SHORT REPORT

Creutzfeldt-Jakob disease with congophilic kuru plaques: CT and pathological findings of the cerebral white matter

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Abstract
In a patient whose Creutzfeldt-Jakob disease with congophilic kuru plaques that was proved at necropsy, the early brain CT showed low-density areas in the cerebral white matter before cortical atrophy and ventricular enlargement became apparent. Subsequently, there occurred diffuse white matter lucency and severe brain atrophy. At necropsy, there was severe white matter destruction which was more prominent than cortical neuronal loss. Serial CT scans were of great value for demonstrating the early and predominant changes in the cerebral white matter.

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Congophilic kuru plaques rarely occur in the brain in sporadic cases of Creutzfeldt-Jakob disease (CJD). The disease CJD with congophilic kuru plaques (CJD-KP), is clinically characterised by an unusually long duration of illness for CJD and ataxia usually preceding dementia, but there is considerable clinical variability. Neuropathological features other than kuru plaques include variable degrees of spongiform changes in the cerebral cortex and frequent involvement of the cerebral white matter. Despite the unique clinicopathological features, there has been very little information about the neuroradiological findings. We report here a necropsy case of CJD-KP, with special reference to the cerebral white-matter changes revealed by CT.

Case report
A 60 year old woman was admitted to our hospital in January 1984 because of impaired memory, disorientation to time, numbness in both legs and truncal ataxia for the previous five months. She had a nine year history of diabetes mellitus which had been well controlled. One month before admission, she showed mental changes with delusions and hallucinations. There was no family history of neurological disease.

Neurological examination on admission revealed a limitation of upward gaze, perceptive deafness, irregular involuntary movements of the tongue and soft palate, areflexia of the lower limbs and truncal ataxia without limb kinetic ataxia. Spinal fluid examination showed a slight elevation of protein (59 mg/dl) without pleocytosis. Five months after admission, she was confined to bed and her condition rapidly deteriorated. In early September, she was in the state of akinetic mutism and showed muscle rigidity, areflexia in the four limbs, myoclonus in the face and four limbs, and a startled response to sounds. An EEG showed a background activity of 3–5 Hz with intermittent synchronous 100 μV θ activity in the fronto-centro-parietal region. The clinical pictures remained relatively unmodified over the next three years; the intensity of myoclonus progressively decreased; the intermittent discharges of the EEG gradually disappeared to be replaced by a slow monomorphous activity by July 1985. On 8 December 1987, she died of respiratory failure. The duration of the disease was 41 months.

The initial CT scan performed in February of 1984 revealed a bilateral pallidal calcification and low attenuation in the cerebral white matter without cortical atrophy and ventricular dilatation (fig 1A, B). The low density of the white matter was accentuated at the cerebral subcortex. A second CT scan obtained six months after the initial CT scan, indicated moderate enlargement of the ventricles with diffuse white-matter lucency (leuko-araiosis). A dramatic enlargement of the ventricles and widening of the cerebral and cerebellar sulci developed within four months after the state of akinetic mutism appeared (fig 1C, D).

At necropsy, focal pneumonia and mild arteriosclerosis were found. The brain weighed 805 grams and was markedly atrophic with cystic softening of the white matter at the cerebral subcortex (fig 2A). There was marked atrophy of the cerebellum and brainstem. Microscopically, there was severe spongy degeneration and less marked diffuse neuronal loss of the cerebral cortex, thalamus and basal ganglia. Diffuse loss of myelinated nerve fibres accompanied with proliferation of hypertrophic astrocytes and fat-laden macrophages were seen throughout the cerebral white matter. At the subcortical white matter of the cerebrum, there were circumscribed necrotic foci. In the cerebellum, both Purkinje and granule cells were diminished. Numerous congophilic kuru like plaques were present in the cerebellar cortex (fig 2B), and to a lesser extent in the cerebral cortex.

Experimental transmission of the disease to mice through inoculation of the brain homogenates was successful. To distinguish the present case from Gerstmann-Sträussler syn-
nine-to-valine change at codon 129 was carried heterozygously.

Discussion
The patient showed clinical features of CJD, but the duration of the illness was longer than that of typical CJD. Neuropathological, molecular genetic and transmission studies confirmed the diagnosis of CJD-KP.

The most prominent CT finding in our patient was the appearance of low-density areas in the cerebral white matter before cortical and central atrophy became apparent. Subsequent enlargement of ventricles progressed more rapidly than the cerebral cortical atrophy, suggesting that the involvement of white matter was more severe than the involvement of the cortex. With regard to the CT finding, previous reports of CJD-KP mention only cortical atrophy and ventricular dilatation of the brain, but no white-matter changes have been described. On the other hand, the CT imagings of CJD without kuru plaques show bilateral cortical atrophy and no apparent white-matter changes. The white-matter abnormalities, if present, develop during the final stage. Recent study with MRI also showed increased signal intensity in the basal ganglia, thalamus and cerebral cortex without significant change in the white matter. The discrepancy in the occurrence of white-matter changes suggests the wide spectrum of pathology in CJD patients.

The neuropathological findings of this patient closely matched the CT images, showing diffuse marked degeneration with necrotic foci in the cerebral white matter. Such white-matter lesions seem to be an exceptional finding in CJD, because the main affected area in CJD has been regarded as cortical and subcortical grey matter. However, extensive degeneration of cerebral white matter which cannot be explained simply as secondary to cortical degeneration has been described in the patients with CJD-KP and a special type of CJD without KP. The outstanding white-matter lesions observed in this patient are not only due to the long clinical course, since the

Figure 1  A, B) CT scans (February 1984) show a bilateral pallidal calcification and low attenuation in the cerebral white matter without cortical atrophy and ventricular dilatation. The low-density is accentuated at the cerebral sub cortex (arrow heads). C, D) CT (January 1985) shows remarkable enlargement of the lateral ventricles with diffuse white matter lucency.

drome (GSS), polymorphic changes in the prion protein gene were examined using the frozen brain tissue. As a result, the patient had neither deletion and insertion nor variant substitution at codons 102, 117, 200 previously reported to be specific to GSS. However, a common polymorphism, methio-

Figure 2  A) Coronal section of the cerebral hemispheres showing overall necrosis with cystic softening (arrow heads) of the white matter. B) Photomicrograph of the cerebellar cortex showing kuru-like amyloid plaques (arrow heads). H-E staining; 206 x.
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