the disease course. As opposed to these multiple white matter lesions seen in MS, in the pseudotumoral inflammatory demyelinating form the onset of disease is characterized by a large isolated mass, typically in the white matter, usually in the brain hemispheres [2]. When these lesions are over 20 mm in diameter, they are defined as tumefactive, or pseudotumoral demyelinating lesions [1]. This form of demyelinating disease is rare, and even more so in the pediatric population. Here we present a case of an adolescent boy with pseudotumoral MS with recurrence several years later, while on no disease modifying therapy with multifocal pseudotumoral lesions. To our knowledge this is the first pediatric case of recurrent pseudotumoral disease occurring within several years.

Case Report

A previously healthy 12 year old boy presented with a persistent and worsening right frontal headache. The parents report a longstanding history of focal headaches, localized to the right frontal region, however in recent days these headaches had increased in frequency and severity. There was no reported photophobia, phonophobia, or visual disturbances reported. The general physical as well as neurological exams were within normal limits.

The MRI of the brain (Figures 1a and 1b) revealed a large mass with a central, well circumscribed cystic area. A ring-like enhancement pattern was seen as well as a mass effect. Spinal imaging did not reveal abnormalities or evidence of similar lesions.

Surgery was attempted to decompress the suspected tumor. However during the surgical procedure, no demarcation could be made between normal and abnormal tissue, therefore a biopsy of the area in question was performed. Fluid from the cystic decompression failed to reveal any evidence of tumor/malignancy. Cytology was negative. Pathological evaluation of the biopsy specimen revealed a markedly hypercellular with somewhat congested vasculature tissue. A perivascular lymphocyte component could not readily be identified. However, the CD68 immunohistochemistry for macrophages diffusely stained the majority of the cells in the lesion and identified clusters of macrophages in the Virchow-Robin spaces as well (Figures 2a and 2b). The differential diagnosis of granular cell astrocytoma and demyelinating process was considered and preference to demyelination given due the major component of the lesion comprising macrophages.

![Figure 1a: Axial FLAIR MRI, revealing a large hyperintense lesion in the right periventricular region extending to the internal capsule.](image1)

![Figure 1b: Axial T1 post-contrast, revealing with patchy contrast enhancement in an incomplete/open ring-enhancement pattern.](image2)
Figure 2a: H&E stained section of biopsy showing the marked hypercellularity of the tissue comprising clear and granular cells. Granular cells surround blood vessels in the Virchow-Robin space.

Figure 2b: CD68 immunohistochemistry demonstrates the diffuse infiltration of the tissue by macrophages. The Virchow-Robin space vascular cuffing is also pronounced.

A visual evoke response (VER) revealed a delayed P100 on the left. As this lesion was seen in isolation with histological evidence of inflammation and demyelination, he was diagnosed as having a pseudotumoral demyelinating process. He was treated with steroids, and the lesion decreased significantly in size. Clinically he returned to his baseline. Repeat MRIs and VERs were performed on a yearly basis and were stable.

Five years later, he experienced a sudden worsening of symptoms, at which time he experienced diffuse right sided hemiplegia as well as a right temporal visual field defect. This occurred three weeks after treatment with amoxicillin for a pharyngeal infection. An MRI (Figure 3a-3f) revealed multiple large white matter lesions with surrounding edema in the left frontal and temporal regions as well as in the right fronto-temporal region. There was evidence of contrast enhancement. Cerebrospinal fluid (CSF) analysis was normal, without oligoclonal bands or NMO IgG antibodies (serum and CSF). His symptoms showed only modest improvement with steroids. He was then treated with plasma exchange followed by intravenous immunoglobulin (IVIG), which aided in the speed of recovery. Because the recurrence and severity of the second demyelinating event, the patient was initiated on monthly IVIG 0.4 gr/kg for 6 months with the consideration of close monitoring. Repeat MRIs improved significantly with no new lesions within a follow up of almost 1 year.

Figure 3a and 3b: Axial FLAIR (3a) and corresponding T1 post-contrast (3b) revealing encephalomalacia in the area of the previous lesion in right periventricular region.

Figure 3c and 3d: Axial FLAIR (3a) revealing a large lesion in the left frontal region with surrounding edema as well as a small lesion in the right parietal region. The corresponding T1 post-contrast (3b) revealing an incomplete/open ring-enhancement pattern in the left frontal lobe as well as enhancement in the small lesion in the right parietal region.
Cognitive functions. In addition, (apraxia, agnosia, aphasia) can occur when the frontal regions are involved and acute memory dysfunction in case of involvement of the temporal regions. Hemiparesis, of variable severity, and of sensorial and visual disorders, are also frequent. Another typical feature is the appearance of seizures when there is a cortical involvement [2,14]. Headaches of varying degrees can also be a common presenting symptoms, as in our patient.

The differential diagnosis can be quite problematic, particularly in the initial stages of the disease. The main entities to keep in mind would be tumor and ADEM, and to a lesser extent progressive multifoccal leukoencephalopathy (PML), abscess or parasitic cysts. The tumor, which can be easily mistaken for a demyelinating lesion, is lymphoma. The brain lymphoma is usually unifocal, but in some cases can be multicentric. ADEM usually appears with small and bilateral lesions, which are generally symmetric, but in rare cases large isolated lesions can be observed [4]. Abscesses and parasitic cysts can have aspects similar to inflammatory pseudotumoral lesions, but the characteristics of enhancement are different as the ring around the cystic cavity is usually complete, whereas in the demyelinating lesion, it is mostly “incomplete” [2].

The long-term evolution of these inflammatory lesions is not well defined. It is estimated that about one third of the patients have a monophasic course with complete recovery, one third have a monophasic course with prolonged evolution and residual deficits, and the remaining third evolves to MS [1].

The relative paucity of reported cases of tumefactive demyelination in children likely represents an underestimate of the actual frequency of cases. In 2002, McAdam et al., [15] reviewed 12 cases in the literature, plus 4 cases observed personally. Morin et al., [16] described 3 additional cases. Puri and associates [17] described a case of recurrent pseudotumoral MS, with two episodes occurring 6 months apart. Our case is unique due to the prolonged interval in between recurrence (5 years). In addition, upon recurrence the symptomatology and clinical severity was considerably worse, requiring plasma exchange as steroids alone were not sufficient. Typically with pseudotumoral MS solitary lesions are found, whereas in our patient there were multiple lesions at recurrence. This raised the concern for the need of a preventative medication, however following six monthly IVIG cycles the patient has shown no evidence of clinical or radiological recurrence of disease for 1 year.

Summary

Pseudotumoral MS can be a diagnostic challenge. The diagnosis in itself can be rare, however our case is unique because there was clinical recurrence after 5 years while on no therapy and because of the early age of presentation. Moreover, upon recurrence our patient exhibited multiple lesions with pseudotumoral features, which is unusual for pseudotumoral MS. Initiation of a preventive disease modifying therapy (DMT) was deferred because the atypical presentation and the 5 year initial remission.

References


