Over-Treated Corneal Abscess May Be Toxic Keratopathy

P. Rubino    J.G. Orsoni    A. Rampini    P. Mora
Institute of Ophthalmology, University of Parma, Parma, Italy

Key Words
Toxic keratopathy · Corneal abscess · Corneal mycosis

Abstract
Background/Aims: Keratitis, especially when long-standing and unresponsive to common antimicrobial treatment, leads to a suspicion of fungal aetiology.
Methods: Photographically documented case report.
Results: A 65-year-old man with diabetes was referred for corneal abscess unresponsive to antibiotic and antifungal treatment lasting 6 weeks. Corneal biopsy was performed following a 72-hour washout for identification of bacteria and fungi. Previously administered drops were withdrawn and only preservative-free artificial tears were maintained. Neither bacteria nor fungi were cultured. After 2 weeks, the clinical situation had conspicuously improved.
Conclusion: Over-treatment of corneal affections fearing mycosis may lead to toxic keratopathy.

Toxic keratopathy due to varying aetiologies has been well known for decades [1–4]. Ophthalmologists, desperate when faced by long-standing ocular inflammation unresponsive to treatment, tend to over-treat corneal affections with every conceivable therapy. Particularly worrying is fungal keratitis. We describe here a typical example of this situation.

Case Report
A 65-year-old Italian farmer with orally treated diabetes was referred to us for a left eye corneal abscess unresponsive to topical antibiotic and anti-inflammatory treatment lasting 6 weeks. The patient had suffered a cow-tail trauma 20 years earlier in the same eye. This gave rise to a central corneal ulcer and poor vision (20/160, measured with the Early Treatment of Diabetic Retinopathy Study (ETDRS) chart). It resulted in central leucoma with mild vascularisation. After a more recent trauma, the patient was hospitalised due to a corneal abscess in the same eye. Owing to the lack of response to various antimicrobial local drugs, i.e. tobramycin 0.3% drops 8 times/day and ampicillin 0.8% drops 4 times/day, even associated with steroid eyedrops, i.e. dexamethasone 0.3% eyedrops, all with preservatives, fungal infection was suspected despite negative corneal scraping. This suspicion was (psychologically) strengthened in the treating ophthalmologist for three reasons: the corneal ulcer had...
been caused by a mild agricultural trauma, the patient was diabetic, and the clinical history and the corneal picture were suggestive of fungal keratitis. Before referral to us, systemic and local ketoconazole 0.4% (3 times/day) and topical levofloxacin 0.5% (4 times/day) had been unsuccessfully administered for two weeks. Thus, we were asked to perform extreme antifungal treatment with corneal intrastromal voriconazole [5]. At admission, severe conjunctival hyperemia with 360° corneal neovascularisation, a central, ulcerative, abscess-like keratopathy with a satellite lesion, and hypopyon were evident (fig. 1). A corneal biopsy was performed following a 72-hour washout for identification of bacteria and fungi. Previously administered drops were withdrawn and preservative-free artificial tears and cycloplegics were given instead. Neither bacteria nor fungi were found. Over the next 2 weeks, the clinical picture improved and resolved 4 weeks later: conjunctival hyperemia was significantly reduced, hypopyon and corneal infiltrates had disappeared, and the corneal stroma appeared more transparent (fig. 2). The clinical picture was similar to the one described clinically after healing from the cow trauma.

Comment

In clinical practice, the ophthalmologist is often tempted to treat ocular inflammation without a working diagnosis and microbiological confirmation, especially if coping with a poorly cooperative patient. In this specific case, very probably, an asymptomatic central corneal epithelial defect persisted after the earlier cow-tail trauma. Clinical signs of toxic keratopathy are usually so nonspecific that a diagnosis can only be confirmed by withdrawing suspected medications and drops with preservatives.

Fig. 1. Aspect of the RE at admission.
Fig. 2. Aspect of the RE four weeks after admission.
References


