Voriconazole in Treatment of Resistant Fungal Keratitis

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ABSTRACT
Purpose: To study the effect of intrastromal Voriconazole for the treatment of resistant fungal keratitis in a tertiary care eye hospital in Lahore, Pakistan.
Study Design: Experimental interventional study.
Place and Duration of Study: Avicenna Medical College Hospital, Lahore, from July 2017 to July 2019.
Methods: Sixty four patients were selected. All patients with fungal keratitis were included. The patients with previous corneal scar, mature cataract, endophthalmitis, Panophthalmitis, scleral involvement, impending or frank corneal perforation and uncontrolled diabetic patients were excluded. Corneal scrapings of all patients were sent for 10% KOH staining. All patients were given intrastromal Voriconazole at 3 to 4 sites in divided doses in one ml syringe with 27-gauge needle. Injection was repeated on 4th and 8th day. It was combined by topical antifungal and antibiotic eye drops six hourly. Patients were followed at day two, five, nine, three weeks and at 3 months.
Results: There were 55 males and 9 females. Average size of ulcer was 6.4 mm ranging from 5.5 mm ± 1.8 mm. Fifty six (88%) patients showed improvement while eight (12%) patients ended up in melting of cornea which was managed with tectonic corneal graft. In three (5%) patients penetrating Keratoplasty was done. Conjunctival congestion and ocular pain improved significantly one week after third dose but final visual acuity was not significantly improved due to scarring.
Conclusion: Intrastromal corneal voriconazole is an effective treatment for fungal keratitis in term of healing of the corneal ulcer, control of infection and prevention of corneal perforation and permanent blindness.
Key Words: Voriconazole, Fungal keratitis, Penetrating Keratoplasty.

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INTRODUCTION
Fungal keratitis is challenging disease, which is difficult to diagnose and treat. Once colonized in the cornea, fungi have the propensity to penetrate deeper corneal layers and if they get access to anterior chamber of eye then control of infection is extremely difficult.¹ Cornea is an avascular structure and restricted defense mechanisms make it easy for fungi to colonize. These are usually found in soil, water and on plants. The important risk factors in developing fungal keratitis are trauma with vegetative material, contact lens wear and immunocompromised status.²³⁴⁵ The prevalence of fungal keratitis is more in warm climate. Commonly implicated organisms in developing warm countries are Fusarium and Aspegillus.⁶⁷ The rapid development of fungal...
keratitis leads to visual loss so early diagnosis is essential to prevent long term ocular complications. Poor outcome of fungal keratitis than bacterial keratitis is due to poor penetration and limited availability of antifungal drugs.

Fungus was first documented in 1879 and accounts for 40 to 50% of all cases of keratitis. There are some 70 different types of fungi but two are more relevant in ophthalmology which are yeast and filamentous fungi (septate and non-septate).

Fungal keratitis is very difficult to treat. Many a times, anti-fungal drugs given in the form of eye drops are not sufficient. The purpose of our study was to find out the effectiveness of intrastromal Voriconazole in the treatment of fungal keratitis.

METHODS
There were 64 patients included in the study. Study duration was from July 2017 to July 2019. All patients with clinical diagnosis of fungal keratitis were included either from outpatient department (OPD) or were referred from other medical centers. Patients’ age, gender, size of ulcer and visual acuity were noted. The patients with previous corneal scar, mature cataract, endophthalmitis, Panophthalmitis, scleral involvement, impending or frank corneal perforation and uncontrolled diabetic patients were excluded. Clinical features of fungal keratitis included ocular pain, photophobia, decreased vision, satellite corneal lesions, corneal edema, haze and hypopyon. Corneal scrapings were sent for 10% KOH staining in all patients. Voriconazole is an antifungal drug available in 200 mg vial, which needs dilution and once reconstituted it needs refrigeration. It can be used for 7 to 10 days after dilution. All patients were given intrastromal injection. The dose was 50 micro-lit/ml at 3 to 4 sites, in divided doses, in clear cornea around the lesion. It produced hydration of corneal stroma around the lesion. One ml syringe with 27guage needle was used for injection, with needle bevel down wards in corneal stroma. It was given at first, 4th and 8th day of presentation. Moderate to severe pain was common complaint by all patients which was treated by oral NSAIDS in all patients at the time of injection. Intrastromal injection was combined by topical antifungal. Voriconazole eye drops 1mg/ml and antibiotic eye drops, Moxifloxacin 0.5% both QID for 4 weeks. Patients were followed-up on day two, five, nine, three weeks and 3 monthly post treatment. Visual acuity was noted at each visit. Patient was asked about pain and slit lamp examination was done to note size of lesion, resolution of hypopyon and epithelization of defect.

No systemic side effects of the drug were detected. No case of endophthalmitis or Panophthalmitis was observed. The patients were followed-up for 6 months.

RESULTS
Sixty four patients were selected, fifty five were males and nine were females. Mean age was 32 ± 8 years and age range was 18 to 60 years. Corneal scrapings were sent to lab for 10% KOH staining. Only 37% showed positive staining and 63% were negative. Fifty six patients (88%) responded well (47 males and 8

Fig. 1: Corneal Ulcer Before (Left) and After Treatment (Right).
females) to intrastromal Voriconazole, in terms of
decrease in size of corneal infiltrates, improvement in
ocular pain, resolution of corneal edema and healing of
corneal ulcer with scarring of cornea. Eight (12%)
patients (7 males and 1 female) did not respond and
ended up in melting of cornea which was managed
with tectonic corneal graft. Out of these eight, only
three patients required Penetrating Keratoplasty for
visual restoration. Conjunctival congestion and ocular
pain improved significantly one week after third dose
of intrastromal Voriconazole but final visual acuity
was not significantly improved due to scarring and
most patients had counting finger vision after
resolution of keratitis. No patient needed evisceration
during 6 months follow-up period.

DISCUSSION
There are different anti-fungal drugs; which include,
Polyenes, Imidazoles, Triazoles and Fluorinated
Pyrimidines. Commonly used antifungal drugs are;
Natamycin, Amphotericin B, and Voriconazole. These
drugs are used as topical and systemic therapy for
treatment of fungal keratitis. We found intrastromal
anti fungal drug delivery most effective as it achieved
targeted drug delivery.

Higher incidence of fungal keratitis in male
patients may be due to our social set up where males
are more exposed to outdoors than women and it was
comparable with other studies done by Al-Hatim et al
and CH Cho et al.

Ideally every corneal scraping should be sent for
PCR and culture for diagnosis. PCR takes only two to
three hours and culture takes up to 35 days. We did
not perform these tests in our study.

Natamycin 5% eye drops belong to polyene group
and it was the first approved antifungal agent. It
inhibits transport of amino acids and glucose across
fungal plasma membrane by binding with ergosterol
leading to cell damage, but it is used only as a topical
drug as negligible oral absorption makes it unfit for
systemic drug. Voriconazole belongs to Triazole
group, fungistatic and fungicidal, available for oral and
parenteral use and metabolized in liver. It inhibits
fungal cytochrome P-450 3A dependent enzymes and
inhibits ergosterol synthesis, which is the principal
sterol in cell wall of fungus and inhibits cell membrane
synthesis. It is effective against Candida, Aspergillus,
Fusarium, Scedosporium and Paecilomyces. It is
effective in fungal keratitis resistant to Polyenes and
first line Triazoles. It is also used as alternative to
Amphotericin B in fungal endophthalmitis.

Recent studies show that intrastromal use of
voriconazole has produced better results in term of
control of fungal infection and healing of ulcer.
Ganapathy K showed that Intrastromal voriconazole
helped to resolve the infection in 18 (72%) patients
and about 15% of these needed more than one
injection. Smaller ulcers responded better to treatment.
Fusarium species were responsible for six of the seven
cases. According to Namrata Sharma, of 12 eyes, 10
eyes healed with scar formation, and the mean
resolution time was 39.75 ± 7.62 days. Two corneas
perforated and required therapeutic penetrating
keratoplasty. Other studies also showed that
Voriconazole has potential to achieve adequate drug

Fig. 2: Corneal Ulcer Before (Left) and After Treatment (Right).
concentration at the site of infection through a targeted drug delivery.\textsuperscript{20–22}

Intrastromal Amphotericin B is used in the same fashion as Voriconazole. Average healing time in our study was 21 days which was comparable with studies in other countries. A study in India by Kalaiasseli et al, showed mean resolution time of resistant fungal keratitis in 25 patients was 17 days.\textsuperscript{18} Another study by Sharma et al. showed healing time of 39 ± 7 days.\textsuperscript{19}

In our study 5\% patients showed no response to treatment and ended up as candidate of penetrating Keratoplasty. Literature shows that 30\% patients with fungal keratitis develop corneal perforation or do not respond to topical antifungal therapy.\textsuperscript{23–25} Penetrating Keratoplasty is expensive treatment but if intraocular contents are not involved it can result in complete eradication of infection.

Limitation of our study are that we did not compare our results with any other antifungal drug, single center study and small sample size.

\section*{CONCLUSION}

Our study showed that intrastromal Voriconazole is effective in fungal keratitis in term of good healing of corneal ulcer, control of infection, saving eye from corneal perforation, evisceration and loss of vision. No systemic side effects of drug were observed.

\section*{Ethical Approval}

The study was approved by the Institutional review board/Ethical review board.

\section*{Conflict of Interest}

Authors declared no conflict of interest.

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Authors’ Designation and Contribution
Khalid Mehmood; Professor: Concepts, Data Acquisition, Literature research, Manuscript preparation, Manuscript review.

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Tariq Khan; Professor: Literature research, Manuscript preparation, Manuscript review.

Mahfooz Hussain; Assistant Professor. Manuscript writing, Critical review.

Sara Riaz; Assistant Professor. Manuscript editing, Manuscript review.