Choreiform Movement Disorder Associated with Subdural Hematoma: A Case Report

Subdural Hematoma Bağlı Koreiform Hareket Bozukluğu Bir Olgu Sunumu

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Dear Editor,

A 68-year-old female was admitted to our clinic with headache and involuntary curling-like movements involving the right upper and lower extremities and trunk, which suddenly started 4 days ago and progressed. These movements were lasting 3-4 seconds and recurring in every 4 seconds. There was no history of trauma or anything else in her medical history. She was conscious, orientated, and cooperated. There was no neck stiffness and cranial nerves were intact. Cerebellar tests were normal, muscle strength was 5/5 in all extremities, Babinski signs were negative bilaterally, and deep tendon reflexes were normoactive. There were choreiform movements involving the right arm and leg. Fasting blood glucose, electrolytes, liver function tests, thyroid function tests, hemogram, complete urine analysis, and urine microscopy resulted as normal. Creatinine was 0.56 mg/dL, prothrombin time was 18.8 s, INR was 1.42, and the erythrocyte sedimentation rate was 30 mm/h.

Cranial magnetic resonance imaging (MRI) with sagittal T2 fast spin echo (FSE), axial T2 FSE, axial T1 fluid-attenuated inversion recovery (FLAIR) and coronal T2 FLAIR sequences were performed. Neuroimaging showed enlarged and deepened bilateral Sylvian fissures and hemispheric cortical sulci, which were suggestive of cerebral cortical atrophy. A smoothly-contoured septum pellucidum cyst that was approximately 18 mm at the widest point was seen hypointense in T1A and hyperintense in T2A-weighted images in the vicinity superior of the third ventricle. A lesion in the right fronto-parietal subdural area, which was seen hyperintense in all MRI sequences and which was approximately 32 mm at the widest point, suggested a subdural hematoma (SDH) (Figure 1). After evaluation of the cranial MRI, the patient was asked if she had had any trauma and she admitted that she had a head trauma 4 days ago. With the data gathered from anamnesis, neurologic examination, laboratory tests, and cranial MRI, a diagnosis of subacute SDH due to the trauma was considered and a surgical operation was planned by the neurosurgery department.

Patients with SDH could be asymptomatic or could present with loss of consciousness due to acute bleeding, hemiparesis or coma. Seizure and headache are the other common findings of SDH (1,2). Choreiform movement disorder caused by SDH was first reported in a child with leukemia in 1977 (3). Neuroimaging has a crucial role in the diagnosis of SDH. Computed tomography, cranial MRI, and diffusion MRI are neuroimaging modalities that are used in diagnosis of SDH (1). Clinically stable patients who have a bleed less than 5 mm in diameter could be followed up for spontaneous resolution. Surgery is advised in clinically unstable patients, patients with 9 or more points in the Glasgow Coma Scale (GCS), patients with a more than 2 points increase in GCS after admission, patients with >10 mm thickness of clot or with >5 mm shift (4). Cranietomy with membranectomy and single or double- “Burr hole” drainage are the surgical procedures used in the treatment of SDH (1,5). The superiority of one method over another is controversial. Right frontal “Burr hole” drainage was performed and the patient’s involuntary movements improved in the postoperative period. The patient was discharged and
followed up. As a result, SDH could rarely result in choreiform movements and these movements could totally improve with treatment of the SDH.

Ethics

Informed Consent: The patient gave informed consent.
Peer-review: Internally peer-reviewed.
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References