Persistent superior oblique paresis as a manifestation of familial periodic cerebellar ataxia

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Abstract
A brother and sister complained of persistent diplopia due to superior oblique palsies. The cause of their symptoms became apparent when they were diagnosed as having familial periodic cerebellar ataxia (FPCA), a rare autosomal dominant condition. Oral acetazolamide (250 mg twice daily) not only prevented all the periodic symptoms but also relieved their diplopia, which had been present between attacks.

Case reports

PATIENT A
Now aged 22 years, patient A, male, began at the age of 10 to have 'episodes of giddiness' about once a week, usually while taking exercise. The episodes would last about 10 minutes and, because he became unsteady, forced him to sit down. On the basis of minor electroencephalographic (EEG) abnormalities he was believed to have 'vertiginous epilepsy and was treated with anticonvulsants, but these were unhelpful and were withdrawn. At 16 he developed diplopia and was referred to an ophthalmologist. Examination revealed visual acuities of 6/6 in the right eye and 6/5 in the left. No field defects were present, and there was no significant astigmatic error. His head was tilted and turned to the left, and a right superior oblique palsy was apparent on testing his eye movements. This was confirmed on a Hess chart (Fig 1). There were no abnormal physical signs and in particular no cerebellar deficit.

Figure 1 Hess chart demonstrating right superior oblique palsy in patient A. Ipsilateral (right) inferior oblique overaction and superior rectus underaction with contralateral (left) superior rectus underaction and inferior rectus overaction are also present, which suggests that the palsies are long standing.
minor EEG irregularities, she was prescribed anticonvulsants, but these merely exacerbated her diplopia and increased the severity of the attacks and were eventually withdrawn. At the age of 13 she was referred for an ophthalmic opinion because of persistent diplopia. On examination her corrected visual acuities were 6/9 bilaterally. She had normal visual fields and no significant astigmatic errors. There was a right superior oblique palsy which was recorded on a Hess chart (Fig 2). Her head was tilted and rotated to the left, but no other abnormalities were evident on general physical examination, and there was no cerebellar deficit.

The father of both patients had experienced similar ‘giddy spells’ since childhood. These episodes recurred weekly, lasted about 20 minutes, and consisted of slurred speech and incoordination of the lower limbs. He was unable to stand during an attack. Examination, between episodes showed ocular dysmetria, a scanning dysarthria, and transient first degree nystagmus on looking to the right. He had never complained of diplopia, and there was no evidence of an extraocular muscle palsy. No other members of the family appeared to be affected.

The Table shows the types of nystagmus found, between attacks, in familial periodic cerebellar ataxia.

<table>
<thead>
<tr>
<th>Type of Nystagmus</th>
<th>(A) In primary position of gaze</th>
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<tr>
<td></td>
<td>(i) Downbeat</td>
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<td></td>
<td>(ii) Oblique downbeat</td>
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<td></td>
<td>(iii) Rotary</td>
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<td>(B) Gaze evoked</td>
<td>(i) Intermittent 1° horizontal</td>
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<tr>
<td></td>
<td>(ii) Persistent 1° horizontal</td>
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<tr>
<td></td>
<td>(iii) Asymmetric horizontal (with increased amplitude of the jerks in the abducting eye)</td>
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<tr>
<td></td>
<td>(iv) Downbeat</td>
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<tr>
<td></td>
<td>(v) Upbeat</td>
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<tr>
<td></td>
<td>(vi) Combinations of horizontal and vertical</td>
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INVESTIGATIONS
Both patients underwent identical investigations, which included a cerebral CT scan, pure tone audiometry, optokinetic nystagmus, blood ammonia and amino acid assays, a full blood count, erythrocyte sedimentation rate, liver function tests, urea and electrolytes, blood sugar, vitamin B12 and red cell folate levels, mid stream urine tests, and urinary amino acid assays; the results were all normal. Repeat EEGs were normal in both patients.

The family’s intermittent symptoms suggested a diagnosis of familial periodic cerebellar ataxia. Treatment with oral acetazolamide (250 mg twice daily) abolished all their attacks, and Hess chart examinations confirmed that the superior oblique palsy had resolved. The effectiveness of acetazolamide has been maintained, without a single relapse, for over a year. Their diplopia has not recurred.

Discussion
Familial periodic cerebellar ataxia was first described by Parker in 1946. Since then several families have been described in which disabling attacks of dysarthria, limb ataxia, and dysequilibrium occur. In the majority of these families the attacks have lasted from 10 minutes to 6 hours, were inherited as an autosomal dominant characteristic, and were preventable by oral acetazolamide. Between ataxic episodes the patients were usually asymptomatic and on examination were completely normal or had mild cerebellar signs, including the following ocular motility disorders: bidirectional square wave jerks on attempted fixation, broken pursuit movements, dysmetric saccades, and the types of nystagmus documented in the Table. In one very atypical case, a 3-year-old with craniofacial
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dysmorphism" (kindred II, patient IV-35), ptosis was observed, and in one other case' (case 3) decreased upgaze was reported.

Our observation of persistent right superior oblique palsies in siblings with FPCA is unique and illustrates the heterogeneity of the eye signs in this condition. The responsiveness of their diplopia to acetazolamide, which also abolished their attacks, suggests that the palsies were not coincidental but the direct result of the underlying defect in FPCA. Evidently this disease can affect systems outside the cerebellum and its connections, in our cases involving the IVth nerve or superior oblique muscleature. This suggestion is supported by Trillet et al., who observed abnormalities in the visual (longer lag period) and somataesthetic (slowed conduction rate in the medial lemniscus) evoked potentials of their patients.

FPCA is easily misdiagnosed or unrecognised and may be more frequent than is generally appreciated. It appears capable of causing a variety of ocular manifestations which can persist between attacks. Once recognised it is treatable, responding impressively to maintenance oral acetazolamide, 250 mg twice daily. This diagnosis should be considered in patients presenting with intermittent ataxia or vertigo.

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