Apparently Ipsilateral Parkinsonism in a Patient with Chronic Subdural Hematoma

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There were a few exceptional cases with hemi-parkinsonism in the ipsilateral side to the lesions [ipsilateral parkinsonism (ILP)] whereas the most patients with secondary parkinsonism had causative lesions in the contralateral hemisphere.1,2 Possible mechanisms for ILP have been discussed but still remain speculative. Herein, we report a patient with ILP. Although the patient had a chronic subdural hematoma (SDH) in the ipsilateral side of parkinsonism, his parkinsonism might be more closely related to hydrocephalus.

Case

A 77-year-old man was admitted for memory loss, clumsiness of the right hand, and gait disturbance. He experienced headache and nausea 4 years before without any history of head trauma, which subsided without specific treatment. His right hand became clumsy around the episode. He had a difficulty in arising from the chair, and began to walk slowly. His symptoms were gradually aggravated afterward.

A neurologic examination showed impaired orientation and memory recall with a 17/30 on Mini-mental status examination. Motor and sensory examination was normal. Deep tendon reflex (DTR) slightly increased in the right knee. His right hand showed a dystonic posture resembling a striatal hand deformity with intermittent resting tremor. Cogwheel rigidity was present in the right arm. He showed a stooped posture and impaired postural reflex. His right leg was less agile with mild external rotation of the foot and mild shuffling.

Brain MRIs were taken four years before and at admission. Both showed a huge SDH with hydrocephalus without significant interval change. The striatum (STR) was compressed and displaced, more severe on the left side (Figure 1). The compressed thalamus and STR on the right side was displaced downwardly and medially and abutted the right midbrain (MB) (Figure 1A and B, arrows). Dorsal MB lost its convexity with widened ventricle (Figure 1C).

To investigate the nigrostriatal dopaminergic system, positron emission tomography with 18F-fluorinated N-3-fluoropropyl-2-beta-carboxymethoxy-3-beta-(4-iodophenyl) nortropane (18F-FPCIT) was done. 18F-FPCIT uptake was decreased in both STR, more severe on the left side (Figure 1). The compressed thalamus and STR on the right side was displaced downwardly and medially and abutted the right midbrain (MB) (Figure 1A and B, arrows). Dorsal MB lost its convexity with widened ventricle (Figure 1C).

Surgical treatment was waived. His cognitive function was improved with donepezil. Levodopa treatment was mildly effective for his parkinsonism.
Discussion

Subdural hematoma could be hardly considered as the primary diagnosis because of protracted course, slow progression, and the scarce signs of increased intracranial pressure. However, fixed dystonia, increased DTR, and the spastic nature of his gait were suggestive of secondary parkinsonism.

The decreased uptake of $^{18}$F-FPCIT in STR could be result-

![Figure 1](image1.png)

**Figure 1.** A brain magnetic resonance imaging shows a huge subdural hematoma with calcification over the right cerebral convexity with prominent hydrocephalus (A: axial, B: coronal, C: sagittal). The compressed right brain is displaced abutting the midbrain (arrow), while the left side of midbrain retains its normal contour.

![Figure 2](image2.png)

**Figure 2.** Positron emission tomography with $^{18}$F-fluorinated N-3-fluoropropyl-2-beta-carboxymethoxy-3-beta-(4-iodophenyl) nortropane ($^{18}$F-FPCIT) shows the decreased uptake of $^{18}$F-FPCIT in both striatum, which is more severe on the left side (A: axial, B: coronal). In the midbrain, $^{18}$F-FPCIT uptake is well preserved on the left side (arrow). The $^{18}$F-FPCIT uptake of the right midbrain is preserved but appears as dispersed in the axial image (arrowhead) mixed with that of overlying thalamus (asterisk).
ed from direct compression of STR or the disruption of the nigrostriatal pathways. As the caudate nuclei retained their convexity, disrupted nigrostriatal pathways may be more likely to be responsible for abnormal \(^{18}\)F-FPCIT uptake. The loss of dorsal convexity of MB would be compatible with the compromised nigrostriatal pathway by hydrocephalus. The anterior-to-posterior gradient of dopamine transporter scan could be a feature of pre-synaptic damage.

In our patient, asymmetric involvement of the contralateral nigrostriatal pathway may result in apparent ILP. It remains enigmatic the absence of parkinsonism in the contralateral side to SDH, in spite of STR abnormality with decreased \(^{18}\)F-FP-CIT uptake.

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