Case report: Microphthalmos associated with cataract, persistent fetal vasculature, coloboma and retinal detachment

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Abstract

We present here a case of left eye microphthalmos associated with cataract, persistent fetal vasculature (previously known as persistent hyperplastic primary vitreous), iris and retinochoroidal coloboma and retinal detachment. No surgical intervention was done and the patient was kept on regular follow-up.

Keywords: Microphthalmos, Cataract, Persistent fetal vasculature, Coloboma, Retinal detachment.

Case Summary

A 12 years old girl was referred to our department with chief complaint of sudden diminution of vision in left eye for last 20 days. She was apparently well 20 days back, and then incidentally she closed her right eye and noticed diminution of vision in her left eye. Antenatal history was uneventful. There was no history of trauma during delivery or during perinatal period. Parents took her to a local practitioner where she was diagnosed with unilateral left eye congenital cataract and referred to our centre.

Her visual acuity at presentation was 6/6 in right eye on Snellen’s chart; left eye visual acuity was only 6/60 on Snellen’s visual acuity chart. Intraocular pressure was 18 mm of Hg in right eye and 14 mm of Hg in left eye by Applanation tonometry. There was no history of trauma, fever, headache, any drug intake before she noticed her decreased vision in right eye. She had normal birth at term.

On ocular examination, Left eye was smaller than right eye and left eye exotropia of about 15 degree was present (Fig. 1).

Left eye corneal diameter (9.50 mm) was also smaller than the right eye (11.03 mm) (Fig. 3).

Fig. 1: Left eye microphthalmos and exotropia

Fig. 2: Left eye inferior iris coloboma, posterior synechiae superiorly and cataract

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Fig. 2: Left eye inferior iris coloboma, posterior synechiae superiorly and cataract

Left eye iris coloboma inferiorly with key hole deformity (Fig. 2) associated with posterior synechiae at 12’o clock hr. position was present. Pupil was non-reacting in left eye. Mature cataract (Fig. 2) was present with fundus details not visible.
Right eye was within normal limits (Fig. 4).

Left eye B-scan Ultrasound with 10 MHz probe revealed cataractous lens, short axial length (13mm) and a V-shaped membrane connecting from optic nerve head to the lens associated with retinochoroidal coloboma at peripapillay area (Fig. 5).

Right eye axial length was 19.4mm. Rest of the posterior segment structures were within normal limits (Fig. 6).
Ocular coloboma can involve any part of the eye and is due to defective embryogenesis. The affected eye is most commonly associated with microphthalmia. The prevalence of coloboma among blind adults is around 0.6–1.9%; and among children, it accounts for a greater proportion of blindness (3.2–11.2%). The common cause of coloboma is failure of fetal fissure to close during the 5th to 7th week of fetal life, at the 7-14mm stage. During this period, there is invagination of the optic vesicle and the closure of the fetal fissure. The coloboma is called “typical”, if the inferonasal quadrant is involved and it is the most frequent type to be found. It may involve any part of the globe from the iris to the optic nerve. If the coloboma is located anywhere other than the inferonasal quadrant of the globe, it is termed “atypical”.

A complete iris coloboma is a full thickness defect, involving both the pigment epithelium and the iris stroma. It may be total, extending to the iris root and giving rise to the “keyhole pupil”, or partial, involving only the pupillary margin and causing a slightly oval pupil. Small strands of mesodermal tissue are occasionally seen bridging the coloboma and forming multiple pupils, or polycoria. Such strands may extend to the lens as a persistent pupillary membrane. An incomplete iris coloboma is usually partial thickness, involving either the pigment epithelium or the iris stroma. It tends to be wedge-shaped and is best demonstrated by iris transillumination.

Retinochoroidal coloboma is more commonly associated with visual prognosis. The defect represents an area of bare sclera devoid of overlying normally differentiated retina or choroid. Some retinal vessels may remain and are visible easily. The sclera in the affected area may bulge posteriorly, forming a staphyloma which may involve the entire posterior pole. Histological findings include absence of the RPE, hypoplastic and gliotic overlying retina. Where retinal tissue is recognizable, the retinal layers are reversed, with the rods and cones facing inward and the nerve fiber layer adjacent to the sclera. But, the RPE at the edge of the defect is hyperplastic. Clinically, they can be seen has leukocoria due to exposed sclera. The coloboma can be involving the macula in which case visual prognosis is poor; or sparing macula where visual prognosis is good. The ocular colobomas may be associated with many chromosomal aberrations and syndromes. Environmental causes and intrauterine insults as thalidomide A exposure, intrauterine Vit. A deficiency, use of anti-convulsants in pregnancy, fetal alcohol syndrome, intrauterine infections can also lead to these complications.

The visual prognosis depends mainly on the fact that the optic nerve, macula, and papulomacular bundle is involved or not. If these structures are involved, severe visual loss is inevitable. Bilateral cases present early in infancy with poor visual function and nystagmus. Unilateral cases may develop a sensory strabismus. Retinal detachment and cataract are the most common complications associated with retinochoroidalcoloboma. Of all cases of retinal detachment, 0.6–1.7% is associated with retinochoroidal coloboma.

Proper detailed evaluation of infants with coloboma is really difficult, due to their lack of cooperation, presence of nystagmus and microphthalmos. In fact, effective examination cannot be done without general anesthesia. Retinal, choroidal and optic nerve involvement can be examined with direct and indirect ophthalmoscopy. Accurate refraction should be done, as amblyopia can be induced by anisometropia or by bilateral high ametropia.

Iris coloboma patients can be given cosmetic contact lenses or in severe cases coloboma can be repaired surgically by non-absorbable sutures (Mc-Canal’s iridoplasty). For retinochoroidal coloboma, as it is associated with non-rheumatogenous retinal detachment, prophylactic laser treatment is applied posteriorly along the edge of the coloboma and cryopexy anteriorly.
surgical intervention is at all required, first laser

treatment is done followed by vitrectomy, buckling and
air fluid exchange.

During the development of the eye in the womb,
there is a blood vessel which runs between the optic
nerve and the back of the lens. This blood vessel carries
nutrients and oxygen to the developing parts of the front
section of the eye. The blood vessel is known as the
hyaloid vessel. The vessel, along with surrounding
embryonic material, is also known as the primary
vitreous. In the 3rd trimester (last third) of pregnancy,
the hyaloid vessels are supposed to dissolve, as they are
no longer needed. However, if the blood vessels do not
disappear, this can result in a particular form of cataract
involving the back part of the lens. The condition is
known as Persistent Hyperplastic Primary Vitreous, or
now named as Persistent fetal vasculature. PHPV is
sometimes further divided into subtypes. Anterior
PHPV occurs when the remnant vascular stalk is seen
attached to the back of the lens but no longer extends
back to the optic nerve. They have best chance of visual
rehabilitation. Here the posterior pole is totally normal
with no retinal folds. They present with leukokoria early
in life. There is characteristic elongation of ciliary
processes and anterior chamber may be shallow leading
to glaucoma.

Posterior PHPV occurs when the remnant vascular
stalk is seen arising off the optic nerve but not reaching
the lens and therefore not usually causing cataract.
Posterior PHPV may be associated with developmental
abnormalities of the optic nerve or surrounding retina.
The surrounding retina can be scarred or even detached.
If there is significant involvement of the optic nerve
and/or retina, good vision may not be possible. Most
often, patients have some element of both Anterior

and Posterior PHPV. PHPV is often associated with a small
eye (microphthalmia). In addition, the pupil often does
not dilate well and there may be traction on the tissue
behind the iris (ciliary processes).

Surgical management of PHPV depends on its
type. In patients with anteroposterior PHPV in which
vision is unsalvageable, a lensectomy may be done. In
posterior PHPV, when rehabilitation of the eye is
deemed possible, a vitrectomy may be performed. In
cases of purely anterior PHPV, a lensectomy-
membranectomy and anterior vitrectomy. The goal of
surgery is to salvage useful vision (in concert with
aggressive amblyopia therapy) and/or to prevent or
alleviate glaucoma and to correct amblyopia. Prognosis
depends on the type of PHPV. Treatment of posterior and
combined anteroposterior PHPV has a less favorable
outcome, with most patients attaining perception only of
motion or light. Conversely, in purely anterior PHPV, a
good visual outcome is achieved when aphakic
rection (contact lenses) and amblyopia therapy are
successful.

However, surgery for PHPV is complicated by a
higher rate of retinal detachment than seen with any
other pediatric cataract surgery. The cause of PHPV is
unknown. It is almost always a disorder that affects one
eye. When PHPV is found in both eyes, this may indicate
a syndrome affecting other parts of the body as well.

Conclusion

Patient with microphthalmos associated with
cataract, iris and retinocochrioid coloboma and
persistent fetal vasculature is a complex situation which
should be handled very carefully. The visual prognosis
already remains poor and can sometimes be worsened
with surgical intervention. So, not all cases with cataract
and related complications should be operated. An expert
decision and approach is required to tackle these cases.

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