Generalised nevus flammeus, episcleral capillary malformation and glaucoma

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DESCRIPTION
A 33-year-old woman presented with no vision, redness, pain and photophobia in the left eye (LE) since 4 months. This was associated with left-sided headache and vomiting too. She had history of poor vision in LE since 2–3 years, and had been treated with topical antiglaucoma medication. There was no significant family history. On examination, visual acuity was 6/6 in right eye (RE) and no perception of light in LE. Intraocular pressure (IOP) was measured as 14 mm Hg in RE and >60 mm Hg in LE. Slit lamp evaluation revealed conjunctival congestion, corneal oedema, fully dilated pupil and white cataract in LE (figure 1). Gonioscopy was within normal limits. RE ocular examination did not reveal any significant anomaly. RE pupil did not show any consensual response, while LE pupil was fixed. Systemic workup revealed nevus flammeus (NF) over the face involving both periorical regions but more prominent on the left side. Further examination showed the NF to be present all over the body (trunk and limbs), and was more prominent over the left side. The capillary malformation was dark purple in colour and non-blanchable to external pressure. Plaque-like changes and underlying soft tissue hypertrophy were noticeable in some areas (figure 1). No other pigmentary changes were seen. On closer ocular examination, the episcleral capillaries around the limbus were found to be engorged, with telangiectasia and focal bulbous dilations (figure 1).

The patient was diagnosed to have a generalised NF with end-stage glaucoma in LE. Hyperosmotic agents and oral acetazolamide were advised while the systemic workup proceeded. Ultrasound of the LE revealed an anechoic vitreous cavity and no signs of choroidal tumours. No anomalies were detected on radioimaging of head, chest and abdomen. She was also evaluated by a dermatologist, physician and gynaecologist, but no other systemic anomalies were found. A 270° transconjunctival cryoablation of the ciliary body was done, and 15 days following the procedure the IOP was 18 mm Hg on a topical combination of dorzolamide and timolol. As the patient was asymptomatic only on topical medication, further surgical interventions were deferred.

NF, also called port wine stain, is a capillary malformation well known to involve the head and neck area. The capillary ectasia may be progressive. Syndromic involvement is well known in the form of Klippel-Trenaunay syndrome, Sturge-Weber syndrome, Cobb syndrome, Proteus syndrome, Rubinstein-Taybi syndrome and others.1 It may simulate vascular tumours in infancy or young children. Glaucoma is also a well-documented association, and may be seen in as much as one-fifth of these eyes.2 The causes include raised episcleral venous pressure due to the capillary malformation, associated intraocular tumours and angle anomalies. A large number of such eyes, close to half, need surgical procedures including cyclodestruction for IOP control as done in our case.2

Capillary malformations may be classified depending on whether they represent naevi or not.3 NF is believed to be a type of malformation representing naevi. It can be generalised and involve the rest of the body apart from the face, and can also have predilection for involvement of one side of the body only. While generalised NF is known to be associated with glaucoma as a part of syndromes or phacomatoses like phacomatosis pigmentovascularis, solitary ocular involvement with generalised NF is rare.

Figure 1 Upper half of the figure shows nevus flammeus to be present on the face, upper limbs and lower limb. The lesions are more prominent over the left side of the body. Lower half of the figure shows the perilimbal episcleral capillary malformation and total cataract in the left eye.
Learning points

► Generalised port wine stain can occur with glaucoma in absence of systemic anomalies. However, systemic involvement must be ruled out as a multidisciplinary approach.
► Cyclodestruction remains a useful technique for managing such painfully blind eyes with absolute glaucoma.

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