Linear Medullary Pericanal Lesion with Longitudinal Myelitis can be Diagnostic of Neuromyelitis Optica

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A 51-year-old woman was admitted to our hospital because of progressive numbness of the trunk and extremities with gait unsteadiness. On admission, she showed an intractable hiccup. Her gait was slightly spastic with hyperactive
deep tendon reflexes in all extremities. She complained of decreased sensation below the C5 spinal level. Although sicca symptoms were not observed, the results of laboratory data and a lip biopsy fulfilled the criteria of Sjögren’s syndrome. The serum anti-aquaporin-4 (AQP4) antibody titre (1) was 256×. A lumbar puncture revealed pleocytosis of 368 cells/μL (357 mononuclear cells) with an increased protein level of 90 mg/dL. T2-weighted MRI showed a high-intensity long cord lesion extending from the C3 to the T6 level and a linear pericranial lesion in the medulla oblongata (Picture 1). These lesions were gadolinium enhanced. She was diagnosed as having neuromyelitis optica (NMO) spectrum disease (2) associated with Sjögren’s syndrome and was successfully treated with IV methylprednisolone with subsequent oral prednisolone.

Misu and colleagues (3) reported six patients with recurrent NMO, who exhibited intractable hiccup with linear medullary lesions as seen in our patient. The present case further suggests that NMO spectrum disorder may be diagnosed in patients having longitudinal myelitis and characteristic medullary lesions, even without optic neuritis or without recurrent episodes. Because NMO requires long-term treatment with immunosuppressants, correct diagnosis should be made by measuring the anti-AQP4 antibody/NMO-IgG in such patients. Another interesting finding was that our patient had Sjögren’s syndrome. Since longitudinal myelitis has been reported in Sjögren’s syndrome (4), we speculate that some patients previously reported as having Sjögren myelitis may also have had NMO spectrum disorder. We emphasize that awareness of these MRI lesions will lead to the successful management of this occasionally devastating disorder.

References