Systemic lupus erythematosus presenting as generalized chorea: case report and review of literature

Mohamed O Abdalla¹, Ahmad O Ahmad¹, Husam A AlTahan², Ibrahim A AlOrairi² and Abdulrahman M AlTahan*³

¹Dallah hospital, Riyadh, Saudi Arabia
²King S bin Abdulaziz University, Riyadh, Saudi Arabia
³King S bin Abdulaziz University, Dallah hospital, Riyadh, Saudi Arabia

Introduction

Neuropsychiatric systemic lupus erythematosus (NPSLE) is one of the main diagnostic criteria of systemic lupus erythematosis (SLE), occurring in 25% - 75% of affected patients [1,2]. Chorea is recognized as one of the clinical manifestations of NPSLE especially in children, however it is very rare to occur as the first presenting feature of SLE [3]. In this report we describe a teenage girl who presented acutely with generalized chorea and was found later on to have SLE. Her case will be described in detail and the topic of NPSLE will be reviewed.

Case report

A 14-year-old female presented with subacute onset of random, unintentional movements involving the neck, trunk, limbs as well as facial muscles. The movements were severe enough to interfere with mobility. Otherwise she did not have any systemic complaints. Had no history of a recent infectious illness and did not use any medication. Her past and family history was unremarkable. On examination, she was alert, fully oriented with intact cognitive functions. The movements were noted as involuntary, rapid and purposeless generalized movements. She could not control it and had difficulty stabilizing her head. Auscultation revealed pansystolic murmur over the apex. She had no skin lesion, joint tenderness or swelling. Blood investigations showed normochromic normocytic anemia with thrombocytopenia (Hb: 8g/dl, MCV: 80 fl, WBCs: 4.5×10⁹ Platelets: 84×10⁹). Had normal renal and liver functions. ANA was positive at 1:180. Coombs test anti-dsDNA Abs were positive. Furthermore her cardiolipin IgG was positive (36.9 U/ml) and urine analysis shows hematuria (20-30/HPF) and proteinuria (100 mg/dl). Brain MRI demonstrated multiple bilateral old and recent ischemic lesions (Figure 1).

On confirming diagnosis of NPSLE, the patient was offered 3 doses of 1000 mg IV methylprednisolone, together with enoxaparin 30 mg BID, hydroxychloroquine 200 mg BID and mycophenolic acid 1g BID. She improved partially over the next few days and another course of 3 doses of methylprednisolone were offered. She showed more improvement and was able to walk unaided by the end of the week. On follow up after discharge two weeks later, involuntary movements was almost absent, and she was able to return to her normal daily routine.

Discussion

NPSLE may present before the diagnosis of SLE is made or during the course of the illness, and can be classified as either primary, when there is direct nervous system involvement, or secondary, when the involvement is a complication of the disease or its treatment [4,5].

NPSLE manifestations are extremely variable and include features of both psychiatric and neurological dysfunction [4-9]. Psychiatric manifestations may include cognitive dysfunction, anxiety, mood and personality disorders, and psychosis [5,6]. The commonest neurological manifestations include headache, stroke, seizure, and peripheral neuropathy [4,7-9]. Other less common features include

*Correspondence to: Abdulrahman AlTahan, King S bin Abdulaziz University, Dallah hospital, Riyadh, Saudi Arabia, E-mail: atahan45@hotmail.com

Received: January 25, 2018; Accepted: February 21, 2018; Published: February 25, 2018
cranial neuropathies, transverse myelitis, meningitis, and movement disorders which occur in less than 5% of patients and include chorea, ataxia, choreoathetosis, dystonia, and hemiballismus [10-12].

Chorea accounts only for about 2% of all NPSLE manifestations, which qualified it as a clinical diagnostic challenge [12]. In practice, the differential diagnosis of chorea is wide and include a long list of hereditary and non-hereditary disorders [13-15]. Acquired causes include drugs and toxins, Sydenham chorea, SLE and antiphospholipid (APL) syndrome, chorea gravidarum, stroke and hyperthyroidism [14,15].

In view of the above described wide NPSLE clinical manifestations, pathogenesis is expected to include multiple mechanisms. However, NLSLE pathogenesis is still controversial, and probably involve immune mediated as well as ischemic mechanisms [16,17]. In our patient, the old and new ischemic changes noted on MRI involve an ischemic vascular process, but these can’t be explained simply by the procoagulant effects of APL antibodies causing thrombotic occlusion of capillaries. It was hypothesized that an immune process may be responsible for a direct injury to the blood-brain barrier allowing autoantibodies to enter the nervous system and attack neurons involved in nigro-striatal pathways [16-19]. One meta-analysis showed that NPSLE patients have higher serum levels of certain antibodies including antiphospholipid, lupus anticoagulant, anti-ribosomal P, and antineuonal antibodies when compared with SLE patients who have no neurological involvement [20]. Other secondary factors that also contribute to the pathogenesis include infections associated with immunosuppressive treatments, hypertension, and renal failure [21].

NPSLE and APL syndrome are very important differential diagnosis of acute chorea, especially in children as apparently APL antibodies are more relevant in this population [21].

A higher index of suspicion should probably promote earlier diagnosis of NPSLE and particularly its most rare manifestations including chorea.

References
13. Cardoso F (2012) Difficult diagnoses in hyperkinetic disorders - a focused review. Front Neurol 3: 151. [Crossref]

Copyright: ©2018 Abdalla MO. This is an open-access article distributed under the terms of the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original author and source are credited.