CASE REPORT

Exofocal Anterograde Transsynaptic Neuronal Death in the Globus Pallidus: Two Case Reports

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Background Exofocal neuronal death in the substantia nigra (SN) is a well-known form of anterograde transsynaptic cell death. Exofocal neuronal death could theoretically also occur in the globus pallidus (GP) after striatal injury.

Case Report Case 1. A 70-year-old woman visited the emergency room because of decreased mentality. On admission, blood-gas analysis indicated that her oxygen tension was 69.1 mm Hg. The caudate nucleus, putamen, and temporoooccipital cortex on both sides of the brain exhibited high-intensity diffusion-weighted magnetic resonance imaging (MRI) signals. At 10 days after admission, new high-intensity signals had developed in the SN and GP on both sides. Case 2. A 48-year-old man visited the emergency room because of right-sided weakness. Lesions were noted in the left caudate nucleus and putamen. At 4 days after admission, newly developed high-intensity MRI signals were observed in the left SN and GP.

Conclusions Exofocal neuronal death can occur in the GP as well as in the SN; these findings need to be clearly distinguished from those of recurrent ischemic injuries, such as recurrent stroke.

Key Words transsynaptic, degeneration, globus pallidus, substantia nigra.
Case 2
A 40-year-old man visited the emergency room because of right-sided weakness. Medical Research Council grade 4 weakness was noted on the right side upon neurological examination. No sensory signs were observed. Diffusion-weighted MRI of the brain performed on the second day of admission revealed high-intensity signal lesions in the left caudate nucleus and putamen. Although there was no change in the symptoms, a follow-up MRI was performed 4 days after admission; this disclosed newly developed high-intensity signals in the left SN and GP (Fig. 2). The patient’s weakness improved without development of any extrapyramidal symptoms.

Discussion
Pallidonigral damage after ischemia has been reported previously, but such reports are not supported by early MRI findings; therefore, the authors of those reports concluded that the pallidonigral damage was the result of the primary ischemic injury. In the cases described herein, both patients underwent both early and follow-up MRI, and changes in signal intensity in the SN and GP were seen only in later MRI scans. Therefore, we believe that the GP lesions were delayed changes, because the high-intensity signals in the GP and SN were not observed during the initial MRI. The timeline of the change in signal intensity in the GP was similar to that in the SN, and both can be attributed to exofocal postischemic neuronal death.

Exofocal postischemic neuronal death in the SN has been studied extensively in rat models; however, these studies have not revealed secondary changes to the rat GP, and therefore many investigators have inferred that GP damage in humans is primary in origin. In nonprimates the GPi is simply referred to as the GP. The nonprimate GPi is not equivalent to the primate GPi; it is small and is buried in the fibers of the internal capsule to form the entopeduncular nucleus. Thus, it is not surprising that exofocal neuronal damage cannot be observed in the rat GPi after striatal damage. Furthermore, the middle cerebral artery occlusion method was used to induce striatal damage in the rat model, which may have primarily damaged the GPi, thereby explaining why exofocal neuronal damage was not observed in the GPi.

Degeneration of the ipsilateral SN in humans following infarction in the striatum has been reported, but those authors did not report secondary changes in the GPi. However, all of the patients in that study underwent MRI only once, and hence the GPi injury might have been considered a primary infarction. Injection of a GABAergic antagonist into the GPi in monkeys causes dyskinesia in the contralateral forelimb. Similarly, isolated GP lesions cause motor disturbances. However, no movement symptoms developed in our two cases: in case 1 we believe that this was due to the patient’s stuporous mentality.

The exact mechanism underlying preservation of the GPe is unknown. One possible mechanism is that although both the...
GPe and GPi receive inhibitory inputs from the striatum, the GPi receives inputs from substance-P- and dynorphin-containing neurons (direct pathway), while the GPe receives both those direct-pathway neurons together with additional inputs from enkephalin-containing neurons (indirect pathway).10

We performed follow-up MRI despite unchanged symptoms in our two cases because we predicted that there would be changes to the MRI findings. Marked astrocyte swelling developed on day 4 following injury in rats.1 Thus, in our case 2, the follow-up MRI was performed 4 days after the symptom onset. In case 1, the follow-up MRI was delayed until 10 days after symptom onset because of the general condition of the patient. Exofocal anterograde transsynaptic neuronal death in the GP was more prominent in case 1. These observations suggest that exofocal neuronal death develops later in humans than in rats.

The GPi can be subjected to delayed damage via the mechanism of anterograde transsynaptic cell death. However, it is important that this is distinguished from recurrent ischemic injuries such as stroke.

Conflicts of Interest

The authors have no financial conflicts of interest.

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