SHORT REPORT

Atypical dopa responsive parkinsonism in a patient with megalencephaly, midbrain Lewy body disease, and some pathological features of Hallervorden-Spatz disease

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Abstract
A 38 year old patient with megalencephaly, mental retardation, and lifelong tremor developed levodopa responsive parkinsonism in his mid-30s followed by the appearance of dyskinesia, motor fluctuations, hallucinations, and dementia. Brain MRI showed, as well as other changes, iron deposition in the globus pallidus, substantia nigra, and the pulvinar of the thalamus. Postmortem examination disclosed depigmentation of the substantia nigra pars compacta with neuronal loss, gliosis, and Lewy body formation. Axonal dystrophic spheroids, neuronal loss, calcification, and iron deposition were found in the substantia nigra pars reticulata. Less severe changes without neuronal loss were seen in the globus pallidus. This combination of megalencephaly with neuroaxonal changes predominantly in the pars reticulata and Lewy body degeneration isolated to the substantia nigra pars compacta has not been previously reported.

Case report
A 38 year old left handed man with congenital macrocephaly and mild mental retardation was referred to us at 35 years of age because of progressive and disabling tremor which had been present since childhood. The tremor predominantly involved his left hand and leg and impaired his ability to write and perform other activities of daily living. He denied slowness in performing activities and other motor impairments. There had been no exposure to neuroleptic medications and no history of encephalitis. His family history was significant in that his mother was diagnosed with "normal pressure hydrocephalus". In her 60s she developed parkinsonism and a few years before her death at the age of 70 she became demented.

No necropsy was performed.

His initial examination disclosed a masked facies, slight hypophonia, and normal eye movements. No Kayser-Fleischer rings were seen. A resting tremor with persistence with posture was prominent in the left arm and was present to a lesser degree in the left leg. A fine action tremor was present in the right arm. Moderate cogwheel rigidity and bradykinesia were found on his left side with mild involvement on the right. There was no dysmetria or ataxia. On ambulation there was decreased arm swing and he had normal postural stability.

Brain MRI T2 weighted images disclosed hypointensities, consistent with iron deposition, in the globus pallidus, substantia nigra, and the pulvinar of the thalamus. T2 weighted hyperintensities were seen lateral to the occipital horns of the lateral ventricles, and to a lesser extent there was involvement of the corticospinal pathways in the internal capsule and in the midbrain cerebral peduncles. No hydrocephalus was noted. 6-[18F] fluoro-L-dopa (F-dopa) PET showed normal caudate uptake and reduced bilateral putamenal uptake more pronounced on the right than the left.

Various medical and surgical treatments were employed in the hope of alleviating his disabling tremor. Propanolol, titrated to 240 mg per day, failed to change the tremor and it "worsened" on 300/75 mg of levodopa/carbidopa per day. Trials of primidone and
levodopa was increased to 1600 mg per day and 50 mg of trihexyphenidyl was added the resting tremor was eliminated for variable periods. With time he developed typical levodopa induced dyskinesiae of the face, trunk, and limbs with greater involvement on the right side of his body. He also developed a fluctuating response to medication and was immobile with a prominent tremor when he was in the “off” state. Severe postural instability eventually resulted in his becoming confined to a wheelchair. This was accompanied by cognitive changes including altered memory and concentration; his mini mental state examination score was 19/30. He developed frequent visual hallucinations and because of severe “off” periods he required continued use of high doses of medication (1300 mg levodopa and 50 mg of trihexyphenidyl per day). As a result he was admitted to hospital and clozapine was titrated to 37.5 mg per day, but he still had persistent hallucinations. He also developed fevers without a change in his white cell count and negative blood cultures. Empirically, antibiotics were started for a presumptive urinary tract infection. On the day of his planned discharge he was found in bed asystolic and failed to respond to resuscitation.

**Pathology**

A complete general necropsy disclosed extensive pulmonary aspiration of vegetative material.

Neuropathological examination was performed on the brain, two peripheral nerves, and muscle specimens using routine stains, immunohistochemistry (neurofilament, β-4 amyloid, synaptophysin, and ubiquitin), and special histological stains (Congo red, Bielschowsky silver, Perl’s, and Von Kossa). The table summarises the pathological findings.

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**Figure 1** MRI of brain: axial section T2 weighted images at the level of the midbrain/diencephalic junction (A) and the level of the midbrain (B). Note the prominent hypointensity of the substantia nigra, which is much darker than the red nucleus (A). Bilateral hyperintensities are present in the region of the corticospinal tract pathways in the cerebral peduncles. (C) MRI of brain: axial section T2 weighted image at the level of the midthalamus. The globus pallidus and the pulvinar of the thalamus are hypointense. Hyperintensities are seen in the internal capsules bilaterally, and lateral to the occipital horns of the lateral ventricular system. Postoperative changes from two prior thalamotomies are visible (hypointense lesion) in the right thalamus.

Trihexyphenidyl also failed to diminish the tremor. He subsequently underwent a right thalamotomy which completely resolved his tremor for six days but then it returned to its preoperative severity. Four months later, a second right thalamotomy did not change the tremor or his rigidity but resulted in left hemiparesis and hemineglect.

Brain MRI performed 20 months after the initial study showed changes similar to the first MRI as well as the right thalamotomy lesion (fig 1A-C).

A second trial with levodopa/benserazide greatly improved his motor function. When...
The brain weight was increased at 2180 g. The brain was enlarged diffusely in a symmetric manner indicative of megalencephaly, and was without associated polymicrogyria. Coronal sections of the left hemisphere showed an incidental grey matter heterotopia in the white matter and a left occipital capillary telangiectasia. There was subtle histological evidence of cortical microdysplasia without other cortical pathology.

The striatum and brainstem structures, other than the substantia nigra, did not show any gross discoloration. The substantia nigra pars compacta, particularly the lateral aspects (fig 2A), showed asymmetric (right > left) depigmentation due to moderate depletion of large neurons containing neuromelanin. This was accompanied by gliosis and prominent Lewy bodies in remaining neurons (fig 2B).

The locus ceruleus, nucleus basalis of Meynert, and dorsal motor nucleus of the vagus were normal. Axonal swellings (dystrophic axonal spheroids) were present primarily in the substantia nigra pars reticulata (fig 2C) and were accompanied by scattered microscopic brown pigment deposition due to iron deposition and calcification. The pallidum was involved to a lesser degree and some of the deposits were present in a perivascular location (fig 2D). The other brainstem structures (pons, medulla), cerebellum, spinal cord, muscles (psoas and gastrocnemius), and peripheral nerve specimens (sciatic and sural) were normal.

**Discussion**

This patient's clinical course was suggestive of primary nigral neuronal loss because of his dopa responsive parkinsonism and subsequent development of motor fluctuations and levodopa induced dyskinesias. However, the MRI showed prominent hypointensity of the pallidum and the substantia nigra on T2 weighted images, which suggested a diagnosis other than idiopathic Parkinson's disease.

The pathological findings reaffirmed that this was not typical Parkinson's disease. To our knowledge, no case of megalencephaly has been reported with either dystrophic axons or...
Lewy bodies. Rather, patients with either or both of these features usually have small, atrophic brains. Thus the association between the megalencephaly (idiopathic?) and these “neurodegenerative” changes in our patient is unique. It is impossible to determine whether there was an aetiological or pathogenetic relation between these features or whether the combination was a mere coincidence.

Megalencephaly is defined as a brain that weighs above the 98th percentile and arises due to “metabolic” or “anatomical” causes. This patient had no evidence of a metabolic cause such as a gangliosidosis, a mucopolysaccharidosis, or a sulphatidosis. Anatomical megalencephaly is attributed to an increase in the number or the size of cells and can be associated with neurological disorders such as Alexander’s and Cunnan’s disease. Inherited (autosomal dominant or recessive) and sporadic (idiopathic) megalencephaly are also included in the anatomical causes and these are usually not associated with other neurological syndromes. As best it could be determined that there was no one else in the family with a large head. The focal cortical dysplasia and focal heterotopia have been reported with megalencephaly. However, they often coexist with evidence of severe neuronal migrational abnormalities—that is, pachygyria or polymicrogyria—which were not present in our patient. The cortical dysplasia does not account for the megalencephaly itself, but may represent the basis of this patient’s mental retardation.

Lewy bodies are eosinophilic cytoplasmic inclusions composed partially of cytoskeletal protein with a high neurofilament content and polyubiquitin chains. Dystrophic axons, which are also ubiquitin-reactive, are composed of accumulated cytosolic and cytoskeletal components that result in the swelling of distal or terminal axons (“spheroids”) probably due to defective axonal transport. The presence of Lewy bodies in the substantia nigra accompanied by gliosis and neuronal loss is consistent with a diagnosis of idiopathic Parkinson’s disease although this inclusion may occur in other cerebral disorders. The nigral pathology suggestive of Parkinson’s disease also correlates with many of the patient’s clinical features. However, Lewy bodies were not found in structures that are often affected in Parkinson’s disease (for example, hypothalamus, septal nuclei, locus ceruleus, dorsal motor nucleus of the vagus), or in diffuse Lewy body disease.

Dystrophic axons have been reported in Parkinson’s disease but not to the extent or severity found in our case. More abundant dystrophic axons are typically associated with neuroaxonal dystrophies (Seitelberger’s disease) and Hallervorden-Spatz disease, although in these disorders, dystrophic axons are usually present diffusely. In Hallervorden-Spatz disease, dystrophic axons are typically found in the spinal cord, the dorsal medullary region, the cuneate and gracile nuclei, the area postrema, the cerebral cortex, the pars reticulata, and the most severe changes are in the pallidum, particularly the internal segment. In our case dystrophic axons primarily involved the pars reticulata with a few scattered dystrophic axons found in the pallidum. The focality of dystrophic axons suggests a different sequence or pathophysiological process than occurs in typical Hallervorden-Spatz disease or neuroaxonal dystrophy, possibly one more closely linked to the pathology in the pars compacta.

In Hallervorden-Spatz disease, the coexistent pathology of the reticulata and the globus pallidus interna is not unexpected as both structures are embryologically related and any genetic or metabolic defect would presumably exist in their primordial cells. However, Kessler et al. have reported a case with selective pallidal involvement, which they include in the more general category of “Hallervorden-Spatz syndrome”. This focal pallidal pathology is, in a sense, enantiomeric to that of our patient with more “selective” reticulata findings.

Although the nature of these combined features in our case is unique, the concurrence of the more typical Hallervorden-Spatz disease pathology and Lewy bodies have been described. In these patients, Lewy bodies have either been limited to the pars compacta or have been more widespread as in diffuse Lewy body disease. In some cases Lewy bodies were accompanied by neuronal loss and gliosis in the pars compacta, features which are not a part of typical Hallervorden-Spatz disease. However, all had accompanying Hallervorden-Spatz disease pathology (both dystrophic axons and mineralisation) involving both the pars reticulata and pallidum, the pallidum typically being most severely affected. None have had pathological changes similar to our patient. Many have had parkinsonism, some presenting during childhood or early adulthood. Some were treated with levodopa and responded whereas others failed to benefit. Theoretically, the extent of the pallidal changes may have differed in these subgroups at the time of levodopa trial, thereby explaining the different levodopa responsiveness. We think that appreciable degeneration of the pars compacta combined with the lack of pallidal neuronal loss and gliosis in our patient explains why there was a response to levodopa.

In summary, our patient presents the unique combination of nigral degeneration with Lewy bodies and dystrophic axons, predominantly involving the substantia nigra, and megalencephaly. Further careful observation and research will be necessary to determine why this combination of cytoskeletal derangements occurs and its relation to the more typical forms of Parkinson’s disease and Hallervorden-Spatz disease. The presence of megalencephaly is also of considerable interest, but is of uncertain relevance.

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Parkinsonism, megalencephaly, Lewy bodies, and Hallervorden-Spatz disease