Bilateral retinal vasculitis in a patient with lichen planus

**Vasculite retiniana bilateral em um paciente com líquen plano**

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**ABSTRACT**

Lichen planus (LP) is an autoimmune, inflammatory disease of unknown etiology that commonly affects the skin and mucous membranes. Retinal vasculitis is a group of vision-threatening disorders, in which autoimmunity is thought to play a role in pathogenesis. We present the case of a patient who was diagnosed with retinal vasculitis and who was followed up for mucosal LP. LP has not been reported as a cause of retinal vasculitis in the literature. We believe that the retinal vasculitis in this case was related to LP because cellular immunity plays a role in the pathogenesis of both entities.

**Keywords:** Retina, Retinal vasculitis, Lichen planus, Fluorescein angiography

**INTRODUCTION**

Lichen planus (LP) is an autoimmune, inflammatory disease of unknown etiology that commonly affects the skin, nails, hair, and mucous membranes in middle-aged adults[1]. Retinal vasculitis comprises a group of disorders characterized by retinal vascular inflammation accompanying by intraocular inflammation, and autoimmunity is thought to play a role in its pathogenesis. Diagnosis is made with ophthalmoscopy and fundus fluorescein angiography (FA) in cases of retinal vasculitis that threaten vision. Retinal vascular sheathing, vitritis, and accompanying cystoid macular edema are among the typical clinical findings, and staining and leakage of the retinal vessels on FA are characteristic in active disease[2-2]. LP has not been linked to the development of retinal vasculitis in the literature. Here, we present the case of a patient with LP with no additional disorder who developed bilateral retinal vasculitis, possibly due to LP.

**CASE REPORT**

A 44-year-old male patient presented to our clinic with symptoms of floaters in both eyes for a week. The best corrected visual acuity (BCVA) was 0.8 on the right eye and 0.9 on the left, and intraocular pressures were normal. The case had been followed up for mucosal LP for the last 10 years at another center. Anterior segment examinations were normal within the first year follow-up; therefore, the vitritis and peripheral vascular sheathing had regressed. BCVA fully recovered bilaterally at the first month of follow-up, and the vitritis and peripheral vascular sheathing had regressed. BCVA and anterior and posterior segment examinations were normal within the first year follow-up; therefore, the immunosuppressive treatment was discontinued (Figure 3).

**DISCUSSION**

LP is an autoimmune, inflammatory disease that affects the skin, skin appendages such as the hairs and nails, and the oral, vulvovaginal, esophageal, laryngeal, and conjunctival mucosa[3]. The res-

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responsible antigen for this T cell-dependent immunologic disease has not been identified. Although the oral mucosa is generally affected by the type of LP that affects the mucosae, esophageal and ocular involvement may remain undetected. Serious eye problems related to the decrease in tear production may develop, and the patients have a relatively low number of conjunctival epithelial goblet cells. Blepharitis, lacrimal duct obstruction, eyelid lesions, keratouveitis, keratoconjunctivitis sicca, punctate epithelial erosions, corneal ulcer/scarring, and ocular surface squamous neoplasms such as dysplastic conjunctival lesions can also be seen in addition to xerophthalmia and cicatrizating conjunctivitis. An isolated ocular form of LP has also been reported, and biopsy and immunofluorescence studies may be required to enable differentiation of this form from keratoconjunctivitis, which can also cause conjunctival scarring.

Many inflammatory and infectious agents are included in the etiology of retinal vasculitis, and a proper diagnostic work-up should be performed in these patients. However, in certain cases, no etiologic reason can be identified, a condition described as primary retinal vasculitis. Some of these patients may later develop a systemic disease known to cause retinal vasculitis. Disorders that can cause retinal vasculitis fall into three groups, including systemic, infectious, and ocular disorders. Behcet’s disease, sarcoidosis, multiple sclerosis, systemic lupus erythematosus, and Wegener’s granulomatosis are the most common systemic diseases that can cause retinal vasculitis. The most common infectious diseases include toxoplasmosis, tuberculosis, Lyme disease, syphilis, cat scratch disease, and herpes simplex and varicella zoster infections, whereas ocular diseases include intermediate uveitis, Birdshot choroidopathy, and Eales disease. CD4+ T cell-dependent immunity is thought to play a main role in the pathogenesis of retinal vasculitis, but humoral immunity and immune complex formation may also be important.

LP has not yet been reported in the literature as a cause of retinal vasculitis. Cellular immunity is known to play an important role in the pathogenesis of both retinal vasculitis and LP. In the present case, LP was suspected as the etiologic reason that the patient developed bilateral retinal vasculitis. To the best of our knowledge, this is the first case report on a patient developing retinal vasculitis due to LP.

REFERENCES