A new form of familial ataxia, deafness, and mental retardation

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Abstract
Conditions causing familial ataxia, deafness, and developmental delay are considered in the context of describing brothers with a new disorder characterised by these clinical features.

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It is uncommon for hereditary ataxia of early onset to be associated with deafness and developmental delay.1 A search using these three criteria in the London Neurogenetics Database yields 20 diagnostic possibilities.2 However, most of these conditions are associated with additional pathological features involving other systems. Accordingly the differential diagnosis in the familial ataxic, deaf patient without other signs is limited to a few distinct entities.

We report brothers with an apparently undescribed form of deafness, ataxia, and mental retardation and contrast their condition with similar, but distinct, disorders.

Case reports
The family comes from Kuwait and the pedigree is shown in the figure. The proband, V-2, was born after a normal pregnancy and term delivery, weighing 2550 g. Initial development was slightly delayed. He sat at 10 months, walked independently at 18 months, and had several single words by the age of 21 years. He continued to gain skills slowly and there was no history of seizures.

On examination aged 4 years his head circumference was on the 50th centile, as was his height, while his weight was on the 25th centile. Tone, power, and sensation were normal. There was no evidence of muscle wasting. Reflexes were difficult to elicit but were present and were thought to be normal. However, there were clear signs of cerebellar incoordination in his upper limbs and in his gait, which was broad based. There were no neurocutaneous stigmata. Clinical evaluation of auditory function suggested an abnormality and this has subsequently been confirmed by formal audiometry. Extensive investigation failed to show a chromosomal or metabolic basis for his developmental problems and CT scan of his brain proved to be normal. The family presented again six years later when V-3 was referred aged 7 years.

He had been a term delivery, weighing 3200 g. Pregnancy had been normal apart from mild hypertension in the latter weeks. Developmental milestones were slightly delayed, with sitting being achieved at 9 months, crawling at 9 to 10 months, and walking at 18 months. His gait was unsteady from the beginning. By the age of 3 years, he was using simple sentences, but his language development made little progress thereafter. Unlike his brother, V-3 has had six generalised seizures, all associated with febrile episodes but none of these episodes has been prolonged beyond a few minutes. On examination, height, weight, and head circumference were all on the 50th centile. There were no dysmorphic or neurocutaneous features. Gait was ataxic; he was unable to perform heel-toe walk or to hop. Finger-nose incoordination confirmed cerebellar involvement in the upper limbs. Tone was reduced, especially in the upper limbs but power and reflexes were thought to be normal, as was muscle bulk. Assessment of hearing confirmed a severe bilateral sensorineural hearing loss with normal middle ear function, as confirmed by tympanometry. As with V-2, extensive investigations, including urinary and plasma amino acids, urinary organic acids, serum lactate, and immunoglobulins failed to show a biochemical basis for the abnormalities and CT brain scan was normal.

Both boys attended a special school, reflecting a mild degree of intellectual delay. Although not assessed, a maternal first cousin, V-4, was said to be similarly affected.

Discussion
These brothers share similar clinical features in all respects apart from the seizures associated with fever in V-3. Essentially, theirs is a condition characterised by cerebellar ataxia,
sensorineural deafness, and global development delay. This triad has been observed in familial conditions by a number of previous authors. Berman et al reported infantile onset progressive ataxia, hearing loss initially noted at the age of 2 to 3 years, and mental retardation in three brothers out of a sibship of four boys and one girl. These patients were described as having a myopathic face and, in addition, had a marked degree of muscle wasting of the lower limbs which was present by the age of 6 years. By this age, hyperreflexia of the lower limbs was observed with extensor plantar responses. Subsequently, heel contractures developed. EMG and sural nerve biopsy were suggestive of a mild peripheral neuropathy so whether the muscle wasting was the result of pyramidal involvement or peripheral neuropathy remains unclear. The hearing loss was progressive resulting, ultimately, in a severe sensorineural deafness. The inheritance was assumed to be autosomal recessive or X linked. Subsequently three affected sisters said to have the same condition were reported as confirmation of autosomal recessivity. As with the original report, there was a myopathic facies and the plantar responses were extensor, although the lower limb muscle wasting noted in the report of Berman et al was not commented upon. The ataxia in this second family was late in onset, being noted at 8 to 10 years, unlike the observation of Berman et al of ataxia from infancy. Thus, it is by no means certain that these two reports refer to the same condition and autosomal recessive inheritance for the condition described by Berman et al has not been proven by the second report. In contrast, the brothers we describe have normal plantar and other reflexes without evidence of pyramidal tract involvement or of peripheral neuropathy. Neither are contractures nor muscle wasting of the lower limbs features in our patients (table).

Richards and Rundle described three affected brothers and two affected sisters, offspring of a consanguineous union, initially of normal intelligence which then gradually deteriorated from the age of 5 or 6 years, with obvious dementia in the second decade of life. A similarly affected sib pair may previously have been described by Koennecke. Ataxia developed early and progressive deafness was initially noted at about 2 years of age but deteriorated to leave a severe sensorineural loss. Tendon reflexes became reduced with age and talipes and claw hand were features of the second decade. Secondary sexual characteristics did not develop and the external genitalia were small. Although the subjects of this report have yet to reach the age of developing secondary sexual features, their clinical course differs from that of the Richards and Rundle syndrome in that there has been no evidence of regression. In addition, there is no evidence of pyramidal tract signs in the lower limbs.

Patients with Viallet-Van Laere syndrome also have deafness, ataxia, and mental retardation. However, in this disorder, which may comprise a number of separate conditions with differing modes of inheritance, bulbar palsy is a key feature. Neither of our patients showed any evidence of bulbar weakness. Similarly, the X linked condition described by Schmideley et al can be ruled out by the absence of optic atrophy in the current report, and the autosomal recessive disorder described by Saul and Stevenson in sibs with deafness, ataxia, and retardation was also characterised by an extreme degree of emotional lability, which we have not observed. Mild incoordination was seen in two of the three cases, but the author states that this "was not suggestive of ataxia".

In summary, we report brothers with ataxia, sensorineural deafness, and global developmental delay, who appear to represent a new genetic entity. Autosomal recessive inheritance seems likely in view of the parental consanguinity, but the presence, by family report, of an identically affected maternal male first cousin leaves X linked inheritance still a possibility.
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