Polypoidal choroidal vasculopathy-associated vitreous haemorrhage presenting as hyphema

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DESCRIPTION
A 59-year-old Indian male patient presented to our clinic with sudden-onset, painless diminution of vision in the right eye for the last 1 week. Records from previous follow-up (2 months back) showed that his vision was 20/200 right eye (OD) and 20/80 left eye (OS). He had no known systemic medical conditions. He was diagnosed as a case of right eye pseudophakia with polypoidal choroidal vasculopathy (figure 1A), as evidenced by his previous fundus fluorescein angiography (FFA) (figure 1B), indocyanine green angiography (ICGA) (figure 1C) and swept source optical coherence tomography (SSOCT) (figure 2A,C). He has been on regular follow-up for the same condition for the past 4 years. During the course, he had received four intravitreal antivascular endothelial growth factor (anti-VEGF, aflibercept 2 mg in 0.05 mL) injections, with the last injection being administered 4 months back.

On ocular examination, his best-corrected visual acuity was hand movement close to face OD and 20/80 OS. Intraocular pressure was 16 mm Hg OD and 14 mm Hg OS. On slit lamp examination, presence of hyphema (figure 3A) was noted in the right eye obscuring the details of the rest of the anterior segment. The anterior segment of the fellow eye was unremarkable. Ocular ultrasonography showed the presence of moderate amplitude spikes in the vitreous area suggestive of haemorrhage in the right eye (figure 3B). Dilated fundus evaluation of the fellow eye showed a sharply delineated area of geographic atrophy (hypopigmentation with apparent absence of retinal pigment epithelium (RPE) (figure 4A). Fundus autofluorescence imaging of the same showed sharply demarcated area of hypopautofluorescence (figure 4B) with window defect on FFA due to atrophy of the underlying RPE (figure 4C). ICGA excluded the presence of any secondary choroidal neovascular membrane in the left eye (figure 4D). SSOCT of the left eye (figure 2B,D) revealed thinning of the hyper-reflective external band due to RPE/Bruch’s membrane complex attenuation. Thus, we made a diagnosis of right eye hyphema with vitreous haemorrhage (VH) secondary to polypoidal choroidal vasculopathy and left eye geographic atrophy.

In view of the normal intraocular pressure and the absence of endothelial staining, the patient was medically managed for hyphema with topical steroids (prednisolone acetate 1%), mydriatic (homatropine 2%) and antiglaucoma (timolol 0.5%), and was given propped-up positioning. The patient is being followed up weekly. A surgical pars plana vitrectomy has been planned as the next step after the clearing of hyphema.

Polypoidal vasculopathy (PCV) is a disease of pachychoroidal spectrum primarily involving the choroidal circulation. It is characterised by the aneurysmal bulge of the inner choroidal vessel endings, which are clinically visible as reddish-orange polyps. Spontaneous rupture of these venules and occasionally arteries is not uncommon, and may lead to massive breakthrough subretinal and/or vitreous haemorrhage causing a sudden, painless...
This can occur as the initial presentation of the disease or during the course of follow-up. PCV-related breakthrough VH can also develop following photodynamic therapy (PDT), intravitreal injection of anti-VEGF agents, pneumatic displacement of submacular haemorrhage, or combined therapy with PDT and anti-VEGF agents. The occurrence of breakthrough VH, subretinal haemorrhage and suprachoroidal haemorrhage is known in PCV, but anterior chamber hyphema secondary to PCV-related VH has not previously been described. We presume that the amount of breakthrough VH in our case was extensive enough to pass through the weakened zonules in pseudophakic eye and appears in the anterior chamber. In this case, the breakthrough bleed was likely a spontaneous event, and not triggered by intravitreal injection that was administered 4 months before the bleed. VH due to intravitreal injection mostly occurs early (within 1–4 days of drug administration). There are variable reports on the visual outcome in patients undergoing vitrectomy for VH in PCV. Visual recovery after pars plana vitrectomy is mainly limited by macular involvement and the presence of subfoveal or juxtapfoveal polyps.

Learning points

- Polypoidal choroidal vasculopathy with vitreous haemorrhage can rarely present as hyphema.
- Thorough work-up is warranted to identify the underlying cause.
- Management of hyphema is usually conservative; subsequent pars plana vitrectomy is needed for clearing of vitreous haemorrhage.

Contributors

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