Myelitis is one of the rare neurological complications of mumps infection. However, there have been few reports of multifocal longitudinally extensive transverse myelitis (LETM) as a complication during the convalescent stage after acute mumps infections. A 14-year-old boy abruptly developed paraplegia, paresthesia of both legs, and urinary retention 4 weeks after acute mumps infection, from which he had recovered with supportive treatment. Spinal magnetic resonance imaging revealed two separate LETM lesions over the thoracic spinal cord. Other radiological and electrophysiological studies revealed no evidence of demyelinating disorders. The results of laboratory studies were all negative except for evidence of recent mumps infection. He was treated with a high-dose intravenous steroid, and recovered from his neurological symptoms about 1 month later. This is the first report of multifocal LETM after acute mumps infection. LETM is thus not only associated with neuroimmunological syndromes, such as neuromyelitis optica, but it can also develop after acute viral infection.
paraparesis during manual motor power examination. Hip flexion/extension and knee flexion/extension were reduced to Medical Research Council (MRC) grade 2/2, and ankle dorsiflexion, planter flexion, and foot inversion/eversion were reduced to MRC grade 1/1. The deep-tendon reflexes on the knee and ankle were absent, and there was hyperesthesia and allodynia below the T6 level. The patient had urinary retention and anal reflex tone was diminished. He was thought to suffer from spinal shock due to an acute cord lesion at around T6, for which emergent spine magnetic resonance imaging (MRI) was performed.

A T2-weighted thoracic spine MRI (Fig. 1) revealed hyperintensity lesion in T2-5 and T9-11 spinal cord with moderate swelling of the cord at these levels. Laboratory studies including complete blood count, and renal and liver function tests were normal. Cerebrospinal fluid (CSF) analysis revealed pleocytosis (a white blood cell count of 130/mm³) with elevated albumin concentration (38 mg/dL; normal range 10-30), but otherwise was normal. Anti-mumps IgG and IgM tests in serum were positive (Platelia Mumps IgM, immunoenzymatic capture method and Platelia Mumps IgG enzyme-linked immunosorbent assay [ELISA]; Bio-Rad, Marnes-la-Coquette, France). Polymerase chain reaction (PCR) for mumps virus, Epstein-Barr virus (EBV), herpes simplex virus (HSV) 1, HSV2, varicella zoster virus (VZV), and enterovirus in CSF yielded negative results. Serum anti-aquaporin-4 (anti-AQP-4) antibody test was negative. CSF oligoclonal band analysed with concurrent serum sample and the result were negative. Visual evoked potential test showed no conduction abnormalities. In addition, brain MRI showed neither a pathological lesion nor an abnormally enhancing pattern. In total, other relapsing demyelinating disorders, such as NMO or multiple sclerosis were excluded. To exclude other autoimmune diseases, we evaluated autoimmune antibodies in serum. Rheumatoid factor, c-antineutrophil cytoplasmic antibodies (ANCA), p-ANCA, anti-ds DNA, and paraneoplastic antibodies including anti-Hu, anti-Yo, and anti-Ri were negative results. Although fluorescent antinuclear antibody (FANA) was positive (titer 1:140, speckled pattern), it did not have clinical significance due to negative anti-extractable nuclear antigen (ENA) panel.

The patient was finally diagnosed as having post- or parainfectious LETM following mumps infection and he was treated with intravenous methylprednisolone (1 g/day for 5 days). A comprehensive rehabilitation program was also prescribed. During the course of treatment the patient exhibited marked improvement of weakness by the 5th day of admission, and on the 7th day began to walk using walking aid. His urinary retention also improved gradually. He had recovered fully from the motor weakness and urinary symptoms 6 weeks after symptom onset. In next 6 months, he showed no suspicious symptoms of relapse.

**DISCUSSION**

Most of the previous reports on the neurological complications after mumps infection have described disseminated syndromes over the entire central nervous system, such as meningoencephalitis or encephalomyelitis. The present case differs from these reports in that our patient presented with only paraparesis and urinary difficulty due to acute LETM. In addition, his recovery from the neurological impairment was considerably faster than has been reported previously.

LETM can be associated with several conditions, of which NMO is the most common, followed by spinal cord infarction and para- or postinfectious myelopathy. NMO is a relapsing-remitting inflammatory disease...
that preferentially involves the optic nerves and spinal cord. Anti-AQP-4 antibodies are highly specific for NMO and its spectrum disorders, and are considered to play a central role in the immunopathogenesis of the disorder. Therefore, both NMO and NMO-spectrum disease can be diagnosed using commercially available antibodies to AQP-4.1,6

Interestingly, laboratory evidence of previous viral infection has been reported in some cases of NMO (e.g., mumps, measles, rubella, human cytomegalovirus, VZV, EBV, hepatitis A virus, human immunodeficiency virus, and dengue virus).7 In this context, some authors claimed that these infectious pathogens are either directly involved or activate a major immunological step in the pathogenesis of NMO. This suggestion raises the need to evaluate NMO or NMO-spectrum disease in the setting of para- or postinfectious myelopathy.

However, AQP-4 antibody testing in cases of mumps-related myelitis has not been reported previously. Whether or not NMO or NMO-spectrum disease can initially present with para- or postinfectious myelopathy has yet to be established.

The patient described herein presented with both serological evidence of mumps infection and previous typical symptoms of mumps infection, and the observed neurological involvement was not disseminated like in previous reports. Therefore, although this patient exhibited only LETM without optic neuritis or brain lesions, the possibility of NMO or NMO-spectrum disease as a cause of the LETM was suspected. However, both CSF analysis with RNA-PCR for mumps and serum testing for antibodies to AQP-4 yielded negative findings. This case does not meet the “2017 McDonald criteria”8 or the “2015 International consensus diagnostic criteria for neuromyelitis optica spectrum disorders”.9 However, we cannot fully exclude the first attack of other demyelinating disease, such as a ‘clinically isolated syndrome (CIS)’. Of further relevance, it is more likely to be post-infectious myelitis due to myelitis occurring after a certain period of mumps infection. Although the sensitivity of reverse transcriptase polymerase chain reaction (RT-PCR) was approximately 70%, the time of sampling is critical for the diagnosis of mumps virus infection in CSF.10 Therefore, we concluded that this patient’s diagnosis was postinfectious LETM as a rare complication of mumps. Despite its rare occurrence, the findings of this case report suggest that acute mumps infection can be complicated by postinfectious LETM, mimicking NMO or NMO-spectrum disease. Clinicians need to be familiar with this rare complication and its differential diagnosis.

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Conflicts of Interest
All authors have no conflicts of interest.

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