LETTER TO JMG

A novel neurodegenerative disease characterised by posterior column ataxia and pyramidal tract involvement maps to chromosome 8p12–8q12.1

P N Valdmanis, A A Simões Lopes, F Gros-Louis, J D Stewart, G A Rouleau, N Dupré

The recent barrage of linkage assignments and gene discoveries has confirmed the clinical and genetic heterogeneity of ataxic diseases. They all share the prototypic feature of difficulty in walking though many additionally present dysarthria, spasticity, retinopathy, and other neurological symptoms. Broad subgroups of ataxias and related diseases exist including spinocerebellar and spastic ataxias, each with their own characteristic features.

The clinical and genetic heterogeneity of ataxias is best represented by the autosomal dominant cerebellar ataxias (ADCAs). Indeed, a minimum of 22 loci have been discovered, including those for the spinocerebellar ataxias (SCA1–8, SCA10–17, SCA19, SCA21, and SCA22), the episodic ataxias EA1 and EA2, and the complex disorder, dentatorubropallidoluysian atrophy (DRPLA). Similarly, Friedreich’s ataxia (FRDA) is an autosomal recessive disease which affects the spinocerebellar and pyramidal tracts. Symptoms are typically noticed before 20 years of age and include dysarthria, nystagmus, areflexia, and a positive Babinski sign. Hereditary spastic ataxia (HSA) is characterised by retinopathy, muscle wasting, nystagmus, and dysarthria. Spastic ataxia (SAX1) is the first described dominant form, while the autosomal recessive spastic ataxia of Charlevoix-Saguenay (ARSACS) is one of a number of diseases associated with ataxia (SMNA) of neuropathic origin. All told, the loci responsible for a significant proportion of hereditary causes of ataxias still have not been elucidated.

A large amount of heterogeneity is also observed within the hereditary neuropathies. These include the hereditary sensory neuropathies (HSNs) and the more common hereditary motor and sensory neuropathies (HMSNs). Sensory ataxia is not present in the HSNs, since they affect mainly the unmyelinated and small myelinated nerve fibres, nor is it present in the HMSNs, where sensory symptoms are seldom the presenting complaint. A few rare families have been described with a hereditary sensory-motor neuropathy associated with ataxia (SMNA) of neuropathic origin. These can include both central and peripheral nervous system involvement and neurogenic muscle atrophy. Similarly, a combination of ataxia originating in the cerebellum with signs of peripheral neuropathy has been reported. Neuropathies that affect large myelinated peripheral nerve fibres or their cell bodies located in the dorsal root ganglia can induce a sensory ataxia. Both preganglionic and postganglionic sensory nerve fibres are implicated in these forms of sensory ataxias.

Here we report an eastern Canadian family from New Brunswick with Anglo-Saxon ancestry. The family has a novel neurodegenerative disease characterised by sensory (posterior column) ataxia and variable pyramidal weakness but with no overt signs of peripheral sensory or motor neuropathy. This suggests a pure sensory ataxia caused primarily by involvement of the preganglionic sensory nerve fibres (posterior column). To describe this distinct diagnostic entity, we propose the term autosomal dominant sensory ataxia (ADSA).

METHODS

Clinical findings

This family has 10 affected living members including six females and four males (fig 1, table 1). This progressive disease does not appear to influence the overall lifespan of those affected. The typical age of onset is in the third and fourth decades ranging from 28 to 55 years, while there is no definite evidence of genetic anticipation. Most patients initially experience a difficulty in gait although some also report instability, especially in the dark. Deep tendon reflexes are usually diminished in the arms and are altogether absent.

Key points

- The ataxias and hereditary sensory neuropathies (HSNs) are clinically and genetically heterogeneous disorders.
- A Canadian family from New Brunswick, of Anglo-Saxon origin, has been identified whose affected members do not display classic signs of sensory neuropathy or spinocerebellar ataxia but who nonetheless have a debilitating sensory (posterior column) ataxia occasionally coupled with pyramidal weakness; as a result, we entitle this disease autosomal dominant sensory ataxia (ADSA).
- Ten affected patients from a single pedigree were subjected to clinical characterisation and linkage analysis. A genome scan performed on the family yielded a locus on chromosome 8 with a maximum LOD score of 4.90 for marker D8S1791. This locus has been designated as sensory ataxia 1 (SNAX1) and maps to a 9.1 cM chromosomal interval from 8p12 to 8q12.1. The candidate genes fibroblast growth factor receptor 1 and glutathione reductase were screened without detection of a coding variation.

Abbreviations: ADCA, autosomal dominant cerebellar ataxia; ADSA, autosomal dominant sensory ataxia; ARSACS, autosomal recessive spastic ataxia of Charlevoix-Saguenay; C-MAPS, compound motor nerve action potentials; DRPLA, dentatorubropallidoluysian atrophy; FGFR1, fibroblast growth factor receptor 1; FRDA, Friedreich’s ataxia; GSR, glutathione reductase; HAS, hereditary spastic ataxia; HMSN, hereditary motor and sensory neuropathy; HSN, hereditary sensory neuropathy; MRI, magnetic resonance imaging; SAX1, spastic ataxia; SMNA, sensory-motor neuropathy associated with ataxia; SNAPs, sensory nerve action potentials; SNAX1, sensory ataxia 1
in the legs. Some affected subjects have extensor plantar responses and weakness of the lower extremities. Sensation to all modalities (touch, pain, temperature, proprioception, and vibration) is invariably decreased distally, but more so in the legs than in the arms. There are never significant bruises or foot ulcerations. Autonomic dysfunction, such as sphincter disturbance or orthostatic hypotension, is not present. Apart from mild jerky pursuit abnormalities on eye movement testing in a few subjects, there are no signs of cerebellar dysfunction such as dysmetria or dysarthria. Brain and spine magnetic resonance imaging (MRI) are normal in all five individuals tested. Similarly, compound motor nerve action potentials (C-MAPs) and sensory nerve action potentials (SNAPs) are normal. To illustrate the variation in phenotypic expression, the clinical evaluation of four affected members conducted by one of the authors (ND) is described here in more detail.

**Case 1 (III:5)**

This subject is the most severely affected. His symptoms began around 45 years of age as progressive gait difficulty. By the age of 49 he needed a walker, and a year later he was wheelchair bound. He was examined at the age of 55. Cognitive function was normal. Cranial nerve examination was normal except for some mild jerkiness of ocular pursuit, while his saccadic eye movements were normal. There was neither nystagmus nor dysarthria. Motor examination was normal in the upper limbs, while the lower limbs showed a

<table>
<thead>
<tr>
<th>Table 1</th>
<th>Clinical findings of ten confirmed affected cases</th>
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<tbody>
<tr>
<td>Patient</td>
<td>Age at onset</td>
</tr>
<tr>
<td>---------</td>
<td>--------------</td>
</tr>
<tr>
<td>IV:4</td>
<td>?</td>
</tr>
<tr>
<td>IV:8</td>
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</tr>
<tr>
<td>III:14</td>
<td>7</td>
</tr>
<tr>
<td>III:12</td>
<td>50</td>
</tr>
</tbody>
</table>

↓↓↓, diminished; 0, abolished; (+), mild; (++), moderate; (+++), severe; EMG, electromyography; eval., evaluation; MRI, magnetic resonance imaging; NA, not applicable; NCS, nerve conduction studies; P, pain; Pr, proprioception; SSEP, somatosensory evoked potentials; T, temperature; V, vibration.

**Figure 1** The chromosome 8p12–8q12.1 haplotype for collected individuals in this study’s family. Males appear as squares and females as circles. Black and white symbols represent affected and unaffected individuals, respectively, while a question mark indicates an individual affected by hearsay. A diagonal line denotes a deceased individual. For disease haplotypes, a thick black bar indicates the area on the haplotype associated with the disease, while a thin black bar indicates a potentially associated area. Alleles in parentheses are inferred. Markers D8S1769 and GATA156H01 were determined to be boundaries of the 9.1 cM segregating region when taking into account the recombinations in individuals III:12 and III:14.
complete flaccid paraplegia. Despite this, there was no significant muscle atrophy. Reflexes were absent in the limbs. Plantar stimulation elicited a triple-flexion reflex. Sensory examination revealed moderately diminished perception of all main modalities distal to the elbows in the upper limbs. In the lower limbs, all modalities were severely diminished distal to the knees. There was no dysmetria on the finger-to-nose test. There was no sphincter disturbance. Electrophysiological studies showed that C-MAPs and SNAPs were both normal. Of particular note, the left sural nerve SNAP was normal despite severely diminished sensation in the leg. The needle electromyographic examinations of distal and proximal muscles were normal. Interestingly, somatosensory evoked responses from upper limb stimulation showed a prolonged latency, while evoked responses from lower limb stimulation were unobtainable. The combination of normal sensory nerve conductions and abnormal evoked responses clearly indicates preganglionic (posterior column) involvement. Cerebrospinal fluid examination was normal, with no oligoclonal bands. Similarly, right sural nerve biopsy and left deltoid muscle biopsy were normal. Levels of vitamin B12, homocysteine, vitamin E, β-lipoprotein, and very long chain fatty acids were all normal. Finally, brain and spine MRI were also normal.

Case 2 (III:7)
This patient developed progressive gait difficulty, especially in the dark, at around 35 years of age. He needed a cane at the age of 58 and was examined at 65. Cognitive function, cranial nerve examination and oculaur pursuit and saccades were all normal. No dysarthria was observed. Motor examination was normal in the limbs, but all reflexes were absent and plantar responses were flexor. Sensory examination revealed a mildly diminished perception of all modalities distal to the elbows. In the legs, all modalities were severely diminished distal to the knees. No dysmetria was seen on the finger-to-nose test. The gait was quite abnormal with a strikingly wide base and an inability of the patient to perform more than two steps on his own. The patient developed marked instability on closing his eyes indicating a positive Romberg sign.

Case 3 (III:12)
This patient developed progressive gait difficulty around the age of 50. By 60, he required a cane, and was examined at the age of 67. Cognitive function was normal. Apart from some mild jerkiness of his oculaur pursuit, his cranial nerve examination and saccadic eye movements were normal. Nystagmus and dysarthria were absent. Muscle tone and strength were normal. Reflexes were diminished in the arms and absent in the legs. The plantar responses were extensor bilaterally. Sensory examination revealed a mildly diminished perception of pain and temperature distal to the elbows while vibration and proprioception were normal. In the legs, pain and temperature perception were severely diminished distal to the knees while proprioception and vibration were absent at the big toes. Vibration was also absent at the ankles. There was no dysmetria on the finger-to-nose test; however, the heel-to-shin test was impaired due to proprioceptive loss. An abnormal gait with a wide base was observed. A Romberg sign was also present.

Pedigree and DNA analysis
Analysis of the pedigree reveals an autosomal dominant mode of inheritance given the evidence of male-to-male transmission and a penetrance that correlates with the age of the patient (table 1). Upon receipt of informed consent, blood samples were obtained from 21 individuals, including nine affected members in two generations of the family, as approved by the Ethics Committee of the McGill University Health Center. DNA was extracted from the peripheral blood by standard methods. Ten samples comprised of eight affected individuals and two normal spouses were subjected to a genome-wide scan. A total of 400 markers spanning all chromosomes at approximately 10 cM intervals were typed. Subsequent marker positions were determined using the Marshfield genetic map (Marshfield Center for Medical Genetics). Primers for each marker were generated from their respective UCSC genome browser sequence (UCSC Human Genome Project Working Draft). Alleles were visualised by incorporating [35S]dATP in PCR products and separated on a 6% polyacrylamide gel. The size and frequency of alleles were based on values from the Fondation Jean Dausset - CEPH database and compared to an M13mp18 sequence ladder. Two-point and multipoint linkage analyses were conducted using the MLINK and LINKMAP programs respectively within the LINKAGE software package. LOD score calculations were based on a disease frequency estimated at 1 in 10 000, a penetrance of 90%, and equal male-female recombination frequencies.

RESULTS
Initially, genetic testing was performed on the proband (case 1 above) to exclude SCA1, SCA6, and SCA7. The subsequent
genomic scan yielded 12 suggestively linked markers based on a LOD score of between 1.5 and 2.0 including two adjacent markers on chromosome 8. No markers yielded an LOD score greater than 2. Each of the 12 suggestive loci was further examined by genotyping other affected and unaffected individuals for nearby markers. This allowed more informative LOD score calculations and haplotype analysis, which confirmed the locus on chromosome 8 (table 2). Significant LOD scores of 4.74 and 4.76 (θ = 0) were obtained at markers DS8110 and DS8601, respectively, while the maximum LOD score was 4.90 (θ = 0) at marker DS81791. Multipoint analysis of the area, increasing the LOD score to 5.07, is shown in fig 2. A disease-related haplotype was constructed such that it incorporated the minimum number of recombinants. Individual III:12 defines the upper recombination fraction that accounts for the ataxia. Genetic analysis has excluded diseases of a similar nature and mapped ADSA to a novel locus on chromosome 8p12–8q12.1. The region corresponds to an interval of approximately 9.1 cM between markers DS81769 and GATA156H01 and contains 21 reviewed and 67 predicted genes. Currently, only one family has been identified in a geographically isolated region making it difficult to obtain more detailed clinical testing. Naturally, the search for other families with ADSA will continue. This will allow a better appreciation of the phenotype and further narrow the critical region in the event that an informative recombinant is found.

Recently, a flood of genetic information has provided insight into the mechanisms and dysfunctions leading to the various forms of ataxias. Of the 12 genes responsible for various SCAs, ten are defective due to trinucleotide repeat expansions.34 The length of the repeat becomes more pronounced with each successive generation leading to an earlier age of onset of the disease and therefore genetic anticipation. This has been shown to occur in the gene for SCA1, ataxin-1, in which the number of CAG repeats encoding glutamine amino acids is directly correlated with the severity of the disease and its age of onset.35 Although the limited number of generations collected may mask its presence, there is no direct evidence of anticipation through this family, suggesting that a triplet repeat expansion is unlikely to be implicated. When queried, the compiled genomic sequence from UCSC revealed no lengthy CAG repeats in the candidate region.

Another major theme for pathogenesis, seen in autosomal recessive ataxias, involves genes implicated in metabolic processes. The pathways involved usually provide hints to the genes implicated. Exemplary of this, a mutation in the microsomal triglyceride transfer protein is responsible for abetalipoproteinemia which presents with ataxia.36 Thus, a gene in the region that displays great promise is one that shares homology with tetrahydrofolate synthase (Celera database, hCG1647345). The folate biosynthesis pathway depends on vitamin B12, and the two have both been linked to neurodegenerative diseases. Gait disturbance can regularly be seen as an initial complaint of vitamin B12 deficiency, with or without haematological defects such as megaloblastic anaemia.37 Many of the phenotypic features of ADSA patients are similar to those with vitamin B12 deficiency.38

Incorporating this information, other genes in the chromosome 8 region for ADSA will also be screened in an order relative to their homology or commonality to genes previously discovered for ataxias and sensory neuropathies. Naturally, priority will be given to genes highly expressed in nerve conduction testing accompanied by ataxia with cerebellar atrophy.39 features which clearly differentiate it from ADSA. That ADSA affects sensory instead of spino-cerebellar pathways and the cerebellum also sets it apart from the SCAs.1 It is the loss of proprioception and not of movement coordination that accounts for the ataxia. Genetic analysis has excluded diseases of a similar nature and mapped ADSA to a novel locus on chromosome 8p12–8q12.1. The region corresponds to an interval of approximately 9.1 cM between markers DS81769 and GATA156H01 and contains 21 reviewed and 67 predicted genes. Currently, only one family has been identified in a geographically isolated region making it difficult to obtain more detailed clinical testing. Naturally, the search for other families with ADSA will continue. This will allow a better appreciation of the phenotype and further narrow the critical region in the event that an informative recombinant is found.

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### Table 2 Two-point LOD scores for markers on chromosome 8

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<thead>
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<th>Marker</th>
<th>Position, cM</th>
<th>Recombination fraction (θ)</th>
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<th>0.01</th>
<th>0.05</th>
<th>0.10</th>
<th>0.20</th>
<th>0.30</th>
<th>0.40</th>
<th>Zmax</th>
<th>θmax</th>
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<td>0.65</td>
<td>0.64</td>
<td>0.59</td>
<td>0.54</td>
<td>0.42</td>
<td>0.30</td>
<td>0.16</td>
<td>0.65</td>
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<tr>
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<td>1.61</td>
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<td>0.89</td>
<td>0.39</td>
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<tr>
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<td>1.85</td>
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<td>0.76</td>
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<td>4.66</td>
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<tr>
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<td>1.81</td>
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<td>1.05</td>
<td>0.49</td>
<td>1.81</td>
<td>0.10</td>
</tr>
</tbody>
</table>
the central nervous system because of the distribution of the sensory deficits evident in this family.

One other locus related to a neurodegenerative disease is present in the same region and thus particular attention must be paid to it. This locus is for the spastic paraplegia SPP5A which extends from 8q11.1 to 8q21.2 and shares one linked marker (D8S601). However, the disease is a recessive one and presents a separate phenotype so the likelihood that the same gene is involved in both diseases is slim. Likewise, the gene for ataxia with vitamin E deficiency (AVED) was excluded because it lies outside the candidate region at 8q12.3 and vitamin E levels were not diminished.

Identification of the gene involved in ADSA would aid in the genetic screening and counselling of subjects. It would further provide an understanding of the pathway(s) involved in a sensory ataxia, show any mechanistic differences between this disease and the SCAs, and perhaps provide a link between sensory ataxia and the peripheral neuropathies. The identification and characterisation of ADSA offers a unique perspective of the pathophysiology of ataxic diseases on the whole.

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ELECTRONIC-DATABASE INFORMATION


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