Introduction

Retinal vasculitis is a poorly characterized, potentially sight-threatening, inflammatory ocular condition that occurs when there is the presence of abnormal blood vessels in the retina. The annual incidence of retinal vasculitis in the United States is estimated at 1-2 per 100,000 with variation between regions. Because it is not well understood, the definition of “retinal vasculitis” is not well established. Currently, the Standardization of Uveitis Nomenclature Working Group considers “perivascular sheathing and vascular leakage/occlusion on fluorescein angiogram as evidence of retinal vascular disease” in classifying retinal vasculitis [1]. There are two main causes of retina vasculitis: primary ocular vasculitis, occurring as an isolated condition or secondary retinal vasculitis, caused by an inflammatory systemic condition. Common systemic diseases associated with retinal vasculitis include Behcet’s disease, sarcoidosis, multiple sclerosis, systemic lupus erythematosus, Wegener granulomatosis, polyarteritis nodosa and other rheumatologic conditions. Infectious agents may also cause retinal vasculitis such as syphilis, tuberculosis, and Lyme disease. Common diseases associated with isolated conditions of retinal vasculitis include primary retinal vasculitis, Eales disease, pars planitis, birdshot retinochoroidopathy, and Fuchs uveitis syndrome. Many autoimmune manifestations have been seen with retinal vasculitis when no systemic diseases are present; however, the exact etiology is unclear [1,2].

Clinically, patients with retinal vasculitis most commonly present with blurred vision, altered color perception, distortion of images, floaters and scotomas. However, sometimes patients may present with no visual symptoms. Retinal vasculitis can be detected using ophthalmologic examinations and fluorescein angiography, which can show perivascular sheathing, vascular leakage, inflammatory cells of the vitreous body, narrowing of the retinal blood vessels, and neovascularization. Late leakage or staining of retinal vessels is also evident on FA. Sometimes, mild anterior or posterior uveitis may be seen. The presence of choroidal inflammation may indicate sarcoidosis or birdshot retinochoroidopathy [2,3]. Active vasculitis causes the formation of exudates around retinal vessels, which results in white sheathing or cuffing of the vascularization. This can lead to vascular leakage, which causes retinal swelling and macular edema that can greatly affect visual acuity and lead to vision loss. Visual acuity prognosis is difficult to anticipate due to complications such as macular ischemia, branch and central retinal vein occlusion, persistent neovascularization and vitreous hemorrhages.
A 27-year-old female from Saudi Arabia presents with decreased vision in her left eye for a year. She denies any recent illnesses, infections or trauma and is currently on no medications. She states that she has had difficulty accessing medical care. A review of systems indicated a cough, fever and night sweats. The patient also presented with joint pain and swelling, along with orogenital ulcerations.

On exam, her vision was 20/200 OD and 20/50 OS with symmetrically reactive pupils and normal intraocular pressures. Anterior segment exam was normal. However, posterior segment exam of the right eye showed peripheral vessel sheathing with peripheral non-perfusion. The left eye showed extensive neovascularization of the optic disc, central and peripheral non-perfusion, along with intraretinal microvascular anomalies (Figure 1). Fluorescein angiography of both eyes revealed significant late leakage of the retina vessels with the left eye to a greater degree (Figure 2b). Spectral Domain OCT showed appropriate foveal contour with no macular edema or subretinal fluid (Figure 3). The following laboratory tests were performed: CBC with peripheral smear, PPD, Quantiferon, ACE, lysozyme, ESR, RPR, ANA, and a chest x-ray. All findings were negative. The patient was also found to be negative for diabetes, hypertension, HIV and TB.

**Discussion**

In diagnosing primary retinal vasculitis, there should be a differential diagnosis based on a detailed history, review of systems, ophthalmic and physical examination. If the patient’s workup suggests an underlying systemic cause, then further examinations and laboratory tests should be performed. The absence of any diagnostic clues from the workup and a negative medical history for systemic diseases associated with retinal vasculitis would suggest primary retinal vasculitis. However, the pathophysiology of primary retinal vasculitis is often closely aligned with Eales disease. Eales disease is also associated with isolated conditions of retinal vasculitis, most often seen in young men. Characteristic features include inflammation of the peripheral retinal vasculature causing retinal phlebitis that can lead to retinal non-perfusion, neovascularization and recurrent vitreous hemorrhages. It is also highly associated with individuals with tuberculosis. The natural progression of Eales disease begins with inflammation, leading to occlusion and then lastly, the presence of neovascularization. Other clinical presentations seen in Eales disease include exudative vasculitis, cotton-wool spots, macular edema, and anterior uveitis. Although the differentiation between primary retinal vasculitis and Eales disease is often unclear, there are certain distinctions seen in each of the diseases. The patient was ultimately diagnosed with primary retinal vasculitis due to the location of her abnormal retinal vasculature being more posterior, sectorial and exudative. Disease presentation is also seen equally in both genders while in Eales disease, it is more prominent in males. The patient also exhibited peripheral sheathing and non-perfusion in both eyes. In individuals with Eales disease, peripheral vascular inflammation only occurs at a minimum while retinal vascular occlusions are more common [3,5]. Eales disease is also often associated with positive TB patients, which was not seen in this case. Nevertheless, the diagnostic workup and treatment for primary retinal vasculitis and Eales disease are similar.

Due to disease course variation, the poor outcome of primary retinal vasculitis is dependent on a number of factors. To control inflammation of the retinal vasculature, the main treatment option for primary retinal vasculitis often involve the use of corticosteroids with or without immunosuppressive therapy. Cyclosporine and
azathioprine have shown to be effective in reducing inflammation. Anti-tuberculosis treatment is given for positive TB patients, which is more often seen in individuals with Eales disease. Laser photocoagulation is often combined with bevacizumab intravitreal injections to treat neovascularization and prevent further retinal ischemia [1,6]. Our patient responded well to panretinal photocoagulation of the left eye and scatter photocoagulation of the right eye. Additionally, the extensive neovascularization of the disc regressed with two treatments intravitreal bevacizumab over a 6-month period. The prognosis of patients with primary retinal vasculitis is variable due to the heterogeneous nature of the disease. Some patients that undergo treatment have full preservation of their ocular function and vision. However, others may have permanent loss of function despite treatment. Current status of patient is unknown due to her return to Saudi Arabia.

References