Autosomal juvenile retinoschisis without foveal retinoschisis

KATSUHIRO YAMAGUCHI AND SATOSHI HARA
From the Department of Ophthalmology, School of Medicine, Tohoku University, Japan

SUMMARY We describe hereditary retinoschisis without foveal retinoschisis in a girl (the product of a consanguineous marriage) and her father. The father had peripheral retinoschisis with dendritic changes of the retina and good visual acuity. The daughter had peripheral retinoschisis complicated by vitreous haemorrhage and retinal detachment. Neither had foveal retinoschisis. The pattern in a scotopic single-flash electroretinogram differentiated this condition from other forms of retinoschisis. We believe that these cases represent the autosomal recessive form of juvenile retinoschisis without foveal retinoschisis.

Autosomal juvenile retinoschisis is a rare vitreo-retinal dystrophy characterised by peripheral retinoschisis without pathognomonic foveal retinoschisis. We describe a father and daughter who had peripheral retinoschisis without foveal retinoschisis, which possibly was transmitted as an autosomal recessive disorder. The results of a scotopic single-flash electroretinogram showed a slightly reduced a wave, a diminished b wave that returned to above baseline levels, and diminished oscillatory potentials, a pattern considered pathognomonic in this disorder.

Case reports

Case 1
A 2-year-old girl suffered a fall on 16 May 1987. Several days later her mother noticed the presence of right esotropia. The child was referred to Tohoku University Hospital because of a vitreous haemorr-
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The left inferotemporal peripheral fundus of the proband shows retinoschisis and a large break in the inner layer of the retina.

Hage in the right eye on 27 May 1987. She was the product of an uneventful pregnancy resulting from a consanguineous marriage (Fig. 1). Her physical and mental development had been normal.

The ophthalmic examination showed 15 prism dioptres of right esotropia and normal eye movements. The pupil was normal in shape and showed a normal light reflex. Visual acuity could not be measured because of the patient’s youth. The right fundus was difficult to see owing to vitreous haemorrhage. The left fundus showed peripheral retinoschisis with inner retinal holes at the equator (Fig. 2). The macula appeared normal, and no foveal retinoschisis was observed (Fig. 3). A scotopic single-flash electroretinogram, which was recorded after 20 minutes of dark adaptation, displayed a slightly reduced a wave, a diminished b wave that returned to above baseline levels, and diminished oscillatory potentials (Fig. 4). Ultrasonography disclosed a strong sheet-like echo that was suggestive of a retinal detachment.

After a week’s observation an ophthalmoscopic examination of the girl’s right eye showed a small, full-thickness retinal detachment that involved the inferotemporal retina and extended from the 5 to 9 o’clock meridian. Also observed were a large grey membrane floating in the inferotemporal vitreous cavity and two holes in the outer retinal layer at the 6 o’clock meridian between the vascular arcade and equator, among the old vitreous haemorrhage. Another lesion of retinoschisis spread across the superotemporal fundus. The optic disc appeared normal, and foveal retinoschisis was not observed.

Surgical treatment consisting of cryoretinopexy and scleral buckling was performed successfully on 9 June 1987. The retina became reattached during the postoperative period of observation.

Control

Patient

Fig. 2 Right: Single-flash electroretinogram, measured after 20 minutes of dark adaptation, shows a normal a wave, a reduced b wave that returned to above the baseline levels, and diminished oscillatory potentials in the proband. Left: An electroretinogram from a normal 2-year-old, measured after 20 minutes of dark adaptation.
displayed a slightly reduced a wave, a diminished b wave that returned to above baseline levels, and diminished oscillatory potentials (Fig. 6). Fluorescein angiography disclosed arborescent figures of retinal vessels in the regions of retinoschisis.

Discussion

Among the various hereditary retinoschisises foveal retinoschisis is an important pathognomonic finding. In X-linked juvenile retinoschisis, which is carried by the X chromosome and is transmitted in a recessive manner with a high degree of penetrance, foveal retinoschisis occurs in 98% to 100% of patients. In approximately 50% of patients foveal retinoschisis is the sole pathological finding. This disease is characterised by a negative electroretinogram, which showed a normal a wave, a b wave that does not return to the baseline values, and reduced oscillatory potentials. In a female carrier of X-linked juvenile retinoschisis radial wrinkling of the inner limiting membrane occurs around the fovea, but no peripheral retinoschisis is seen. The electroretinogram is normal. Favre-Goldmann disease, which is inherited in an autosomal recessive manner, is not only characterised by peripheral retinoschisis and foveal retinoschisis but also includes an optically empty vitreous, chorioretinal atrophy, cataract, loss of peripheral vision, glial bands, eventual blindness, and an extinguished electroretinogram. In autosomal recessive foveal retinoschisis the lesion is limited to the macular area. The electroretinogram is normal. Autosomal dominant juvenile retinoschisis sometimes affects the macula. In eight patients described by Yassur and colleagues three presented with foveal retinoschisis and five had macular pigmented changes. Six patients had a normal single electroretinogram; the remainder had anormal a wave and a reduced b wave that returned to above the baseline levels.

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Retinoschisis is diagnostic for autosomal juvenile retinoschisis. Cibis described a family in which female monozygotic twins and their grandmother had peripheral retinoschisis without foveal retinoschisis. Their disorder was considered an autosomal form of retinoschisis with recessive or possibly even dominant transmission of inheritance. The father and daughter described here have retinoschisis without foveal retinoschisis. All members of the family live in a small, remote village. At least two consanguineous marriages have been recorded in the pedigree. Because the girl described in this report is also the product of a consanguineous marriage, the most likely mode of transmission in this family is autosomal recessive. The absence of foveal retinoschisis and the electroretinographic pattern displayed by the father and daughter also differentiate this condition from other forms of vitreoretinal dystrophy.

References


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K. Yamaguchi and S. Hara

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