Cogan Reese syndrome

Síndrome de Cogan Reese

Anelise de Medeiros Lago¹, Ricardo Henrique Goulart Bittar¹, Larissa Rossana Souza Stival¹, João Jorge Nassaralla Junior²

ABSTRACT

Study conducted to report a case of Cogan Reese syndrome. Female patient, 55 years old with diagnosis of Cogan Reese syndrome. Best visual acuity of 0.67 in the right eye and 0.2 in the left eye. By biomicroscopic examination there was no abnormalities in the right eye. In the left eye there were pigmented nodules on the anterior surface of the iris, corneal irregularities and iris hole (pseudopolycoria). The intraocular pressure was 18 mmHg in the left eye and there was glaucomatous optic atrophy of the optic disk. The patient had been subjected to trabeculectomy three years ago. Recently medical treatment allowed the relative control of intraocular pressure. Gonioscopy revealed peripheral anterior synechiae. Corneal specular microscopy showed ICE-cells and low cell count. Glaucoma filtering surgery is usually successful when done early, but it may fail due to endothelialization of the fistula by the abnormal corneal endothelium. Medical treatment was effective despite a fail trabeculectomy.

Keywords: Glaucoma; Corneal diseases/diagnosis; Iris diseases/pathology; Endothelium, corneal/pathology; Syndrome; Case reports

RESUMO


Descritores: Glaucoma; Doenças da córnea/diagnóstico; Doenças da íris/patologia; Endotélio corneano/patologia; Síndrome; Relato de casos

¹ Ophthalmology Resident of Instituto de Olhos de Goiânia, Goiânia – Goiânia, GO, Brazil.
² Doctor, Department of Retina and Vitreous, Instituto de Olhos de Goiânia – Goiânia, GO, Brazil.

Study conducted at the Instituto de Olhos de Goiânia - Goiânia, GO, Brazil.

The authors declare no conflicts of interests.

Received for publication 05/09/2012 - Accepted for publication 28/01/2013
INTRODUCTION

This study was conducted to describe a case of Cogan Reese syndrome, which is a rare disease classified as one of the variants of the iridocorneal endothelial syndrome, which affects young, healthy individuals and is not correlated to family history. The rapid evolution courses with persistent glaucoma. It is often progressive, although it has already been reported the regression of a case. The corneal abnormalities found are features such as polymorphism and endothelial polymegathism. The specular microscopy shows pathognomonic endothelial alterations with the presence of abnormal cells, characterized by a dark area with a clear central spot and often a clear peripheral zone. The size of the cells varies, but typically they are larger than normal endothelial cells. They appear in areas that are presented with a metal aspect beated to specular reflection at the slit lamp examination. They are called ICE-cells. Some histological studies and electron microscopy showed ectopic Descemet’s membrane on the posterior corneal surface and before the iris, and the presence of cell necrosis and mild chronic inflammation, with loss of contact inhibition associated to the formation of multiple endothelial layers. These changes may or may not affect the entire endothelium. Nodes representing spots of normal iris stroma surrounded by endothelial cells are characteristic, besides the iridian dystrophic changes such as atrophy, pseudopolycoria and uveal ectropion. This syndrome is rare and usually unilateral.

CASE REPORT

Patient diagnosed with Cogan Reese syndrome in ophthalmological follow-up in the Eye Institute of Goiânia since February 2010. A.P.S., female, 55, from Paraúna - Goiás. The patient presented only hypertension under control as pathological background. Patient reports history of sudden low visual acuity (LVA) in the left eye (LE) held in July 2009. A trabeculectomy was performed in the LE 15 days after the episode of low visual acuity. Follow-up in this hospital began in February 2010. In November 2010 she was prescribed eyedrops of timolol maleate 0.5% and brimonidine tartrate 0.2% every 12 hours due to surgical treatment failure. A corneal specular microscopy was performed in June 2011, which showed the usual pattern in examination of the RE, and endothelial cell count decreased in the LE, as well as characteristic changes confirming the diagnosis of Cogan Reese Syndrome.

In March 2012 she was making use of the following eyedrops: dorzolamide hydrochloride 2% in the dosage of every 12 hours, timolol 0.5% every 12 hours, and brimonidine tartrate 0.2% every 12 hours in both eyes. The eye examination showed: visual acuity with best correction in right eye (RE) (+8.75): 20/30 and LE (+4.25 -1.50 95º): 20/100. The biomicroscopy presented: in the RE - clear cornea, nuclear cataract 1+/4+, wide anterior chamber, reagent pupil; and in the LE - cornea with central and peripheral discrete leukomas, diffuse superficial punctate keratitis, nuclear cataract 2-3+/4+, non-reactive mydriasis , accessory pupil, nevus iridian.

Gonioscopy of the left eye revealed high iris insertion, narrow angle and synechiae of 5 to 7 hours and 9 to 12 hours. The eye fundus examination showed: RE - retinal applied, papillary excavation 0.5, nerve with usual staining and aspect, retinal pigment epithelium in the usual pattern, vessels with normal path and size, macula with no visible changes.
Eye fundus: LE - applied retina, papillary excavation 0.9 deep, slightly pale nerve, retinal pigment epithelium within the usual pattern, slightly tapered arteries in relation to the right eye, veins with usual path and size, macula with no visible changes.

Figure 6 - Left eye fundus reveals excavation 0.9 with discretely pale nerve in comparison to the right eye.

The intraocular pressure (IOP) was 11 mmHg in the RE and 18 mmHg in the LE. The visual field exam showed glaucomatous-type injury, and a progression was observed when compared to the same exams held in 2010, 2011 and 2012. There was a reduction of sensitivity in the eye affected consistent with the damage caused by glaucoma in the left eye.

DISCUSSION

Iridocorneal endothelial (ICE) syndrome is rare and predominant in leukodermic females with age of onset usually from the third decade of life(2,9). It is usually progressive and unilateral(2,8-11), and its etiology is attributed to chronic endothelial inflammation probably of viral etiology(1,10-11). Cogan Reese syndrome may present some degree of the characteristics present in other variants. However, its main finding is the nodules on the surface of the iris representing a remaining iris stroma surrounded by endothelial cells(8,9,11). Multiple delayed goniosynechias extend in all directions, and may block the camerular sinus. Patients with Cogan Reese syndrome often complain of pupil changes(2). It is believed that glaucoma is caused by the proliferation of corneal endothelium, with the formation of a membrane that progressively covers the trabecular meshwork, and not due to goniosynechias that emerge lately(1,4,11).

The clinical treatment of glaucoma should be initially tried with agents that reduce the production of the aqueous humor, from the third decade of life(3,9). It is usually progressive and predominant in leukodermic females with age of onset usually from the third decade of life(2,9). In the difficult cases of corneal edema, penetrating keratoplasty can be carried out 2 years afterwards.

A case of Cogan Reese syndrome was reported, in which the patient presented corneal endothelial changes, iridian nodules, pseudopolycoria and secondary glaucoma. Early trabeculectomy was performed, there was an increase in the intraocular pressure two years afterwards, and currently reasonable pressure control is presented with the use of antiglaucoma eye drops. The patient will be submitted to a new filtering surgery combined with facetectomy with implantation of IOL.

Based on the results and the case discussed, it was concluded that knowledge of the syndrome and its prognosis are essential for its proper management. In case of failure of drug therapy, a filtering surgery shall be suggested before there is glaucomatous damage to the affected eye, since the disease is progressive with endothelial injury and risk of corneal decompensation.

This study aims to highlight the importance of the general ophthalmologist in identifying and better conducting rare and specific cases as the one described, since with early diagnostic and therapeutic it will be possible to decrease the morbidity of the syndrome.

ACKNOWLEDGEMENT

To Dr. João Jorge Nassella Junior for the methodological orientation.

REFERENCES


Corresponding author:
Anelise de Medeiros LagoRua 19, número 90 apartamento 303 - Setor Oeste
ZIP Code: 74120-100 - Goiânia (GO), Brazil
Fax: (62) 32202557
E-mail: aneliselago@hotmail.com