Coexistence of optic pit and coloboma of iris, lens, and choroid: a case report

Coexistência de fosseta óptica e coloboma de íris, cristalino e coroide: um primeiro relato de caso

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ABSTRACT

A 42-year-old woman was admitted to our clinic with a complaint of glare in both eyes. Biomicroscopic examination of both the eyes revealed iris and lens colobomas in the inferior quadrant. Fundus examination of the right eye revealed an oval and gray inferotemporal optic pit and two choroid colobomas in the inferior quadrant. In the left eye, two choroid colobomas were detected that were inferior to the optic nerve head. Furthermore, a 21-year-old man presented to our clinic for a routine ophthalmologic examination. Bilateral biomicroscopic examination was normal. Fundus examination of the left eye revealed an oval and gray inferotemporal optic pit and a choroid coloboma that was inferior to the optic nerve head. Here we describe optic pits co-occurring with iris, lens, and choroidal colobomas. On the basis of these cases, a defect in the closure of the embryonic fissure is the most plausible etiology of the optic pit.

Keywords: Coloboma; Eye abnormalities; Optic nerve abnormalities; Iris abnormalities; Choroid abnormalities; Lens, crystalline abnormalities

INTRODUCTION

Optic pit is a congenital anomaly of the optic nerve head that may cause acquired or congenital vision loss and serious macular detachment[1]. It usually occurs unilaterally and temporally on the optic disc, is round and gray, and may coexist with the cilioretinal artery. In the early phase, the optic pit is hypofluorescent on fundus fluorescein angiography (FFA); however, it is hyperfluorescent in the late phase[2]. Here we report two cases of co-occurring optic pit and uveal coloboma, including the first published case of an optic pit accompanied by a unilateral iris, lens, and choroid coloboma.

CASE 1

A 42-year-old woman was admitted with a complaint of glare in both the eyes. On ophthalmic examination, her best corrected visual acuity (BCVA) was 20/20 in both the eyes. Biomicroscopic examination of the right eye revealed iris and lens colobomas in the inferior quadrant, pigment precipitates on the lens, and posterior subcapsular cataract (Figure 1 A). In the left eye, there was an iris coloboma in the inferonasal quadrant and iris septa near the limbus that connects the iris tissue, pigment precipitates on the lens, and posterior subcapsular cataract (Figure 1 B). No retinal detachment was found, and the bilateral maculae were normal. In the right eye, the optic pit revealed hypofluorescence in the early phase and relative hyperfluorescence in the late phase of FFA, as did the choroid coloboma anteriorly located one was 8-disc size, extending to the equator in the inferior quadrant. Under the left optic disc was a 4-disc-sized choroid coloboma that was covered with a membrane with a 1-optic disc-sized hole. A second 7-disc-sized coloboma was detected at the anterior of the coloboma, and pigment alterations were observed around both (Figure 2 B). No retinal detachment was found, and the bilateral maculae were normal. The right eye, the optic pit revealed hypofluorescence in the early phase and relative hyperfluorescence in the late phase of FFA, as did the bilateral choroid colobomas (Figure 3). Results of optical coherence tomography (OCT) of the optic pit are shown in figures 3 C and 3 D.

CASE 2

A 21-year-old man presented at our clinic for a routine bilateral ophthalmologic examination. His BCVA was 20/20 in both the eyes, with a refraction of -4.00. Bilateral biomicroscopic examination was normal. Intraocular pressure was 18 mmHg. Fundus examination of the right eye revealed an oval and gray inferotemporal optic pit and two choroid colobomas (one was 2-disc size and an anteriorly located one was 8-disc size, extending to the equator in the inferior quadrant (Figure 2 A). Under the left optic disc was a 4-disc-sized choroid coloboma that was covered with a membrane with a 1-optic disc-sized hole. A second 7-disc-sized coloboma was detected at the anterior of the coloboma, and pigment alterations were observed around both (Figure 2 B). No retinal detachment was found, and the bilateral maculae were normal. In the right eye, the optic pit revealed hypofluorescence in the early phase and relative hyperfluorescence in the late phase of FFA, as did the bilateral choroid colobomas (Figure 3). Results of optical coherence tomography (OCT) of the optic pit are shown in figures 3 C and 3 D.
choroid coloboma with half the size of an optic disc that was inferior to the optic nerve head. Additionally, two cilioretinal arteries were detected, one of which transited through the optic pit (Figure 4). The macula was normal, and there was no retinal detachment in the left eye. The fundus of the right eye was normal.

DISCUSSION

Two hypotheses have been proposed for optic pit development. The first proposes developmental anomaly of the primordial optic nerve[3], whereas the second attributes optic pit development to the incomplete closure of the embryonic fissure[4-7]. The coexistence of optic pits and uveal colobomas supports the latter hypothesis. Because of the defective closure of the embryonic fissure, colobomas may arise anywhere from the optic nerve head to the pupil. After invagination of the primary optic vesicle, the distal portion approaches the proximal part, thus creating the optic cup. Abnormal closure of the inner layer of the optic cup leads to chorioretinal coloboma, and the closure defect of the distal end of the embryonic fissure causes iris coloboma[8].

Cases of optic pit co-occurring with serous macular detachment, chorioid coloboma, or retinochoroidal coloboma have been reported, both unilaterally[6,7] and bilaterally[5]. One report described a 2-year-old patient with aniridia and with an optic pit in one eye and a morning glory anomaly in the other[6]. The optic pit and iris coloboma developing in three generations of the same family have also been reported[4]. Here we described a case of optic pit co-existing with iris, lens, and choroid colobomas and a case of optic pit co-occurring with chorioid coloboma. Chorioid coloboma in the first patient had a hole in it, but retinal detachment was absent. Overall, a very small number of cases have been published regarding the co-occurrence of optic pits and chorioid colobomas[4-7], and to the best of our knowledge, this is the first published description of optic pit accompanied by iris, lens, and choroid coloboma. On the basis of these cases, a defect in the closure of the embryonic fissure appears to be the most plausible etiology of the optic pit.

REFERENCES