Devic’s Disease: A MRI Finding Case

E. V. Acko-Ohui1*, A. Setchéou2, I. Garba2, U. V. Acko3, A. Konan2, P. Yapo4

1Department of Radiodiagnosis and Medical Imaging, University Hospital (UH) of Treichville, Abidjan, Côte d’Ivoire
2Department of Radiodiagnosis and Medical Imaging, University Hospital (UH) of Yopougon, Abidjan, Côte d’Ivoire
3Department of Internal Medicine and Geriatrics, University Hospital (UH) of Treichville, Abidjan, Côte d’Ivoire

Email: *ohuestelle@yahoo.fr

Abstract

Introduction: Devic’s neuromyelitis optica is an autoimmune and central nervous system demyelinating disease. It mainly affects the spinal chord and optic nerves. Considered for a long time as a special form of Multiple Sclerosis (MS), it is distinguished clinically by its rarity and poor prognostic, biologically by NMO (IgG) antibody, radiologically by extended myelitis. Treatment include mega dose of methylprednisolone during relapses and immunosuppressive drug in prevention of recurrences.

Observation: Devic’s disease manifests itself in outbreaks of neuritis optic or myelitis. We report the case of a patient who had no particular antecedents. She presented a spastic tetraparesis, accompanied by a decreasing left visual acuity (4/10 th); the all evolving by thrust since one year. Spinal chord lesions on MRI, second stage brain damages appearance affecting the white substance and the discovery in the patient serum of anti-aquaporin 4 antibodies (anti-AQP4 Ab) allowed the diagnosis of Devic’s disease.

Conclusion: Diagnosis of Devic’s neuromyelitis optica that was initially based on clinical examination and radiological assessment is currently confirmed by the discovery in the patient serum of the NMO immunoglobulin G directed against Aquaporin 4.

Keywords

Optic Neuritis, Spinal Chord, Transverse Myelitis, Aquaporin 4, Devic’s Disease, NMO, MRI

1. Introduction

The neuromyelitis optica (NMO) has been described for the first time by Eugène Devic [1] [2] [3]. Devic’s disease is an inflammatory and demyelinating pathology of the central nervous system. It mainly affects the spinal chord and the optic nerves. Long regarded as a special form of sclerosis (MS), it is different. MS is
a cell-mediated autoimmune disease in which the Lymphocytic infiltrate is predominant [4]. Devic’s disease is an autoimmune disease with humoral mediation. It implies the presence of an antibody directed against the Aquaporin 4. This antibody, recognized under the name of Aquaporin 4 antibody (anti-AQP4 Ab) [4] [5], was originally called immunoglobulin G of the NMO (NMO - IgG). Devic’s disease manifests itself in outbreaks of optic neuritis or myelitis. The mode of entry into the disease is, in more than two thirds of the cases, an optical neuritis. Optic neuritis may be bilateral but is more often unilateral [6]. It is generally of acute installation; the decrease in visual acuity is profound, and the prognosis is severe. Myelopathy during Devic’s disease is also most often acute and associated with back pain [7]. It usually leads to severe disability. Repetition of flare-ups can lead to permanent disability. We report the case of 35 years old patient in whom the MRI allowed to evoke the diagnosis of Devic’s disease.

2. Observation

Miss TY, 35 years old, was admitted to a neurology department in our town. She had a spastic paraparesis associated with a decrease of the left eye visual acuity. She had no particular antecedents. The symptomatology had evolved for about a week. Three days after her hospitalization, left eye blindness was installed, associated with urinary retention. The thoracic spine Magnetic Resonance Imaging (MRI) involved highlighted an extended intra medullar signal anomaly from T9 to T12. This anomaly appeared in hypo intensity on EST1 sequences, in hyperintensity on EST2 sequences [Figure 1] and in hyper intensity on STIR sequences. This hypersignal also extended at the level of the dorsal spine from T2

Figure 1. From T9 to T12. Intra medullar hyperintensity on EST2 and FLAIR sequences.
to T7 on EST2 [Figure 2]. These lesions were not affected by the gadolinium injection. Encephalic MRI was normal. The analysis of the cerebro-spinal fluid (CSF) highlighted a rise in its protein content and a rise in its content in cells which reached more than 60 leukocytes/mm3. All of these data was in favour of active transverse myelitis. A corticosteroids intravenous bolus has been initiated at the dose of 1g of methyl prednisolone a day for three days. The relay was taken orally with prednisone, 1 mg per kilogram and per day for a month. This treatment was followed by an incomplete recovery of motor function and left eye visual acuity.

Two months later, she introduced a new thrust. This time, there was a bilateral reduction in visual acuity and the reappearance of sphincter disorders. She was again hospitalized and under corticosteroid therapy at the same doses as above. A week later, her general condition had improved. Six months after this last episode, she introduced a third thrust marked by the worsening of the unrest. It was about quadriplegia, binocular blindness and sphincter disorders. The MRI of cervical spine, thoracic spine, skull and brain were conducted. At neck and chest, the lesions described above were increased, revealing themselves in the form of an extension of the intra spinal chord hyper intensities from C2 to T10 [Figure 3]. Myelitis involved less than 50% of the transverse diameter of the cervical chord [Figure 4]. The brain, in this third thrust, had signal abnormalities. Were observed bilateral protuberantial and sub-ependymal demyelinization lesions [Figure 5]. The discovery of the anti-AQP4 antibody in the patient’s serum helped bring certainty diagnosis of Devic’s neuromyelitis optica [Figure 6]. The same treatment was renewed, but it has been spread over a longer period of time (one week of high-dose treatment and three months of maintenance treatment).

![Figure 2](image_url). MRI sagittal EST2 of the dorsal spine. Intra medular hyperintensity on EST2 from T2 to T7.
3. Discussion

NMO is a central nervous system demyelinating disease. It preferentially affects the spinal chord and optic nerves [2] [5]. The NMO is also known under the name of Devic’s disease, by the name of Eugène Devic who was the first one who identified the symptoms in 1894 [3] [6]. Devic’s disease is a rare pathology and epidemiological data are lacking. We can however estimate that it represents less than 1% of central nervous system inflammatory pathologies [7] [8]. Authors describe a female predominance [5] [6] [9] with a peak incidence at the end of the third decade. Our study concerns a woman of 35 years old. This disease affects Caucasians but also West Indians [10] and Japanese [10]. In one-third of relapsed forms, a personal history of autoimmune diseases is found [9] [10].
Some authors state that these associations justify the non inclusion of these cases in the nosological framework of Devic’s syndrome [11].

This entity is not a hereditary disease, but there is a reported case of 2 homozygous sisters [10] [11].

The viral and bacterial infections preceding or accompanying the NMO are a known phenomenon. The influenza-like illness that precedes the onset of neurological disease has been reported in about 25% - 30% of cases [12]. Varicella zoster virus (VZV) and tuberculosis have been the most common infectious triggers; however, other bacterial and viral agents such as chlamydia, cytomegalovirus, HIV and Epstein Barr virus (EBV) have been reported [12].

Clinically, Devic’s disease is characterized by acute episodes of optic neuritis and myelitis. Optic neuritis is a retrobulbar-like optic neuritis, unilateral or bilateral, often more severe than that described in MS. Cases of oculomotor abnormalities, Claude Bernard Horner’s syndrome and nystagmus have been reported by some authors [13].

The myelitis is transverse, acute and is clinically expressed by the association of motor, sphincter and sensitive disorders. Its onset is on average of 2 years [14]. Classically the installation of symptoms is acute reaching acme in a few
hours or in a few days. The topography is readily cervico-dorsal. The table can be dramatic in the type of flaccid tetraplegia associated with respiratory distress requiring ventilatory assistance. Medullary sprays may also adopt a more limited topography, such as Brown-Sequard syndromes [14] [15], single-paresis, or even isolated sensory disorders. In the state phase, paralysis in a pleural mode of at least one member and its flaccid character would be frequent.

Spinal chord MRI is characteristic. This imaging technic highlights an intra medullar hyper intensity on the EST2 and STIR sequences. Signal abnormalities extend to more than 3 metameres in the sagittal plan [11]. Cerebral MRI is normal outside the extended hypersignal of the two optic nerves dating back to the chiasma. However, most authors admit the presence of brain white hypersignals on the T2-weighted sequences especially correlated with the duration of evolution, without calling into question the diagnosis of NMO [16].

During the third thrust, the neck and thoracic spine MRI performed in our patient have highlighted the hyper intensity lesions on T2 and STIR sequences that stretched to 5 metameres for the cervical spine and 10 metameres for the thoracic spine. Brain MRI was normal at the beginning.

At the last thrust, it demonstrated bilateral protuberantial and sub-ependymal demyelinization lesions.

The analysis of CSF and blood makes it possible to differentiate on the one hand between the infectious and systemic inflammatory etiologies and on the other hand between NMO and MS [1] [4] [12] [13]. In the NMO, levels of CSF in cells (greater than 50 leukocytes/mm3) and in proteins are high [2] [7] [9]. These were observations we made in our study. The NMO is secondary to the production of an antibody directed against the Aquaporin 4, abbreviated AQP4 (anti-AQP4 antibody). Aquaporins are membrane proteins that provide exclusively the passage of water from the extracellular sector to the intracellular environment. The AQP4 differs from other aquaporins by its tropism for the central nervous system and the retina [3] [10] [14]. In our study, the research of anti-AQP4 antibody came back positive.

The current criteria of Devic neuromyelitis are based on the combination of mandatory criteria (optic neuritis, acute myelitis and absence of signs in favor of an attack other than that of the optic nerve and marrow) to a secondary criterion (Normal MRI, extensive medullary lesion on more than 3 vertebrae, plethocytosis of the cerebrospinal fluid) or two minor secondary endpoints (bilateral optic neuropathy, severe optic neuropathy with 1/10th visual acuity and severe muscle deficiency At least one member) [17] [18].

However, these criteria do not allow the diagnosis of Devic neuromyelitis subjects with neurological symptoms in regions of the central nervous system other than the optic nerve or spinal cord, and those with cerebral MRI showing lesions identical to Those of multiple sclerosis [19] [20].

Thrust treatment starts classically with daily infusions of 500 mg to 1000 mg of methyl prednisolone for five to seven days. Secondarily the relay is orally tak-
en by the corticosteroid at the dose of 1 mg/kg/d [5] [7] [15]. Attacks of myelitis may not stop under this treatment and may even worsen. In this case, plasma transfusions must be considered [15]. One day out of two, 5 to 7 sessions of plasmapheresis are performed [10] [12] [14]. Failing to plasmapheresis, intravenous immunoglobulin infusions are often considered as an alternative. Most often used immunosuppressive therapy is an oral combination of azathioprine – prednisone [13] [15]. In our study, thrust gave way under corticosteroid therapy. We didn’t need to resort to plasmapheresis or immunosuppressive therapy.

4. Conclusion

Devic’s disease is an extremely rare pathology. It is a rare clinical syndrome involving a unilateral or bilateral optic neuritis with a transverse myelitis. Devic’s neuromyelitis optica Clinically similar to multiple sclerosis, albeit with a more rapid and more severe, it is a distinct clinical and pathological entity. The diagnosis of NMO should be evoked facing clinical symptoms of increased aggravation of spinal and ocular neuritis. On MRI, lesions are localized first to the spinal cord and then to the brain. The diagnosis of certainty is made by the discovery of the anti-aquaporin4 antibody in the patient’s serum.

Conflict of Interest

None.

References


