Case report

Herpes simplex virus bullous keratitis misdiagnosed as a case of pseudophakic bullous keratopathy with secondary glaucoma: an unusual presentation

Sreedharan Athmanathan⁎1, Mittanamalli S Sridhar2, Raj Anand2, Anil K Mandal3 and Gullapalli N Rao2

Address: 1Jhaveri Microbiology Center, Hyderabad Eye Research Foundation, LV Prasad Eye Institute, Hyderabad, India, 2Cornea services, LV Prasad Eye Institute, Hyderabad, India and 3VST Center for Glaucoma Care L.V. Prasad Eye Institute, Hyderabad, India

E-mail: Sreedharan Athmanathan⁎ - sreedhar@lvpeye.stph.net; Anil K Mandal - mandal@lvpeye.stph.net; Gullapalli N Rao - gnrao@lvpeye.stph.net

⁎Corresponding author

Abstract

Purpose: To report an unusual case of herpetic bullous keratitis misdiagnosed as a case of pseudophakic bullous keratopathy with secondary glaucoma.

Results: A retrospective analysis of the case record of a 60-year-old man who had earlier undergone bilateral cataract surgery, was done. He presented with a complaint of decrease in vision in the right eye of 20 days duration. On examination, cornea showed epithelial bullae all over the surface with stromal and epithelial edema. Intraocular pressure was 30 mm of Hg in RE. He was treated with anti-glaucoma medications. Two dendritic lesions were seen in the cornea during a subsequent visit four days later. Virological investigations confirmed a diagnosis of Herpes simplex keratitis. He was treated with topical acyclovir.

Conclusions: This case highlights the fact that herpes simplex keratitis can present initially as a more diffuse corneal stromal and epithelial edema with epithelial bullae mimicking bullous keratopathy. Herpetic bullous keratitis, although unusual, should be considered in the differential diagnosis under such circumstances.

Introduction

Herpes simplex keratitis (HSK) is a sight threatening ocular infection and is a leading cause of corneal blindness [1]. Clinical presentation of HSK is often protean. While a typical and common presentation of HSK is usually a dendritic or geographic ulcer, atypical presentations are not uncommon [2]. We report here an unusual presentation of HSK. The patient presented to us with bullous keratitis, which was misdiagnosed as a case of pseudophakic bullous keratopathy (PBK) with secondary glaucoma.

Case Report

A sixty-year-old male presented to our cornea services with a complaint of progressive diminution of vision, in the right eye, of 20 days duration. There were no other ocular or systemic complaints. He gave a history of having undergone extracapsular cataract extraction with
posterior chamber intraocular lens implantation 4.5 years earlier in the RE and phacoemulsification with posterior chamber intraocular lens implantation 2 years earlier in the LE. Surgery and postoperative period were uneventful on both occasions with a final visual acuity of 6/6 in both eyes. On examination, visual acuity in RE was restricted to perception of light with accurate projection of light in all the four quadrants. Conjunctiva was congested and cornea showed epithelial bullae all over the surface with mild to moderate epithelial and stromal edema. Anterior chamber was deep and quiet. Intraocular lens was in place and fundus details were within normal limits. Intraocular pressure in the RE was 30 mm of Hg and LE was 14 mm Hg. A diagnosis of PBK with secondary glaucoma, was made. Patient was immediately treated with intravenous mannitol (350 cc) and a single oral dose of 250 mg acetazolamide, followed by 250 mg three times daily and 0.5% timolol eye drops twice daily for the right eye. Intraocular pressure was 16 mm of Hg following mannitol administration and he was discharged. Patient presented to us four days later. Visual acuity in the RE had improved to counting fingers at 1 m and 6/6 in LE. On examination, cornea showed few epithelial bullae with mild to moderate epithelial and stromal edema. The intraocular pressure in the right eye was 16 mm Hg. Two dendritic lesions were seen (Figure 1) in the cornea. A clinical diagnosis of PBK with HSK was made. Corneal scrapings were collected for virological investigations. The patient was treated with 3% acyclovir eye ointment five times daily and cyclopentolate eye drops twice daily for the RE. The patient was lost to follow up.

Papanicolaou stained smear of the corneal scraping showed multinucleated giant cells and intranuclear eosinophilic inclusion (Figure 2). HSV-1 antigen was detected in the epithelial cells of the corneal scraping and the smear revealed multinucleated giant cells (Figure 3). HSV-1 was isolated in culture and PCR was positive for HSV DNA using primers, which amplified a 179 bp region of the DNA polymerase gene of HSV 1/2.

**Discussion**

This case was initially diagnosed as PBK with secondary glaucoma based on a history of cataract surgery, presence of diffuse corneal stromal and epithelial edema, epithelial bullae and a raised intraocular pressure in the affected eye. The subsequent appearance of typical dendritic lesions and virological investigations confirmed a diagnosis of herpetic bullous keratitis. Further, the present event occurred after 4 years after the cataract surgery. Since corneal latency of HSV has been described [4], it is perfectly reasonable to assume that herpes rather than the previous endothelial damage caused the whole syndrome.

The presence of glaucoma possibly suggests HSV trabeculitis in the affected eye. It is difficult to ascertain this finding since we did not perform any investigation. A PCR assay for the detection of HSV DNA using aqueous humor would have provided supporting evidence.

This event was possibly a syndrome of HSV stromal and epithelial keratitis with trabeculitis, which explains the signs of a diffuse corneal stromal and epithelial edema.
Figure 3
Immunoperoxidase assay of the corneal scraping: Corneal scraping showing multinucleated giant cells and the presence of HSV-1 antigen (Seen as brown precipitate) (×500).

and an acute rise in the intraocular pressure in the affected eye. The formation of epithelial bullae due to corneal edema could have predisposed the cornea for the development of dendritic ulcers. It has earlier been shown that the presence of corneal epithelial bullae has a statistically significant effect on the rate of ulcer development [3].

This case highlights the fact that HSK can present initially as a more diffuse corneal stromal and epithelial edema with epithelial bullae mimicking bullous keratopathy.

To the best of our knowledge and based on a MEDLINE search, such a presentation of HSK has not been documented.

Herpetic bullous keratitis, although unusual, should be considered in the differential diagnosis under such circumstances to prevent further complications and for the prompt institution of specific antiviral therapy.

Declaration of competing interests
None declared

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