Bacterial

Etiology
Approximately 25,000 Americans develop bacterial keratitis annually. Of patients with bacterial keratitis, 19-42% are contact lens wearers. Bacterial keratitis is one of the most important potential complications associated with contact lens wear. While even daily wear of contact lenses carries an increased risk of infection, extended or overnight contact lens wear is the greatest risk factor for infectious keratitis in patients choosing to wear contact lenses. Large-scale epidemiological studies have confirmed that the absolute and relative risk of microbial keratitis is unchanged with overnight use of silicone hydrogel materials. The key findings included the following: (1) The risk of infection with 30 nights of silicone hydrogel use is equivalent to six nights of hydrogel extended wear; (2) Occasional overnight lens use is associated with a greater risk than daily lens use. Microbial contamination of CL storage cases is a great risk for gram-negative bacterial infection among soft CL wearers. Many bacteria have been identified in contact lens related microbial keratitis, with the most common organisms cultured from bacterial ulcers being *Staphylococcus, Streptococcus, Pseudomonas* and *Moraxella*. The gram-negative rod *Pseudomonas aeruginosa* is commonly associated with microbial keratitis in soft contact lens wear but a higher incidence of gram-positive organisms than gram-negative organisms are recovered by culture from infections associated with contact lens wear. It is important to remember that the organisms *Neisseria gonorrhoeae, Listeria, Corynebacterium* and *Haemophilus aegyptius* do not require damage to the cornea and may invade directly through intact corneal epithelium. Up to 20% of cases of fungal keratitis (particularly candidiasis) are complicated by bacterial coinfection.

Presentation
Patients with microbial keratitis present with symptoms including decreased vision, photophobia, moderate to severe ocular pain, redness, swelling and discharge. On slit lamp examination, the critical finding is a focal white opacity in the corneal stroma with an overlying corneal epithelial defect that stains with fluorescein. Additional findings include diffuse epithelial edema, stromal infiltration surrounding the ulceration, and mucopurulent exudation. An anterior chamber reaction and hypopyon may be present. It is important to document the depth and location of the epithelial defect and stromal infiltration. The anterior chamber should be evaluated for cells and flare and examined for a hypopyon, and the intraocular pressure should be checked.

Much has been made regarding culture and sensitivity testing in cases of microbial keratitis. In general, consider cultures in ulcers greater than 1mm to 2mm, defects in the visual axis, ulcers unresponsive to initial therapy, or if an unusual organism is suspected. Remember that just approximately 40% of corneal cultures identify causative pathogens. For this reason, culturing the contact lens and contact lens storage case in addition to scrap-ing the corneal ulcer can aid in identifying the causative organism.

Treatment
Ulcers need to be considered infectious until proven otherwise. Therapy begins with immediate, intensive, aggressive treatment with fourth-generation fluoroquinolones while awaiting lab results. In vitro studies and prospective clinical trials have shown that fourth-generation fluoroquinolones are better than the older generation fluoroquinolones and are as potent as combined fortified antibiotics against common pathogens that cause bacterial keratitis. Besifloxacin 0.6% is a topical fluoroquinolone that has similar potency against ocular pathogenic bacteria as the fourth-generation agents. Dosage is every 30 minutes for the first six hours, followed by hourly administration around the clock until improvement is noted. The application of fourth-generation fluoroquinolones in the treatment of corneal ulcers is an off-label use of these medications, but routinely performed. Cycloplegic drops are valuable for patient comfort and to prevent synchiae formation in accompanying iritis.

Avoid steroids initially. Corticosteroid treatment resulted in a statistically significant delay in corneal re-epithelialization and steroid use did not translate to a significant difference in visual acuity or infiltrate/scar size. Infectious keratitis may worsen with topical steroid use, especially when caused by fungus, atypical *Mycobacteria* or *Pseudomonas*. Once the cornea has re-epithelialized and the causative organism has demonstrated sensitivity to the antibiotic (usually after 72 hours of treatment), a steroid may be added to the therapeutic regimen to control persistent inflammation and reduce tissue damage. However, the results of the Steroids in Corneal Ulcers Trial (SCUT) study showed that adjunctive topical corticosteroid did not improve three-month vision in patients with bacterial corneal ulcers. The patient with an infectious keratitis needs to be followed daily, with careful monitoring of the findings. The antibiotic regimen should be reduced depending on the response, but should never be tapered below the minimum dose (usually q.i.d. to t.i.d.) to prevent the possibility of bacterial resistance.

Collagen cross-linking has been shown to improve healing of infectious corneal ulcers in treatment-resistant cases or as an adjunct to antibiotic treatment.

ICD-9 Codes
- 370.00 Corneal ulcer, unspecified
- 370.01 Marginal corneal ulcer
- 370.02 Ring corneal ulcer
- 370.03 Central corneal ulcer
- 370.04 Hypopyon ulcer
- 370.05 Mycotic corneal ulcer
- 370.06 Perforated corneal ulcer
- 371.00 Corneal opacity/scar, unspecified (upon resolution)

References
Viral Infections

ADENOVIRAL KERATOCONJUNCTIVITIS

Etiology
Adenoviruses produce the most common viral conjunctival infections.1 The condition is quite contagious and is transmitted readily in respiratory and ocular secretions, eye drops and mascara bottles, and contaminated swimming pools. Eye care professionals are often guilty of spreading adenovirus because it is highly contagious and has 53 serotypes with variable morphology.2 The incubation period is usually five to 12 days, and the clinical illness is present from five to 15 days. Most cases of viral conjunctivitis resolve spontaneously, without sequelae, within days to weeks. Adenovirus is often difficult to diagnose based on clinical appearance and, in the early stages, is associated with a red eye or superficial keratitis common to herps and other infections.3

There are four forms of adenoviral conjunctivitis: follicular conjunctivitis, pharyngoconjunctival fever, epidemic keratoconjunctivitis (EKC) and acute hemorrhagic conjunctivitis.4 Follicular conjunctivitis is the mildest form of adenoviral conjunctivitis. Pharyngoconjunctival fever is the most common ocular adenoviral infection and is characterized by a combination of pharyngitis, fever, and conjunctivitis. EKC is a more severe form of conjunctivitis and typically lasts for seven to 21 days. EKC can affect the cornea with coarse keratitis and sub-epithelial infiltrates (SEIs). SEIs may last for months, affecting visual acuity. Acute hemorrhagic conjunctivitis produces a severe, painful follicular conjunctivitis with the development of tiny subconjunctival hemorrhages.

Presentation
In general, viral infections present with redness, irritation, itching, foreign body sensation, tearing and photophobia. The condition starts in one eye and then progresses to the other a few days later. Signs include conjunctival injection and swelling. The lids may be swollen. Inferior palpebral conjunctival follicles are seen. Pinpoint subconjunctival hemorrhages and membrane formation over the palpebral conjunctiva are occasionally seen. In some cases multiple, focal infiltrates in the cornea anterior to mid-stroma may be seen. A pre-auricular lymphadenopathy is present. A rapid, in-office immunodiagnostic test using antigen detection is available for adenovirus conjunctivitis. In a study of 186 patients with acute conjunctivitis, this test had a sensitivity of 88% to 89% and a specificity of 91% to 94%.5

Treatment
Palliative therapy is often sufficient for most cases of adenoviral conjunctivitis: cold compresses, artificial tears and topical decongestants/antihistamines. Betadine 5% is now available as an FDA-approved product that can be used “off-label” to treat EKC. This agent has minimal toxicity when used properly and followed by saline irrigation. In vitro, povidone iodine is effective against adenovirus as well as many other infectious agents. Topical steroids are indicated when the visual axis is involved or membrane or pseudo-membrane formation is noted. Patients should discontinue contact lens wear. Avoid the use of topical and oral antibiotics or antiviral agents as these will not help resolution and may promote antibiotic resistance. Patients with adenoviral conjunctivitis need to understand that the condition is highly contagious and should be informed of appropriate measures to reduce the risk of spreading the infection to their other eye or to other people. Recent data also indicate ganciclovir 0.15% ophthalmic gel may hold promise as a treatment for adenoviral keratoconjunctivitis.6

HERPES SIMPLEX KERATITIS

Etiology
The herpes simplex virus (HSV) is the leading cause of vision loss in the United States. Keratitis caused by HSV is also the most common cause of cornea-derived blindness in developed nations.7 The HSV is a DNA virus that resides latent in the trigeminal ganglion, only to resurrect during periods...
of intense stress, illness, irritation and phototoxic exposure. The disease can be present as superficial lesions, neurotrophic disease, or with deep stromal involvement.

**Presentation**

Patients with HSV infection present with rapid onset unilateral pain and redness, watering and light sensitivity. Diagnosis of HSV infection is primarily based on clinical findings. The disease starts as a punctate epithelial keratitis, coalescing into the classic branching epithelial ulceration with terminal end bulbs within 24 to 48 hours. The dendrites stain with rose bengal or lissamine green. Corneal sensitivity may be decreased. The neurotrophic form of HSV disease is characterized by areas of intense punctate change or epithelial denudement, and can result in corneal scarring. Deep stromal lesions appear as a round, fluid filled circle. Scarring can develop in later stages with loss of stromal thickness and corneal thinning.

**Treatment**

Treatment of active HSV keratitis consists of topical trifluridine 1% solution every one to two hours until no sign of active infection (lack of dendrite patterns), then five times a day for an additional seven to 10 days. Zirgan (ganciclovir) ophthalmic gel 0.15% is indicated for the treatment of acute herpetic keratitis. Four international, randomized, multicenter clinical trials have demonstrated that ganciclovir gel is at least as effective as acyclovir ointment for the treatment of HSV keratitis. Ganciclovir gel was better tolerated, with lower rates of blurred vision, eye irritation, and punctate keratitis. The recommended dosing regimen for Zirgan is one drop in the affected eye five times per day (approximately every three hours while awake) until the corneal ulcer heals, and then one drop three times per day for seven days. Topical acyclovir 3% ointment (no longer commercially available, but can be obtained from specialized compounding pharmacies) can be used five times a day as an alternative in patients with a known sensitivity to the above medications.

Oral antivirals are gaining use in the treatment of epithelial disease. Acyclovir 400mg five times a day is the most common oral dosage. Alternatively, oral valacyclovir (Valtrex), a prodrug of acyclovir, can be given 500mg three times a day. Since these drugs are cleared through the kidneys, it is critically important for the patient with renal disease that the patient’s nephrologist or primary care physician and/or pharmacist be consulted regarding dosing. For a patient on long-term oral antiviral therapy for recurrent disease, check creatinine levels to ensure there is no kidney damage. Normal levels of serum creatinine are approximately 0.8mg to 1.4mg per deciliter in adults. Oral antiviral prophylaxis has been associated with a decreased risk of recurrence of epithelial keratitis, stromal keratitis, conjunctivitis, and blepharitis due to HSV.

**HERPES ZOSTER OPHTHALMICUS**

**Etiology**

Herpes zoster ophthalmicus (HZO) is a recurrent infection of the varicella zoster (chickenpox) virus in the ophthalmic division of the trigeminal dermatome, most often affecting the nasociliary branch. HZO can affect any of the ocular and adnexal tissues. One in four people will contract herpes zoster in their lifetime, with this risk rising markedly after 50 years of age, especially in very elderly individuals. Ophthalmic herpes zoster represents 10% to 20% of all zoster cases.

**Presentation**

HZO usually begins as an influenza-like illness characterized by fatigue, malaise, nausea and mild fever accompanied by progressive pain and skin hyperesthesia. A diffuse erythematous or maculopapular rash appears over a single dermatome three to five days later. The skin of the forehead and upper eyelid is commonly affected and strictly obeys the midline with involvement of one or more branches of the ophthalmic division of the trigeminal nerve. Involvement of the tip of the nose (Hutchinson’s sign) has been thought to be a clinical predictor of ocular involvement. Although patients with a positive Hutchinson’s sign have twice the incidence of ocular involvement, one third of patients without the sign develop ocular manifestations. HZO conjunctivitis is a common ocular finding and the conjunctiva appears swollen and injected, with occasional vesicles and petechial hemorrhages.

HZO can cause either a nongranulomatous or granulomatous anterior uveitis with keratic precipitates and posterior synechiae. The diagnosis of herpes zoster disease is generally based on clinical findings.

**Treatment**

All patients with ophthalmic zoster, irrespective of age or severity of symptoms, should be prescribed oral antiviral drugs at the first sign of disease. Patients with HZO are treated with oral acyclovir (800mg, five times daily) for seven to 10 days. Early treatment with acyclovir (within 72 hours after rash onset) reduces the percentage of eye disorders in ophthalmic zoster patients from 50% down to 20% to 30% and also lessens acute pain. Famciclovir 500mg three times daily for seven days or valacyclovir 1,000mg three times daily are alternatives to acyclovir. In general, the dosages for HZO are twice that for HSV infections. Palliative therapy including cool compresses, mechanical cleansing of the involved skin and topical antibiotic ointment without steroid are helpful in treating skin lesions. Conjunctival defects associated with HZO keratitis may be treated with nonpreserved artificial tears, eye ointments, punctal occlusion, pressure patching, or therapeutic soft contact lenses. Topical steroids are useful in the management of kerouveitis, interstitial keratitis, anterior stromal infiltrates, and disciform keratitis. Topical cycloplegics prevent ciliary spasm associated with HZO keratitis.
Acanthamoeba Keratitis

Background

Acanthamoeba keratitis can be severe and vision-threatening. It was first recognized in contact lens wearers in the early 1970s, and contact lens wear is thought to be associated with 80% of the cases.1 Symptom onset is greatest during the summer months. Acanthamoeba species are found in virtually every environment. These protozoa are resistant to killing by desiccation, freeze/thaw cycles, irradiation, chlorination levels and antimicrobial agents. Co-infection with bacteria or fungi is common, providing food and antimicrobial agents. Co-infection with herpes zoster ophthalmicus: predictors of postherpetic neuralgia and ocular involvement. Br J Ophthalmol 1987;71:353–58.

Presentation

Acanthamoeba keratitis presents with pain (ranging from mild foreign body sensation to severe pain), photophobia, decreased vision, injection, irritation, tearing and a protracted clinical course. The patient often presents with a unilateral red eye, where the pain is disproportionately worse than one would surmise from the clinical appearance. Early corneal findings include irregular epithelium, punctate epithelial erosions, microcystic edema, perilimbal injection and dendritiform epithelial lesions. The dendritiform lesions often resemble those of herpes simplex keratitis; however, the Acanthamoeba lesions appear edematous and necrotic rather than frankly ulcerated.

A ring infiltrate is classically thought of as the defining sign of Acanthamoeba keratitis; however, it tends to form four to eight weeks after onset of symptoms and is rarely the presenting sign. Radial perineuritis (perhaps explaining the intense pain) may be seen on slit lamp examination or confocal microscopy. Unchecked, there may be progressive corneal thinning and risk of perforation. Up to 40% of patients may have mild to severe anterior uveitis. Scleritis has been reported in patients with Acanthamoeba keratitis; however, the scleral inflammation was attributed to an immune-mediated response to necrotic organisms and was not believed to be the result of active infection.14

Severe glaucoma has been associated with Acanthamoeba keratitis secondary to an inflammatory angle-closure mechanism, apparently without direct infiltration of the organism.15

Etiology

There are more than 20 different species, several of which are known to cause infections in humans, including A. castellanii, A. polyphaga, A. castellani, A. nailey, A. astronyxis, A. hatchetti and A. rhyodes. A. castellani is the most common amoeba associated with corneal infection. The life cycle of these organisms is comprised of two stages: trophozoite and cystic forms. Trophozoites bind to and desquamate the corneal epithelium. They secrete a variety of proteases, which facilitate the dissolution of the corneal stroma.1 When environmental conditions become unfavorable, the organism converts to a dormant cystic form, which is able to survive many years. These double-walled cysts are highly resistant to killing by desiccation, freeze/thaw cycles, irradiation, chlorination levels and antimicrobial agents. Co-infection with bacteria or fungi is common, providing food for amoeba.

ICD-9 Codes

- 372.00 Unspecified conjunctivitis
- 372.02 Acute follicular conjunctivitis
- 372.03 Other mucopurulent conjunctivitis
- 372.11 Simple chronic conjunctivitis
- 077.1 Epidemic keratoconjunctivitis
- 077.3 Adenoviral (acute follicular)

877-94.

References


with herpes zoster inflammatory disease. Aqueous suppressants and topical corticosteroids should be used to treat glaucoma associated with HZ disease. Consider Zostavax vaccine for your patients over 60 years of age. Aqueous suppressants and topical corticosteroids should be used to treat glaucoma associated with HZ disease. Consider Zostavax vaccine for your patients over 60 years of age.

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Fungal Keratitis

Fungal keratitis is relatively rare in the United States (approximately 5% to 10% of reported cases), although it accounts for up to 50% of ulcerative keratitis elsewhere in the world. Fungal keratitis is usually associated with a history of ocular trauma, ocular surface disease or topical steroid use. There has been a lot of attention focused on the recent epidemic of fungal keratitis in soft contact lens wearers in 2005 and 2006; however, a recent review indicates the number of fungal keratitis cases associated with contact lens wear has been steadily increasing the past 20 years.

Etiology

Fungi require an epithelial defect for corneal penetration. Once the epithelium has been violated, the present fungi can multiply and cause severe tissue damage. Up to 30% of fungal keratitis cases may be associated with bacterial co-infection. Risk factors for the development of fungal keratitis include ocular trauma, topical corticosteroids, systemic immunosuppression, penetrating or refractive surgery, chronic keratitis (vernal/atopic keratitis and neurotrophic ulcers) and contact lens wear with certain lens solutions.

Presentation

Patients present with pain, photophobia, injection, tearing and possible discharge; however, the degree of symptoms may vary. In some cases, the progression of symptoms may be slow, while in others it may move very quickly. Corneal infiltrates tend to have feathery borders, are generally grayish-white and may reveal hyphae in filamentary fungal disease. Up to some cases, the progression of symptoms may be slow, while in others it may move very quickly. Corneal infiltrates tend to have feathery borders, are generally grayish-white and may reveal hyphae in filamentary fungal disease. Up to some cases, the progression of symptoms may be slow, while in others it may move very quickly. Corneal infiltrates tend to have feathery borders, are generally grayish-white and may reveal hyphae in filamentary fungal disease. Up to some cases, the progression of symptoms may be slow, while in others it may move very quickly. 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Epithelial

The corneal epithelium serves a variety of roles, including the primary mode of protection for the corneal surface and as the interface tissue between the corneal stroma and the tear film. Derived from surface ectoderm, it can be best described as a nonkeratinized, stratified squamous layer possessing many of the same properties of regeneration after injury and, fortunately enough, recovers in rapid fashion. For purposes of brevity, the corneal epithelium consists of deep columnar cells attached to the anterior limiting lamina (or Bowman’s membrane), midlayer polygonal cells, and surface wing cells all bound together through a variety of tight adhesions. The typical epithelial layer is approximately 50µm thick. During cell division, the epithelial cells migrate and flow more anteriorly towards the surface, ultimately losing their nuclei and becoming “wafer” thin.

Fungal corneal ulcer

Epithelial basement membrane dystrophy

ICD-9 Codes

• 370.05 Mycotic corneal ulcer
• 370.04 Hypopyon corneal ulcer

References

Corneal abrasions are one of the most common forms of ocular trauma presenting to an optometric clinic. Damage to the corneal epithelium in corneal abrasion and its subsequent repair is influenced by a variety of comorbidities, including the presence of conditions such as diabetes, Sjögren’s or other tear chemistry altering illness, corneal denervation and neuropathy, lagophthalmos, and basement membrane degenerations or dystrophies. Most epithelial trauma will recover within three to five days by a coordinated migration of cells as an epithelial sheet. Large or total abrasions can take significantly longer to heal. Careful attention needs to be paid to the size, position, and depth of the abrasion and whether or not any signs of infiltrate are present. Fluorescein dye should be instilled to measure the abrasion and monitor its improvement; for a deeper abrasion it is imperative to assess for risk of perforation and check for Seidel’s sign. Should either of these be detected, prompt medical management should be initiated. Though any abrasion can predispose to future erosions, those caused by paper, fingernails or tree branches are more commonly associated with recurrent erosions and the patient should be treated accordingly. Sodium chloride eyedrops and ointments, such as Muro 128 5%, are advised.

There are a variety of corneal epithelial dystrophies. Almost all are considered recessive in nature and can cause the patient symptomatic visual disturbance in the form of irregular astigmatism or intermittent pain secondary to recurrent erosions.

Epithelial basement membrane dystrophy (EBMD), also known as map-dot-fingerprint dystrophy or Cogan’s dystrophy, is described as hereditary in nature, presents bilaterally and is progressive. Typically, EBMD can be visualized in white light at the biomicroscope, by using sodium fluorescein dye and a watten filter and looking for areas of discrete negative staining, as well by irregularity on corneal topography. Typically these patients have reduced acuity that can be improved with a gas permeable trial lens.

Subepithelial mucinous corneal dystrophy is a very rare autosomal dominant condition characterized by frequent recurrent erosions in the first decade of life followed by progressive vision loss. The lesions involve the entire cornea, but are typically found centrally.

Meemans’s corneal dystrophy is a bilateral autosomal dominant disorder that generally appears early in life as epithelial microcysts in the first decade of life. Patients may remain asymptomatic for years, until epithelial cysts produce symptoms, such as impaired visual acuity, pain and photophobia.

Lisch epithelial corneal dystrophy (LECD) is a relatively new disorder that was first described in 1992. It is unusual because it is linked to the X chromosome. It is a gelatinous, whoof-like corneal dystrophy associated with surface deposition and appears as epithelial microcysts on retroillumination. It typically begins in childhood, sparing the center of the cornea. As the opacities progress toward the center they can decrease acuity but are not associated with recurrent erosions. Gelatinous, drop-like corneal dystrophy is a rare corneal presentation association with surface deposition thought to be caused by a mutation of the M1S1 gene. Typically associated with Japanese heritage, the condition has been noted in other parts of the world and characterized by severe visual impairment.

Treatment—Abrasion
The treatment will depend upon the extent and presentation of the injury. In cases where there are irregular, ragged edges of epithelial tissue it is essential to debride that tissue. Antibiotic drops can be dosed from q.i.d. to q.h. depending on the size and depth of the injury, along with a cycloplegic agent b.i.d.-q.i.d. for photophobia. A bandage contact lens can be extremely beneficial to promote reepithelialization and improve patient comfort.

Treatment—Dystrophies and Degenerations
EBMD treatment depends on the level of visual performance and patient discomfort. For mild presentations, the patient can use artificial tears and hyperosmotic drops and/or ointments. Ointment at bedtime is especially helpful for those patients having difficulty with overnight erosions. Moisture chamber goggles for nighttime use is part of the palliative treatment. For patients with blurred vision due to irregular astigmatism, a soft, rigid or hybrid contact lens can be utilized. If these treatments are unsuccessful, the faulty epithelial tissue can be debrided by diamond burr keratectomy, phototherapeutic keratectomy (PTK) or anterior stromal micropuncture (ASM). Treatment of the other epithelial dystrophies need to be considered on a case-by-case basis where treatment ranges from symptomatic relief with lubrication, Muro 128, and bandage contact lenses to more advanced treatments including debridement, PTK, and penetrating and lamellar keratoplasty. However, there is a risk of the dystrophy recurring within the graft. All tests should be medically necessary and be important for the management of the case.

ICD-9 Codes
- 918.00 Corneal abrasion
- 367.22 Irregular astigmatism
- 371.42 Recurrent erosion of cornea
- 371.51 Meemans’s
- 371.52 EBMD

New CPT Code: 92071 Fitting of a contact lens for treatment of ocular surface disease

**Tests to Help Delineate Structure and Function**
- All tests should have an interpretation of findings.
- Corneal topography—useful for cases of epithelial irregularity involving visual axis.
- Anterior segment photography—useful for documenting presence and extent of surface changes.
- Tear film performance tests—Shirmer’s I or II, tear break-up time, and phenol red test.

**Pearls**
- Never use a bandage contact lens with corneal trauma that might be at risk for fungal infection. For instance, caution should be used with a patient presenting to the office with a tree branch injury to the cornea.
- Consider using a stiffer modulus soft contact lens for corneal irregularities in the corneal visual axis. Stiffer designs can drape over irregular zones and dampen the effects of irregular astigmatism.
- For patients considering cataract surgery AND possessing significant EBMD in the visual axis, consider a targeted epithelial debridement, particularly if located near the patients’ visual axis.
- For patients with EBMD, Meemans’s or Reis-Bucklers (Bowman’s layer), consider the use of bandage soft lenses for episodes of recurrent erosion. Furthermore, encourage the use of nightly hyperosmotic ointments to limit intermittent nightly erosions.

**References**
Bowman’s

Reis-Bucklers (Corneal Dystrophy of Bowman Layer Type 1 and Granular Dystrophy Type III)

Gray, hazy, reticular deposits (crystallization) typify this Bowman’s layer dystrophy.\(^2\) \(^3\) The deposits will generally form symmetrical patterns in each eye and can easily be detected at the slit lamp. The opacities cause an elevation of the surface epithelium and often create a visual disturbance by affecting surface epithelium as well as underlying stromal areas of the cornea.\(^1\) It is believed to be an autosomal dominant inherited corneal dystrophy associated with mutations in the TGFBI gene and in some cases with R124L and G623D mutations.\(^1\) \(^3\) Onset generally occurs by the first decade of life, typically four to five years of age. This dystrophy is quite rare and its incidence is unknown.\(^1\)

Patient history and symptoms will invariably include recurrent corneal erosion (traumatic or spontaneous), pain, light sensitivity and visual disturbance/distortion. The episodes of recurrent erosion tend to occur multiples times each year.\(^2\) A severe disruption of the basal epithelial cells and their stromal attachments accounts for the frequent erosions. The epithelium is loosely attached since a basement membrane is focally absent.\(^2\) After 30, the erosions tend to become infrequent but acuity continues to decline due to increased superficial corneal opacification.\(^2\)

Early reticular superficial opacity will advance to a honeycomb, gray-white opacity fishnet pattern of the central and mid-peripheral cornea with prominent corneal nerves. An increased corneal thickness, irregular astigmatism and decreased corneal sensation are hallmarks of the corneal malady.\(^2\) \(^3\) Recurrence of this anterior corneal dystrophy is not uncommon after corneal transplantation. Over-the-counter pain relievers, lubricants, NSAIDs and cycloplegics can be used for pain management when necessary.\(^4\) There are no systemic treatments for this dystrophy.

Recommendations

This condition must be differentiated from among several anterior stromal conditions. To add to the confusion, anterior membrane dystrophy of Gayson and Wilbrandt and the honeycomb dystrophy of Thiel and Behnke may represent clinical variants of Reis-Bucklers dystrophy.\(^2\)

Management of the epithelial erosion poses the major problem earlier in life. Lubrication and appropriate supportive therapy for erosions is helpful. When opacification of the cornea impairs vision significantly, superficial keratectomy may improve vision. Advanced cases require lamellar or full thickness grafting but recurrence is common. Over the past decade, researchers using molecular genetics have redefined dystrophies of Bowman’s layer to include two distinct entities (Reis-Bucklers and Thiel-Behnke).\(^1\)

Stroma

The stroma accounts for 90% of the corneal thickness and is composed mainly of water, collagen and keratocytes. It is the specific arrangement of the tightly bound collagen fibers in the stroma that allows for the transparency and mechanical strength of the cornea. Descemet’s membrane is a thick basement membrane that is located at the posterior portion of the stroma. Four corneal dystrophies including granular, lattice, Avellino and Reis-Bucklers have been linked to a mutation in the TGF-1 gene, also known as the BIGH3 gene.

Types of Stromal Dystrophies

Lattice corneal dystrophy (LCD) is the most common of the stromal dystrophies. It has an autosomal dominant pattern of inheritance and appears after the first decade of life as a linear branching pattern that affects the central cornea and can increase over time. LCD is associated with a genetic mutation in the BIGH3 gene, resulting in deposits of amyloid within the anterior corneal stroma. Five subtypes of LCD have been identified. LCD Type I is the classic form of LCD.

Symptoms include decreased vision and recurrent corneal erosions.

Granular corneal dystrophy (GCD) is a bilateral, autosomal dominant disease associated with a mutation in the BIGH3 gene that leads to the deposition of a hyaline material in the corneal stroma. It typically presents within the first decade of life with focal granular deposits between the anterior to mid stromal regions. These opacities are discrete deposits located centrally, with clear cornea located in the periphery and clear cornea between deposits. The disease is typically asymptomatic early on, but with time the opacities can coalesce and lead to decreased vision. Recurrent corneal erosions can occur in GCD but at a lower incidence than in LCD. Three types of GCD have been described. GCD Type I is the classic form of GCD.\(^1\) Avellino

References


ICD-9 Codes

- 371.52 Other anterior corneal dystrophy
- 371.42 Recurrent corneal erosion
- 371.0 Corneal scar and opacity code range

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Recurrent lattice dystrophy after corneal transplant

corneal dystrophy is GCD Type II. It is linked to a mutation in the BIGH3 gene that leads to a deposition of both hyaluronic acid and amyloid in the corneal stroma. Typically, patients present in their second decade with granular opacities like in GCD, but later in the disease process develop lattice lines as well. The disease was thought to have originated from a family in Avellino, Italy. However, GCD Type II has now been reported in patients from many other countries as well.2,3

Reis-Bucklers is GCD Type III (Bowman’s layer). It is characterized by faint, gray-white superficial scarring patterns that are often greatest in the central cornea. The patient may experience corneal erosions, photophobia and irritation that can occur in early childhood. Analysis of the area with confocal microscopy will often reveal the absence of Bowman’s layer. Vision can vary and is diminished by the severity of superficial scarring and irregular astigmatism.

Macular corneal dystrophy (MCD) is the least common, but the most severe, of the stromal corneal dystrophies. Three subtypes of MCD have been described based on the presence or absence of immunoreactive keratan sulfate within various tissues. Type I does not have immunoreactive keratan sulfate in the corneal stroma, keratocytes, sera or cartilage, and is the most common variant of MCD worldwide. It is an inherited autosomal recessive condition. It is typically noted during the first decade of life, characterized by gray-white anterior stromal lesions similar to GCD. There is severe stromal haze throughout the entire stroma and limbus to limbus, with patients typically developing severe visual loss by the second to third decade of life.

Schnyder corneal crystalline dystrophy (SCCD) is a slowly progressive autosomal dominant dystrophy that appears early in life, but may not cause vision loss until the fifth decade of life. It is linked to a metabolic defect of corneal keratocytes that leads to crystalline lipid deposition. Clinically the disease presents with a ring-shaped accumulation of fine needle shaped polychromatic crystal deposits within Bowman’s layer and the anterior stroma, and is often associated with a pre-senile peripheral lipid arcus.4

Congenital stromal dystrophy is an autosomal dominantly inherited condition that is caused by mutations in the DCN (decorin) gene, leading to corneal haze and reduced visual acuity. Strabismus is common, and corneal thickness is increased.

Corneal fleck dystrophy (CFD) is a rare autosomal dominant dystrophy with that is often asymptomatic. Photophobia, reduced vision and recurrent erosions may occur. It is characterized by bilateral irregular shaped “flecks” of grayish matter in the posterior stroma.5 Posterior amorphous corneal dystrophy is a rare condition characterized by bilateral sheet-like opacification of the posterior stroma in association with corneal flattening and thinning. It appears to be non-progressive and patients are asymptomatic.6

Treatment
The treatment for the stromal corneal dystrophies is observation and lubrication for corneal erosions if they occur. Phototherapeutic keratectomy (PTK) and corneal transplants are options as vision becomes impaired, but the dystrophy can recur in the graft.

Tests to Help Delineate Structure and Function

• Corneal topography—useful for cases of stromal thinning causing irregularity involving the corneal surface.
• Anterior segment photography—useful for documenting presence and extent of stromal changes.
• Pachymetry—useful for documenting thickness changes or variations.

Pearls

• Monitor visual acuity with contrast changes. While Snellen acuity might be satisfactory, loss of contrast will affect visual quality more and result in patient complaints of performance.

ICD-9 Codes

• 371.53 Granular corneal dystrophy
• 371.54 Lattice corneal dystrophy
• 371.55 Macular corneal dystrophy
• 371.56 Crystalline corneal dystrophy

References


Endothelium

The endothelium is the posterior layer of the cornea, consisting of a single layer of cells, about 5μm thick, bound together and predominantly hexagonal in shape. Anteriorly, it is in contact with Descemet’s membrane and posteriorly with the aqueous humor. It is the structure responsible for the relative dehydration of the corneal stroma. In the normal adult eye the cell density varies from around 3,000 cells/mm² in the central cornea to about 2,000 cells/mm² in the periphery. With age, disease or trauma, the cell density decreases. But with disease or trauma, this reduction may affect corneal transparency, as some fluid then leaks into the cornea.1,2 The endothelial corneal dystrophies, which result from primary endothelial dysfunction, include Fuchs’ endothelial corneal dystrophy (PECD), posterior polymorphous corneal dystrophy (PPCD or PPMD) and congenital hereditary endothelial dystrophy (CHED).

Fuchs’ dystrophy is an autosomal dominant inherited disease that affects women greater than men. It often presents in the fifth to sixth decade of life as multiple central corneal guttata (exccrescences of Descemet’s membrane) associated with pigment dusting on the endothelium. The condition spreads from the center toward the periphery. As the endothelial cells fall, the remaining cells enlarge to cover the gap. With the reduced number of endo-
thelial cells, the pump function suffers. This leads to stromal and epithelial edema, and loss of visual acuity. Vision is typically worse upon awakening because of the swelling induced by nighttime lid closure. In more advanced stages, the epithelial microcysts later coalesce and form bullae, which can rupture, causing foreign body sensation and pain, as well as exposing the cornea to the danger of infectious keratitis.

Posterior polymorphous corneal dystrophy (PPCD) is an autosomal dominant disorder with extremely variable expression. Three genes have been implicated in PPCD (VSX1, COL8A2, TCP8), but the evidence implicating VSX1 and COL8A2 is questionable. PPCD presents earlier than Fuchs' and is typically more benign. It is characterized by the early appearance of vesicle-like lesions, bands or diffuse opacities. These opacities represent more diffuse thickenings in Descemet’s membrane. PPCD can result in peripheral anterior synechiae so these patients must be monitored for increased intraocular pressure. Corneal edema is also a feature of PPCD. Congenital hereditary endothelial dystrophy (CHED) presents at or shortly after birth with bilateral corneal edema. The pathology of CHED is attributed to endothelial cell degeneration during gestation. There are two types: Type I (CHED 1) is inherited as an autosomal-dominant trait that presents with clear corneas at birth. Type II (CHED II) is more common but more severe. It is inherited as an autosomal-recessive trait, associated with nystagmus and corneal edema from birth.

X-linked endothelial corneal dystrophy (XECED) was first described in 2006. The course in XECED is slowly progressive with intermittent corneal clouding in the form of ground glass and moon crater-like changes of the corneal endothelium. The corneal opacification may be severe and associated with nystagmus. In advanced cases, a subepithelial band keratopathy develops.

Treatment
Early treatment includes hypertonic solutions during the day and nighttime hypertonic ointment. With extreme epithelial edema, bullae may form and cause pain and photophobia. Bandage soft contact lenses may provide temporary relief. However, these patients require careful follow-up care to reduce the risk of stromal neovascularization or infectious keratitis. If at some point vision degrades to a disabling level, surgery should be considered.

Penetrating keratoplasty had been the gold standard for treatment of complications relating to Fuchs’ dystrophy and other endothelial disorders. Long term, the results have been impressive with graft survival rates. Because of the complications associated with the surgery, including healing time, graft failure and visual variability, the surgery has largely been relegated to second choice behind Descemet’s stripping automated endothelial keratoplasty (DSEK).

DSEK is a surgery designed to replace the endothelium alone without violating any of the overlying structures, including stroma and epithelium. The surgery has an even higher rate of success compared to PK as long as graft adhesion to the recipient cornea is successful. Newer techniques of reducing graft dislocation have made the surgery safer. Graft rejection can be difficult to spot, so it is prudent to look for keratic precipitates or diffuse corneal edema.

Tests to Help Define Structure and Function
- Anterior segment photography—useful for documenting presence and extent of stromal thickening or other disease changes.
- Endothelial cell photography—allows tracking of endothelial cell density as well as imaging for pleomorphism and polymegathism.
- Pachymetry—useful for documenting thickness changes or variations in cornea.

Pearls
- Caution should be given for use of mitomycin C for procedures in Fuchs’ patients as studies have raised the potential for further damage.
- Monitor corneal grafts with pachymetry to look for signs of rejection. Typically, the graft will thicken over time.
- Fuchs’ dystrophy can be transmitted to the recipient of a corneal transplant. Do not rule out this possibility if gutter, stromal, and epithelial edema are observed in a graft.

The “A” for “automated” in DSAEK refers to the recent practice of the eye-bank preparing the tissue in advance of the procedure instead of the surgeon preparing the tissue. This practice reduces handling problems with eye-bank tissue in the operating room.

ICD-9 Codes
- 371.57 Corneal guttata/Fuchs’ dystrophy
- 371.58 Posterior polymorphous dystrophy

References

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Degenerations

Keratoconus and pellucid marginal degeneration are the two most common noninflammatory corneal ectasias. Though they have very distinct features, early cases of either can be difficult to distinguish from the other. Keratoconus is seen in adolescents and young adults. Its onset is usually at puberty. There may be a family history in 10% of patients. It may be associated with atopic disease, Down syndrome, retinitis pigmentosa, Leber’s congenital amaurosis, Marfan syndrome, Eilers-Danlos, osteogenesis imperfecta, or other noninflammatory connective tissue diseases. Patients present with blurred vision and exhibit progressive myopia with irregular astigmatism. Slit lamp examination reveals central or paracentral corneal thinning with protrusion of the cornea at the area of thinning. Usually, the apex of protrusion is just below the center of the cornea. The base of the cone is often outlined by a Fleischer ring, or an epithelial iron line. Vogt’s striae are stress lines in Descemet’s membrane at the apex of the cone that disappear with digital pressure. Breaks in Bowman’s membrane can lead to scarring and subepithelial fibrosis. Acute ruptures in Descemet’s membrane can lead to swelling or hydrops. Corneal topography reveals inferior steepening with paracentral thinning and elevation of the posterior and anterior corneal surfaces.

The onset of pellucid marginal degeneration is usually similar to keratoconus, in the second through fourth decades of life. It is also bilateral, but in contrast to keratoconus, it is not inherited and does not have other typical ocular or systemic associations. The corneal ectasia is located peripherally, usually in the inferior cornea extending from 4 to 8 o’clock positions. The thinning is typically 1 to 2mm wide, and located 1 to 2mm from the limbus. Maximal corneal protrusion occurs just superior to the area of thinning, in contrast to keratoconus. The name “pellucid” means “clear.” The cone is clear without any iron rings or striae. Typically there is no scarring; however, acute hydrops can occur as in keratoconus, which can then leave scarring. Topography reveals vertical flattening with horizontal steepening centrally, while inferiorly there is vertical steepening with horizontal flattening. This results in the classic bowed or “bent bowtie” appearance on topography. Acute hydrops is caused by a break in Descemet’s membrane with subsequent influx of aqueous into the stroma. The break is self-repairing with resolution over weeks to months. Hyperosmotics may hasten recovery. Rarely does the cornea thin enough to perforate. A penetrating keratoplasty (PKP) may be indicated if central scarring is severe, although frequently Cl. fitting is easier after a hydrops episode. PKP is contraindicated during the acute phase of the disease.

In contrast to keratoconus and pellucid marginal degeneration, Terrien’s marginal degeneration usually occurs in middle age to elderly males. The thinning usually starts superiorly and may occasionally be accompanied by inflammation. Neovascular vessels may cross the area of thinning and there may be corneal lipid deposition.

Treatment

Treatment options for keratoconus include collagen crosslinking, INTACS, full thickness penetrating keratoplasty or deep anterior lamellar keratoplasty, and contact lenses.

Pearls

- Keratoconus should be considered any time visual acuity does not meet expected norms and other pathology is not obvious.
- Not all inferior steepening seen on corneal topography is keratoconus and diagnosis should not be made solely on an axial topographical map. Comprehensive diagnosis, treatment, topography interpretation and fitting videos are available at GPLI.org.

Mechanical

ABRASION

Corneal abrasions are a common form of injury most often resulting from ocular trauma or inherently poor adhesion of the epithelium to the underlying basement membrane. When the cornea is injured or damaged, sight-threatening consequences are always a possibility, especially in the contact-lens wearing patient. Contact lenses can compromise the corneal epithelium and may facilitate the development of infectious or non-infectious keratitis. It is important to recognize the signs and symptoms of corneal abrasion in order to ease the patient’s pain and provide proper medical management to hasten visual recovery.

Etiology

Damage to the corneal epithelium results in loss of the connections of the surrounding epithelial cells, but Bowman’s layer generally

ICD-9 Codes

- 371.60 Keratoconus unspecified
- 371.61 Keratoconus stable
- 371.62 Keratoconus acute hydrops
- 371.00 Corneal scar unspecified
- 367.22 Irregular astigmatism
- 371.48 Terrien’s marginal degeneration
- 371.10 Iron deposits cornea

References

remains intact. Deeper involvement affecting stroma is rare and generally due to trauma from a sharp or abrasive object. Often corneal flaps of various sizes and thickness can be seen. The speed by which the abrasion resolves can be affected by a variety of factors including whether the patient has diabetes, corneal denervation, dry eye, lagophthalmos, previous recurrent corneal erosions or basement membrane changes. In general, an epithelial insult will recover within 24-48 hours and is accomplished by having the neighboring epithelial cells slide over and begin covering the wound. Common causes of mechanical abrasions are fingernail, paper, foreign body, curling iron, mascara brush, plant or other vegetative source, and contact lens.

**Presentation**

Slit-lamp biomicroscopy of the injured cornea reveals epithelial disruption and diffuse corneal edema. In severe cases, when edema is excessive, folds and inflammation in Descemet’s membrane may be visible. A drop of topical anesthetic may aid in the examination of the ocular structures. The corneal abrasion should be documented for location, size, shape and depth with either a drawing or photograph, making any notation of an infiltrative process. This visual documentation will aid in follow-up to chart the healing progress. Fluorescein dye can be instilled to identify the corneal defect. The newly created wound appears bright green compared to the rest of the cornea because the dye accumulates in the defect where cell loss and disruption have occurred. The anterior chamber should be observed and any anterior chamber reaction should be noted.

Patient symptoms include eye pain, foreign-body sensation, photophobia, blepharospasm and tearing. The time, place and activity surrounding the injury should be noted. For medical and legal purposes, the visual acuity and any residual foreign body should be documented. Differences include recurrent corneal erosion, herpes simplex keratitis, and confuent superficial punctate keratopathy.

**Treatment**

There are several treatment options for corneal abrasions, including bandage soft contact lenses, topical antibiotic ointment and drops, topical non-steroidal and steroid anti-inflammatory drops, cycloplegic preparations, and hypertonic drops and ointments. Topical antibiotics are the mainstay of corneal abrasion therapy. There is a risk of infection with any open wound or defect involving the basal lamina. Topical antibiotics—such as polymyxin B/trimethoprim, aminoglycosides or fluoroquinolones—are all reasonable for providing prophylactic antibiotic coverage.

Depending on the extent and severity of presentation, dosing might be more aggressive at q2h for 24 hours or more conservative at q.i.d. Antibiotic ointments can be liberally applied during the day or augment drops by using them at bedtime. Ointments tend to provide better barrier and lubricating function, but will temporarily blur vision. Debridement of loose or hanging epithelium is necessary to enhance healing. Therapeutic soft contact lenses can be used in abrasion management. Large abrasions have been found to heal more quickly under the protection of a bandage contact lens. The use of bandage contact lenses, however, does not come without risk, including additional ocular irritation, epithelial and stromal edema, sterile infiltrates and hypopyon, infection and ulceration. Bandage contact lenses should be avoided in all situations where plant injury or false fingernails is suspected because of the risk of fungal keratitis.

Bed rest, inactivity and over-the-counter analgesics can be used to manage pain. Topical non-steroidal anti-inflammatory drops can be a useful adjunctive therapy for the management of pain from corneal abrasion and often obviates the need for oral pain medications. These medications provide patient comfort and do not adversely affect corneal healing time.

Oral analgesics may be necessary for pain not controlled by topical medications. These include aspirin, ibuprofen, toradol and tramadol. These medications provide both analgesic as well as anti-inflammatory effects. Analgesics such as acetaminophen lack the anti-inflammatory component. The above anti-inflammatory medications can be prescribed in combination with narcotic analgesics such as codeine or hydrocodone. Cycloplegics should be prescribed to paralyze the ciliary body and thus decrease ocular pain. Mild abrasion can be managed with cyclopentolate 1%, while more severe presentations may require homatropine 2% or 5%.

Fortunately, the cornea heals very rapidly, but keep in mind that recurrent corneal erosion is possible even months to years later. Lesions that are purely epithelial often heal quickly and completely without scarring. The presence of subepithelial infiltration may be a sign of infection. Lesions such as these should be considered vision-threatening and may warrant culturing and the use of fortified antibiotic therapy.

To prevent recurrent erosion and reduce corneal edema, a hypertonic solution or ointment may be prescribed as long as the cornea has re-epithelialized. Examine both eyes carefully for any evidence of corneal dystrophy. Prevention includes wearing safety glasses for athletes and industrial workers. Eyelids should be adequately taped for any patient undergoing general anesthesia.

**Recommendations**

- If the patient’s blepharospasm is intense and visual acuity cannot be obtained, instill one drop of anesthetic onto the bulbar conjunctiva. This should allow you to immediately record visual acuity. A new (sterile) bottle of anesthetic should be used if a penetrating injury is suspected.
- Always remember to evert the upper lid to check for residual foreign substance.
- Cycloplegics will make the patient more comfortable and decrease the likelihood of traumatic iritis developing. Steroids are NOT always necessary and should be avoided initially.
- If using a bandage lens, make sure that you are using a lens of appropriate Dk/t to reduce the likelihood of corneal swelling.
- Avoid bandage lenses if injury involves plant material or false fingernails.
- Patching is rarely necessary. Never patch contact lens related abrasions or injuries involving vegetative material or false fingernails. Patients should be re-evaluated within 24 hours.
- Encourage the use of nighttime hyperosmotic ointments once the cornea has re-epithelialized up to three-six months post insult in cases involving a significant abrasion.

**ICD-9 Codes**

- 371.42 recurrent corneal erosion
- 37.82 corneal disorder due to contact lens wear
- 918.1 significant injury to the cornea

**References**

1. Arunagiri G, Trikha R. Corneal Abrasions, Contusions, Lacerations, and Perforations. In:Roy FH, Fraunfelder FW and...
EXPOSURE KERATOPATHY

Exposure staining of the ocular surface appears to be associated with incomplete blinking with improper lid closure and movement and can have several different causes. 

Whenever possible, correcting any underlying problem will often provide immediate relief and prevention is critical in many cases to maintain normal corneal clarity.

Etiology

Major causes of exposure keratopathy relate to lid malposition or deformity (eyelid scarring from trauma or herpes zoster), ectropion, or chemical burns. Other causes stem from lid surgery or acquired defects/ deformities (especially pteryi repair and blepharoplasty), nocturnal lagophthalmos, sedation and altered mental status, proptosis or cranial nerve palsies that affect the lid or eye. Exposure keratopathy is commonly seen in Parkinson’s disease, Bell’s palsy, neurotrophic and ectropion.

Treatment

A careful medical history may uncover previous Bell’s palsy, lid surgery, anterior segment trauma or thyroid disease. Careful evaluation should include assessment of eyelid closure/laxity and corneal exposure. Always check to be certain there is no evidence of reduced corneal sensation increasing the risk of corneal complications. A slit lamp examination will often uncover any tear film abnormalities, corneal integrity issues or other external or anterior chamber reaction. Lubrication is essential for any seated or obtunded patient. Artificial tears, lubricating ointments and gels are the mainstay of treatment. Punctal plugs can also be used when necessary for neurotrophic states or other dry eye related complications. Eyelid taping and patching can be tried when the condition is believed to be temporary. In severe, recalcitrant cases, with progressive corneal deterioration, eyelid reconstruction, partial tarsorrhaphy and amniotic membrane transplants can be beneficial. Orbital decompression for proptosis and eyelid gold weights for seventh nerve palsies can be helpful.

Recommendations

• Close observation is necessary when an infection is present or a high probability of infection exists.
• Floppy eyelid syndrome and Parkinson’s disease can also cause a poor blink and result in exposure staining.
• Petroleum jelly preparations applied to the periorcular skin can be therapeutic in certain cases of ectropion and may obviate the need for surgery.

ICD-9 Codes

• 370.34 Exposure keratitis
• 374.2 Lagophthalmos
• 374.5 Floppy eyelid syndrome
• 374.0 Entropion
• 374.1 Ectropion

References


FOREIGN BODY

A variety of materials in the environment, often metallic or organic, resulting in foreign body of the eye can be found on the surface of the cornea or conjunctiva, or even intraocular/intraorbital. Fortunately, the majority of foreign bodies encountered in clinical practice lodge on either the cornea or conjunctiva and do not enter the eye or orbit. A dilated eye examination must be performed to rule out any posterior segment involvement and appropriate testing such as B scan ultrasonography, computed tomography (CT scan) of the orbit or ultrasonographic biomicroscopy (UBM) to rule out an intraocular or intraorbital foreign body should also be done when necessary.

Presentation

Patients will generally provide a history of trauma and foreign body sensation with any foreign body of the eye can be found on the surface of the cornea and/or conjunctiva. Additional symptoms include photophobia and lacrimation. A slit lamp examination will reveal a single or multiple foreign bodies. Metallic corneal foreign body will produce rust. The eye is injected with eyelid edema and punctate keratopathy is seen with a foreign body under the lid. An anterior chamber reaction is possible with any significant involvement.

ICD-9 Codes

• 930.0 Corneal foreign body
• 930.1 Foreign body in the conjunctival sac
• 360.60 Foreign body-intraocular
• 930.8 Foreign body in other and combined sites on external eye

Treatment

A careful history will determine the mechanism of injury. A lid speculum may aid in the evaluation. Some attempt should be made to determine the size, shape, weight, velocity, force and composition of the object. Visual acuity assessment should be performed before any procedure is attempted. Topical anesthetic agents can control pain and blepharospasm to aid in examination.

Slit lamp examination will determine the location, depth and whether there are any self-sealing lacerations. Evert the eyelids and carefully inspect the fornices. With conjunctival laceration be certain Tenon’s membrane is intact to rule out scleral laceration or perforation of the globe.

Corneal foreign body can be removed by use of a fine forceps or foreign body spud at the slit lamp after topical anesthetic is applied to the eye. Multiple superficial foreign bodies may be irrigated. Broad spectrum topical antibiotics should be used to treat the resultant defect after assessing the size of the defect (see Corneal Abrasion section, p. 16-17, for a more complete discussion).

Similar treatment employed for any epithelial defect or abrasion can be used for pain management. Conjunctival (and sometimes corneal) foreign body can usually be removed successfully using fine forceps or a cotton-tipped applicator soaked in topical anesthetic. Residual and not easily accessible conjunctival foreign bodies may sometimes be left without undue harm unless they are infectious or pro-inflammatory. Deep injury to the posterior cornea may perforate the cornea and require sutures, glue and or a bandage contact lens.

Recommendations

• Carefully examine the anterior chamber and iris for any signs of intraocular foreign body. A low intraocular pressure or anterior chamber shallowing effects may indicate corneal penetration.
• Any infiltrate suggesting an infectious process must be managed with appropriate antibiotic therapy.
• Remove any rust as completely as possible. This might require allowing time for the rust to migrate to the surface of the cornea with removal of the rust after several days in order to minimize scaring.
• Routine dilated examination, gonioscopy and intraocular pressure checks may be necessary depending on the extent of the injury.

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Chemical

TOXIC/SOLUTION KERATITIS

Chemical toxicity can result from any contact lens solution, and care products are an under-recognized cause of intolerance to contact lens wear. A wide range of signs and symptoms are possible, ranging from mild sensitivity from a micro-punctate keratopathy to significant discomfort as a result of diffuse corneal staining. The patient may experience a foreign body sensation and photophobia. The condition is generally bilateral and disappears after removing the offending agent.

Etiology

Virtually any contact lens care product has the potential to create a toxic effect to the ocular surface, especially inadvertent application of non-neutralized hydrogen peroxide directly in the eye. A less severe adverse ocular response of diffuse punctate staining and conjunctival redness has been associated with wearing contact lenses, especially silicone hydrogel lenses and using certain multipurpose disinfecting solutions.1 Solution-induced staining should not be classified as a toxic response as this finding (what the fluorescein staining actually represents) is being debated.

Presentation

Patients experiencing a toxic response often present with symptoms of stinging, tearing, burning, dryness and a decreased wearing time. Signs include redness, conjunctival chemosis, follicular response, variable corneal staining patterns and infiltrates.

Treatment

Palliative therapy should include the use of artificial tears (non-preserved) or gels/oointments and antibiotic prophylaxis when indicated for more severe forms of coalesced corneal staining. Topical corticosteroids or antibiotic/steroid combinations may be warranted especially when there is an infiltrative response, assuming the risk of corneal infection has been carefully considered and ruled out. A change in solution is advised to remove the offending agent, and some clinicians will switch from a multipurpose solution to an oxidative hydrogen peroxide system.

Recommendations

• Some mild forms of toxicity/hypersensitivity are rarely detectable and the use of vital stain is essential in evaluating patients who are symptomatic with lens wear.
• Obtain a careful history in contact lens wearers including solution used, lens type and replacement frequency.

ICD-9 Codes
• 370.21 Punctate keratitis
• 370.3 Certain types of keratoconjunctivitis
• 370.4 Other and unspecified keratoconjunctivitis

References

CHEMICAL BURNS

Chemical injuries following exposure to solid, liquid or gas forms of corrosives have the potential to permanently damage the ocular surface (unilateral or bilateral). Pathophysiological cascades that may influence the final visual outcome include: 1) ocular surface injury and repair, 2) stromal matrix repair and or ulceration and 3) corneal inflammation.1-3 Ocular burn severity correlates to exposure duration and noxious agent.1 Specifically, chemical burn severity relates to pH, duration, solution quantity and permeability.2 Ocular burns caused by acids are generally less severe compared to alkali burns due to the natural buffering capacity of the corneal stroma and the barrier to penetration formed by coagulated epithelial cells.1 Immediate therapy should include prompt irrigation and removal of any remaining reservoir of chemical contact.1,2 Ocular burns represent 7-18% of ocular trauma. The vast majority are chemical burns that occur in the industrial/occupational setting.4

Etiology

Any noxious agent or radiant energy (thermal or ultraviolet) has potential to irritate/injure the ocular surface and produce other more severe forms of anterior segment abnormality. This includes alkali (i.e., lye, cements, plasters, airbag powder), acids, solvent, detergents and irritants (i.e., mace).3 Burns damage tissues primarily by denaturing and coagulating cellular proteins and through vascular ischemic changes.1,4
• Thermal burns: Injury from radiant energy results from contact with hot liquids, gases or molten metal. Cell death from thermal burns is generally limited to the superficial epithelium, but thermal necrosis and deeper penetration can occur.4
  • Ultraviolet burns: Punctate keratitis results from an epithelial injury. Delayed pain is secondary to actinic keratitis.1
  • Alkali burns: Since alkali substances are more lipophilic, they penetrate more rapidly than acids.1,3 The damaged tissues stimulate an inflammatory response that damages the tissue further by release of proteolytic enzymes (liquefactive necrosis) and alkali substances can pass into the anterior chamber rapidly, exposing the crystalline lens, ciliary body and trabecular meshwork.2,5 When the pH value is above 11.5, irreversible damage occurs.7
  • Acid burns: Acid burns cause protein coagulation of the epithelium, thereby limiting in most cases further penetration and progression beyond the superficial cornea.3

Presentation

Patients will often give a history of a liquid or gas being splashed or sprayed into the eye or of particles falling into the eye.4 The local Poison Control Center may be an invaluable resource in determining the nature of the chemical when unknown.1

Acute alkali burn

Alkali burn status post stem cell transplant with clear visual axis

Alkali burn status post stem cell transplant with clear visual axis

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For each condition, the review of inflammation, extent of involvement, and management is presented.

Inflammatory Corneal Conditions

STAPH MARGINAL KERATITIS/CLPU
Etiology
A contact lens-induced peripheral ulcer (CLPU) typically presents as a single well-circumscribed, circular, dense focal corneal infiltrate involving the anterior corneal layers. It is sometimes difficult to distinguish between a CLPU and microbial keratitis (MK), so MK is always considered part of the differential diagnosis. Differentiating a CLPU from early stage MK is primarily based on clinical judgment rather than on microbiologic or histopathologic investigations. However, there is often an overlap in the signs and symptoms, which can complicate the diagnosis. A critical sign, however, is the response of the presentation immediately after lens wear is discontinued.

CLPUs are different from peripheral marginal ulcers caused by S. aureus exotoxins that are often found in the corneal periphery, which are more often oval and may be associated with corneal vascularization. However, a recent study showed that bacterial carriage on contact lenses during extended wear predisposes the wearer to the development of corneal inflammatory events including
contact lens-associated red eye (CLARE), contact lens peripheral ulcers, and asymptomatic infiltrates. Any diagnosis of CLPU must be monitored carefully to ensure it is not MK.1

Presentation

Patients may complain of redness, discomfort, light sensitivity, tearing and foreign body sensation. These symptoms are milder than in microbial keratitis; however, suspicion must always be high in contact lens-wearing patients of a possible underlying microbial cause.

Slit-lamp examination will demonstrate mild to moderate conjunctival injection near the corneal infiltrate and a round, focal peripheral infiltrate approximately 2mm or less. The overlying small epithelial defect stains with fluorescein. Anterior chamber reaction, if present, is very mild. A significant anterior chamber reaction raises the suspicion of microbial keratitis.

Treatment

The initial management of CLPU involves discontinuing contact lens wear. Patients are often started on topical antibiotics such as a fourth-generation fluoroquinolone and followed with serial examinations for evidence of either improvement or worsening of the condition to rule out microbial keratitis. Topical steroids should generally be withheld until the epithelial defect is healed and there is no evidence of fungal, protozoan, or herpetic infection. In staph-induced marginal keratitis where severe inflammation exists, antibiotic and corticosteroid combination drops or ointments can be rubbed into the lid margins following lid scrubs.

Oral therapy with tetracycline 250mg four times a day or doxycycline 100mg twice a day or minocycline 50mg twice a day may be needed for more severe infections. The lesion generally heals with a small, mid-stromial scar and recurrences are common.

ICD-9 Codes

- 370.00 Corneal ulcer, unspecified
- 370.01 Marginal corneal ulcer

DRY EYE

Etiology

From the 2007 DEWS Report: “Dry eye is a multifactorial disease of the tears and ocular surface that results in symptoms of discomfort, visual disturbance and tear film instability with potential damage to the ocular surface. It is accompanied by increased osmolarity of the tear film and inflammation of the ocular surface.” Dry eye disease is a common disorder that especially affects adults and women and can cause quality of life impairment comparable to migraine, shortness of breath and chronic renal insufficiency, depending on its symptoms or complications.1 There are two etiopathogenic distinctions of dry eye: aqueous deficient dry eye and evaporative dry eye. There are numerous causes for each. Dry eye is the single most common complaint among contact lens wearers. It is the number one reason patients discontinue contact wear. Approximately 34% of patients discontinue contact lens wear at least once, most frequently because of dry eye symptoms.

Presentation

Physical examination includes visual acuity measurement, external examination, and slit-lamp biomicroscopy.2 Additional diagnostic tests may be performed to assess tear film stability, ocular surface damage, and aqueous tear flow. Tear film instability is commonly evaluated by performing a tear breakup time (TBUT) test. Ocular surface damage is commonly assessed by staining with rose bengal, lissamine green or fluorescein dye. Abnormal corneal and/or conjunctival staining patterns, observed on slit-lamp examination, are a sign of damage. Aqueous tear flow is commonly assessed by performing a Schirmer test. While helpful in making the diagnosis, diagnostic test results generally correlate poorly with symptoms.3 There are two new diagnostic tests available commercially: The TearLab Osmolarity System is intended to measure the osmolarity of human tears to aid in the diagnosis of dry eye disease in patients suspected of having dry eye disease. Advanced Tear Diagnostics has just launched a Tear Assay system that measures the quantity of lactoferrin in dry eye patient populations. Lactoferrin is a marker that will indicate the secretory function of the lacrimal gland and aid in determining if the condition is aqueous deficient dry eye or evaporative dry eye. The system also tests IgE levels that can mask as dry eye.

Treatment

There is no single method for determining if a patient is a candidate for dry eye therapy. Patient symptoms and clinical signs should be considered when deciding on therapeutic intervention. The management of dry eye disease encompasses both pharmacologic and non-pharmacologic approaches, including avoidance of exacerbating factors, eyelid hygiene, tear supplementation, tear retention, tear stimulation, and anti-inflammatory agents.4 Artificial tears are the mainstay of dry eye disease therapy but, although they improve symptoms and objective findings, they do not resolve the underlying inflammation. Topical corticosteroids are effective anti-inflammatory agents, but are only used short-term because of their adverse-effect profiles. Topical cyclosporine—currently the only approved pharmacologic treatment for DED—is safe for long-term use and is a disease-modifying therapy.

Treatment selection is guided primarily by DED severity.5 Replacement of tear volume with nonpreserved wetting agents and standard typical anti-inflammatory corticosteroid and/or cyclosporine A continues to be the current central conventional therapy for dry eye.6 Autologous serum eye drops have been reported to be effective for the treatment of severe dry eye-related ocular surface disorders (Sjögren’s syndrome).7 With appropriate management of the ocular surface conditions that produce dry eye, careful selection of contact lenses and solutions, and vigilant follow-up, successful CL wear should be achievable for the dry eye patient.8

ICD-9 Codes

- 370.33 Keratoconjunctivitis sicca, not Sjogren’s
- 370.23 Filamentary keratitis

UVEITIS

Etiology

Anterior uveitis is the most common form of uveitis encountered in general ophthalmic practice...
practice with acute anterior uveitis (AAU) accounting for 90% of all cases of uveitis. Half of all cases of AAU are HLA-B27 positive. It is often self-limiting, but can, in some cases, lead to complications such as posterior synechia, cataract, glaucoma and chronic uveitis.

**Presentation**
Symptoms of unilateral pain, photophobia, redness and watering develop over one to two days, with little or no effect on vision. Pain and photophobia may precede slit lamp signs, due to unseen ciliary body inflammation. Signs depend on the severity of the inflammation but typically include circumlimbal flush, miosis, anterior chamber cells and flare, and small, fine keratic precipitates (KP). The pupil may appear sluggish and may be small and irregular. In more severe attacks, posterior synechia, peripheral anterior synechia, corneal edema, spill-over vitritis and macular edema may be seen. The intraocular pressure (IOP) generally falls slightly with attacks but occasionally it may increase.

**Treatment**
In the majority of cases of AAU, especially those of mild to moderate intensity, topical applied steroids are the mainstay of treatment. Dosage regimens for topically applied steroids depend on the severity of AAU but all strategies must include aggressive initial treatment with the goal of bringing the inflammation under control quickly. Hourly instillation for the first few days is typical. A loading dose to achieve a therapeutic concentration at an early stage is often recommended.

Mydriatic/cycloplegic agents may also be added to reduce pain and prevent the development of posterior synechiae. Topical corticosteroids and cycloplegic agents remain the cornerstone of treatment for AAU.

**ICD-9 Codes**
- 364.00 Acute and subacute iridocyclitis, unspecified
- 364.01 Primary iridocyclitis
- 364.02 Recurrent iridocyclitis
- 364.05 Hypopyon

**CONTACT LENS SUPERIOR LIMBIC KERATOCONJUNCTIVITIS**
**Etiology**
Contact lens superior limbic keratoconjunctivitis (CLSLK) and superior limbic keratoconjunctivitis (SLK) are inflammatory conditions that affect the superior bulbar conjunctiva and adjacent corneal surface. CLSLK is recognized as a completely separate event from Theodore’s SLK. CLSLK is identified in patients wearing soft lenses by superior corneal staining combined with tarsal and superior limbal hypertrophy. There may be an association with atopy, contact lens-related trauma, hypoxia, hypersensitivity responses to proteins deposited on contact lens and chemical preservatives, especially thimerosal in lens care systems. SLK on the other hand is seen in older populations of women and might be related to thyroid eye conditions or other autoimmune diseases.

**Presentation**
Both conditions will manifest with punctate staining of the superior corneal epithelium, although CLSLK is often a more aggressive presentation. SLK is typically a bilateral condition and is clearly evident upon visualization. CLSLK is usually monocular with tremendous variability of presentation. Patients with CLSLK may complain of contact lens intolerance, light sensitivity, burning, redness, watering and pain. Upon lifting the upper lid, an intense area of localized injection at 12 o’clock with associated loose and boggy bulbar conjunctiva can be observed. An irregular epithelial surface, punctate staining with fluorescein, and subepithelial infiltrates may be found on the superior aspect of the cornea in association with hyperemia of the superior bulbar conjunctivae. There may be significant neovascularization and fibro-vascular pannus extending into the superior cornea. Evert the eyelids on all contact lens patients, paying careful attention to the bulbar and tarsal conjunctivae at the superior limbus.

**Treatment**
Case management for CLSLK is straightforward. Discontinuing contact lens wear and treatment with frequent preservative-free topical lubricants is the initial therapy. In most cases, this palliative treatment will suffice. In more severe presentations where the patient is experiencing significant discomfort, it might be necessary to concurrently treat the patient with steroid drops during the day and steroid ointment in the evening. Dosing schedule will vary depending on the case presentation. The treatment for SLK is generally more involved and can require therapeutic, conjunctival resection or chemical cautery with silver nitrate solution.

**ICD-9 Codes**
- 370.21 Punctate keratitis
- 371.82 Corneal disorder due to contact lens
- 372.10 Chronic conjunctivitis, unspecified

**References**
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