Bilateral striatal necrosis, dystonia and optic atrophy in two siblings

V Leuzzi, E Bertini, A M De Negri, M Gallucci, B Garavaglia

Abstract
Two siblings developed a neurological disorder in the first decade characterised by generalised dystonia, hypokinesia, and subacute visual loss. CT and serial MRI examinations showed bilateral lesions of the striatum, mainly in the putamen. The classification of these patients is discussed in relation to infantile bilateral striatal necrosis (IBSN), Leigh's disease, and Leber's optic neuropathy. The literature shows a clinical and aetio-pathogenetic overlap between these syndromes. In our cases parental consanguinity and the involvement of a single generation suggest a new clinical condition with autosomal recessive transmission.

With the development of non-invasive diagnostic techniques, there has been an increase in reported cases of a progressive dystonic syndrome beginning in childhood and associated with selective lesions of the basal ganglia. Improvement of CNS investigations has brought new criteria for the differential diagnosis between idiopathic and secondary dystonias. We report two siblings with dystonia, hypokinesia, symmetrical striatal necrosis, and visual failure. These cases, together with others previously reported, may represent a new clinical entity.

Patients
The parents were first cousins. The paternal grandfather had suffered from left hemiparesis, which started at the age of 8 years during an unspecified exanthematic illness. The couple had four children, and the two patients were the product of the second and third pregnancy respectively (the latter was a dizygotic twin pregnancy).

Case 1
This 20 year old woman had had a normal birth and development. At the age of 6 years she contracted measles. Four days after exanthematic eruption she had transient (a few seconds) amaurosis which disappeared spontaneously without further effects. Six months later she experienced a painless rapid bilateral deterioration in visual acuity. At the age of 7 she began to invert and plantar flex the left foot with dorsiflexion of the big toe when walking; dystonia then slowly progressed to involve her left arm. At the age of 10, she underwent left Achilles tendon and extensor hallucis longus tendon lengthening procedures. At the age of 11, the left dystonic hemiparesis worsened: the hip and the knee were flexed, the foot was equinovarus, the toes plantar flexed. The arm exhibited athetoid postures and movements. Both optic discs were pale. Visual acuity was correctable to VR 1/10, VL 10/10. A CT scan showed an area of low attenuation in the right putamen and in the head of the right caudate nucleus with slight enlargement of the homolateral ventricle.

The patient was re-examined in April 1987 at the age of 18. In addition to dystonic postures and movements of the left limbs, there was distinct generalised hypokinesia and bradykinesia. Both arms presented high frequency distal postural tremor; there was a plastic rigidity of limbs and tendon reflexes of the left leg were weak. The left kneecap was dislocated upward because of the shortening of the rectus femoris, which showed regular, almost rhythmic (2-3 Hz), myoclonic jerks either in the resting condition or during leg movement. She had an external squint. Ophthalmoscopy revealed a bilateral optic atrophy; the vessels were unusually tortuous. On examination, the visual acuity was correctable to VR 3/50, VL 2/10; the study of visual fields (Goldmann) showed an absolute centrocecal scotoma with a small area of normal central vision on the right eye and an absolute centrocecal scotoma, multiple absolute paracentral, and peripheral scotomas on the left eye. The right pupil was larger than the left, but light and near reflexes were normal. Visual evoked responses to flash stimuli showed normal voltage and latency; no cortical responses were evoked by patterned stimuli. Brain CT and MR images showed bilateral putaminal lesions and involvement of the head of the right caudate nucleus (fig 1). Fasting resting blood lactate concentration was normal, but blood pyruvate and alanine concentrations were elevated (8.8 mg/l, normal range 3-6-5.9 mg/l, and 1061.3 μM/l, normal range 273-449 μM/l, respectively). She was re-examined in October 1989. Visual acuity had worsened, and she complained of episodes of sudden transient amaurosis for about 10 minutes several times a day. Neurological, neuro-ophthalmological, and neuroradiological (MRI) examinations were unchanged. Her mental function remained unaffected.

Case 2
This 11 year old boy had been born preterm (eight months, weight 2550 g) after an uncomplicated twin pregnancy and delivery. Growth and development were normal until the age of 7.


Correspondence to: Dr Leuzzi, Istituto di Neuropsichiatria Infantile, Via dei Sabelli, 108. 00185 Roma, Italy. Received 10 September 1990 and in final revised form 1 March 1991. Accepted 8 March 1991.
years and 10 months, when his parents noticed that his right arm irregularly adopted bizarre postures (extension and adduction of the arm, pronation of the forearm and hand). Despite abnormal postures, the child could perform skilful movements (such as writing). When 9 years old, he developed an increasingly severe rigidity of the left leg which disturbed walking. The homolateral upper limb was flexed with a clenched fist. He became anorexic and asthenic. On examination (in April 1987), at the age of 9 years and 5 months, he was a pale boy with normal somatic growth, poor muscle development, and dystonic movements and postures of the limbs (left more than right). The most considerable neurological disorders of the arms, however, were hypokinesia and bradykinesia, which were the cause of clumsiness and inadequacy in intentional skilled movements. There was plastic rigidity and mild weakness (left more than right). Tendon reflexes were brisk and polyphasic with bilateral Babinski’s signs. Speech was quiet, slow, monotonous, and somewhat slurred. Sensation was intact. Mental function remained unaffected.

Both optic discs showed temporal pallor, and the retinal vessels were narrowed. On examination, the visual acuity was correctable to VR 5/10, VL 8/10; perimetry showed a distinct concentric constriction of visual fields and enhanced blind spot. Colour vision was impaired in the red-green region (Ishihara test). Non-sustained horizontal nystagmus to the right was present. Visual evoked responses to flash stimuli showed normal voltage and latency; pattern reversal VEPs showed low amplitude, delayed P2 component (140 ms) in O2, and a normal response in O1. A brain CT scan and MRI examination showed bilateral selective lesions of the putamina (fig 2). His fasting resting blood lactate concentration was normal, but on four separate occasions the blood pyruvate concentration was elevated one and a half to three times the normal values. Pyruvate oxidation and respiratory chain function of cultured fibroblasts were normal. Histological and ultrastructural studies of left deltoid muscle biopsy specimens showed no abnormalities.

The patient was re-examined when 10 years old. He walked with more difficulty. Intentional movements of the arms were slow and clumsy; his speech was slurred and writing much slower. There was generalised muscular hypotrophy and weakness. Visual acuity was correctable to VR 4/10, VL 6/10. A second brain MRI examination was unchanged. He underwent clinical assessments every six months until June 1990. Neurological, neuro-ophthalmological, and neuroradiological (MRI) examinations revealed no change from his previous state. He had no intellectual impairment or learning difficulties. His somatic development, however, was arrested (height below 10% and weight below 3%).

The analysis of mitochrondrial DNA obtained by PCR-amplification on total DNA from muscle specimens showed a normal Sfa NI digestion of the 316 bp PCR product, that includes the nt 11778 mutation associated with Leber’s hereditary optic neuropathy3 (Dr C Gellera, Istituto Neurologico “Besta” Milano). In both patients values were normal for the following analyses: standard blood and urinary studies, VDRL, serum copper and caeruloplasmin, 24 hour urinary copper, serum lipid, serum protein electrophoresis, plasma and urinary aminoacid chromatography, urinary mucopolysaccharides, urinary organic acids, urinary aroylsulfatase-A, and an assay of lysosomal enzymes in white blood cells and fibroblasts in case 2. EMG and nerve conduction studies, EEG, BAEP, SEP, ERG, and slit lamp examination for Kayser-Fleischer ring were also normal. The twin of case 2 has been surveyed for four years. He is an intelligent, normal child with no motor or neuro-ophthalmological disability. Three brain MRI examinations performed at intervals of six months gave normal results.
Discussion
The two siblings reported here show an uncommon clinical disorder characterised by dystonic movements and postures associated with generalised hypokinesia, subacute visual loss of varying degrees, and bilateral and selective lesions of the striatum detected by neuroradiological techniques (CT and MRI). Mental deterioration was absent. We suggest that three clinico-pathological conditions should be considered; infantile bilateral striatal necrosis (IBSN), Leigh's disease, and Leber's optic neuropathy (LHON). With the term of infantile bilateral striatal necrosis, Friede classified a group of infantile encephalopathies in which the main pathological finding was bilateral, symmetrical, spongy degeneration of putamina, caudate nuclei, and, less commonly, pallidum. Modern imaging techniques such as brain CT and MRI can detect basal ganglia abnormalities. Though they cannot substitute neuropathological examination, they do enable new cases of IBSN to be observed in vivo, and serial studies can assess the evolution of the disease.

Cases reviewed by Friede, as well as those more recently described, are clinically heterogeneous. Age at onset varies from infancy to adulthood; some cases present a progressive disorder, others an acute or subacute neurological disorder, sometimes preceded by an acute febrile illness. Clinical features, with various combinations, include dystonia, hypokinesia, spasticity, bilateral optic atrophy, abnormal eye movements, seizures, mental retardation or mental deterioration, disturbed behaviour, and failure to thrive. The disorder has a poor prognosis: death usually occurs, but at intervals from a few weeks to some years after onset of symptoms. Cases with complete or partial remission, however, have also been reported. All previously reported familial cases were in one generation or the same sibship, suggesting autosomal recessive transmission. The family described by Barghavan-Farkas et al is the only published example of parental consanguinity we know of.

Confining the discussion to slowly progressive cases, those described by Miyoshi et al and Peenito et al differ from ours in that mental retardation was prominent in the former and visual function was unaffected in all. Both clinical and neuroradiological findings of our patients are similar to the clinical observations of Marsden et al, who described seven patients in two families. The patients exhibited a slowly progressive generalised dystonic syndrome, which was typically associated with subacute visual loss and striking bilateral symmetrical lucencies on CT, particularly affecting the putamen. The patients showed no obvious mental deterioration. In one case the visual deficit was isolated and in another a diagnosis of Leigh's disease was suggested. Symmetrical areas of low attenuation in the basal ganglia on CT scan, dystonic movements, optic atrophy, and increase of blood pyruvate and alanine are often observed in patients with Leigh's disease.

This constitutes a heterogeneous group of disorders associated with a disturbance in mitochondrial energy metabolism. The clinical features are variable and are related to the multiplicity and site of the neuropathological lesions. The coexistence of multiple neurological deficits is also found in late onset cases which usually progress more slowly. Recently, Van Erven et al reported three sporadic patients with a slowly progressive condition like Leigh's disease in which hypokinesia and rigidity were the most prominent neurological disorders. Nevertheless, other signs and symptoms in these cases suggest the multi-systemic aspect typical of Leigh's disease: deterioration during intercurrent infections followed by a slow recovery, exercise intolerance, poor somatic growth, abrupt changes in respiratory and cardiac rates,
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strabismus, nystagmus, dysarthria, ataxia, tremor, and dysmetria.

Neuroradiological data derived from MRI, particularly if serial studies are performed, offer a valuable tool for differentiating in vivo selective lesions of the striate nuclei, as in IBSN, from the multifocal involvement of the brain seen in Leigh's disease.20-29 In the absence of a neuropathological report, we believe that Leigh's disease is unlikely in our patients as no damage was present outside the striatum (as detected repeatedly by CT and MRI) and no clinical signs of brainstem, cerebellar, and peripheral nerve involvement were ever noted. Moreover, we did not find any pyruvate oxidation defects and respiratory chain abnormalities in fibroblasts of case 2.

In our patients, subacute painless visual failure suggests the diagnosis of Leber's optic neuropathy (LHON). This is a maternally inherited form of optic nerve atrophy associated with neurological and psychiatric symptoms in a high percentage of cases, both in affected patients and in their collateral.30 A few reports exist in which LHON is associated with IBSN. Novotny et al31 studied a single large pedigree (79 members in five generations) of patients affected by Leber's disease. Eight members had neuroretinopathy, 14 had a progressive, generalised dystonia attributed to striatal degeneration, but only one had both disorders. In six out of 14 neurologically affected patients CT examination demonstrated low-density in the putamen as the earliest finding. Caudate nuclei were affected later, and one subject had low density lesions in the centromedian nucleus of both thalami. Recently, Wallace et al32 and Singh et al33 identified a mitochondrial DNA point mutation (at nt 11778) that correlated with Leber's disease in nine out of 11 tested pedigrees. The authors did not find the same mutation, however, in cases of LHON plus IBSN, which may be the result of the presence of a mixture of mutant and normal mitochondrial DNA sequences (heteroplasmasy).34

Our two patients were not seen in the acute phase of the optic neuropathy, when the peripapillary microangiopathy gives a pathognomonic appearance of Leber's disease.35 The clinical picture, some ophthalmoscopic findings (arterial attenuation and persistence of tortuous vessels in case 1), andvisual field deficits may suggest this disease. Although the restriction-fragment-length polymorphism negativity after digestion of mitochondrial DNA with Spha NI enzyme might accord with the hypothesis of different genetic origins of classical LHON and LHON associated with IBSN, the parental consanguinity and the apparent involvement of a single generation suggest a new clinical condition with an autosomal recessive transmission.