Case Reports

Anti-CV2/CRMP5 Paraneoplastic Chorea Effectively Managed with Intravenous Amantadine

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Abstract

Background: Paraneoplastic chorea is typically a subacute progressive hyperkinetic movement disorder. The mainstay of treatment is managing the underlying neoplasm. However, the clinical course may be variable, and effective symptomatic management can precede the start of cancer treatment.

Case report: A 63-year-old man presented with insidious onset, slowly progressive generalized chorea for 1 year, later diagnosed as anti-CV2/CRMP5 autoantibody positive paraneoplastic chorea. His chorea was markedly improved with intravenous amantadine.

Discussion: In patients with anti-CV2/CRMP5 autoantibody-related chorea, sequential follow-up of brain magnetic resonance imaging reveals progression from active inflammation to atrophy. Our report highlights the efficacy of intravenous amantadine in paraneoplastic chorea.

Keywords: Paraneoplastic, autoimmune, CRMP5, CV2, chorea

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Introduction

Paraneoplastic chorea is a rare hyperkinetic movement disorder caused by remote effects of the underlying neoplasm on the basal ganglia.1 Anti-CV2/CRMP5 autoantibody is the most commonly detected anti-neuronal autoantibody.2 Patients with typical paraneoplastic chorea show fully developed chorea in the course of weeks to months with acute inflammation in the striatum.1,2 Median survival period depends on the underlying cancer and related antibody but generally ranges from 10 to 20 months.3,4 Regarding the treatment approach, the best option for symptomatic management remains unclear. We reported a case of anti-CV2/CRMP5 autoantibody positive paraneoplastic chorea presenting with insidious onset and slow progression, decreased striatal volume on serial follow-up magnetic resonance imaging (MRI), effectively managed with intravenous amantadine prior to anti-cancer management.

Case report

A 63-year-old man presented at our clinic with slowly progressive chorea starting from the neck of 1-year duration. The patient was a 60 pack-year smoker with hypertension and was undergoing a statin drug treatment for dyslipidemia at the time of presentation. In anamnesis, there was no family history of movement disorder or stroke, or evidence of recent infection or weight change; slow development of chorea that had spread to the right arm and affected gait as a consequence was noted. Initially, chorea was managed with clonazepam (1 mg/d) and...
haloperidol (1.5 mg/d). At post-treatment, mild improvement of symptoms was observed initially with gradual worsening over the next 6 months. After increasing the dose of haloperidol, the improvement of chorea was achieved, but the development of Parkinsonism as a side effect was noted. Therefore, haloperidol was switched to quetiapine (400 mg/d), but the relapse of chorea was observed. When the patient was referred to our clinic 12 months after the symptom onset (Video 1), his previous medical records were not accessible.

On the brain MRI images acquired at 12 months after the symptom onset, marked bilateral striatal atrophy was observed (Figure 1B). Peripheral blood smear, fasting glucose, and glucose tolerance test were unremarkable. Tumor markers, including carcinoembryonic antigen, prostate-specific antigen, and carbohydrate antigen 19-9 were normal. Genetic tests for spinocerebellar ataxia type 17 and Huntington’s disease were negative. In the results of detailed neuropsychiatric cognitive assessment (Seoul Neuropsychological Screening Battery, 2nd edition), mild cognitive impairment was revealed especially regarding frontal lobe function. In addition, mild depression was noted in the abbreviated version of Geriatric Depression Scale (6/15).

After initial work-up at our clinic, we were able to assess the previous brain MRI scans acquired 4 months after the symptom onset. T2-hyperintensities were present in the bilateral caudate nucleus and anterior putamen (Figure 1A). On the \[^{18}F\]N-(3-fluoropropyl)-2β-carbomethoxy-3β-(4-iodophenyl) nortropane (FP-CIT) positron emission tomography (PET); DAT, Dopamine Active Transporter; FLAIR, Fluid Attenuated Inversion Recovery; MRI, Magnetic Resonance Imaging.

Figure 1. Structural and Functional Imaging of the Patient’s Brain. Brain MRI nonenhanced T2 FLAIR images acquired 4 months (A) and 12 months (B) after initial symptom showed marked striatal hyperintensity and striatal atrophy, respectively, and FP-CIT PET scan showed a decrease in DAT binding in the bilateral striatum (C). Abbreviations: \[^{18}F\]N-(3-fluoropropyl)-2β-carbomethoxy-3β-(4-iodophenyl) nortropane (FP-CIT) positron emission tomography (PET); DAT, Dopamine Active Transporter; FLAIR, Fluid Attenuated Inversion Recovery; MRI, Magnetic Resonance Imaging.
Amantadine in Paraneoplastic Chorea

Few randomized pla...ation using serum and cerebrospinal fluid mixture. Meanwhile, the diagnosis of small cell lung cancer with metastasis in the lymph node, first lumbar spine, and left ureter was made based on the malignancy workup.

Before any treatment plan for the lung cancer was implemented, intravenous amantadine (200 mg in 500 cm³ of normal saline given over a 3-hour period, twice per day for 3 days) was administered to manage chorea; in response, remarkable improvements in chorea, especially of the limbs and trunk were attained, and consequently, the patient’s gait was improved (Video 2). We confirmed the efficacy and safety of drug treatment in our patient and made the switch from intravenous amantadine to oral amantadine (200 mg/d). At the outpatient clinic, the dose of oral amantadine was increased from 200 to 300 mg/d, and beneficial effect of the treatment was maintained at 3 years’ follow-up.

**Discussion**

We reported a case of slowly progressive generalized chorea associated with metastatic small cell lung cancer. We demonstrated a change from active inflammatory state to atrophy through serial MRI and suggested the effectiveness of IV amantadine in treatment of the patient’s chorea.

Oroolingual chorea and retrocollis may resemble tardive dyskinesia. However, at an initial visit with the attending physician (JH Lee), the patient reported no prior exposure to any antiemetic or antipsychotic medication and was experiencing milder form of chorea that was compatible with hyperkinetic movements observed in our clinic 12 months later (Video 1). Subsequently, the patient underwent treatment with haloperidol, which initially showed improvement of chorea. Based on these facts, we considered that chorea in our patient was due to paraneoplastic syndrome and not tardive dyskinesia.

Brain MRI performed 12 months after the symptom onset revealed severe atrophy in the striatum, most prominently involving bilateral head of the caudate nuclei. Subsequently, we assessed the brain MRI scans acquired previously at other hospitals and observed T2-hyperintensities in the identical regions, which agree with the findings of another report of cases of paraneoplastic chorea with positive anti-CV2/CRMP5 autoantibody that included changes of T2-hyperintensity in the basal ganglia. Moreover, FP-CIT PET revealed decreased uptake of dopamine transporter which may be associated with severe striatal atrophy (Figure 1B), a finding most likely explained as a remnant of active inflammation (Figure 1A). Persistence of atrophic landmarks after inflammation suggests pathophysiology involving irreversible cytotoxic process. Previous studies have reported presynaptic dopaminergic deficit in patients with hyperglycemic chorea and Huntington’s disease, but not in those with paraneoplastic chorea with anti-CV2/CRMP5 autoantibody. To the best of our knowledge, this is the first case report to present a time-series follow-up brain MRI, with conclusive FP-CIT PET scan finding of decreased presynaptic dopaminergic uptake in the bilateral striatum in a patient with paraneoplastic chorea.

Immunomodulatory therapy with intravenous immunoglobulin and steroids achieved some success in the control of chorea in patients with paraneoplastic chorea; however, there is no consensus on symptomatic management in such cases. Few randomized placebo-controlled trials on treatment with amantadine are reported for Huntington’s disease. Studies have indicated that the pathophysiology of chorea in Huntington’s disease can be explained by imbalance between indirect and direct pathways due to selective involvement of indirect pathway. We considered that the effectiveness of amantadine on chorea of anti-CV2/CRMP5-related chorea was due to modulation of the imbalance between direct and indirect pathways in the basal ganglia by two possible mechanisms: (1) decrease of glutaminergic input from the neocortices and consequently, decrease of total motor output, and (2) sensitization of the globus pallidus pars interna (GPi) to glutaminergic input and augmentation of inhibitory response at the thalamus. Clinicians should consider IV amantadine as a low-cost alternative treatment to control activity-limiting severe chorea in patients with paraneoplastic chorea before initiating other immunomodulatory therapy or cancer treatment.

One of the limitations of our case report was that our antibody testing was performed only in a qualitative manner. Nevertheless, the method was already validated in previously published literature. Future studies focused on changes of autoantibody titers before and after treatment with amantadine are needed.

In conclusion, the findings of time-series follow-up brain MRI in anti-CV2/CRMP5 autoantibody-related chorea indicated the change from active inflammation to atrophy according to the clinical course, which may reflect the mechanisms involved in pathophysiology of the disease. Moreover, IV amantadine showed potential as a viable option for symptomatic management in anti-CV2/CRMP5 autoantibody-related paraneoplastic chorea.
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