Bilateral primary hyperplastic persistent vitreous: report of two cases

Abstract

Primary hyperplastic persistent vitreous (PHPV) or persistent fetal vasculature is a rare clinical entity that presents with leucocoria, microphthalmos, and cataract. It is mostly unilateral. Here we present a report of two cases of bilateral PHPV. One of the patients had associated Peters’ anomaly. The entity closely mimics retinoblastoma and should be kept as a differential diagnosis of bilateral leucocoria. Examination under anesthesia, ultrasound B-scan, and aqueous lactate dehydrogenase levels helped us reach the diagnosis and differentiate it from the more serious entity retinoblastoma.

Keywords: PHPV, leucocoria, Peters’ anomaly, B-scan

Introduction

Primary hyperplastic persistent vitreous (PHPV) is the result of the anomalous development of the primary vitreous as it persists into the time of formation of the secondary vitreous [1]. It is a rare clinical entity that presents with leucocoria, microphthalmos, and cataract [2]. It can cause complications like tractional retinal detachment, secondary glaucoma, strabismus, and ambylophia [2]. It is mostly unilateral, and bilateral PHPV is rare [3]. Here we present a report of two cases of bilateral PHPV. One of the patients had associated Peters’ anomaly. The entity closely mimics retinoblastoma and should be kept as a differential diagnosis of bilateral leucocoria.

Case description

Case 1

A male child of 1 month presented with the inability to open the left eye from birth. The child had a full-term normal vaginal delivery with a birth weight of 3,500 grams. On clinical examination, the left eye appeared smaller and there was a bilateral white reflex raising the clinical suspicion of retinoblastoma.

A detailed examination was done under general anaesthesia. In the right eye, horizontal and vertical corneal diameters were 9 mm and 7 mm, respectively; and in the left eye 7 mm and 6 mm, respectively. IOP was 8 mm Hg and the anterior chamber was shallow in both eyes. Fundus examination showed membrane extending from the optic disc in the right eye. In the left eye, no posterior details could be visualized due to total cataract. Aqueous samples were collected and sent for Lactate dehydrogenase (LDH) levels which came to be normal.

A B-scan revealed an ill-defined hyperechoic soft tissue lesion of size 10.3×7.4 mm in the right and 12.6×6.6 mm in the left posterior chamber (Figure 1). The lesion also shows hyper reflection echogenic linear strands with few debris. The bilateral lens appeared normal but the left lens capsule was thickened and compressed by the lesion.
anteriorly. On color Doppler the lesion showed mild vas-
cularity (Figure 2). Based on these clinical findings and
investigations, the diagnosis of bilateral PHPV was made.
The patient’s parents were informed about the pathology,
the visual prognosis and possibility of subsequent com-
plications. The parents refused any further intervention.

Case 2

A three-month-old otherwise healthy male was referred
to our center with suspicion of bilateral retinoblastoma.
Birth history was not significant. On ophthalmological
examination, central corneal opacity was noted in the
right eye. White pupillary reflex was present in both eyes.
A detailed examination under general anesthesia revealed
horizontal and vertical corneal diameters of 9 mm and
8 mm respectively in both eyes. Central corneal opacity
was noted in the right eye (Figure 3a). A clinical diagnosis
of Peters’ anomaly was made in the right eye. Intraocular
pressure was 10 mm Hg in both eyes. Fundus details
could not be visualized due to media haze in both eyes.
Aqueous samples were collected for LDH which came out
to be normal.

A B-scan was done for posterior segment assessment
(Figure 4). It showed dense immobile echoes filling the
posterior segment of the right eye and thickening of
chorioretinal layers. No obvious calcifications were seen.
The left eye showed echogenic membrane extending from
the posterior surface of the lens to the optic nerve head.
Based on these clinical findings and investigations, a
diagnosis of bilateral PHPV with Peters’ anomaly in the
right eye was made. The patient’s parents were made
aware of the nature of the disease and associated
complications. However, they were subsequently lost to
follow-up.

Discussion

PHPV is the result of the anomalous development of the
primary vitreous as it persists into the time of formation
of the secondary vitreous [1]. The primary vitreous is
formed during the 1st month of intrauterine life and starts
regressing during the formation of the secondary vitreous
at the 9th week. By the end of the 3rd month, the secondary
vitreous fills most of the vitreous cavity, and the primary
vitreous condenses into a narrow band (Cloquet’s canal)
running from the optic nerve to the posterior aspect of
the lens [1].

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 [4]. 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 sur-
rounding 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 [2].

Bilateral leucocoria in a child may be due to a number of
causes, of which congenital cataract, retinoblastoma,
retinopathy of prematurity (ROP), and retinal dysplasias
are the most frequent [5]. Cataract is a common cause
of leucocoria in children. It carries a good visual prognosis
if not associated with any retinal pathology and when
treated early enough. However, a primary congenital
cataract was not suspected in our case as the cataracts
in our patients were of the posterior capsular type.
Moreover, a B-scan revealed pathology in the posterior
segment.

PHPV is usually unilateral [5], [6]. Pollard [3], in his study
of 83 cases of PHPV, reported only two cases (2.4%) that
had bilateral PHPV. Bilateral presentations are commonly
associated with systemic and syndromic associations
such as trisomy 13, 15, or 18, Norrie’s disease and
Warburg’s syndrome [7]. However, no systemic anomalies were present in our patients. The differential diagnosis in our cases included retinoblastoma, ROP and PHPV. Normal gestational age, birth weight and uneventful birth history eliminated the possibility of retinopathy of prematurity. Presence of microphthalmia, cataract and normal intraocular pressure made the possibility of retinoblastoma less likely. It was supported by normal aqueous LDH levels, ultrasound B-scan and color Doppler.

One of our patients had Peters’ anomaly in the right eye. This association has been reported twice in the literature in the past. Matsubara et al. [8] described two cases with bilateral Peters’ anomaly and PHPV. They postulated that migratory disorder of neural-crest cells from 4 to 7 weeks of gestation was responsible for those ocular anomalies. Muslubas et al. [9] reported a case of unilateral PHPV with Peters’ anomaly and morning glory syndrome. They suggested the possibility of mutation in the PAX 6 gene for this association.

Conclusion
To conclude, younger age at presentation, presence of microphthalmos, cataract and associated congenital ocular anomalies make the possibility of PHPV more likely in cases of bilateral leukocoria. Typical findings in ultrasonography, color Doppler and normal aqueous LDH levels help in differentiating PHPV from the more serious entity retinoblastoma.

Notes
Competing interests
The authors declare that they have no competing interests.

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