Port-wine stain and glaucoma in a 29-year-old male

Karen Kate Quilat,1 Eshir A Ismael1,2

A diagnosis of Sturge-Weber Syndrome (SWS) is made when two out of three criteria—facial port-wine birthmark, increased ocular pressure, and leptomeningeal angiomatosis—are present.1 The facial lesion is a hamartoma that arises from vascular tissue, producing the characteristic port-wine hemangioma of the skin along the trigeminal nerve distribution.2 3

Glaucoma occurs in up to 70% of patients with SWS and is usually diagnosed during infancy, but it can develop later during adolescence or adulthood.4 For late-onset glaucoma, the initial management consists of topical aqueous suppressants and miotics. If topical medications fail, trabeculectomy is the procedure of choice. A cyclodestructive procedure targeting the secretory epithilium of the ciliary body may be performed on eyes with failed medical and surgical interventions.4

A 29-year-old man came to our clinic due to eye pain and redness of four years’ duration, associated with occasional episodes of headache. There were no accompanying seizures or other neurological symptoms reported. Physical examination revealed a left-sided, flat, well-defined violaceous red patch within the dermatome distribution of the ophthalamic branch of the trigeminal nerve, with irregular borders extending from the left upper eyelid inferiorly to the hairline above the frontal area superiorly, and from one centimeter medial to the left inner canthus medially to the left outer canthus laterally (port-wine stain; Figure 1A). The patient had visual acuity of 20/20 on both eyes.

We found more significant findings on the left eye. Intraocular pressure was 30 mmHg. Slit lamp biomicroscopy revealed dilated and toruous perilimbal vessels (Figure 1B). Gonioscopic examination revealed open anterior chamber angles on all quadrants. On funduscopy, the optic nerve had a cup-to-disc ratio of 0.7 (Figure 1C). Retinal vessels were noted to be dilated and toruous. The rest of the ophthalmologic findings were unremarkable. The patient, having port-wine stain and glaucoma, was diagnosed to have Sturge-Weber syndrome.

The patient was initially given timolol eyedrops to control the intraocular pressure (IOP). However, IOP ranged from 24-30 mmHg over a 1-month period. Automated perimetry revealed a temporal quadrantanopsia on the left eye. Both the increase in cup-to-disc ratio and temporal quadrantanopsia were highly suggestive of progressing optic nerve damage and visual field defect on the affected eye.

The patient underwent trabeculectomy on the left eye. On the first postoperative day, the IOP went down to 13 mmHg, the conjunctival bleb was formed and located superonasally, the anterior chamber was shallow, and visual acuity was 20/100. Two weeks after trabeculectomy, the anterior chamber deepened and visual acuity returned to 20/20. A repeat automated perimetry after trabeculectomy revealed no progression of the scotoma (Figure 1D).

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REFERENCES
Figure 1  Flat, well-defined, violaceous red patch with irregular borders (port-wine stain) on the left frontal area (A). Dilated and tortuous perlimbal vessels in the left conjunctiva (B). Cup-to-disc ratio of 0.7 and tortuous and dilated vessels on funduscopy of the left eye (C). Report of automated perimetry test showing left temporal quadrantanopsia (D).