Diagnosis and Management Protocol of Acute Corneal Ulcer

Vaishal P Kenia¹, Raj V Kenia², Onkar H Pirdankar³

¹Kenia Eye Hospital, Mumbai
²Kenia Foundation, Mumbai
³Kenia Medical and Research Foundation, Mumbai

Corresponding Author: Onkar H Pirdankar

ABSTRACT

Corneal ulceration is one of the leading causes of corneal blindness. Various pathogens are responsible for corneal ulceration. Accurate and quick diagnosis and prompt treatment is a key to improve clinical and visual outcomes in cases of corneal ulceration. However there are no specific guidelines or protocols are available for managing the corneal ulcers. Sometimes even an experienced clinician struggle to predict the course of the disease in most of the cases. Here we make an attempt to provide an overview on diagnostic approach and management protocol of acute corneal ulcer.

Key words: Acute corneal ulcer, corneal ulceration, corneal blindness

INTRODUCTION

Corneal ulceration is one of the major ocular emergencies causing ocular morbidity. It is considered a leading cause of corneal blindness especially in the developing countries. It has been estimated that globally, corneal ulceration with ocular trauma are resulting in 1.5 -2 million cases of corneal blindness annually ¹. It affects males more than the females, can be seen at any age and mostly the patients belong to a low socio-economic strata ²,³. In developing countries like India, the major reason for corneal ulcer is ocular trauma whereas in the developed countries, the most common reason is contact lens wear.

Etiology and Epidemiology:

The causes of a corneal ulcer can be infections like bacteria, viruses, fungi or parasitic (Acanthamoeba). Whereas the non-infectious causes are autoimmune, neurotrophic, toxic, allergic keratitis, chemical burns, keratitis secondary to entropion, trichiasis, blepharitis, lagophthalmos.

Infectious causes of corneal ulceration vary based on geographic location. The common microorganism are Fusarium species, Pseudomonas aeruginosa, Aspergillus spp., S. Pneumoniae, Staphylococcus Spp. Fungal organisms are common in Countries like India, China. In other countries such as Philippines, Taiwan, Thailand and Singapore bacterial organisms such as Pseudomonas aeruginosa is common. Table 1 describes the common microorganisms, by counties ⁴.

<table>
<thead>
<tr>
<th>Common organism</th>
<th>Total</th>
<th>IND</th>
<th>CH</th>
<th>SG</th>
<th>PH</th>
<th>JP</th>
<th>TH</th>
<th>KR</th>
<th>TW</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bacterial</td>
<td>38.0%</td>
<td>38.3%</td>
<td>15.3%</td>
<td>41.3%</td>
<td>53.2%</td>
<td>47.8%</td>
<td>50.5%</td>
<td>42.8%</td>
<td>61.8%</td>
</tr>
<tr>
<td>Fungal</td>
<td>32.7%</td>
<td>45.6%</td>
<td>30.2%</td>
<td>0.7%</td>
<td>27.0%</td>
<td>6.3%</td>
<td>9.1%</td>
<td>10.0%</td>
<td>8.2%</td>
</tr>
<tr>
<td>Parasitic</td>
<td>2.4%</td>
<td>2.1%</td>
<td>0.9%</td>
<td>1.3%</td>
<td>5.7%</td>
<td>5.0%</td>
<td>4.0%</td>
<td>1.2%</td>
<td>6.9%</td>
</tr>
<tr>
<td>Viral</td>
<td>12.6%</td>
<td>6.9%</td>
<td>46.2%</td>
<td>7.0%</td>
<td>2.0%</td>
<td>16.6%</td>
<td>6.1%</td>
<td>6.0%</td>
<td>8.2%</td>
</tr>
<tr>
<td>Infectious Keratitis (Not Specified)</td>
<td>14.5%</td>
<td>7.1%</td>
<td>4.4%</td>
<td>49.7%</td>
<td>12.1%</td>
<td>24.3%</td>
<td>30.3%</td>
<td>40.0%</td>
<td>15.0%</td>
</tr>
</tbody>
</table>
Table I describes common organism found in Asian countries (IN-India, CH-China, SG-Singapore, PH-Philippines, JP-Japan, TH-Thailand, KR-South Korea, TW-Taiwan)

Pathogenesis:

Bacterial:
Bacterial corneal ulcers are results from the penetration of bacteria after a breach in the corneal epithelial barrier except organisms like gonococcus can penetrate an intact epithelium to cause ulcer. Factors predisposing the epithelium like corneal edema, prolonged contact lens usage, dry eyes, and trauma make it vulnerable to corneal infection. The most common microorganism is Pseudomonas Aeruginosa which utilizes glyocalux to adhere to epithelium and then invades into the stroma through breach in epithelial. Inflammatory cells (PMNs) reach the site of corneal breach from the tears and limbal vessels which releases cytokines and interleukins resulting in progressive invasion of cornea and increase in size of the ulcer. Phagocytosis of the organism releases the free radicals and proteolytic enzymes leading to necrosis and sloughing of the epithelium, bowman’s membrane and stroma. In addition, process is facilitated by proteases and exotoxin that are produced by multiplying bacteria and endotoxin that are produced by organisms after their death. The endotoxins are polysaccharides within the cell wall of gram negative bacteria and are responsible for ring infiltrates.

Viral:
In viral ulcer, the virus reaches the cornea from within via terminal branches of ophthalmic division of the trigeminal nerve. It has been postulated that in case of herpes simplex there is an involvement of the sub basal nerves which results in epithelial swelling whereas in case of herpes zoster there is an involvement of deep stromal nerves. So without epithelial breach the virus reaches the eye via nerve endings and the inflammation of nerve causes neurogenic pain. The virus actively replicates in corneal epithelium. The virus in the epithelium form raised lesion forming superficial punctate keratitis and then slough to form large epithelial defect and eventually stromal ulceration.

Parasites such as acanthamoeba:
Acanthamoeba keratitis is most commonly associated with soft contact lens use. Once it is adherent to the contact lens, it survives in the space between the contact lens and the ocular surface and later gets attached to the glycoproteins on the corneal villi. The microtrauma to the corneal epithelial surface due to contact lens use promotes the entry of the organism into the epithelium, the invade Bowman's layer and enter the stroma. The infection then moves along the corneal nerves, produces acute inflammation and radial deposits (radial keratoneuritis). The acute inflammation produces metalloproteases that digest collagen fibrils and allows deeper penetration into the stroma. As the disease progresses, it may penetrate the anterior chamber and can cause endophthalmitis.

Symptoms:
- Reduced visual acuity,
- Tearing,
- Discharge,
- Redness are the common symptoms presented by the patients.
- Pain (Disproportionate pain can be seen in Herpes and Acanthamoeba. Fungal ulcers are quieter whereas pseudomonas are fast growing)
- Diagnostic Approach:

Careful History:
It is very important to keep in mind the TRIAD of ocular trauma, Lowered immune status (either the ocular surface or the individual as a whole) or extremely virulent organisms that penetrate the intact ocular surface. A corneal ulcer cannot develop in a healthy individual with a healthy ocular surface, in the absence of ocular trauma. In this respect, a detailed history focussed on finding the cause of an ulcer in the patient is very important so as to ensure an appropriate management. A history of ocular trauma, ocular surgery, long-term use of ocular medications (topical steroids, Anti-glaucoma medications), contact lens wear (age of contact lens and lens cleaning solution), and previous ocular infections is important as all these factors alter the ocular surface milieu and promote microbial invasion of the cornea in the absence of trauma. Similarly, systemic diseases such as diabetes, rheumatoid arthritis, hepatitis, auto-immune diseases and their therapy, tuberculosis, malignancy impair the natural immune status of an individual and predisposes to opportunistic infections, unusual microbes, fungi or viruses.

Thorough Slit lamp Biomicroscopy:
A thorough slit lamp examination is useful to evaluate the clinical signs may be helpful to confirm the probable diagnosis. Figure 1 briefly describes the diagnostic approach for acute corneal ulcer.

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**Diagnostic Approach with Corneal Ulcers Patients**

**Presentation: History**
- Reduced Visual acuity, tearing, discharge, redness
- Pain (disproportionate in case of Herpes and Acanthamoeba)

**Risk Factors:**
- External: Corneal trauma, Contact Lens (CL) wear, contaminated CL solution
- Ocular Factors: Ocular surface and adnexal disorders, Dacryocystitis, corneal epithelial disease
- Systemic Factors: Long term steroid use, diabetes Mellitus, Kidney Failure, HIV

**Slit Lamp Biomicroscopy**

**Infiltrate Features**
- Bacterial ulcer: Necrotic stroma, purulent discharge and hypopyon
- Fungal Ulcer: Stromal Infiltrate with feathery borders
- Viral Ulcer: Dendritic pattern with progressive geographic and Amoeboid configuration
- Acanthamoeba Ulcer: Stromal infiltrates with ring shaped configuration and hypopyon

**Severity Factors**
- Size of infiltrate (<2mm or >2mm)
- Location of Infiltrate (central Versus peripheral)
- Depth of Infiltrate (<50% or > 50%)
- Involving limbus, sclera
- Associated with AC reaction
- Hypopyon

**Initial Medical treatment Approach**
- Severe ulcers need intense therapy and follow ups
- Anti fungal and Acanthamoeba treatment after smear report and strong clinical judgement

**Laboratory Investigation**

<table>
<thead>
<tr>
<th>Procedure</th>
<th>Sample</th>
</tr>
</thead>
<tbody>
<tr>
<td>Conventional Superficial Scraping and Culture</td>
<td>Gienstain, KOH, Calcofluor White,</td>
</tr>
<tr>
<td></td>
<td>Reduced AFB stain eg Nocardia</td>
</tr>
<tr>
<td>Culture</td>
<td>Blood Agar, Sabouraud agar, Special</td>
</tr>
<tr>
<td></td>
<td>Non nutrient Agar with E. Coli</td>
</tr>
<tr>
<td>Confocal Microscopy</td>
<td>For Deep Infiltrate</td>
</tr>
<tr>
<td>For Acanthamoeba and Fungal Infection</td>
<td></td>
</tr>
</tbody>
</table>

**Figure 1: Diagnostic Approach with Corneal Ulcers Patients**
Slit lamp examination starts with:

1. Eyelid assessment for Blepharitis, meibomian glands dysfunction, ectropian/entropian, lagophthalmos.
2. Eyelashes assessment for trichiasis/distichiasis
3. Lacrimal apparatus system assessment for punctal abnormalities, dacryocystitis
4. Conjunctiva assessment for discharge, inflammation, foreign body, papillae, follicle, cicatization, symblepharon, pseudomembrane, filtering bleb, tube erosion
5. Sclera assessment for any nodule, thinning
6. Cornea assessment for epithelial defects, punctate keratopathy, stromal edema, ulceration, thinning, perforation, infiltrate characteristics (size, shape, location, depth), foreign body, sign of previous corneal surgeries. Fluorescein or rose Bengal staining allow clinicians to identify some organism or underlying cause. For example in cases of viral infections dendritic ulcers are stained with fluorescein and rose Bengal stain.
7. Anterior chamber assessment for presence of any inflammation, look for cells and flare, hypopyon, hyphema

**Signs:** 6,7

Although there are no specific signs to identify the responsible organisms, a careful slit lamp assessment with clinical experience help reach probable diagnosis. Various factors such as size, shape, location of Infiltrate, involvement of limbus, sclera, associated with AC reaction and hypopyon gives information about how aggressive the infection is.

**Bacterial:**
- Gram positive infection (Figure 2):
  - Localized infiltrate with distinct borders
  - Minimal stromal haze

**Gram negative infection (Figure 3):**
- Dense stromal suppuration/ Ring Infiltrate
- Hazy surrounding cornea with a ground glass appearance.

**Fungal:**
- Dry raised slough, with a dried appearance of surrounding cornea which is clear.
- Stromal infiltrate with feathery edges,
- Satellite lesions and thick endothelial exudates, (Figure 4)
- Sometimes presents with ring infiltrate
- Hypopyon usually forms which is convex upwards and may wax and wane.

Viral:
- Viral ulcers can result from herpes simplex or herpes zoster infections
- Viral ulcer can be seen in the form of dendritic pattern (linear branching) due to central desquamation.
- The end of the branches manifest a characteristically swollen appearance.
- It generally gets stained with fluorescence.
- Anterior stromal infiltrate appear under the ulcer but resolves spontaneously.
- Corneal sensation is reduced.
- Progressive centrifugal enlargement may result in larger epithelial defect with a geographical and amoeboid configuration. (Figure 5)

Parasitic
A. Acanthamoeba:
- Acanthamoeba keratitis can be contact lens or non contact lens related
- Characterized by epithelial irregularities, corneal edema, with single or multiple stromal infiltrates which has classic ring shaped configuration (Figure 6)
- However diffuse and satellite infiltrates are also common.

- Hyopopyon is also a common finding in acanthamoeba.
- In late cases radial keratoneuritis can also be noted and can be identified as whitish outline of the corneal nerves 6,7.

Figure 5 showing three phases of lesions: epithelial dots (9 o'clock), dendritic pattern (6 o'clock) and a geographic epithelial keratitis (12–2 o'clock), suggesting herpes simplex virus epithelial keratitis.(Image From Gurav P et al 2015 5)

Microbiological Culture and Light Microscopy: 8,9
Traditionally clinicians were heavily dependent on light microscopes, corneal smears and cultures. Conventional smear and culture for bacteria, fungus and Acanthamoeba can be prepare by scraping the base and leading edge of the corneal ulcer using flame sterilized Kimura spatula or sterile surgical blade no 15 on Bard Parker Handle. Every scraping can be use for direct microscopic examination, culture and antibiotic susceptibility testing. These scraping are immediately placed on glass slides for light microscopy and agar plates for culture (Blood agar, chocolate agar, Potato dextrose agar (PDA), Sabouraud agar.
The slides for light microscopy are stained with 10% potassium hydroxide or gram stain or Giemsa stain to aid in the visualization of fungal filament, bacterial or Acanthamoeba cyst growth respectively. Special staining such as modified Ziehl Neelsen for nocardia, microsporadia and KOH or calcoflour white staining for acanthaomeba and fungus can be use. For culture the agar plates are inoculated at 25-27degree C for 7 days in case of (PDA) whereas in case of other media it is inoculated at 35-37 degree C (2 days for blood agar) and microorganism growth is assessed on daily basis. Cultures of contact lens, lens case and contact lens solution can also be done in case of contact lens wearers.

**Corneal Biopsy and deep Stromal Culture technique:**

Corneal biopsy is indicated if the infiltrate is located in the mid or deep stroma with overlying uninvolved tissues. Corneal biopsy can be performed at the slit lamp biomicroscope or operating microscope. After instillation of topical anaesthetic, a small trephin e or blade is used to excise a small piece of stromal tissue at the edge of the infiltrate which can be sent for culture as well as histopathology.

**Antimicrobial Susceptibility:**

Antimicrobial susceptible testing of the isolates is performed by Kirby Buaer Disk Diffusion method using ciprofloxacin (5 µg), ofloxacin (5 µg), gatifloxacin (5 µg), tobramycin (10 µg), chloramphenicol (30 µg), amikacin (30 µg), gentamicin (10 µg), moxifloxacin (5 µg) as per Clinical and Laboratory Standards Institute Guidelines. Disk diffusion method assesses antibiotic sensitivity of bacteria. It uses antibiotic discs to evaluate the extent to which bacteria are affected by those antibiotics. Antibiotic susceptibility does not necessarily directly reflex clinical susceptibility.

**Confocal Microscopy:**

Confocal microscopy has played a crucial role in the diagnosis of microbial keratitis, fungal and acanthamoeba keratitis 8,10,11. With advancement in technology, direct visualization of pathogens within the patients cornea is possible. In Vivo confocal microscopy is non invasive technique available in clinical settings. To best of our knowledge there are presently two modalities available for clinical use are scanning slit IVCM (Confoscan, Nidek Technology, Fremont, CA) and laser scanning IVCM (HRT3 with Rostock corneal module, Heidelberg Engineering, Heidelberg, Germany). On confocal microscopy acanthyamoeba cyst can be identified as double walled ovoid bodies and fungal bodies were seen as bright linear filamentous structures with bright borders that appear as parallel lines (double walled linear bodies)10.

**Treatment protocols**

In majority of the cases the infection is resolved without any acute surgical need. However surgical intervention is required irrespective of infection is resolved or not resolved.2. Initiation of treatment is based on clinical judgement, smear report and the treatment is modified according to culture report and clinical response. It has also been reported that use of topical cortico steroid is controversal hence they are best avoided.

**Medical Treatment:**

Antibiotic, antifungal or antiviral eye drops are the treatments of choice however Antifungal and Acanthamoeba therapy started only after microbiological evidences, in most cases. The line of medical treatment and the route of treatment is decided based on the depth, size and location of infiltrates. Central infiltrate would require more aggressive treatment as compared to peripheral, superficial<2mm infiltrate. Deep intrastromal infiltrate would require intrastromal injections as it gives good drug availability at deeper layer.

**Bacterial Keratitis:**

Topical antibiotics (Monotherapy) can be in given in cases of superficial peripheral infiltrates< 2mm. For deep stromal involvement or an infiltrate larger than 2 mm with extensive suppuration a loading dose every 5-15 minutes followed
by frequent applications such as every hour is recommended. In case of monotherapy, Levofloxacin 1.5% is preferred over Gatifloxacin and Moxifloxacin due to emerging resistance with Gatifloxacin and Moxifloxacin and easier availability of Levofloxacin. In cases of large or visually significant infiltrate or severe infection in the presence of a hypopyon, topical fortified antibiotics is preferred. Systemic antibiotics are rarely required, however they can be considered in severe cases where infection involves limbus and sclera. Role of corticosteroid in treating the bacterial ulcer is controversial. The SCUT treatment study found no benefit of concurrent topical corticosteroid therapy using prednisolone sodium phosphate 1% in conjunction with broad spectrum topical antibiotics. A pervious study have reported no benefits of corticosteroids in managing corneal scars.

Table 2 describes the various antibacterial drugs.

<table>
<thead>
<tr>
<th>Gram Positive Cocci</th>
<th>Gram Negative Cocci</th>
<th>Gram Positive Bacilli</th>
<th>Gram Negative Bacilli</th>
</tr>
</thead>
<tbody>
<tr>
<td>Regular antibiotic</td>
<td>Regular Antibiotics</td>
<td>Regular Antibiotics Amikacin</td>
<td>Regular antibiotic</td>
</tr>
<tr>
<td>Cefazolin</td>
<td>Ceftriaxone</td>
<td>Fluoroquinolone</td>
<td>Fluoroquinolone</td>
</tr>
<tr>
<td>4th Generation Fluoroquinolone</td>
<td>Ceftazidime</td>
<td>Clarithromycin</td>
<td>F. Tobramycin</td>
</tr>
<tr>
<td>Higher antibiotics</td>
<td></td>
<td></td>
<td>Higher antibiotics</td>
</tr>
<tr>
<td>Vancomycin</td>
<td></td>
<td></td>
<td>Amikacin</td>
</tr>
<tr>
<td>Linezolid</td>
<td></td>
<td></td>
<td>Ceftazidime</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Piperacillin</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Meropenem</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Colistin</td>
</tr>
</tbody>
</table>

Fungal Keratitis:
Fungal ulcers are difficult to treat since the diagnosis is delayed. Mycotic ulcer treatment trial (MUTT) I compared natamycin and voriconazole revealed that Natamycin had showed significant clinical improvement as compared to voriconazole. MUTT II compared oral voriconazole and oral placebo which did report benefits of oral voriconazole in treating Fusarium Ulcer. Steroids are contraindicated in fungal ulcers. Subconjuctival antifungals should be avoided since they result in severe pain and induce tissue necrosis to some extent. Since Intrastromal Voriconazole has a good penetrating capacity it can be considered to treat deep and larger ulcers. Intrastromal Voriconalzole can be used as an adjunct to Natamycin in eyes not responding to topical natamycin. Table 3 describes Antifungal drugs. Figure 7 describes the algorithm for managing severe bacterial keratitis.
Table 3 describes Antifungal drugs

<table>
<thead>
<tr>
<th>Polyenes</th>
<th>Pyrimidines</th>
<th>Imidazoles</th>
</tr>
</thead>
<tbody>
<tr>
<td>e.g. Nystatin,</td>
<td>e.g. Fluconazole</td>
<td>e.g. Clotrimazole</td>
</tr>
<tr>
<td>Amphotericin B</td>
<td></td>
<td>Miconazole</td>
</tr>
<tr>
<td>Natamycin</td>
<td></td>
<td>Ketoconazole</td>
</tr>
</tbody>
</table>

Acanthamoeba Keratitis:
A combination of propanodine isethionate (Brolene) 0.1% and polyhexamethylene biguanide 0.02% can be prescribed in initial stage. A combination of brolene and neomycin or monotherapy with chlorhexidine also gives good results. Steroid should be avoided however in case of deep vascularisation steroids and cysticidal agents combination can be used. Oral Miltefosine can be considered as adjunctive treatment for Acanthamoeba keratitis. Table 4 describes Antiameoboid drugs.

Table 4 describes Antiameoboid drugs

<table>
<thead>
<tr>
<th>Anticeptic Biocides</th>
<th>Aminoglycosides</th>
<th>Diamidines</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chlorhexidine</td>
<td>Neomycin</td>
<td>Dibromopropamidine</td>
</tr>
<tr>
<td>PHMB</td>
<td>Paromomycin</td>
<td>Hexamidine</td>
</tr>
</tbody>
</table>

Viral Keratitis:
Usually, about 50% of active epithelial lesions heal spontaneously without treatment. The cure rate of antiviral therapy is 95%. In most of the cases healing occur by day 10. After healing has occurred medication should be quickly tapered and discontinued by day 14. Steroids are contraindicated in viral ulcers. Table 5 describes the anti viral drugs.

Table 5 describes anti viral drug

<table>
<thead>
<tr>
<th>Topical</th>
<th>Systemic</th>
</tr>
</thead>
<tbody>
<tr>
<td>Acyclovir 3% ointment</td>
<td>Oral Acyclovir 400mg</td>
</tr>
<tr>
<td>Ganciclovir 0.15% Gel</td>
<td>Famciclovir 500mg</td>
</tr>
<tr>
<td>Trifluoroxyamide 1%</td>
<td>Valacyclovir 1gm</td>
</tr>
<tr>
<td></td>
<td>Valganciclovir 900mg</td>
</tr>
</tbody>
</table>

Supplementary treatment:
Cycloplegic agents such as atropine sulphate 1%, homatropine 1%, or cyclopentolate 1% can be prescribed for three times a day to reduce the ciliary spasm and produce mydriasis, thus help relieve pain and prevent the formation of synechiae. Glaucomatous drug can be prescribed to lower the IOP.

Causes of Medical treatment failure
- Wrong diagnosis
- Resistance to Antibiotic especially (Ciprofloxacin, moxifloxacin etc.)
- Non compliance from patients

Surgical treatment
Surgical treatment depends on various factors such as size, location and causes of the ulcer.

Corneal Gluing to manage perforations:
For managing corneal perforations less than 2mm cyanoacrylate glue is applied. Healing of the cornea occurs as fibrovascular tissue grows under the glue and dislodges the glue. If the perforation is larger than 2mm then either a tenon patch or multilayer amniotic membrane graft is considered. It has been reported that Gunderson flap have also been use to treat perforation however it was found to be non effective.

Amniotic membrane transplant (AMT):
AMT can provide structural support in areas of corneal ulceration. The use of a single layered AMT may be sufficient to treat ulcers lacking depth and significant stromal thinning however non effective in cases of deep ulcers and multilayered AMT can be considered. It has been reported that AMT is also efficacious in treating neurotrophic Ulcer.

DALK using the big bubble technique seems to be effective and safe in the treatment of deep fungal as well Acanthamoeba keratitis unresponsive to medication. Also, therapeutic Keratoplasty have shown good results in Acanthamoeba keratitis. Penetrating keratoplasty (PKP) is performed in advance cases where medical treatment fails or perforation too large to treat by other options.

Novel treatment approaches
Topical Povidone-Iodine 1.25% has shown to be as effective as topical antibiotics for bacterial keratitis. Collagen shields or contact lenses soaked in antibiotics are used in some cases which may enhance the drug delivery. Nano drug
particles as it has better penetration and drug availability.

Newer treatments such as photoactivated chromophore for infectious keratitis - corneal collagen cross-linking (PACK-CXL)\(^{22,23}\) have been used in therapy resistant cases of melting corneas and also in novel cases of bacterial keratitis. It has been reported that Dresden protocol technique is found be efficacious by damaging the DNA and RNA in microbes and thus help in improving and reducing inflammatory response to bacteria. Pack CXL has a very good healing rate in cases of bacteria as compared to fungal. However it works better in superficial infiltrate and future work is required to explore its use in other cases with consideration of treatment parameters as well as pathogen types.

**Prevention**

Since it is vision threatening condition, it is important to increase awareness about eye care. Many causes of corneal ulcers can be prevented by using protective eye wear during travelling or work. Educating the patients about care and maintenance of contact lens can help prevent ulcers related to contact lens wear.

**REFERENCES**


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