SHORT REPORT

Autosomal dominant optic atrophy with asymptomatic peripheral neuropathy

R M Chalmers, A C Bird, A E Harding

Abstract

The association between hereditary motor and sensory neuropathy (HMSN) and optic atrophy has been termed HMSN type VI. The autosomal dominant inheritance of this syndrome is reported. Three generations were affected with optic atrophy, which differed in some respects from classic dominant optic atrophy, and an asymptomatic, mainly sensory, neuropathy.

(Trans Neurosurg Psychiatry 1996;60:195-196)

Keywords: hereditary motor and sensory neuropathy; optic atrophy; peripheral neuropathy

An association between optic atrophy and hereditary motor and sensory neuropathy (HMSN) was first described by Vizoli in 1889. Further examples have been described in families showing varied patterns of inheritance and in sporadic cases and the syndrome has been termed HMSN type VI. These patients have developed distal muscle wasting and weakness and, in some reports, lancinating pain suggesting involvement of peripheral sensory neurons. We report a three generation family that shows autosomal dominant inheritance of progressive optic atrophy and an asymptomatic peripheral, mainly sensory, neuropathy.

Case reports

The figure shows the pedigree. Four family members in three generations were affected. The proband (III-2), a 31 year old woman, developed impaired vision at the age of 5 years. By the age of 16 visual acuity was 6/18 bilaterally; visual field examination showed bilateral central scotomata and fundoscopy showed bilateral optic atrophy. Examination of colour vision using Hardy-Rand-Ritler plates and the Farnsworth-Munsell 100 hue test showed red-green deficiency of medium extent. Vision gradually deteriorated and by the age of 31 acuities were 3/36 bilaterally. Neurological examination showed the previously documented ophthalmological signs and bilateral gaze evoked nystagmus. In addition, she was areflexic with impairment of pain and vibration sensation distally in the lower limbs. There was no muscle wasting or weakness. Hearing was normal. In the upper limbs motor conduction studies were normal, with median and radial sensory action potential (SAP) amplitudes of 13 μV and 18 μV respectively. In the legs, tibial nerve muscle action potentials (MAPs) were normal, with a reduced MAP amplitude in the right peroneal (3-6 mV) and left peroneal nerves (2.7 mV) without significant reduction in motor conduction velocities (MCVs) (39-43 m/s). Sural and superficial peroneal SAP amplitudes were 1 μV and undetectable respectively. Needle EMG showed evidence of chronic partial denervation in distal muscles of both upper and lower limbs.

Her father (II-1), a 65 year old man, developed progressive visual problems from the age of 10 and by the age of 40 he was only able to perceive light. Examination at the age of 65 showed bilateral optic atrophy; there was no nystagmus. He was areflexic with impaired vibration sensation distally. There was no muscle wasting or weakness. Hearing was normal. In the upper limbs motor conduction studies were normal, but median and ulnar SAP amplitudes were 6-5 μV and 2 μV respectively. In the legs, tibial nerve MCVs were normal; there was a reduction in MAP amplitude of 0-6 mV) and MCV (35 m/s) in the peroneal nerve. The sural SAP was small (3 μV). Electromyography was normal.

The following family members were considered affected on the basis of history provided by relatives. Case I-1 developed progressive visual loss at the age of 10 and was only able to perceive light from the age of 30; case II-3, a 60 year old woman, developed optic atrophy at the age of 7 and was only able to perceive light from the age of 30. Neither patient had any symptoms of muscle wasting, weakness, or impaired hearing.

Discussion

This family presented with optic atrophy and the autosomal dominant mode of inheritance is shown by the presence of both maternal and paternal transmission. None of the affected members had symptoms related to peripheral nerve disease but they had diminished tendon
reflexes and impairment of sensation on examination. Nerve conduction and EMG confirmed the presence of an axonal sensorimotor neuropathy.

Two features are atypical for the classic form of dominant optic atrophy (“Kjer type”) as defined by Smith. Firstly, the extent of visual loss—to perception of light only—is more severe than the moderately reduced visual acuity of 20/40 to 20/400 suggested by Smith and others. Secondly, this family has red-green dyschromatopsia and, while some report a range of colour defects, a blue-yellow colour defect is thought to be characteristic of dominant optic atrophy.

The association of neuropathy and optic atrophy was first described in 1889 by Vizoli in a father and son; another son had neuropathy without optic atrophy. The only other reports in which autosomal dominant inheritance is possible is of a male with progressive optic atrophy and neuropathy, whose father had distal lower limb wasting but no visual symptoms. Autosomal recessive inheritance is likely in two further reports of affected siblings with normal parents (with parental consanguinity in one instance). X-linked recessive inheritance is proposed for the case of a grandfather with neuropathy and optic atrophy whose grandson had a neuropathy but no visual complaints. There have been some reports of sporadic cases of optic atrophy and neuropathy.

The classification of these reports is complicated by the need to differentiate progressive optic atrophy, as seen in classic “Kjer-type” optic atrophy, from other forms, such as Leber’s hereditary optic atrophy (LHON). Four of the reported patients, with subacute visual loss, were young adult men, suggestive of LHON. One family has been described in which LHON and HMSN segregate independently.

Optic atrophy and neuropathy have also been found in association with other features. In particular, the triad of optic atrophy, neuropathy, and deafness has been described in autosomal dominant, recessive, and X-linked recessive forms.

This report is the first clear description of autosomal dominant inheritance of progressive optic atrophy and neuropathy. The optic atrophy differed in some respects from classic dominant optic atrophy and the neuropathy was an asymptomatic, mainly sensory, axonal degeneration. The aetiology of these syndromes of optic atrophy and neuropathy is unknown. Recent evidence suggests that some families with dominant optic atrophy show linkage to markers on the telomeric portion of the long arm of chromosome 3. Analysis of pedigrees with HMSN VI and other syndromes that include optic atrophy and neuropathy will determine whether they share a similar molecular basis.

Professor A E Harding died on 11 September 1995.

7 Gordon A. Remarks on primary neurotropic atrophy (Charcot-Marie-Hoffman type) with report of a case in which there was excessive indulgence in tea and coffee. J Neurol Ment Dis 1903;30:354–9.