Case Report

Encephalopathy, Hypoglycemia, and Flailing Extremities: A Case of Bilateral Chorea–Ballism Associated with Diabetic Ketoacidosis

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Abstract

Background: Hypo/hyperglycemia is a known cause of chorea and hemiballism. The temporal lobes, hippocampus, basal ganglia, and substantia nigra are most susceptible to hypoglycemic changes.

Methods: We present a case of bilateral chorea and bi-ballism accompanied by encephalopathy in the setting of severe hypoglycemia and diabetic ketoacidosis. The patient had brain MRI changes involving both caudate nuclei, temporal lobes, and hippocampi.

Discussion: This case demonstrates the basal ganglia’s vulnerability to hypoglycemia and the need for cautious evaluation of involuntary movements when they occur in the setting of encephalopathy.

Keywords: Encephalopathy, hypoglycemia, chorea, ballism, diabetic ketoacidosis, hyperglycemia

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Introduction

We present a case of bilateral chorea and ballism (bi-ballism) accompanied by encephalopathy in a patient with severe hypoglycemia and diabetic ketoacidosis. This case was significant because the movements were initially mistaken as simple flailing in the setting of hypoglycemic encephalopathy. Although there have been case reports of chorea and hemiballism associated with hypo/hyperglycemia, the findings in this case manifested bilaterally and were associated with brain magnetic resonance imaging (MRI) changes involving both right and left caudate nuclei, temporal lobes, and hippocampi. The clinical manifestations of hypoglycemia can be broad and may include neurological manifestations such as amnesia, seizures, and hyperkinetic movement disorders. The temporal lobes, hippocampus, basal ganglia, and substantia nigra appear most susceptible. This case emphasizes the sensitivity of the basal ganglia to hypoglycemia, and the need for careful and thoughtful evaluation of involuntary movements when they co-occur with encephalopathy.

Case report

A 19-year-old Caucasian male with insulin-dependent type 1 diabetes developed involuntary movements of all four extremities after several episodes of prolonged hypoglycemia. These movements were initially described as “flailing” movements associated with diabetic encephalopathy. He had no previous history of chorea or kernicterus, and he had no family history of Huntington’s disease, benign hereditary chorea, neuroacanthocytosis, or other neurodegenerative disorders. He was admitted with diabetic ketoacidosis with an anion gap of 24. On day 5 of his hospitalization, he suffered two episodes of severe hypoglycemia with blood glucose levels recorded in the range of 37–44 mg/dL. The episodes of hypoglycemia lasted approximately 2 hours. Within hours of these episodes, the patient developed encephalopathy and involuntary movements of his arms, legs, and face.

During his examination, he appeared mildly agitated and in some distress due to his hyperkinetic movements. He was alert and oriented...
to self, time, and place. He had scanning dysarthric speech, but was easily understood. He demonstrated deficits in recall and attention. His language was intact. Cranial nerves were normal. He had full strength and his reflexes were 2+ throughout with plantar flexor responses bilaterally. His involuntary movements included facial chorea and chorea of the fingers, hands, arms, and legs. He also had prominent intermittent ballistic movements of the arms and leg. His electroencephalogram showed evidence of mild background slowing. His ceruloplasmin was normal at 26 mg/dL and his cerebrospinal fluid analysis revealed eight red blood cells and two white blood cells with a normal range protein and negative bacterial cultures. His electroencephalogram showed evidence of mild background slowing. His brain MRI demonstrated abnormal T1 and T2/FLAIR (fluid attenuation inversion recovery) signal in the caudate nucleus and basal ganglia bilaterally, right greater than left (Figure 1). There was also edema in both hippocampal gyri. Diffusion imaging showed no restriction on diffusion.

He was treated with olanzapine 5 mg twice a day for 3 days without change in his hyperkinetic movements. Eventually he was started on clonazepam 1 mg twice a day with a significant decrease in involuntary movements after 24 hours of treatment. In addition, he was placed on an insulin drip for improved blood glucose control. Within 24 hours of these changes his involuntary movements had largely resolved. Coincident with improvement in his involuntary movements his encephalopathy also improved.

Discussion

Repetitive episodes of hypoglycemia can lead to chorea and hyperkinesia. These symptoms can be a temporary or a permanent sequela. Although there are several case reports of transient chorea in the setting of hypoglycemia, to our knowledge there have been no cases in which bilateral chorea and bi-ballism have been reported with hypoglycemia and associated closely with the co-occurrence encephalopathy.

In our patient we observed abnormal T2 and T1 hyperintensities in the basal ganglia and hippocampi bilaterally without evidence of infection or an inflammatory process. At the cellular level, both glucose deprivation and ischemia/anoxia in the brain can result in decreased energy reserve, alterations in membrane potentials, and overall cell stress. However, in cases of hypoglycemia, there is a topographical preference for the hippocampus, temporal lobe, and basal ganglia. This topographical preference is well correlated with the T2 changes on brain MRI in hypoglycemic individuals. In addition, consistent with previous reports, our patient did not demonstrate restricted diffusion on MRI, making ischemia/anoxia less likely.

Although the exact pathophysiology of chorea in hypoglycemia is still not well understood, based on MRI and single photon emission computed tomography (SPECT), it has been postulated that hypoglycemia results in temporary striatal dysfunction. SPECT imaging in patients with hypoglycemia and chorea has shown decreased blood flow in the basal ganglia and increased perfusion of the thalamus contralateral to the side of the body manifesting chorea. These findings have led to speculation that decreased pallidal inhibitory input to the thalamus, resulting in increased thalamocortical outflow, might explain the chorea and the hyperkinesia, although the exact mechanism remains unknown.

Interestingly, several case reports of hyperglycemia-induced chorea–ballism associated with striatal lesions on MRI have been reported. It is thought that depletion of gamma-aminobutyric acid (GABA) and acetylcholine, which are needed as an alternative energy source during
non-ketotic hyperglycemia, can lead to a decreased inhibitory signal to the thalamus, resulting in a hyperkinetic movement disorder. However, this theory has been disputed as chorea–ballism can also manifest in ketotic patients where acetoadipate is abundant and can be utilized as a source of GABA. Furthermore, there have been other cases of chorea–ballism in patients with hypoglycemia.

Another hypothesis has been that of cerebrovascular insufficiency. SPECT and positron emission tomography suggest the presence of regional hypoperfusion of the striatum of patients with hyperglycemia induced chorea–ballism. Furthermore, MR spectroscopy in these same patients has shown neuronal loss and damage in the striatum. Abe et al. recently reported six patients with hyperglycemia-induced movement disorder who had biopsies of the striatum performed. They found patchy necrotic tissue, severe thickening of all layers of arterioles, and marked narrowing of vessel lumens. Hyaline degeneration of the arteriolar walls, extravasation of erythrocytes, and prominent capillary proliferation were also noticed together with lymphocytic infiltration and macrophage invasion. They coined this finding diabetic striatopathy. However, whether the relationship between ischemic insult and changes on MRI is associated with striatal vasculopathy is still to be elucidated. To date, there have been no cases of hypoglycemia-induced movement patients with biopsies performed.

Treatment of chorea and ballism with low-dose olanzapine monotherapy was ineffective, but the addition of clonazepam notably resulted in rapid improvement of symptoms; however, this improvement occurred coincident to tighter glucose control. In the current case, tight regulation of blood glucose may have resulted in a rapid improvement of the transient metabolic dysfunction of the basal ganglia, and also improvement of the encephalopathy. Interestingly, the recovery was more rapid than that in previously reported cases. In a previous case of hypoglycemia-induced choreoathetosis, the patient was treated with tiapride for 4 weeks with improvement, but residual hyperkinetic issues still remained. The same patient continued to experience symptoms even after blood glucose normalization. Takahai and Ohkawa described a patient with hypoglycemia-induced bilateral ballism whose symptoms disappeared after controlling serum glucose with insulin treatment. A recent report of 25 patients from Korea with hyperglycemia induced chorea–ballism found that all patients recovered within 1 month with glycemic control and administration of neuroleptics.

We propose that correction of the metabolic derangement and rapid glucose control with an insulin drip if necessary should play a critical role in the best management of these cases. Furthermore, this current case suggests that the addition of clonazepam may be useful in some cases for control of hypoglycemia-induced chorea–ballism. Finally, patients with encephalopathy, hypoglycemia and flailing extremities should prompt clinicians to obtain rapid neurological consultation, MRI scanning, and glycemic control, and this syndrome should not be mistaken for a simple mental status change in a patient trying to get out of bed.

References


