Hemichorea and Hemiballismus Associated with Cerebral Vascular Malformation Induced by Hyperglycemia: Case Report

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Summary

Hemichorea, hemiballismus, and hyperglycemia associated with vascular malformation is rare. We report a patient who presented with involuntary movements in the left-side with concurrent hyperglycemia. The patient had type 2 diabetes mellitus (DM) and a venous angioma in the basal ganglia on the cranial magnetic resonance imaging (MRI). A woman aged 49 years presented with flinging and throwing movements of the left upper and lower limb, which she had had for one month. Her neurologic examination confirmed involuntary, irregular, wide amplitude movements of the left limbs consistent with left hemiballismus. She had hypertension and type 2 DM. Her glucose level was 400 mg/dL. Cranial MRI showed a cavernoma in the right subependymal area of the lateral ventricle and a venous angioma in the right nucleus lentiformis, which was confirmed on digital subtraction angiography. Hemiballismus improved after blood glucose level had been regulated in the follow-up period. Especially in the elderly secondary causes should be investigated in patients with acute or subacute onset of chorea-ballismus. We think that our patient’s clinical presentation was induced by unregulated DM. Venous angioma with chorea-ballismus is rarely stated in literature and we presume the mass effect of venous angioma could be responsible of our clinical findings.

Keywords: Chorea, ballismus, vascular malformation, hyperglycemia


Anahtar Kelimeler: Kore, ballizm, vasküler malformasyon, hiperglisemi

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Introduction

The term chorea, which is included in hyperkinetic disorders, means ‘dance’ in Greek and is used for irregular muscle contractions that are clinically random, arrhythmic, short-lived, low amplitude, sudden and jerky in nature, and involving distal parts of extremities especially, and sometimes the tongue (1). If severity and amplitude of these involuntary movements increase and involve proximal parts of extremities they are called ballismus (1). Chorea and ballismus may sometimes co-occur or follow each other (2). If they involve only one half of the body they are called as hemichorea or hemiballismus (1). Hemiballismus may sometimes occur together with other involuntary movements such as dystonia, myoclonus, or orofacial tics (2). These movements rarely occur intermittently; they are usually continuous in nature and can be interrupted by patients only for short periods of time (1). Their disappearance during sleep is characteristic. Putamen and globus pallidus are important for choreic and ballistic movements and subthalamic nucleus pathologies are especially important for ballismus (2). They can be classified as primary (neurodegenerative diseases) or secondary (autoimmune, structural, vascular, metabolic, and iatrogenic) according to the etiology (2). Late-onset cases are secondary and vascular causes are usually prominent (2,3,5,6,7,8,9,10,11,12,13,14). Infarct, hemorrhage, cavernoma, and arteriovenous malformation (AVM) are among the possible vascular causes (2,3,5,6,7,8,9,10,11,12,13,14). Metabolic disturbances including hyperglycemia, particularly non-ketotic hyperglycemia may be associated with many neurologic disorders (4). Chorea-ballismus is among the most common of these neurologic disorders (4). Co-occurrence of hemiballismus, hemichorea, and hyperglycemia due to vascular malformation is rarely seen (5,15,16). We present a woman who presented with left-sided involuntary movements. Hyperglycemia was detected in a blood analysis and she had a history of type 2 diabetes mellitus (DM). A venous angioma was detected at the level of the basal ganglia in cranial magnetic resonance imaging (MRI).

Case Report

A woman aged 49 years presented with involuntary jerks in her left arm and leg, which she had had for nearly a month. She had never been admitted to a center for these symptoms before. Her medical history was not significant except for hypertension and type 2 DM. The patient’s type 2 DM was diagnosed 5 years earlier but she did not use her diabetes drugs and had not adhered to her diet for the last 6 months. With the exception of the left-sided severe hemiballismus, which started from the hip and shoulder, her physical and neurologic examinations were normal. Her laboratory analysis revealed a blood glucose of 400 mg/dL. Cranial MRI showed a 10x13-mm cavernoma located subependymally on the lateral side of the ventricular corpus in the right hemisphere, which had heterogeneous signal intensity. Its periphery was hypointense, especially in T2-weighted and FLAIR sequences, and susceptibility-weighted imaging (SWI) suggested a vascular lesion (Figures 1, 2, 3). Digital 4-vessel cerebral angiography evaluation revealed findings consistent with venous angioma at the level of the left nucleus lentiformis. The patient’s blood sugar was regulated in approximately 1 month. Haloperidol was stopped after 3 weeks because of a high level of somnolence. At the end of the month, near-total improvement was seen in choreiform movements. She had no active symptoms during her 2 years of follow-up. Then her left-sided choreiform movements restarted and analysis again revealed hyperglycemia. Her HgbA1c value was 11.2%. Repeated cranial MRI revealed no significant change.

Discussion

Secondary causes should be kept in mind in acute or subacute onset chorea-ballismus in patients at advanced ages (2). These causes include drugs, vascular diseases, metabolic causes, autoimmune diseases, neoplasia, senility, and hereditary causes.
(2,3,5,6,7,8,9,10,11,12,13,14). Hemichorea/hemiballismus is used to state presentation in on half of the body. They form approximately 0.7% of all movement disorders. They are usually associated with contralateral structural pathologies such as vascular lesions, tumors or tuberculomas, sometimes they may be associated with diabetic ketoacidosis, or non-ketotic hyperglycemia (5). Vascular etiologies that may cause chorea and ballismus include ischemic or hemorrhagic cerebrovascular events and vascular malformations (1,2,3,5,6,7,8,9,10,11,12,13,14). Vascular malformations consist of venous angiomas, AVM, and cavernous angiomas (6). Chorea due to vascular anomalies such as cerebral angioma, or cavernoma is rarely seen (7). The etiopathogenesis of chorea was tried to be explained by the disruption of balance between direct and indirect pathways. The impairment of striatal outputs that project to the globus pallidus externa (indirect pathway) increases inhibition of neurons at this region, and this decreases output of the globus pallidus externa by increasing inhibition on the globus pallidus interna. As a result, thalamocortical activity which is known to be the basic mechanism in hyperkinetic disorders develops. In the presented case, a venous angioma was detected at the level of the right nucleus lentiformis. Previous cases of chorea and ballismus associated with cavernoma have been reported (8,9,10,11,12,13,14). Very rarely cases have been associated with developmental venous anomalies (15,16). In these cases, venous hypo-perfusion associated with venous anomalies at the contralateral putamen level were detected. In the present case, possible hemodynamic changes due to the venous anomaly or a direct effect of the mass were thought to be responsible for the patient’s clinical situation.

The second most common cause of hemiballismus after stroke is hyperglycemia (17,18). Hyperglycemia associated with ballismus/chorea (HABC) is commonly seen in patients with DM aged between 50-80 years, whose blood glucose levels cannot be regulated (19,20). Typically, the clinical picture settles within hours. During this period the patients’ blood glucose levels are between 400-1000 mg/dL and involuntary movements regress after control of hyperglycemia. Some patients have normal computed brain tomography (CBT) findings although others may have hyper-dense lesions in the basal ganglia. Cranial MRI evaluations may also be normal or they may demonstrate hyper-intense lesions in the putamen or caudate nucleus in T1-weighted images and variable intensity lesions in T2-weighted images (21,22,23). The pathophysiology of HABC has not yet been completely understood. Hyperglycemia may lead to tissue edema and this may increase vascular resistance and viscosity, decrease metabolic rate of neurons, inactivate Krebs cycle, and lead to consumption of gamma-aminobutyric acid (GABA) by the brain to produce energy (24). Moreover, Chang et al. (21) evaluated cranial MRI and MR spectroscopy findings of 18 patients with HABC and suggested that transient ischemia triggered by hyperglycemia may be responsible for the clinical picture. Shan et al. (24) suggested that ischemia triggered by hyperglycemia damages GABAergic neurons in particular, this affects the indirect pathway more prominently and thereby inhibits inhibition on the subthalamic nucleus. Additionally, hyperosmolarity decreases the epileptic threshold in the direct pathway, which increases firing of this pathway. At the end of all these processes, reactive swollen astrocytes and gliosis occur and these may cause petechial hemorrhages. This explains the hyper-density in acute CBT. We observed no such finding in the CBT of our patient. Oh et al. (22) performed a meta-analysis of 53 cases of HABC and demonstrated hyperintense lesions at the level of the putamen in T1-weighted images in all cases. Enhancement was also seen in T2-weighted images. Additionally, in our case a hypointense lesion in T2-weighted image was detected to be a venous anomaly. In the same study by Oh et al. (22) in 19 of 22 patients evaluated with MRI, cranial MRI findings disappeared with improvement in chorea; blood sugar regulation was sufficient in most patients and drug treatment was given very rarely. However, in the present case diabetes regulation was combined with haloperidol.

Conclusion

In conclusion, although the present case did not fully meet the HABC criteria, especially with imaging findings, we believe that poorly-regulated DM precipitated the clinical picture and regulation of diabetes formed the basis of our treatment. Moreover, we suggest that the mass effect and hypo-perfusion due to venous anomaly may be responsible for the clinical picture in rare cases of comorbid venous angioma and chorea-ballismus.

Ethics

Informed Consent: Consent form was filled out by this patient.
Peer-review: Internal peer-reviewed.

Authorship Contributions

Surgical and Medical Practices: Ferda Selçuk, Süha Akpınar, Concept: Ferda Selçuk, Mine Hayriye Sorgun, Design: Ferda Selçuk, Mine Hayriye Sorgun, Data Collection or Processing: Ferda Selçuk, Mine Hayriye Sorgun, Süha Akpınar, Analysis or Interpretation: Ferda Selçuk,

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References