Cerulean cataract: a case report

Catarata cerúlea: relato de caso

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KEYWORDS:
Cataract; Ophthalmologic surgical procedures; Congenital abnormalities.

ABSTRACT
Cerulean cataract is a rare subtype of cataract that is primarily caused by an autosomal dominant gene, which causes low visual acuity in children and adolescents. Herein, we report a patient with a rare type of cataract to raise awareness of its occurrence and allow its prompt diagnosis and more efficient treatment. Cerulean cataract is a rare form of congenital cataract; of them 25% are of hereditary and genetic etiology. Neonates should undergo a proper eye test (red reflex examination) while at the nursery for an early congenital cerulean cataract diagnosis. Presently, several surgical techniques have been described for congenital cataract treatment, considering the challenges faced during the intraoperative period, as it deals with crystallins that are softer than usual. Cerulean cataract has rare occurrence, with 25% being hereditary and needing no treatment, making observation the best course of action. Surgical treatment was recommended in this case to improve the patient’s visual acuity and reduce the risks of amblyopia.

INTRODUCTION
Cerulean cataract is a rare subtype of cataract that causes low visual acuity in children and adolescents¹. Low visual acuity does not significantly occur before the age of 2 years; thus, patients are usually asymptomatic during early childhood¹-⁴. However, opacities may grow and cause lost red reflexes and decreased vision in both eyes because both child and crystallin evolve; thus, parents often seek ophthalmological care.

Herein, we report a patient with a rare type of cataract to raise awareness of its occurrence and allow prompt diagnosis and more efficient treatment.
Ophthalmic examination of the patient, including examination under cycloplegia, biomicroscopy of the anterior segment, retinography, and ocular ultrasound, were documented through their medical record and additional tests.

CASE REPORT

J.B.R. is a 9-year-old boy from Rio de Janeiro who came to our clinic with complaints of progressive loss of visual acuity in both eyes. He reported a family history of congenital cataracts (mother and uncle). The corrected visual acuity was 20/50 in the right eye and 20/80 in the left eye, and the pinhole visual acuity potential was 20/30 in both eyes. The biomicroscopy of the anterior segment showed bluish, punctate, and scattered opacities in the adult, fetal, and embryonic nuclear regions of the crystalline, respectively, in both eyes (Figure 1). Optical coherence tomography of the anterior chamber (Figure 2) was conducted, which revealed no abnormality.

DISCUSSION

Several scholars linked some genes, such as nucleotide 12 deletion associated with the lens βB12 gene and γ-crystalline pseudogene activation; however, out of all the genes studied in animals, only CRYBB2 was highly expressed in the lens of adults5. Harder particles that once could not be aspirated can now be easily broken down by this device with the advent of phacoemulsification techniques. Among the most common complications is posterior capsule opacification.

Cerulean cataract has a rare occurrence, and 25% of them are hereditary and need no treatment, making observation the best course of action. In this case, surgical treatment was recommended to improve the patient’s visual acuity and reduce the risks of amblyopia.

REFERENCES


