Association of an ataxia indistinguishable from Friedreich's ataxia and congenital glaucoma in a family: a new syndrome

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SUMMARY An inbred family is described in which three sibs have congenital glaucoma and two of them also have an ataxia indistinguishable from Friedreich's ataxia. The association between these two disorders has not previously been reported. The genetic mechanisms of this association are discussed.

Impairment of vision often occurs in patients with spinocerebellar degenerations. The most frequent ocular disorders are optic atrophy or pigmentary retinal degeneration, but instances of cataract have also been described. Congenital glaucoma is not known to be associated with the hereditary ataxias. We report on three sibs, the offspring of consanguineous parents, who had congenital glaucoma in conjunction with an ataxia indistinguishable from Friedreich's ataxia in two cases.

Case reports

The pedigree of the family showed obvious consanguinity between the parents of the affected subjects (fig 1). The parents of our patients were examined and found to be normal. Four subjects in this family were affected by history. There is good evidence that all four had progressive ataxia in their first or second decade of life and severe visual failure in infancy. The paternal grandmother and the parents of the proband were convinced that the clinical picture of the affected members by history was the same as described below.

Case VI. 6

The proband was an 18 year old man whose motor development was normal until the age of four when progressive imbalance of gait and clumsiness of the hands were noted. At the age of seven, examination revealed a wide based, ataxic gait. He was unable to walk heel to toe without losing his balance and was unsteady on standing with the feet close together. Muscle tone was decreased. All deep tendon reflexes were absent except for the jaw jerk, and plantar responses were flexor. A slight degree of intention tremor on finger to nose testing was noted and he had difficulty performing rapid alternating...
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movements. There was nystagmus on lateral gaze to both sides. Strength and sensation were normal. The following investigations were either normal or negative: complete blood count, glucose tolerance tests, VDRL, serum protein and lipid electrophoresis, urine screen for amino acids, chromosome analysis, routine CSF studies, X rays of the thorax and spine, and CT scan. Nerve conduction studies showed an absence of sensory action potentials, with normal motor conduction in both upper and lower extremities. Electrocardiogram showed left ventricular hypertrophy and extensive T wave inversion.

On neurological examination at the age of 14 there was axial and appendicular ataxia which was so severe that the boy was unable to walk or stand unassisted. He had dysarthric speech and moderate ptosis of the left eyelid. There was mild thoracolumbar scoliosis, flat feet, and slight wasting of the interosseous muscles of the hands, but there was no weakness. The plantar response was extensor bilaterally. Vibration sensation was diminished up to the iliac crests and there was loss of position sense in the toes. A diastolic third heart sound was present on auscultation and a systolic ejection murmur was audible along the left sternal border. Echocardiogram showed an increase in interventricular septal thickness. There had been a progression of the previous ECG abnormalities. The patient’s condition continued to deteriorate steadily and two years later he was completely wheelchair bound. His mental state was normal.

Ocular findings (fig 2)

At birth he was noted to have large eyes. The right cornea was clear, but the left showed an edematous opacity with tears in Descemet’s membrane. Intraocular pressures were 32 mmHg OD and 45 mmHg OI (Schiotz). A goniotomy was performed on the left eye. Because of the continued raised pressure in the left eye, additional goniotomies and trabeculotomy were necessary. A year later, the left eye was found to be developing first a cataract and then a spontaneous and persistent hyphema. Finally, phthisis bulbi and blindness developed in this eye. When the boy was two, the right cornea measured 14 mm and the left 15 mm. The pressure in the right eye was 37 mmHg. The anterior chamber was quite deep. When viewed gonioscopically the angle was wide open, but there was increased opacification of the structures. A goniopuncture was performed on the right eye and within the next four years the patient underwent seven additional filtering procedures. Over the next years frequent examinations failed to show an increase in the intraocular pressure above 20 mmHg in the right eye. When the boy reached 10, a high myopia developed. The refractive error was −12·00 diopters. Vision was 20/200 with myopic correction. Glaucomatous cupping of the optic disc and contraction of the visual field were noted.

CASE VI.5

The proband’s 22 year old sister was normal until the age of 10 years. When progressive unsteadiness of gait with frequent falls was noticed. When she was first examined two years later, her gait was broad based and ataxic, and Romberg sign was positive. Her performance of the heel-knee-shin test was moderately impaired, with less marked incoordination on the finger-nose testing. Tone was decreased in the upper and lower limbs but there was no wasting or weakness. Deep tendon reflexes and vibratory sensation were absent in the toes. She had bilateral pes cavus with hammer toes. Routine laboratory investigations and CT scan were normal. ECG showed wide spread T wave inversion. Electrocardiogram was not carried out. Electrodagnostic studies showed normal conduction velocity in the.

**FIG. 2 The eyes in case VI.6.**

*Note megalocornea of the right eyeball and phthisis bulbi of the left eyeball.*
motor fibres of the median and peroneal nerves, but
the sensory action potentials could not be recorded.

The disease progressed slowly, forcing her to walk
with a stick. At the age of 22, examination revealed
bilateral horizontal nystagmus on lateral gaze and
mild dysarthria. The gait was markedly ataxic and
she could not walk without support. There was mild
weakness in both lower limbs and plantar responses
were extensor bilaterally. Vibratory sensation
and proprioception were diminished distally in all extre-
mities, more in the legs than in the arms.

CASE VI.7

The proband’s 15 year old brother was neurologi-

cally asymptomatic. At the age of five, examination
revealed generalised areflexia and pes cavus. At the
age of 15, neurological examination showed the same
findings together with hypopallæsthesia in the
feet. Plantar responses were equivocal. There was
no cerebellar ataxia. Electrophysiological studies,
including motor and sensory conduction velocities
of median and peroneal nerves, brainstem auditory
evoked potentials, and somatosensory potentials
from tibial and median nerves, were normal. No
anomalies were noted on echocardiogram and
repeated ECGs.

Ocular findings in cases VI.5 and VI.7 were
similar but less severe than in the proband.

Discussion

The three patients reported here showed the ocular
hallmarks of congenital glaucoma. This is an un-
usual, inherited anomaly of the trabecular network
and anterior chamber angle which leads to obstruc-
tion of aqueous outflow, increased intraocular pres-
sure, enlarged globes (buphthalmos), corneal en-
largement, oedema, and optic nerve damage.2 As in
our family, autosomal recessive inheritance is con-
sidered to be the genetic basis for most cases of
congenital glaucoma.3 4

In addition, two members (cases VI.6 and VI.5)
of the affected sibship fulfilled all the
clinical diagnostic criteria of Friedreich’s ataxia,5 such
as onset at an early age, autosomal recessive inheri-
tance, ataxia of gait, dysarthria, generalised areflexia,
extensor plantar responses, loss of vibration and
position sensation, skeletal involvement, and car-
diac disease. Moreover, in both cases there was
absence of distal sensory potentials, an electrophys-
iological abnormality consistently present in Freid-
reich’s ataxia.6 7 By contrast, in case VI.7, neuro-
logical signs were minimal and non-progressive and
there was absence of electrophysiological abnor-
malities. Even though these features militate against
the actual diagnosis of Friedreich’s ataxia,5 we think
that a more prolonged follow up would be necessary
to establish whether the patient is affected or not
with this form of hereditary ataxia.

The present concurrence of ataxia and congenital
glaucoma, as described in our family, could be
explained by the pleiotropic effects of one gene.
Friedreich’s ataxia is often accompanied by non-
neurological manifestations, for example, cardiac
involvement, endocrine dysfunction, or skeletal
abnormalities, and it is likely that these features are
due to the effects of a pleiotropic gene.5 This could
theoretically explain the findings in our kindred, but

our knowledge there is no previous evidence that
this gene can cause congenital glaucoma. It is
possible that more than one mutant gene is segre-
gating in this family, especially since the likelihood
of two rare mutant genes segregating together in
one family is enhanced by consanguinity.5 However,
the fact that virtually all affected subjects within this
inbred family exhibited both the neurological and
ophthalmological defects might imply the transmis-
sion of a single and new mutant recessive gene
rather than two closely linked genes.

The authors thank the Department of Ophthalmo-
gy for their assistance in evaluating the patients. We
would also like to thank Mrs Marta de la Fuente for
secretarial help and Mr John Hawkins for stylistic
revision of the manuscript.

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