Tuberculous corneal ulcer with hypopyon: A case report

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This case report represents an unusual presentation of Mycobacterium tuberculosis-associated corneal ulcer with hypopyon. A 64-year-old man presented with a hypopyon corneal ulcer in the right eye. His corneal scrapings were negative, and the ulcer was not responding to empirical antimicrobial therapy. A real-time polymerase chain reaction from an aqueous aspirate of the right eye detected 422 copies of M. tuberculosis. Subsequent investigations revealed positive Mantoux and interferon-gamma release assay, supportive radiology. The patient was successfully treated with a course of antitubercular therapy and topical steroids. Hypopyon corneal ulcer, though a rare manifestation of ocular tuberculosis, should be considered in cases not responding to standard antimicrobial therapy.

Key words: Antituberculosis therapy, corneal ulcer, hypopyon, ocular tuberculosis, tuberculosis, uveitis

The clinical manifestations of ocular tuberculosis are diverse and protean. The diagnosis of ocular tuberculosis always remains challenging because of the difficulty in ocular sample collection considering the delicate structure of the eye. Due to the paucibacillary nature of the infection, live Mycobacterium tuberculosis is rarely isolated from tissue samples. Thus, hypotheses on immune-mediated inflammation and direct bacteria-mediated inflammation in the pathogenesis of intraocular tuberculosis have been advocated. Over the last few years polymerase chain reaction (PCR) has emerged out as a valuable tool for the diagnosis of infections from various tissue samples and has been able to fill this gap in the diagnosis of ocular tuberculosis partially.

Corneal involvement in tuberculosis is rare. Peripheral corneal ulceration, limbal nodule, and stromal keratitis have been described in the literature. In the majority of the cases with corneal involvement even in an endemic country like India, possibility of M. tuberculosis infections comes last in the list of differential diagnoses. Similarly, in a patient presenting with hypopyon corneal ulcer, ocular tuberculosis is never considered, though there are several reports of ocular tuberculosis presenting as hypopyon uveitis. TB-causing immune stromal keratitis with hypopyon has been reported once in the past. We report a case of hypopyon corneal ulcer, where an association with M. tuberculosis infection was established with the help of real-time polymerase chain reaction (RT-PCR) and subsequent investigations revealed positive Mantoux test, interferon-gamma release assay (IGRA), and pulmonary involvement on radiography.

Case Report

A 64-year-old male presented to our outpatient clinic with a sudden increase in redness, pain, and watering in the right eye for the past 1 month. He was a known diabetic under antidiabetic treatment for the past 8 years. He had received a diagnosis of infective keratitis elsewhere and underwent a series of preliminary investigations, which also included microbiologically negative corneal scraping. His blood investigations were within normal limits, except for raised erythrocyte sedimentation rate (ESR). The ocular condition got worse in spite of being on intensive topical antimicrobial medications. On examination, best-corrected visual acuity (BCVA) in the right eye was 6/24 and 6/6 in the left eye. Slit-lamp examination of the right eye revealed mid-stromal infiltrate with overlying epithelial defect along with the adjacent superficial limbal vascularization. Examination revealed cells 1+, flare 1+ with trace hypopyon [Fig. 1a]. Slit-lamp and fundus examination of the left eye were unremarkable. View of fundus in right eye was hazy, but grossly it looked normal. A repeat corneal scarping was not helpful in isolating any microorganism. An anterior chamber paracentesis was done, and his aqueous aspirate was sent for microbiological evaluation and PCR was done for eubacterial, pan fungal genus, herpes group viruses, and M. tuberculosis. RT-PCR detected 422 copies/mL DNA of M. tuberculosis. Based on his PCR report, patient was thoroughly evaluated for systemic tuberculosis. Mantoux test was positive with induration of 15 mm and IGRA was also positive. High-resolution computerized tomography (HRCT) of chest was suggestive of granulomatous sequelae. His serum angiotensin enzyme and serum lysozyme were within the normal limit and serology for syphilis and human immunodeficiency viruses (HIV) were negative. We considered a provisional diagnosis of immune stromal keratouveitis secondary to ocular tuberculosis. The patient...
Phlycten, the corneal edema responded to topical steroid and ATT in consultation with chest physician. The patient was closely monitored and there was a complete resolution of the corneal ulcer within a month of the initiation of the treatment [Fig. 1b]. The patient underwent phacoemulsification with the implantation of the intraocular lens in the right eye after 6 months of resolution of inflammation [Fig. 1c]. A repeat RT-PCR on aqueous aspirate, collected during cataract surgery, did not detect any copies of \textit{M. tuberculosis} genome. His BCVA improved to 6/9 following surgery. He is in follow-up with us for last 1 year and after the treatment, till now, he did not have any recurrence of inflammation.

**Discussion**

The involvement of conjunctiva, cornea, and sclera are rare though but not uncommon in tuberculosis.\cite{1,3} Phlycten, interstitial keratitis, and conjunctival granulomas have been described as hypersensitivity diseases of cornea and conjunctiva.\cite{4,5} A hypersensitivity reaction to tubercular protein has been implicated primarily rather than direct involvement of \textit{M. tuberculosis}. However, direct involvement by the bacilli is not uncommon and has been reported widely in the literature.\cite{1,4} Corneal involvement in ocular tuberculosis includes interstitial keratitis,\cite{7} disciform keratitis,\cite{3} and stromal keratitis.\cite{9} Similarly, the reports on hypopyon uveitis in association with tuberculous infection are increasing in number.\cite{3,10,11} The present case illustrates the protean corneal manifestations of tuberculous corneal ulcer. It was RT-PCR report that raised our suspicion toward tuberculous etiology in our case and was confirmed subsequently by positive Mantoux test, positive IGRA, pulmonary involvement in HRCT of chest.

Bilateral interstitial keratitis and granulomatous anterior uveitis were reported to be the initial manifestation in a 17-year-old girl. A presumptive diagnosis of ocular tuberculosis was achieved on the basis of raised ESR, positive Mantoux test and IGRA.\cite{7} The corneal edema responded to topical steroid and ATT. Delayed type of hypersensitivity reaction to the probable tubercular antigen in aqueous humor was implicated by the authors. Arora \textit{et al}.\cite{3} described bilateral disciform keratitis as presenting manifestation of extrapulmonary tuberculosis in a 11-year-old girl. Subsequent investigations revealed tubercular Pott’s spine, and treatment with topical steroid and ATT shows considerable resolution of corneal lesion.\cite{9} In another separate case report, multiple disc-shaped stromal infiltrates with overlying corneal ulcers with non-granulomatous uveitis in both eyes and hypopyon were reported in an 11-year-old female by Singhal \textit{et al}. The patient had raised ESR, positive Mantoux test, and radiographic evidence of pulmonary tuberculosis, and finally her ocular lesions responded to topical steroid and ATT. Similar to this case, our case initially did not respond to anti-infective treatment and initial microbiological investigations were negative. There are many reports of hypopyon uveitis associating ocular tuberculosis in the literature.\cite{3,10,12} RT-PCR was able to detect and quantify the presence of \textit{M. tuberculosis} genome, which we believe was not reported previously.

To conclude, the possibility of ocular tuberculosis should be considered in the cases with corneal ulcer not responding to the standard antimicrobial therapy, and PCR can be added as a valuable tool in the diagnosis of such cases.

**Declaration of patient consent**

The authors certify that they have obtained all appropriate patient consent forms. In the form the patient(s) has/have given his/her/their consent for his/her/their images and other clinical information to be reported in the journal. The patients understand that their names and initials will not be published and due efforts will be made to conceal their identity, but anonymity cannot be guaranteed.

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**Conflicts of interest**

There are no conflicts of interest.

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Strabismus associated with Crouzon syndrome is common and often complex. V-pattern strabismus is most commonly reported in this condition and is mainly thought to be due to an excyclorotation of the orbits and rectus muscle pulleys. Various theories have been proposed as causes of the V-pattern. V-pattern horizontal strabismus has been reported as the most common deviation in Crouzon syndrome. V-pattern strabismus with Crouzon syndrome, including dysmorphic orbits and misalignment. He had undergone an expansion cranioplasty and trans-suture distraction osteogenesis at 3 years of age.

A 4-year-old boy with Crouzon syndrome and a confirmed heterozygous mutation in the fibroblast growth factor receptor 2 (FGFR-2) gene (c.1030G>C) was referred for ocular misalignment. He had undergone an expansion cranioplasty and also considered as a contributing cause of strabismus. Various extraocular muscles in Crouzon syndrome with V-pattern exotropia who had rectus muscle heterotopy on orbital imaging and were also found intraoperatively to have anomalous insertion. A computerized tomography (CT) scan of the orbit confirmed the presence of all rectus muscles and superior oblique muscle and also considered as a contributing cause of strabismus.

Case 1

A 9-year-old boy presented with a long history of right exotropia and misdirection, and fibrosis of extraocular muscles were noted. This highlights the various causes of V-pattern strabismus associated with Crouzon syndrome, including dysmorphic orbits and also considered as a contributing cause of strabismus. The right eye had an exotropia of 35 prism diopter (PD) and a left hypotropia of 6 PD in primary position with mild left inferior oblique overaction. On presentation, his visual acuity was 20/30 in each eye, and +1.25 diopters of hyperopia in both eyes. Motility examination revealed bilateral inferior oblique overaction. He had undergone an expansion cranioplasty and trans-suture distraction osteogenesis at 3 years of age.

He had an exotropia of 35 prism diopter (PD) and a left hypotropia of 6 PD in primary position with mild left inferior oblique overaction. On presentation, his visual acuity was 20/30 in each eye, and +1.25 diopters of hyperopia in both eyes. Motility examination revealed bilateral inferior oblique overaction. He had undergone an expansion cranioplasty and trans-suture distraction osteogenesis at 3 years of age.

However, at the time of surgery, a bifid left lateral rectus muscle insertion was found and a thin left inferior rectus muscle was also planned to correct the left hypotropia. A recession of lateral rectus muscle and inferior oblique myectomy in both eyes were planned via limbal incisions to correct the V-pattern exotropia, and a recession of left inferior rectus muscle was also planned to correct the left hypotropia.

Postoperatively, he had a left hypotropia of 6 PD in primary position and +1.25 diopters of hyperopia in both eyes. Rectus muscle heterotopy on orbital imaging and were also found intraoperatively to have anomalous insertion. A computerized tomography (CT) scan of the orbit confirmed the presence of all rectus muscles and superior oblique muscle and also considered as a contributing cause of strabismus.

In Case 2, a 7-year-old boy presented with right exotropia and misdirection, and fibrosis of extraocular muscles were noted. This highlights the various causes of V-pattern strabismus associated with Crouzon syndrome, including dysmorphic orbits and also considered as a contributing cause of strabismus. The right eye had an exotropia of 35 prism diopter (PD) and a left hypotropia of 6 PD in primary position with mild left inferior oblique overaction. On presentation, his visual acuity was 20/30 in each eye, and +1.25 diopters of hyperopia in both eyes. Motility examination revealed bilateral inferior oblique overaction. He had undergone an expansion cranioplasty and trans-suture distraction osteogenesis at 3 years of age.

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