BILATERAL COLOBOMA OF IRIS AND CHOROID WITH UNILATERAL MICROCORNEA AND ASTIGMATISM – A CASE REPORT

Dr. Arun kumar S. Bilodi1*, Dr. Muthu Ramalingam2, Dr. Vijayalakshmi3, Dr. Vikas Shantaram Kakade4 and Dr. P. Arunkumar5

1Professor and Head, Department of Anatomy, Velammal Medical College Hospital and Research Institute, Madurai – 625009.
2Professor and Head, Department of Ophthalmology, Velammal Medical College Hospital and Research Institute, Madurai – 625009.
3Assistant Professor, Department of Ophthalmology, Velammal Medical College Hospital and Research Institute, Madurai – 625009.
4Senior Resident, Department of Ophthalmology, Velammal Medical College Hospital and Research Institute, Madurai – 625009.
5Senior Resident, Department of Ophthalmology, Velammal Medical College Hospital and Research Institute, Madurai – 625009.

ABSTRACT

A 22 years old female came to outpatient department of ophthalmology at Velammal Medical College Hospital and Research Institute, Madurai with history of defective vision since birth. She had no systemic diseases. No family history of similar complaints. On examination she had bilateral coloboma of iris and choroid with microcornea in right eye. She was examined thoroughly and treated. Her vision improved on wearing glasses. Her vision improved on wearing glasses which is rare. Hence this case made us interesting to study in detail, and reported.

KEYWORDS: Defective vision, coloboma, microcornea, iris, choroid, astigmatism.

INTRODUCTION

Occurrence of ocular coloboma is due to incomplete closure of neuroectodermal optic cup that occurs between 5-8 weeks of intra uterine life.1 The coloboma affects the posterior segment of the eye which may be unilateral or bilateral. It occurs bilaterally in 60% of cases.2 The coloboma affects the retinal pigment epithelium (RPE),
neurosensory retina, or choroid due to failure of fetal fissure to close posteriorly.[3]

Chorioretinal coloboma may affect other systems like cardiovascular, nervous, musculoskeletal, gastrointestinal and genitourinary systems.[4]

Incidences of coloboma
Studies have shown various incidences in different countries. They are
1) In Spain, it is 0.5/10,000 live births. [5]
2) In France it is 1.4/10,000 live births. [6, 7]
3) In United States it is 2.6/10,000 in the USA. [8]
4) In China, it is 7.5/10,000 live births. [9]
5) In Canada incidences of coloboma in blind adults ranges from 0.6%-1.9%. [10]

CASE REPORT
A 22 years old female from Madurai came to the outpatient department of ophthalmology with history of defective vision since birth. She had no systemic diseases. No similar complaints in her family.

On general examination
She was underbuilt and moderately nourished. She had no cyanosis, jaundice, clubbing, and lymphadenopathy.

On local examination
1) Microcornea of right eye
2) Bilateral coloboma of iris and choroid
3) Astigmatism in both eyes (right eye 3.0x15 degrees and in left eye -1.0x180 degrees).

She was prescribed glasses to correct her astigmatism and refractory errors. Her vision improved to 6/9 after wearing the glasses which is of rare sequel. Optic discs were normal in both eyes.
DISCUSSION

Chorioretinal colobomas are congenital anomalies present since birth.\(^{[12]}\) Vitamin A is very essential for development of eye.\(^{[13]}\) The retinal detachment is the most common complication that occurs in 8.1 to 43% of cases.\(^{[14-20]}\) As per Warburg M. and Stoll C. et al. incidence of coloboma ranges from 0.5 to 2.2 cases per 10,000 live births.\(^{[21,22]}\) In some cases, it may be associated with systemic disorders which may be sporadic or inherited.\(^{[23]}\) Rarely isolated microcornea has been reported having horizontal diameter of less than 11 mm without any significant systemic disorders.\(^{[24, 25]}\) Ribhi Hazin and Arif O. Khan, have reported a case of bilateral small eyes in a boy of 12 years present since birth. Visual acuity with his current glasses (-3.75-3.50\times020 right eye [OD], -4.25-1.25\times165 left eye [OS]) was 20/40 in both eyes. No history of strabismus. Pupillary examination was normal, By Goldman tonometry, intraocular pressure was 14mm in both eyes. Slit lamp examination showed small but normal eyes having horizontal diameter of less than 10 mm of normal cornea. Anterior chamber was within normal limits.\(^{[26]}\) It has reported by. Schubert, H.D. that in 23-42% of cases risk of retinal detachment occurs in coloboma of posterior pole.\(^{[27]}\)
Present study
Microcornea of right eye associated with bilateral coloboma of iris and choroid with astigmatism in both eyes associated with defective vision was seen in a young female aged 22 years. These multiple anomalies were not associated with cataract, retinal detachment, or any systemic disorders. No history of strabismus, glaucoma. Both optic discs were normal. Her vision improved after wearing glasses which is very rare.

CONCLUSION
In spite of multiple congenital defects of the eye present in this patient, her vision improved after wearing glasses which is a rare sequel. Hence it has been studied and reported.

Take home message
Any ocular congenital anomalies should be well examined, should be treated carefully and proper counseling has to be done and reassurance to be given so as to boost their morale.

REFERENCES


