Clinical and imaging features of intracranial demyelinating pseudotumor: report of one case and literature review.

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Abstract

Demyelinating pseudotumor mainly attacks in teenagers and more commonly seen in female compared with male counterparts. Comprehensive understanding of medical history, clinical manifestations, imaging and pathological characteristics contribute to clinical diagnosis and management of demyelinating pseudotumor. Demyelinating pseudotumor in the central nervous system is mainly characterized as nerve fiber demyelination and inflammatory cell infiltration surrounding the minute vessels. However, the clinical and imaging characteristics of demyelinating pseudotumors mimic those of brain tumors, especially the brain glioma. It is likely to make a misdiagnosis of demyelinating pseudotumors. Herein, we reported one case of intracranial demyelinating pseudotumors and described the diagnosis and treatment of this patient in this study.

Keywords: Demyelinating pseudotumor, Misdiagnosis, Imaging characteristics, Clinical symptoms.

Introduction

At present, the pathogenesis underlying demyelinating pseudotumor remains elusive, probably affected by hereditary factor, virus infection or medication use etc., [1-3]. Demyelinating pseudotumor mainly attacks in teenagers, more commonly seen in female compared with male counterparts [4]. The tumor size is >2 cm in diameter. No evident symptoms occur upon the onset. It is mainly characterized with acute or sub-acute episode. In this report, we have reported a male patient, aged 56 y, presented with sub-acute onset due to left limb weakness.

Case Presentation

Baseline data

The male patient, aged 56 y, was admitted to hospital due to left limb weakness for 2 weeks. Physical examination upon admission revealed that body temperature was 36.6°C and blood pressure was 140/80 mmHg. The patient remained mentally conscious and could communicate normally. The Glasgow Coma Scale score was assessed as 15, suggesting that the patient suffered from minor brain injury. Bilateral pupil size was measured as 3.0 mm in diameter with sensitive direct and indirect light reflex. His left limb muscle strength was assessed as grade IV and grade V for the right limb muscle strength. The limb muscle tension was evaluated as normal. No edema was documented in bilateral lower limbs. Physiological reflex was normal and pathological reflex was not noted.

Twenty years ago, he underwent bilateral fronto-temporal decompressive craniectomy due to craniocerebral trauma. Written informed consents were obtained from the family members of the patient.

Imaging characteristics

Head CT of the outpatient: Multiple hypointensity lesions and suspected isointensity nodes were noted within the brain, which were considered as space-occupying masses. Following decompressive craniectomy, the signs of multiple encephalomalacias were observed (Figure 1). Plain head MRI and enhanced MRS upon admission: multiple masses of slightly long T1 and long T2 signals were noted in the right frontal lobe, left frontal lobe proximal to corpus callosum and right temporoparietal lobe. Hypointense signals were observed on Diffusion Weighted Imaging (DWI). No contrast-enhanced signals were noted on the enhanced MRI, indicating no incidence of cerebral ischemia. Nodular long T1 and T2-signal was noted in the right centrum semiovale and brain stem. Hypointense signals were observed on Diffusion Weighted Imaging (DWI). No contrast-enhanced signals were noted on the enhanced MRI, indicating no incidence of cerebral ischemia. Nodular long T1 and T2-signal was noted in the right centrum semiovale and brain stem. T2 flair hypointense signal was detected on the MRI, as illustrated in Figure 2. Single and multi-voxel 1H-Magnetic Resonance Spectroscopy (MRS) were carried out (TE=135 ms). The right temporal lobe, focal and peripheral lesions in the basal ganglia region were located as the region of interest and statistically compared with the imaging findings of the contralateral normal region, hinting decrease in NAA peak, increase in Cho peak, decrease in Cr peak, decrease in NAA/Cr ratio and increase in Cho/Cr and Cho/NAA ratio in the right lesions. Inverted lactate peak was detected. In the center of the lesions, decrease in...
NAA, Cho and Cr peak was observed (Figure 3). Inverted lactate peak was found. No other abnormality was documented.

**Figure 1.** CT scan of the head.

**Figure 2.** MRI of the head.

**Laboratory test**

**Conventional cerebrospinal fluid and biochemical examination:** A relatively high level of protein was detected in the cerebrospinal fluid. Blood routine examination found a slightly high level of leukocyte. The examination outcomes indicated the possibility of intracranial demyelinating lesions. The final diagnosis was confirmed by subsequent biopsy.

**Surgical biopsy**

Percutaneous brain tissue biopsy was performed via the temporo-occipital junction behind the ear. The blood loss was 100 ml during surgery. Intraoperatively, yellow colloid-type lesions with soft texture were noted in the subcortical brain tissue (Figure 4).

The deep layer of the lesions was excised and no explicit margin was observed, significantly differing from the brain tissue from healthy counterparts. After tissue sampling, the incisional wounds were sutured.

Immunohistochemical staining revealed positive for CD68, GFAP, Vim, S-100, MBP and Olig-2 and negative for CK, EMA, NSE, NF, Syn and CD34 (Figure 5).

**Discussion**

In this report, the male patient, aged 56 y, presented with sub-acute onset due to left limb weakness. He reported no family history, virus infection or use of chemotherapy agents.

Therefore, the exact pathogenesis remained unclear. Demyelinating disease, dominantly involved with the frontoparietal lobe, is mainly characterized with dysfunction and refractory high intracranial pressure, leading to a high risk of relevant symptoms, such as limb and trunk dyskinesia, epilepsy, headache, nausea, vomiting and optic disk edema, etc. These symptoms can be aggravated due to the edema severity.

It presents as a solitary contrast-enhanced mass and poses a diagnostic challenge for both radiologists and surgical pathologists to distinguish demyelinating tumors from astrocytic gliomas.

Initially, CT plain scan hinted that the possibility of metastatic tumor was suspected due to multiple lesions with hypo-intensity accompanied by severe edema. MRI findings demonstrated multiple abnormal signals in the white matter of bilateral cerebral hemispheres. The pathological changes presented with long T1 and T2 signals.

Since the blood-brain barrier was destroyed, significant transient enhancement was noted within the lesions and decreased along with the mitigation of inflammatory response.

Masdeu et al. have first proposed that open-ring sign is often presenting large, contrast-enhancing demyelinating lesions and contributes to differentiate demyelinating pseudotumors from neoplasms and infections [5].

In this report, we also identified open-ring signs in the contrast-enhancing demyelinating lesions with unclear margins, which were surrounded by edema signs with hypointense signals.

White matter of the brain is abundant with myelinated nerve fiber, whereas no myelinated nerve fiber is present in the brain cortex and gray matter, probably leading to a higher risk of demyelinating pseudotumors occurring in the white matter of brain.
The patient in this report had sub-acute onset. He was pathologically characterized with myelin sheath injury, whereas the nerve axon was intact. A high quantity of lymphocyte infiltration in a cuffing pattern was observed surrounding the vessels under light microscopy. However, a large amount of monocyte and froth-shaped macrophage infiltration was noted in the myelinolysis region, accompanied by proliferative spongiosity. Lymphocyte and macrophage infiltration is a common manifestation of demyelinating pseudotumor, which is consistent with the patient in this report. Demyelinating pseudotumor is highly sensitive towards the adrenal cortex hormonotherapy and yields relatively favourable prognosis [6-8]. Plasma exchange is recommended for patients who are untreated with glucocorticoid administration. To prevent the progression into multiple sclerosis demyelinating pseudotumor, interferon administration is highly considered to alleviate the injury, interfere the progressive development and improve clinical prognosis [9,10]. In this report, the patient was administered with methylprednisolone after surgery and clinical symptoms were significantly mitigated. Nevertheless, the long-term clinical efficacy remains to be elucidated by subsequent follow-up.

Comprehensive understanding of medical history, clinical manifestations, imaging and pathological characteristics contribute to clinical diagnosis and management of demyelinating pseudotumor. For juvenile patients with sudden onset, accompanied by a history of multiple sclerosis, vaccine inoculation or virus infection and lesions with open-ring signs vertical lateral ventricle, the possibility of demyelinating pseudotumor should be highly considered. Prior to surgery, hormone treatment or brain tissue biopsy can be performed initially to avert the serious damages and complications to the nerve function induced by excessive surgery and/or radiotherapy, etc.

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Conflicts of Interest
None.

References


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