Short report

Sandhoff disease mimicking adult-onset bulbospinal neuronopathy

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SUMMARY A 32 year old male is described with an onset of upper limb postural tremor in adolescence followed by muscle cramps. Progressive proximal amyotrophy and weakness in the limbs developed late in the third decade. Examination disclosed, in addition, bilateral facial weakness and mild dysarthria. Enzyme studies revealed hexosaminidase A and B deficiency, indicating a diagnosis of Sandhoff disease. Intra-axonal membranocyttoplasmic bodies were present in a rectal biopsy. The presentation, which resembled that of X-linked bulbospinal neuronopathy, widens the clinical spectrum for disorders related to G\textsubscript{M2} gangliosidosis.

The G\textsubscript{M2} gangliosidoses result from a deficiency either of the hexosaminidase enzyme which hydrolyses the terminal N-acetylgalactosamine from the ganglioside G\textsubscript{M2}, or of the activator protein for the enzyme. Two major isoenzymes of hexosaminidase are found in human tissues, namely hexosaminidase A and hexosaminidase B (hex A and hex B). Hex A is composed of \(\alpha\) and \(\beta\) subunits coded for on chromosomes 15 and 5 respectively whereas hex B is composed only of \(\beta\) subunits. Thus a mutation at the \(\alpha\) locus gives rise to deficiency of hex A and a mutation at the \(\beta\) locus to a deficiency of hex A and hex B.

Hex A deficiency occurs in Tay-Sachs disease and has also been reported in adult G\textsubscript{M2} gangliosidosis.\textsuperscript{1} Hex A and B deficiency occurs in Sandhoff disease. Sandhoff disease usually presents before the age of 9 months with mental and motor regression, visual failure with cherry-red spots, and macrocephaly. Cases of Sandhoff disease with a later onset and protracted course have been described, presenting with progressive ataxia, spasticity and psychomotor retardation in late childhood,\textsuperscript{2,3} as a progressive ataxic syndrome in adult life,\textsuperscript{4} or with spinal muscular atrophy that resembles Kugelberg-Welander disease.\textsuperscript{5} We report a new presentation in early adult life, mimicking that of X-linked bulbospinal neuronopathy.\textsuperscript{6,9}

Case report

This 32 year old male had a normal early developmental history. He received speech therapy for a mild articulatory disturbance of dyslalic nature during childhood. He was seen by a neurologist at the age of 15 years because of upper limb tremor and a diagnosis of essential tremor was made. This has persisted. He has been prone to muscle cramps for some years. From the age of 28 or 29 years, slowly progressive proximal lower limb weakness has become evident, accompanied by wasting of the thighs. He has noticed no upper limb weakness and has had no symptoms referable to the cranial nerves. Bladder, bowel and sexual function has been normal.

He has no children or siblings. His father is neurologically normal. His mother has a mild lower limb sensory abnormality related to spinal cord involvement secondary to midthoracic vertebral crush fractures sustained in a car accident. There is no other family history of neurological disorder. His parents are both Caucasian and nonconsanguinous.

Neurological examination revealed normal cranial nerve function except for mild bilateral facial weakness and slight dysarthria. There was no contraction fasciculation of the face. His tongue was normal. In the upper limbs there was mild bilateral proximal muscle weakness and wasting and a postural tremor of the outstretched hands, present during movement. In the lower limbs, there was bilateral quadriiceps wasting and moderately severe weakness of hip flexion and abduction, lesser weakness of quadriiceps, and mild weakness of the anterolateral muscles in the lower legs. There was no lower limb tremor or ataxia. His tendon reflexes were normal.

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Table 1  Hexosaminidase activity in the patient and his parents

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<th>Total hexosaminidase</th>
<th>% hexosaminidase A</th>
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|                        | Plasma (μmol/ml/h)   | WBC (μmol/mg ptn/h)
|                        | Skin fibroblasts     | Skin fibroblasts   |
| Patient                | 0-02                 | 0-06               | 0-69               |
| Mother                 | 0-3                  | 0-9                | —                  |
| Father                 | 1-0                  | 1-9                | —                  |
| Normal range           | 0-5-2-0              | 0-7-3-6            | 2-4-13-9           |
| Sandhoff disease (infantile) | 100              | 100                | —                  |
| n                      | 0-0-04               | 0-02-0-20          | 0-07-0-09          |
| Obligate heterozygotes | 0-3-0-9              | 0-4-1-1            | —                  |
| Sandhoff disease       | 18                   | 22                 | —                  |

Fig 1  Semithin section of rectal biopsy specimen showing multiple macrophages containing lipid inclusions (arrows). Araldite section, thionin and acridine orange stain, × 550.

Fig 2  Electron micrograph from rectal biopsy specimen showing an axon filled with multiple lamellar inclusion bodies. Bar = 1 μm.

apart from sluggish knee jerks. His plantar responses were flexor and there was no sensory loss. Examination of other systems was negative. There was no gynaecomastia.

Routine haematological and biochemical studies were normal. There were no vacuolated lymphocytes. His serum creatine kinase activity was slightly increased (219 IU/l; normal < 195 IU/l). Electromyography showed chronic partial denervation in the right deltoid, biceps brachii, first dorsal interosseous, vastus medialis and tibialis anterior muscles. Motor nerve conduction velocity was normal (median 63 m/s; ulnar 55 m/s; peroneal 43 m/s). Median, ulnar, radial and sural sensory nerve action potentials were absent. Central motor conduction time was normal (cortex—C8/T1, magnetic stimulation: right 6-0 ms, left 5-6 ms) as was an electrocardiogram.

His karyotype (fibroblast culture) was normal.

Special investigations
Lyosomal enzyme studies revealed a profound deficiency of hexosaminidase activity (table 1) compared with other

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Table 2 Clinical spectrum of $G_{M2}$ gangliosidoses

- **a locus mutations (deficiency of hex A)**
  - Tay-Sachs disease
  - Juvenile $G_{M2}$ gangliosidoses with hex A deficiency resembling Batten-Spielmeyer-Vogt disease
  - Atypical spinocerebellar degeneration
  - Cases resembling juvenile spinal muscular atrophy (Kugelberg-Welander) and amyotrophic lateral sclerosis
  - Normal adults

- **b locus mutations (deficiency of hex A and B)**
  - Sandhoff’s disease
  - Juvenile Sandhoff’s disease
  - Progressive ataxia ± dementia
  - Chronic $G_{M2}$ gangliosidoses with hex A and B deficiency
  - Multisystem disorder with ataxia, dementia, pyramidal signs and amyotrophy in varying combinations
  - Juvenile onset spinal muscular atrophy
  - Neuromuscular disease resembling X-linked bulbospinal muscular atrophy
  - Normal adults

**Activator factor mutations (normal hex A and B activity)**
- Resembling Tay-Sachs disease, but milder
- Seizures, dementia and normal pressure hydrocephalus
- Normal adults

Lysosomal enzymes which had activities within the normal ranges, indicating a diagnosis of Sandhoff disease. Hexosaminidase was measured in plasma, leucocytes and cultured skin fibroblasts using the synthetic 4-methylumbelliferyl substrate as described previously. Urinary oligosaccharides were normal.

Rectal biopsy

Moderate numbers of macrophages containing PAS-positive material were present in the lamina propria (fig 1). Small numbers of neurons related to the muscularis mucosa contained rounded sudanophilic eosinophilic inclusions which were PAS-negative but which showed positive staining with Luxol fast blue. The appearances and staining reactions were those of a gangliosidosis and consistent with Sandhoff disease (Prof B Lake). Electron microscopy confirmed the presence of multiple concentric lamellar structures within axons (fig 2).

Discussion

The clinical features in this patient bore a distinct resemblance to X-linked bulbospinal neuronopathy with the early onset of upper limb postural tremor, followed by muscle cramps, limb girdle and predominant proximal limb muscle weakness, mild facial weakness and dysarthria. X-linked bulbospinal neuronopathy was initially considered to represent a “spinal muscular atrophy”, but sensory nerve action potentials are depressed or absent and mild sensory loss may be present at times. Sensory nerve action potentials were lost in the present case, although there was no sensory loss. Gynaecomastia is present in approximately 50% of patients with X-linked bulbospinal neuronopathy but was not shown by our patient.

Cases of hexosaminidase A deficiency of delayed onset have been described in recent years in whom lower motor neuron involvement has been a prominent feature. In some of these it has constituted the sole neurological manifestation, such as resembling juvenile-onset spinal muscular atrophy. Our patient most closely resembles that of Case 1 of Parnes et al in whom postural tremor and muscle cramps preceded limb muscle weakness but in whom sensory nerve conduction was normal. The three cases of Mitumoto et al has a multisystem degeneration with amyotrophy, ataxia, spasticity and mental retardation, one having an axonal sensorimotor neuropathy. The six described by Argov and Navon also showed a multisystem degeneration, as did those reported by Adams and Green and Harding et al.

The enzyme studies in our patient indicate a deficiency of hexosaminidase A and B. The case is thus classifiable as an example of delayed onset Sandhoff disease. An asymptomatic individual has been reported who possessed the homozygous Sandhoff phenotype, but the presence in the rectal biopsy from our patient of intraaxonal membranocyttoplasmic bodies of the type that occur in $G_{M2}$ gangliosidosis makes it highly probable that the hexosaminidase deficiency is responsible for his neurological disorder. The higher residual activity in his fibroblasts as compared with patients with the classical infantile form of Sandhoff disease may explain the later onset of symptoms. A case has been described with the Franceschetti syndrome and a significant reduction in hexosaminidase B in which there was a balanced chromosomal translocation (5:13) (q11:p11). The karyotype in the present case was normal.

Of the 23 obligate heterozygotes for infantile Sandhoff disease that we have tested, 21 had either a total hexosaminidase activity or percentage hex A outside the normal range in either plasma, leucocytes or both. The remaining two obligate carriers had both total hexosaminidase activity and percentage hex A in the overlap range of heterozygotes and normal controls. The mother of our patient had enzyme levels consistent with a heterozygote for Sandhoff disease whereas the levels in his father were within the normal range. Although confirmatory serological studies were not undertaken, the physical resemblances between father and son were so close as to leave no doubt that the stated paternity was correct.

It is possible that the parents possess different gene mutations, only one of which is reflected in altered enzyme activity with the synthetic 4-methylumbelliferyl substrate. The patient might therefore be a compound heterozygote. If so, this could be relevant to his atypical presentation.

Measurement of the hexosaminidase activity in the parents against the natural $G_{M2}$ substrate may be more
informative, as may DNA analysis. At present, however, no material from the parents is available for further study.

This case widens the already extensive spectrum of clinical presentations that have been shown to be attributable to $G_{M2}$ gangliosidoses (table 2).21

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