Case Report

Parkinson-Like Symptom as a Rare Manifestation of Systemic Lupus Erythematosus: A Case Report
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Abstract

A 65-year old Bangladeshi woman with Systemic Lupus Erythematosus (SLE) developed Parkinson-like movement disorder. Steroid pulse therapy followed by prednisolone was most effective in this case. Psychosis, seizure and meningitis are common central nervous system (CNS) manifestations in SLE patients, and Parkinson-like rigidity or tremors are rare.

Keywords: Systemic Lupus Erythematosus, CNS lupus, Parkinson-like rigidity, Steroid pulse therapy.

INTRODUCTION

Patients with SLE, CNS involvement may have varied presentations like headache, cognitive dysfunction, seizure, cerebrovascular disease, acute confusional state and psychiatric disturbance. Movement disorders like chorea, hemiballismus, cerebral ataxia and Parkinson-like rigidity or tremors are rare manifestations1. We came across a woman with SLE and Parkinson-like symptoms, an extremely rare form of CNS lupus manifestation.

CASE REPORT

A 65-year old Bangladeshi woman with Diabetes Mellitus (DM) and Hypothyroidism was diagnosed as a case of SLE on the basis of polyarthritis, photosensitive malar rash, and alopecia, positive anti Nuclear Antibody (ANA) and anti ds DNA in 2009. She was treated with hydroxyl chloroquine (HCQ), methotrexate (MTX) and prednisolone on a tapering course along with thyroxin. She improved with these medications. Later MTX was stopped due to haemolysis and azathioprine was added. At the same time, course of prednisolone was completed but the patient developed Jaccoud’s like deformities in metacarpophalangeal joints of right hand and interphalangeal joints of right foot. She also developed Coombs’ positive autoimmune haemolytic anemia. Subsequently she developed difficulty in walking, slowness of speech with behavioural alteration (like apathy, mutism and irritability), bradykinesia and rigidity. She was considered as a case of Parkinson Disease (PD) and put on Levodopa 150 mg along with Carbidopa 15 mg daily for four years. There was gradual progression of symptoms, there by Ropinirole was added but no improvement was seen. She had a single episode of convulsion in 2011. In 2012 she experienced low trauma fracture of neck of left femur and pulmonary embolism. She had no history of unconsciousness, nasal regurgitation, swallowing difficulties and bowel-bladder abnormalities.

She had mild splenomegaly (12.4cm), no ascites, GIT bleeding or sign of CLD. There were no abnormalities detected in heart and lungs. Her consciousness was clear. She had expressionless face with slow and monotonous speech. There was cogwheel and lead pipe rigidity of upper and lower limbs. Muscle power was 4/5 in both proximal and distal groups with diminished deep reflexes, equivocal plantar reflexes bilaterally with short step gait.

Laboratory examination showed haemoglobin 5.3gm/dl, CRP was 6, ANA and anti ds DNA were positive, liver function, renal function and urine R/E were within normal limits. Anti cardiolipinAb positive but Lupus anti-coagulant, β2 Glycoprotein-1 IgM and IgG were negative. Vitamin D total was 32.1ng/ml. Rheumatoid factor, anti CCP, anti SmAb, anti SSA, Anti SSB, HBs Ag, anti HCV, ICT for Malaria and Kala azar, anti Mitochondrial Ab, Anti LKM Ab were negative. CT scan and MRI of the brain showed age related mild cortical atrophy and brain SPECT revealed hypo perfusion of fronto-parietal region and left basal ganglion.

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Diagnosis of CNS lupus with Parkinsonian-like symptoms was added, stopped all anti Parkinson drugs and steroid pulse therapy (800 mg Methylprednisolone daily for 3 days) was given, followed by 40 mg daily oral prednisolone. After steroid therapy along with pulse cyclophosphamide, there were improvement of rigidity, facies, Speech, walking and reduced aggressiveness of behavior. Finally she was discharged after one month of hospital course.

DISCUSSION
Our patient was initially diagnosed as a case of SLE on the basis of polyarthritis, photosensitive malar rash, alopecia, positive ANA and anti-ds DNA in 2009. Later she developed seizure, Jaccoud’s like joint deformities, rigidity, bradykinesia, expressionless facial features, slowness of speech with behavioral alteration (as irritability, aggressiveness, apathy, mutism). Other causes of secondary Parkinsonism was not considered as the patient had never been given responsible drugs and never exposed to toxic agent like CO or Manganese. Cranial CT and MRI scan showed only degenerative changes. Brain SPECT shows hypo perfusion of fronto-parietal region and left basal ganglia. Anti-phospholipid antibodies like Anti-cardiolipin (aCL) Ab, lupus anti-Coagulant (LA) and B2 glycoprotein-1 IgM were positive. Therefore, in this case, Parkinsonian like symptoms were considered to be manifestations of CNS lupus. After a high dose of methyl prednisolone pulse therapy, Parkinsonian-like symptoms were improved dramatically and all anti Parkinsonian drugs were withdrawn.

Seizures, mental disorders and cranial neuropathy are the more commonly observed symptoms of CNS lupus. Chorea is also common as extrapyramidal involvement of SLE. In contrast, Parkinsonian-like symptoms are extremely rare. Willoughby et al described a 30-year-old male SLE patient with cogwheel rigidity. Despite 40 mg daily of prednisolone, complications due to meningitis and endocarditis followed and he died. Autopsy revealed multiple areas of encephalomalacia in the basal ganglia of the brain. In a study Yancey et al, 2 out of 37 children with SLE had Parkinsonian-like symptoms and in one child had cogwheel rigidity. The clinical outcome was not
described. In another case, Parkinsonian-like symptoms and coma was developed in a 16-year-old girl; a nearly complete recovery was achieved. Although the authors did not describe how they had treated that particular patient, they favoured combination of high dose prednisolone (2mg/kg/day) and a cytotoxic agent as initial therapy of CNS lupus, and steroid pulse therapy for non responding patients. Nagaoka et al. reported SLE in a 35-year-old woman with akinesia, muscle rigidity and expressionless facial features. Her Parkinsonian-like symptoms were controlled by 40mg daily methylprednisolone and anti-Parkinsonian drugs. For this patient, anti-Parkinsonian drugs were necessary since symptoms relapsed during the course of steroid-tapering when amantadine was discontinued. In our patient the Parkinsonian-like symptoms were improved dramatically after Methylprednisolone pulse therapy and the patient remained in clinical remission after discharge. In many cases, CNS lupus can be effectively treated with high dose corticosteroids including methylprednisolone pulse therapy. About 25% of CNS lupus episodes are not responsive to steroid. In such cases, cyclophosphamide pulse therapy and plasma exchange may be effective.

Although the putative pathogenic mechanism of CNS lupus includes vasculitis in the CNS, functional disorder of nerve cells due to auto-antibodies against neuronal cells (eg. Anti-Asialo GMI Antibody, Anti-Ribosomal P protein Antibody, Anti-Neuronal cell Antibody), and thrombi formation in the CNS by Anti-Phospholipid Antibodies, precise mechanisms are yet to be defined.

CONCLUSIONS

We reported a rare case of SLE, in which Parkinsonian-like symptoms were noticed as manifestation of CNS lupus. The patient’s symptoms were effectively managed with high dose prednisolone with anti Parkinsonian drugs ropinirole only.

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