Scheimpflug Imaging in Unilateral Acute Pupillary Block Glaucoma in a Weill-Marchesani Syndrome in an Adult Female

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ABSTRACT
We report an interesting case of Weill-Marchesani syndrome presenting as unilateral acute pupillary block glaucoma after dislocation of crystalline lens into anterior chamber in a middle-aged woman. Scheimpflug imaging demonstrated accurately the dimensions of spherophakic crystalline lens and its corneal lenticular touch. Glaucoma resolved completely after lensectomy and anterior vitrectomy and she achieved best corrected visual acuity of 20/30.

Keywords: Brachydactyly, Pupillary block glaucoma, Scheimpflug image, Spherophakia, Weill-Marchesani syndrome.

INTRODUCTION
The syndrome of Weill-Marchesani (WMS) is a rare hereditary affection, retrieved for the first time by Weill in 1932,1 then described in 1936 by Marchesani.2 Alternatively, it has been named spherophakia-brachymorphia syndrome, or congenital mesodermal dysmorphodystrophy. Diagnostic criteria of WMS includes: (1) Short stature, (2) brachydactyly, (3) microspherophakia and/or ectopia lentis.3 The principal complication is the secondary glaucoma that is frequent in the absence of treatment. Spherophakia is an uncommon condition in which the small, spherical lens may leads to pupillary block and secondary angle closure glaucoma.4,5 We present a case of WMS presenting as unilateral acute pupillary block glaucoma in a middle-aged woman documented with Scheimpflug image.

CASE REPORT
A 40-year-old Sikh woman presented with acute pain and decreased vision in right eye since 5 days. She had a history of treatment with oral acetazolamide, 4% pilocarpine drops in right eye for last 5 days. There was no light perception in the left eye for last 13 years following similar complaint and history of lensectomy in left eye 6 years back following which symptom got relieved. At presentation, the right eye had a visual acuity of counting finger close to face (CFCF) and no light perception in left eye; intraocular pressure (IOP) was 50 and 34 mm Hg in the right and left eye respectively. There was diffuse epithelial corneal edema. Spherical dislocated lens was present in the anterior chamber touching corneal endothelium in the right eye (Fig. 1A). The left eye had steel suture at the superior limbus, vascularized leucomatous corneal opacity inferiorly, few epithelial bullae and stromal edema. There was aphakia with anterior capsular opacification, retained lens matter. Retinoscopy was not possible owing to the poor fundal glow due to corneal edema in right eye and vascularized corneal opacity in left eye. The axial length was 24.58 and 26.19 mm in the right and left eye respectively. B-scan ultrasonography showed normal posterior segments in both eyes. The optic disk and posterior segment was normal in right eye and advanced glaucomatous cupping was present in left eye.

Scheimpflug image obtained using Oculus Pentacam of right eye showed spherical lens touching corneal endothelium as well as subluxated from patellar fossa. The lens thickness was 5,000 microns and distance between posterior lens surface and patellar fossa was 1,470 microns (Fig. 1B). Scheimpflug imaging showed corneal thickness varying between 653 and 865 micron. Right eye keratometry was 45.8 D at 90º and 50 D at 180º.

Systemic Findings
Patient’s height was 142 cm (150 ± 12 cm) and weight 60 kg. Electrocardiogram was normal. There was brachydactyly (Fig. 1C) and no joint hypermobility.

The patient was given intravenous injection mannitol 20%, 350 ml stat followed by systemic acetazolamide 250 mg four times a day, topical timolol maleate 0.5% twice-a-day and brimonidine 0.15% thrice-a-day. The IOP reduced to 28 and 20 mm Hg in the right and left eye respectively after treatment.

Patient underwent right eye lensectomy with anterior vitrectomy with peripheral iridectomy from limbal route with vitreous cutter and no IOL implanted.

At first postoperative day, the best corrected visual acuity (BCVA) in right eye was 20/40 and IOP was 24 mm Hg.
There was mild corneal edema, few Descemet’s folds. Pupil was mid dilated and there was a peripheral iridectomy at 9 o’ clock (Fig. 1D), the optic disk was normal with cup-disk ratio of 0.2 and healthy neuroretinal rim. Twelve weeks later BCVA of right eye was 20/30 [+2.5 DS with rigid gas permeable (RGP) contact lens and base curve of 7.2 mm and diameter 9 mm] and IOP was 17 mm Hg with topical timolol maleate 0.5% twice-a-day with CD ratio of 0.2 and healthy optic disk. Left eye IOP was 24 mm Hg on topical timolol maleate 0.5% twice-a-day and brimonidine 0.15% twice-a-day and patient was symptom free in the left eye. Postoperative gonioscopy showed the open angles in right eye and left gonioscopy not possible due corneal opacity.

DISCUSSION

This is the first reported case of microspherophakia presenting as unilateral acute pupillary block glaucoma in an adult female and Scheimpflug image demonstrated accurately the dimensions of spherophakic crystalline lens and its corneal lenticular touch.

Microspherophakia is usually associated with systemic disorders, such as WMS, homocystinemia, Marfan’s syndrome, Alport’s syndrome and Klinefelter’s syndrome. Our patient had features suggestive of WMS with brachydactyly, short stature and spherophakia with acute pupillary block glaucoma. Glaucoma in spherophakia can result from several mechanisms: Pupillary block by the spherical lens, irritation of the ciliary body by the dislocated lens, or by complete luxation of the lens into the anterior chamber. Unrelieved pupillary block may lead to peripheral anterior synechiae (PAS) formation and irreversible trabecular damage. Chronic pupillary block without complete angle closure may lead to crowding of the trabeculae by the spherophakic lens. Our patient presented with unilateral acute pupillary block glaucoma, which was worsened by miotics and relieved to some extent by cycloplegic treatment. Our patient had right eye acute glaucoma secondary to pupillary block which was relieved after lensectomy and anterior vitrectomy. The IOP in our patient remained controlled with timolol maleate medication postoperatively and patient achieved good visual outcome.
REFERENCES


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