X-linked juvenile retinoschisis

Retinosquise juvenil ligada ao X

Cesar Gomes da Silveira¹, Gabriela Hertz Soares², Jacqueline Provenzano³

ABSTRACT

Report the case of a male 28 years-old patient, who was referred to Centro de Estudos e Pesquisas Oculistas Associados – RJ, to retina assessment and presented the typical findings of the X-Linked Juvenile Retinoschisis: a cystoid maculopathy with formation of foveal cysts and schisis of the peripheral retina. It was carried out the conservative management, with attention to the complications.

Keywords: Retinoschisis/diagnosis;  Genetic diseases, X-linked/diagnosis; Case reports

Resumo

Relatar o caso de um paciente masculino de 28 anos, que foi encaminhado ao Centro de Estudos e Pesquisas Oculistas Associados – RJ, para avaliação de retina e apresentou os achados típicos de Retinosquise Juvenil Ligada ao Cromossomo X: maculopatia cistóide com formação de cistos na fóvea e retinosquise periférica. Foi realizado o manejo conservador, com atenção para as complicações.

Descritores: Retinosquise/diagnóstico;  Doenças genéticas ligadas ao cromossomo X/diagnóstico; Relatos de casos
INTRODUCTION

X-linked juvenile retinoschisis (XJR) is an inherited recessive vitreoretinal degeneration more common in young men characterized by bilateral maculopathy with associated peripheral retinoschisis in 50% of patients. A prevalence of 1:5000 to 1:28000 is estimated.

There are descriptions of mutations in the XLRS1 gene, responsible for coding of retinoschisis - a protein that provides adhesion and interaction among cells and among the retinal layers. Defects or absence of its secretion may reduce the adherence between the layer of nervous fibers and the remainder of the sensory retina, forming cystic cavities.

XJR manifests with reduced visual acuity (VA) between the ages of five and ten years of age, and evolves with progressive visual loss during the first two decades of life. There is the possibility of stabilization of the VA until the fifth or sixth decade, when it can worsen due to macular atrophy, no longer showing the cystic aspect.

The visual acuity may be less than 20/200, keeping an average of 20/70 in young adults. Abnormalities in the angle of the anterior chamber such as occlusion angle, strabismus and nystagmus are described as other findings. Vitreous hemorrhage and retinal detachment are the main complications.

The foveal retinoschisis is characterized by the “wagon wheel” standard of radial streaking due to changes in the layer of nerve fibers related to a defect of the Müller cells. Despite the cystic appearance, the exam of fluorescein angiography does not evidence the progressive extravasation characteristic of cystoid macular edema and contributes to the differential diagnosis between pathologies. Peripherial retinoschisis predominantly involves the inferior temporal quadrant, often with large detachments of the inner retinal layers. These changes of the inner layers of the retina predispose vitreous hemorrhage and retinal detachment.

CASE REPORT

MJS, male, 28, white, single, born and raised in Rio de Janeiro - RJ was referred to the Study and Research Center Opticians Associates - RJ (CEPOA) in February 2011, for retinal evaluation with the diagnosis of XJR. When asked about his family history, he mentioned a brother with whom he is no longer in contact, but who suffers from the same pathology diagnosed.

He brought additional exams from 2008, in which the red-free photography showed the standard of foveolar retinoschisis that extends circumferentially to the parafoveal area. Fluorescein angiography demonstrates cystic changes in the macula without fluid leakage through the subeyoid space (Figure 1). Optical coherence tomography (OCT) (Figure 2) shows foveal cysts with retinal cleavage between the outer and inner plexiform layers in both eyes (BE).

In the eye exam held on 02/01/2011 the visual acuity was 20/60 in BE, with the best correction. The biomicroscopy of the anterior segment shows no changes. The intraocular pressure was 17 mmHg at 04pm in BE. The retinal mapping performed showed a macular image in foveolar retinoschisis and inferior temporal retinoschisis in the right eye. The left eye showed the same pattern of macular and peripheral retinal images. These findings corroborate the diagnosis of X-linked juvenile retinoschisis.

A new fluorescein angiography (Figure 3) was made in 2011 showing similar appearance to that found in the previous exam. The exam of optical coherence tomography (OCT) (Figure 4) also held in 2011 showed an image of subfoveolar cystic confluence, further changing the fovea architecture and justifying the visual impairment.

DISCUSSION

X-linked juvenile retinoschisis is a relatively rare disorder which determines low visual acuity, mainly by foveolar retinoschisis.

The inferior temporal bilateral presentation is considered the classic pattern of disease, and the most frequent complications and of worst prognosis are vitreous hemorrhage and retinal detachment.
The fluorescein angiography reaffirms the diagnosis, ruling out the possibility of cystoid macular edema, since there is no extravasation of dye through the macula. The optical coherence tomography (Figures 2 and 4) shows changes in the evolving foveolar architecture (foveal thickness changes in BE), resulting from cysts on the layer of nerve fibers. These cystic formations are also found in the retinal periphery, where they may break to the vitreous and the retina, causing retinal ruptures and hemorrhages.

The electroretinogram is important in the diagnosis of this disease. The wave amplitude A is normal or close to normal, but wave B has a very reduced amplitude. This reduction in the wave of the B is justified by the change in the initiation of the electrical impulse in the Müller cells.

The treatment aims to prevent complications that increase visual loss, since nothing has been proven effective in preventing the development of the disease. The possibility of treatment with topical dorzolamide is being studied, but still remains unproven.

In cases of vitreous hemorrhage, some authors suggest the use of photocoagulation of the vessels responsible for the bleeding. But in cases of retinal detachment, a tamponade on the rupture and vitrectomy via pars plana are indicated.

Children who have concomitant strabismus deserve attention, and factors that can induce amblyopia should be minimized. Aid for low vision, and adaptive skills training can help individuals who develop visual loss. Genetic counseling can help identify family members who are carriers of the gene XLRS1, in order to avoid further damage.

REFERENCES