SEZ6L2 Antibody–Associated Cerebellar Ataxia Responsive to Sequential Immunotherapy

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Abstract

Objectives
Seizure-related 6 homolog like 2 (SEZ6L2) antibody–associated ataxia is an extremely rare disease. Six patients have been reported and none of them improved significantly with immunotherapy. Herein, we present the case of a patient with cerebellar ataxia and SEZ6L2 antibodies who benefited from immunotherapy, which dramatically altered the course of her disease.

Methods
We present a case report of a 73-year-old woman with progressive balance problems. Her condition had rapidly deteriorated in the 2 weeks before the admission to our hospital leading to repeated falls and eventually left her bed-ridden.

Results
She presented with severe trunk ataxia, bidirectional nystagmus, dysarthric speech, and persistent nausea. With the exception of cerebellar atrophy, extensive imaging studies revealed no pathology. SEZ6L2 antibodies were found in both CSF and serum. Over a period of 9 months, our patient received immunotherapy consisting of steroid pulse therapy, IV immunoglobulin infusions, rituximab, and cyclophosphamide. Consequently, her condition improved markedly, and she was discharged home from the neurologic rehabilitation unit.

Discussion
Our case report shows that intense sequential immunotherapy may considerably improve level of functioning in some patients with SEZ6L2 antibody–associated cerebellar ataxia.

Classification of Evidence
This provides Class IV evidence. It is a single observational study without controls.
Neuroimmunologic advances have recently led to the identification of autoantibodies in patients with progressive cerebellar ataxia. Among these are antibodies against seizure-related 6 homolog like 2 (SEZ6L2), a type 1 transmembrane protein. The symptoms associated with these immunoglobulin (Ig) G4 subclass antibodies also include dysphagia, cognitive impairment, mild extrapyramidal symptoms, retinopathy, apraxia, and spastic paraparesis.\(^1\) Six patients\(^1,2\) with SEZ6L2 autoimmunity have been reported, and they have all had little or no treatment response. Here, we report a patient who markedly improved following sequential immunotherapy.

A 73-year-old woman was admitted to our hospital. She had been using escitalopram for depression for 19 years. Otherwise, her medical history was unremarkable, and she had no previous functional impairment. Her balance problems began 6 months before the admission leading eventually to several falls. A persistent nausea had also developed. Nevertheless, she had been able to walk unaided even outdoors until her condition began to deteriorate more rapidly a couple of weeks before the admission. Eventually, she was found lying in bed, with severe back pain. She said that she had fallen a couple of days earlier and had to crawl to bed because of severe postural instability.

On admission, she had persistent bilateral gaze-evoked nystagmus, dysarthric speech, and severe truncal ataxia. There was dysmetria in the finger chase test on the right. She was unable to stand or even sit without support. Mild bradykinesia and hypophonia were also observed (Videos 1 and 2). The disease course along with relevant clinical assessments and therapies is presented in the Figure.

Brain and spinal cord MRIs were unremarkable with the exception of cortical cerebellar atrophy (eFigures 1 and 2, links.lww.com/NXI/A680). Whole-body CT and PET-CT investigations revealed no pathology. Lumbosacral radiography showed a compression fracture at L1. Vitamin B1, B6, and B12 levels were within the normal range. Antibodies against tissue transglutaminase, glutamic acid decarboxylase, HIV, and Treponema pallidum were negative. A cell-based indirect immunofluorescence assay conducted at Stöcker Clinical Immunological Laboratory revealed SEZ6L2 antineuronal antibodies in both CSF (titer 1:3.2) and serum (1:100).

Over a period of 9 months, the patient received sequential immunotherapy. Steroid pulse therapy and 2 IVIg infusions resulted in slight improvement, but despite markedly improved scores on clinical scales, her overall condition remained poor with persistent nausea and poor mobility even after rituximab (RTX). Consistent improvement commenced following cyclophosphamide (CTX) initiation with steady recovery thereafter (Videos 3 and 4). Repeat brain MRI was unchanged (eFigures 3 and 4, links.lww.com/NXI/A680).

**Figure** June and July 2021 Were Not Included in the Timeline as No Specific Medication Was Administered in That Time

CDT = Clock Drawing Test; CTX = cyclophosphamide; IVIg = intravenous immunoglobulin infusion; Karnofsky = Karnofsky Performance Status Scale (0–100); MMSE = Mini-Mental State Examination; mRS = modified Rankin Scale (0–6); RTX = rituximab; SARA = Scale for the Assessment and Rating of Ataxia (0–40).
She was discharged from the neurologic rehabilitation ward to her own home 5 months after starting CTX. At a follow-up visit 5 months after the final CTX infusion, she reported slow but continuous improvement, which was also clinically noticeable (Videos 5 and 6), despite the fact that she had sustained a hip fracture in the meantime (Figure). She now lives at home requiring only 1 daily visit by home care.

SEZ6L2 antibody–associated cerebellar ataxia is an ultra-rare disease, therefore our case is interesting because of the substantial treatment response.

The symptoms caused by SEZ6L2 antibodies stem from their known predilection against cerebellar and hippocampal neuronal cell surface antigens. This is in accordance with our patient’s initial symptoms, consisting of ataxia, oculomotor disturbances, dysarthria, mild bradykinesia, and mild cognitive impairment, which were also manifested in previously published cases. Contrary to prior reports, our patient showed substantial improvement following immunotherapy, culminating in her return home.

Four of the 6 previously published patients did not benefit from immunotherapy, whereas the remaining 2 patients stabilized and showed mild improvement. Notably, immunotherapy was initiated within 2 months of symptom onset in only 1 of these patients and more than a year after the onset of symptoms in 2.

In our patient, clinical improvement cannot be solely attributed to CTX because it was started only 2 months after RTX. It is feasible that the sustained response was due to combined medication. At some point, 2 previously published patients had received CTX without clear treatment response. Two other patients had been given RTX with treatment response in only 1, whose symptoms (less severe than in our patient) stabilized. The reasons for these differences in treatment response are unknown.

Previous reports on SEZ6L2 antibody–associated syndrome have suggested that even without treatment, the symptom progression may slow down or cease, but spontaneous improvement has not been reported. Our case report shows that intense sequential immunotherapy may considerably improve level of functioning in some patients.

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Disclosure
A. Mehdiyeva and A. Hietaharju report no disclosures relevant to the study from the past 2 years. J. Sipilä has received honoraria (Merck, Pfizer, and Sanofi), consultancy fees (Rinnekti Foundation and Medafficon), and travel grants and congress sponsorship (AbbVie, Orion Pharma, Merck Serono, Sanquin, Lundbeck, and Novartis) and holds shares (Orion Corporation). Go to Neurology.org/NN for full disclosures.

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