INTRODUCTION

Uremia is a syndrome of clinical and metabolic abnormalities that develops in parallel with deteriorating renal function and brain involvement. Uremic encephalopathy (UE) is a well-known complication of uremia (1). The movement disorder seen in uremic patients typically consists of asterixis and myoclonus, which may be related to cortical dysfunction. Several recent studies have been reported with acute movement disorder accompanying akinetic, parkinsonian syndromes associated with bilateral basal ganglia (BG) lesions in uremic patients (2). However, to the best of our knowledge, cerebellar lesions are extremely rare, with only one case reported to date. We describe the findings from computed tomography and MRI for typical BG and cerebellar vermis lesions.

CASE REPORT

A 50-year-old man presented with dysarthria and hand trembling, that had persisted for 3 days, and uncontrollable involuntary swinging or dancing-like arm movements, which had been noted for 1 day. Examination of his medical history showed that he had a 20-year history of hypertension and chronic renal failure and had been diagnosed with diabetes mellitus 2 months previously. He had received regular hemodialysis treatment 3 times a week for 20 years.

The patient’s vital signs were stable; his blood pressure was 130/90 mm Hg; body temperature, 36.6°C; heart rate, 72 bpm; and respiration rate, 20 breaths/min. An examination of his mental status showed drowsiness, and neurologic examination showed slurred speech, negative cerebellar function test, and negative myoclonus without motor weakness. Laboratory findings revealed metabolic acidosis (pH 7.17), elevated levels of blood urea nitrogen (BUN; 30 mg/dL) and serum creatinine (Cr; 5.1 mg/dL), as well as normal blood glucose levels (124 mg/dL).

Initial brain MRI showed symmetrically increased signal intensities involving the entire BG region, and T2-weighted, and the fluid attenuated inversion recovery (FLAIR) images showed a brightly hyperintense rim extending to the internal...
and external capsules. Further, three-dimensional FLAIR images showed increased signal intensity involving the CV. Diffusion-weighted imaging (DWI) showed localized hyperintensities with reduced apparent diffusion coefficient (ADC) values in the globus pallidus (GP) on both sides, suggestive of cytotoxic edema. However, in the CV, it showed hyperintensity without signal change on ADC (Fig. 1).

Intensified daily hemodialysis combined with supportive therapy was administered, but the symptoms did not improve. The patient was in a confused state 3 days after admission, immediately after hemodialysis. At this point, laboratory findings revealed further elevated BUN (74.7 mg/dL) and Cr (10.1 mg/dL) levels and reduced blood glucose level (54 mg/dL). Follow-up brain computed tomography (CT) was performed, and the extent of the hypodense areas corresponding to the lesions shown on initial MRI seemed to have increased.

The patient suddenly presented with weakness on the right side and a semi-comatose mental status 9 days after admission.

Brain CT performed immediately showed massive intracerebral hemorrhage (ICH) in the left BG and mild ICH in the right BG (Fig. 2). The patient was transferred to a long-stay institution but showed no improvements on hospital day 28.

**DISCUSSION**

In uremia, brain involvement is a common complication (1). In cases of this complication, the imaging findings are very characteristic, and there are two types of radiological manifestations of UE: cortical (2) and BG involvement (1, 3-6).

A case of UE with reversible bilateral BG involvement was first reported by Okada et al. in a patient with chronic glomerulonephritis (4). They presented the MRI findings, which demonstrated decreased signal intensity on T1-weighted images and increased signal intensity on T2-weighted images; the lesions resolved after dialysis.

Wang and Cheng (5) presented a group of 12 patients with

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**Fig. 1.** Uremic encephalopathy in a 50-year-old man.

Three-dimensional FLAIR images (A-D) show hyperintensities in the entire BG extending to the internal and external capsules, representing the “Lentiform fork” (arrowheads) and CV (arrows). DWI (E) shows hyperintensities with reduced ADC values (F) in bilateral globus pallidus. In the CV, DWI (G) shows hyperintensities without signal change on ADC (H).

Note: ADC = apparent diffusion coefficient, BG = basal ganglia, CV = cerebellar vermis, DWI = diffusion-weighted imaging, FLAIR = fluid attenuated inversion recovery, GP = globus pallidus
end-stage renal disease who developed acute movement disorders. They also found bilateral BG lesions ranging from focal putaminal and pallidal lesions to widespread involvement of the entire BG with perifocal edema.

Although no definite pathophysiology has yet been established, metabolic acidosis and long-term hyperglycemia in uremic patients are believed to play an important role as risk factors in the development of BG lesions. The cellular functions of the BG in such uremic patients are already compromised by long-term diabetes mellitus through microangiopathic changes or energy utilization failure. Since these cells are particularly susceptible to toxic and metabolic insults, when they were further exposed to markedly elevated levels of uremic toxins, their regional cellular metabolism and/or vascular autoregulation collapse, and tissue damage and focal edema ensue (1, 5).

Bilateral symmetrical BG lesions are typically caused by diffuse systemic or metabolic disturbances. Bilateral symmetrical BG T2/FLAIR hyperintensities on MRI are non-specific and remain non-diagnostic in the absence of a clinical setting. Therefore, the differential diagnosis includes hypoxic injury, mitochondrial disease, vasculitis, carbon monoxide intoxication, hypoglycemia, extrapontine myelinolysis, Wilson’s disease, Creutzfeldt-Jakob disease, deep venous thrombosis and so on (7). Kumar and Goyal (8) assert that the “Lentiform Fork” sign is a unique radiologic picture presented by metabolic acidosis in a patient with UE. The lateral arm of the fork, constituted by the edematous external capsule, extended from the anterior end of the putamen at the rostral end of the frontal horn of the lateral ventricle to the “stem” of the fork, which is formed by the fusion of edematous external and internal capsules at the infero-posterior end of the putamen. The medial arm extended from the stem up to one-third of the medial edge where it split into two branches engulfing the GP. In our case, a bright hyperintense rim extending into the internal and external capsules surrounding the swollen BG was seen on T2-weighted and FLAIR images, representing the “Lentiform Fork.” The “Lentiform Fork” sign is not pathognomonic but a specific imaging finding in UE, so it is important with regard to differential diagnosis (8).

The main responsibility of the CV is proprioception, which is the ability to recognize the relative positioning of body parts used for movement. And, the other function of the CV is the control of muscle tone and level of force. Also, the CV affects ocular movement, although, in our case, T2-weighted and FLAIR images showed hyperintensity in the CV, and the patient presented without disturbance of ocular movement, including nystagmus, ocular motor apraxia, rapid eye movement disorder, or hypotonia.

A and ADC map showed diffusion restriction (hyperintensity on DWI and hypointensity on ADC map) only in the GP, but not in the putamen, external capsule, internal capsule and CV. This finding indicates that cytotoxic and vasogenic edema may simultaneously affect the BG and CV. Previously reported cases also showed that diffusion restriction was observed only in the GP on DWI (6, 8). Thus, DWI findings may play a key role in distinguishing UE from BG lesions caused by other conditions.

Ertl-Wagner et al. (9) suggested that BG hemorrhage may be caused by the “breakthrough” mechanism. Loss of autoregulation due to acidosis leads to severe BG edema. Correction of the metabolic derangement results in a decrease in the intracranial pressure and an increase in vascular reactivity. The initial loss and subsequent increase in reactivity probably disrupts the blood-brain barrier, which is the so-called “breakthrough” mechanism, and this leads to bilateral edema of the BG and finally to hemorrhagic transformation (9). In our case, bilateral but asymmetrical ICH in the BG (massive on the left and mild on the right, as shown by CT) developed, as was reported in 2 previous cases (9, 10). Considering that the patient’s blood pressure was relatively stable in the range of 130/80-

![Fig. 2. Follow-up brain CT performed on hospital day 9.](image)
150/90 mm Hg when he was in the hospital, we believe the hemorrhage was caused not by hypertension but by the hemorrhagic transformation due to UE.

In summary, bilateral BG lesions are the most common radiologic finding in UE, but they are not specific or pathognomonic. Because of their differentiating ability, DWI findings and the “Lentiform fork” sign are important for differential diagnosis. Lastly, clinicians and radiologists should be well-aware of the rare features of UE including CV involvement and hemorrhagic transformation.

REFERENCES