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Eyelid and Orbital Infections

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1. Introduction

The major infections of the ocular adnexal and orbital tissues are preseptal cellulitis and orbital cellulitis. They occur more frequently in children than in adults. In Schramm's series of 303 cases of orbital cellulitis, 68% of the patients were younger than 9 years old and only 17% were older than 15 years old.

Orbital cellulitis is less common, but more serious than preseptal. Both conditions happen more commonly in the winter months when the incidence of paranasal sinus infections is increased. There are specific causes for each of these types of cellulitis, and each may be associated with serious complications, including vision loss, intracranial infection and death. Studies of orbital cellulitis and its complication report mortality in 1-2% and vision loss in 3-11%. In contrast, mortality and vision loss are extremely rare in preseptal cellulitis.

1.1 Definitions

Preseptal and orbital cellulites are the most common causes of acute orbital inflammation. Preseptal cellulitis is an infection of the soft tissue of the eyelids and periorbital region that is localized anterior to the orbital septum outside the bony orbit. Orbital cellulitis (3.5 per 100,000) is an infection of the soft tissues of the orbit that is localized posterior to the orbital septum and involves the fat and muscles contained within the bony orbit. Both types are normally distinguished clinically by anatomic location.

1.2 Pathophysiology

The soft tissues of the eyelids, adnexa and orbit are sterile. Infection usually originates from adjacent non-sterile sites but may also expand hematogenously from distant infected sites when septicemia occurs. Preseptal cellulitis usually originates from skin infection with or without local trauma. It may also originate from structures inside the eyelid that are connected to the surface and become infected such as external and internal hordeolom. Chalazion is an example of internal hordeolom and these are all infected glands with surface connections. Glands with even partial preseptal location such as the lacrimal gland, in which the palpebral lobe is located preseptally, may also cause preseptal cellulitis.

Orbital cellulitis occurs in the following three situations:

- Spreading of an infection from the periorbital structures, usually from the paranasal sinuses, but also from the face, the globe and the lacrimal sac.
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- Direct inoculation of the orbit from surgical trauma.
- Hematogenous spread from distant sites (bacteremia).

In case of local cutaneous infection, preseptal cellulitis can arise from the spread of a contiguous anterior eyelid infection such as a chalazion, local trauma resulting in infection such as insect bite, or a foreign body. The skin and, in some instances, the sinuses and lacrimal mucosa, are colonized by various microorganisms. Orbital cellulitis following trauma is the consequence of a direct exposure of the orbital contents to these microorganisms. Open periorbital fractures, as well as closed fractures involving the sinuses or the nasal bone, may be a risk factor for orbital infections.

Orbital cellulitis, in contrast, usually arises from spread of infection from the paranasal sinuses. The ethmoid sinus is the most common source that extends to the orbit in children. In adults, pansinusitis is often accompanied by orbital cellulitis and its spread is believed to be caused through the ethmoidal or frontal sinuses. The ethmoid sinus separeated from the orbit medially by the thinnest orbital bone – lamina papyracea. Often, the lamina contains congenital dehiscences through which sinus infections can easily spread into the orbit. This may support the frequency of orbital cellulitis secondary to ethmoiditis. The anterior and posterior ethmoidal foramina may also serve as potential passages for infection.

The orbital roof borders the frontal sinus. It is a diploeic bone and is thicker than the lamina papiracea. Infection may spread more easily through the valveless facial veins. Since the frontal sinus is adjacent to the anterior cranial fossa, it may serve as an intermediary for the spread of infection.

The orbital floor that borders the maxillary antrum also contain congenital dehiscences through which infection from the maxillary sinus can enter and facilitate infection spreading.

The posterior medial wall of the orbit borders the sphenoid sinus. Isolated sphenoiditis is rare. The sphenoid may be involved secondary to ethmoiditis. Sphenoethmoidal sinusitis has distinct clinical characteristics. Marked visual loss with or without ophthalmoplegia usually precedes the findings of proptosis and inflammatory orbital signs. This condition is rare due to the thick bony barrier and firm attachment of periorbita to the posterior orbital wall.

One dehiscence or more is often present in orbital walls, particularly in the thin-walled lamina papyracea and this facilitates the spread of infection to the orbit. Posteriorly, the optic nerve within the optic canal is adjacent to the lateral wall of the sphenoid sinus. Dehiscence can also be found in the lateral wall of the sphenoidal sinus adjacent to the optic canal. The free valveless venous communication between the orbit and the sinus is another reason predisposing the orbit to the spread of adjacent sinus infection.

However, not all orbital cellulitis infections caused by sinus disease are secondary to acute sinusitis. Orbital fracture involving sinuses may allow spreading of an existing chronic sinus infection. Open fractures may also result in orbital cellulitis due to direct contact with the environment. Foreign bodies, such as a glass, wood or orbital floor implants, can cause orbital cellulitis. Infection can extend to the orbit from the eye, teeth, middle ear, or in neglected cases of preseptal cellulitis, from the eyelids and face.

Uncomplicated eyelid, strabismus, cataract surgery, glaucoma valve and retinal surgery may all expose the orbit to infection. Orbital implants that are imbedded and other foreign bodies,
such as Molteno valve implant, may carry a risk for infection, especially if they are colonized or exposed. Orbital cellulitis secondary to keratitis may develop after radial keratotomy.

Advanced carries with secondary infection, poor dental work or an infected root or dental cyst can cause orbital cellulitis. Extraction of maxillary premolars, molars or canines exposes the patient to orbital infection. The most common pathway for odontico-orbital infections is through the paranasal sinuses. The apices of maxillary molars and premolars are in close proximity to the floor of the maxillary sinus and are, in fact, in direct contact with the maxillary mucosa. Direct fistula to the antrum may be caused by a floor fracture during dental extraction or maxillary mucosa disruption. Infection spreading from the sinus to the orbit can occur through congenital dehiscences in the medial orbital wall or through communication between the venous plexus of the maxillary mucosa and the ophthalmic veins, thereby causing thrombophlebitis.

Another pathway for the spread of infection is the thin buccal cortical plate of the alveolar processes.

Finally orbital cellulitis can occur secondarily from embolic spread in subacute bacterial endocarditis and from other distant organs.
1.3 Classification

In 1970, Chandler described a spectrum of progressive infectious changes in orbital cellulitis.

Chandler’s Classification:

Stage I – Preseptal cellulitis

Occasionally, edema may spread secondarily to preseptal cellulitis posterior to the septum without infection. In these cases chemosis may be present, but the extraocular movements and visual acuity remain intact.

Stage II – Orbital cellulitis

Diffuse edema of orbital contents, with leukocytosis, fever, proptosis and impaired extraocular motility without discrete abscess formation.

Examples are shown in the images below:

A male with left orbital cellulitis presented with proptosis, ophthalmoplegia, and edema and erythema of the eyelids. The patient also exhibited pain on eye movement, fever, headache, and malaise.
A male with left orbital cellulitis with proptosis, ophthalmoplegia, and edema and erythema of the eyelids. The patient also exhibited chemosis and resistance to retropulsion of the globe.

**Stage III - Subperiosteal abscess**

The globe is often displaced and limited in the field of gaze of the abscess.

An axial computed tomography scan in a patient with a right orbital infection caused by *streptococcus pneumoniae* and a right superior orbital subperiosteal abscess that resulted in blindness.

A coronal computed tomography scan of a child with pansinusitis as well as a left orbital and subperiosteal abscess.
A coronal computed tomography scan in a patient with sickle cell disease. In this image, the patient has a left subperiosteal bleeding that mimicked the appearance of an infectious subperiosteal abscess.

**Stage IV – Intraorbital abscess, purulent collection:**

These patients have severe proptosis, chemosis, ophthalmoplegia and often visual loss.

Frontal view of the patient with a right orbital abscess showing periorbital redness, swelling and proptosis.

Coronal computed tomography scan in a pediatric patient with pansinusitis sinusitis and left orbital abscess.
Axial computerized tomography scan shows a right classic proptosis associated with an abscess of the orbit, as well as displacement of the medial orbital tissues and tenting of the posterior.

**Stage V – Cavernous sinus thrombosis (septic abscess)**

In these instances, the orbital signs evolve in the fellow eye (bilateral) and other central nervous system signs supervene.

A T1 weighted coronal MRI demonstrating asymmetry between the cavernous sinuses with obliteration of the right cavernous sinus.

2. **Preseptal cellulitis**

Preseptal cellulitis is more common than orbital cellulitis. It can present with swelling and erythema of tissues surrounding the orbit, with or without fever. Preseptal cellulitis most commonly occurs from a contiguous infection of the soft tissues of the face and eyelids.
secondary to local trauma, foreign bodies or insect or animal bites. It is rare for untreated preseptal cellulitis to progress to orbital cellulitis by local extension. Defining the exact location of inflammation is essential for proper diagnosis and treatment.

2.1 Symptoms and signs
Patients with preseptal cellulitis often have a short history (days) of painless swelling of the eyelids. A history of early upper respiratory tract infection, trauma, insect or animal bite, conjunctivitis, or chalazion may be disclosed. Fever is an inconstant feature. The eyelid characteristically is erythematous, edematous, tender and warm. Vision and pupillary response are always unaffected and proptosis, globe displacement and limitation in ocular motility are never present. Concurrent preseptal cellulitis was discovered in the presence of many systemic diseases including, varicella, asthma, nasal polyposis and neutropenia. Preseptal cellulitis is more common among children than in adults.

2.2 Microbiology
The most common inciting microorganisms include *Streptococcus pneumoniae*, *Staphylococcus aureus*, other Streptococcus species and anaerobes. *Haemophilus influenzae* type B was the most common cause in children under four years old. However, routine vaccination with conjugate *Haemophilus influenzae* vaccines since 1985 has dramatically decreased this infection in young children. Less commonly implicated microorganisms include *Acinetobacter* spp., *Nocardia brasiliensis*, *Bacillus anthracis*, *Pseudomonas aeruginosa*, *Neisseria gonorrhoea*, *Proteus* spp., *Pasteurella multocida*, *Mycobacterium tuberculosis* and *Trichophyton* spp.

2.3 Differential diagnosis
Conditions that might masquerade as preseptal cellulitis include allergic edema (anaphylactoid reaction) of the eyelids, severe blepharitis with scruffs (seborrheic),
collarettes (staphylococcal), sleeves (demodex) with or without erythema but no local warmth. The meibomianitis is characterised with eyelid swelling, pouting of meibomian gland orisices and discharge from orifices but no local warmth.

In addition dacryoadenitis, blunt trauma, thyroid eye disease, leukemic infiltrates, blepharochalasis syndrome and autoimmune inflammatory disorder such as lupus. Other disorders of less resemblance include orbital tumors/pseudotumours, orbital vasculitis, necrotising fascitis and others.

2.4 Management

Treatment regimens cover the most likely organisms to cause infection in this setting and according to case series. Outpatient treatment should include broad-spectrum oral antibiotics and close observation. The author came to a conclusion that most cases of preseptal cellulitis can be safely managed as outpatients with oral antibiotics and follow-up until improvement is documented. If the condition does not improve or deteriorates 48 hours or more after oral antibiotic treatment, the patients should be admitted for intravenous antibiotic treatment and close observation. The average time of hospitalization is four days.

The appropriate antibiotics in adults include Amoxicillin–clavulanate (Augmentin) 875 mg every 12 h, and in children 90 mg/kg/day amoxicillin and 6.4 mg/Kg/day of clavulanate divided to two doses. Another option includes Cefpodoxime (Vantin) 200 mg every 12 h in adults, and 10 mg/kg/d divided every 12 h in children with maximum daily dose of 400 mg. Cefdinir (Omnicef) 600 mg daily in adults, and 14 mg/kg/d divided every 12 h in children with maximum daily dose 600 mg is another option.

Pediatric patients under 1 year of age and all more severe cases require the same approach as patients with orbital cellulitis, namely, intravenous broad-spectrum antibiotics and hospital observation. Blood cultures should be obtained if the systemic fever increases. The recommended duration of antimicrobial therapy is for 7-10 days. Occasionally, patients will continue to have local signs of cellulitis at end of treatment. Oral antibiotic therapy is recommended to be continued in these patients until resolution of the erythema. Children with preseptal cellulitis, no orbital involvement, and who do not appear toxic can be treated with intramuscular or oral antibiotics on a daily basis as outpatients. Preseptal cellulitis in adults can be managed on outpatient basis with oral antibiotics with frequent monitoring for progression.

3. Orbital cellulitis

Orbital cellulitis occurs in three settings:

i. Extension of infection from periorbital structures, such as the face, lacrimal sac and globe, but particularly from the paranasal sinuses. Acute sinusitis is complicated by orbital cellulitis in 1-3% of cases and the coexisting sinusitis is present in 73-94% of patients with orbital cellulitis. Ethmoid sinusitis and pansinusitis are most likely to progress to orbital cellulitis. The first ocular sign of sinusitis may be preseptal inflammation only. This can then quickly progress to the classic clinical picture of orbital cellulitis. Other causes include – orbital trauma, with fracture or a foreign body, dacryocystitis, and infection of teeth, middle ear or face.
ii. Direct inoculation from accidental trauma or from surgery. Orbital cellulitis is an uncommon complication of ophthalmic surgery, being reported after strabismus surgery, blepharoplasty, radial keratotomy, retinal surgery and following peribulbar anesthesia. A special case is fungal orbital cellulitis, a relatively rare condition occurring in two principal forms - 1. Subacute infection due to genera of Zygomacetes (mucormycosis). 2. A more chronic orbital infection caused by species of Aspergillus. The distinction between these two may be difficult on clinical observation alone.

iii. Hematogenous spread (endogenous from bacteremia). The microorganisms responsible for most cases are aerobic non-spore forming bacteria.

3.1 Symptoms and signs
Orbital cellulitis as preseptal cellulitis can present with swelling and erythema. The cardinal signs and symptoms are proptosis, ophthalmoplegia and globe displacement. Pain on eye movement, vision loss (indicates orbital apex involvement), diplopia, conjunctival chemosis and elevated intraocular pressure are common but variable accompanying signs.

3.2 Microbiology
*S. aureus* and Streptococci are the most commonly identified organisms in culture-positive orbital cellulitis. Less common causes are *H. influenza* and non-spore forming anaerobes. However, many other common and rare bacterial pathogens include *Eikenella corrodens, Aeromonas hydrophila, P. aeroginosa*, fungal and mycobacterial pathogens, including *Scedosporium apospermum, M. tuberculosis* and *Mycobacterium avium* complex. The prognosis of aspergillosis is poor. More than 80% of reported patients have died from this causative.

3.3 Differential diagnosis
Conditions that may mimic orbital cellulitis include:

- Anaphylactoid reaction can be characterised with eyelid swelling and erythema but no local warmth, no proptosis, limited ocular motility or optic nerve involvement.
- Cavernous sinus syndrome that is characterised by proptosis, complete ophthalmoplegia, optic nerve involvement, V2 involvement, evolve to bilateral condition and usually in debilitated patients (diabetics, drug abusers, HIV).
- The orbital apex syndrome is characterised with complete external ophthalmoplegia, optic nerve and V1 involvement with or without proptosis.
- Superior orbital fissure syndrome is a condition that is characterised with complete external ophthalmoplegia, V1 involvement with or without proptosis.
- The orbital compartment syndrome and the orbital tumors can present with proptosis, limited ocular motility, optic nerve involvement and increased intraocular pressure but no local warmth.

3.4 Diagnosis
There have been no controlled trials examining the utility of radiologic studies (e.g. computed tomography scanning, orbital ultrasound, or magnetic resonance imaging) in the diagnosis of orbital cellulitis or in distinguishing preseptal from orbital cellulitis.
Computed Tomography scanning (CT)

CT can confirm extension of inflammation into the orbit, detect coexisting sinus disease, and identify an orbital or subperiosteal abscess. Whether every patient with suspected orbital cellulitis needs a CT scan is controversial. Some experts suggest that a CT scan be performed only in those patients who deteriorate or fail to respond to 48 hours of IV antibiotics, as the majority of patients with orbital cellulitis do well with conservative medical management. It is suggested that patients with suspected orbital cellulitis – those with proptosis, globe displacement, limitation of eye movements, double vision, vision loss, and those patients in whom the physician cannot accurately assess vision – usually patients less than one year of age, at presentation have a baseline CT scan.

Orbital Ultrasoundography (US)

US provides higher-resolution details of orbital contents and is useful when sequential follow-up of an abscess or drained abscess is required. However, orbital sonography is not widely available and is dependent on the expertise of the sonographer.

Magnetic Resonance Imaging (MRI)

MRI is superior to CT in the resolution of soft tissue disease. However, it is not usually performed because of the need for sedation in pediatric patients and because MRI is rarely immediately available.

Microbiologic studies

The causative microorganism in orbital cellulitis may be difficult to identify due to normal flora contaminants, mixed infection, and prior antibiotic therapy. Cultures for aerobic and anaerobic organisms may be obtained from blood, sinus aspirates, and abscess. Because blood cultures are usually negative, some clinicians obtain cultures of eye secretions or pharyngeal culture. However, these cultures are likely to be contaminated with normal oropharyngeal flora and should not guide the choice of antibiotic therapy. Microbiologic data limited to microorganisms recovered by surgical drainage from orbital abscesses or involved sinuses and/or positive blood culture are the most reliable information. It is recommended that in patients with suspected orbital cellulitis, blood cultures should be obtained before the initiation of antibiotic treatment. If surgery is performed, culture of abscess material or sinus contents should be sent for aerobic, anaerobic and fungal cultures.

3.5 Complications

The complications of bacterial orbital cellulitis may be orbital or intracranial. Orbital complications of orbital cellulitis include subperiosteal or orbital abscess formation in 7-9%, permanent globe displacement, limited ocular motility that may cause diplopia, and vision loss in 1%. Orbital abscess may be clinically indistinguishable from orbital cellulitis. Proptosis and globe displacement tends to be more severe with orbital abscess than in orbital cellulitis, and patients are more likely to be systemically ill. The diagnosis of orbital abscess is confirmed by imaging or at surgery. The identification of an orbital abscess on the baseline CT scan is important since these patients almost always require surgery. Intracranial complications, which are encountered in 4% of orbital cellulitis secondary to sinusitis include meningitis in 2%, cavernous sinus thrombosis in 1%, intracranial abscess
formation, epidural or subdural abscess or parenchymal brain abscess in 1%, and carotid artery occlusion. Intracranial involvement may be heralded by ophthalmoplegia, changes in mental status, contralateral cranial nerve palsy, or bilateral orbital cellulitis. Cavernous sinus thrombosis has become relatively rare in developed countries because of prompt and adequate treatment of most cases of acute sinusitis, but it still poses a major threat. The mortality rate of cavernous sinus thrombosis may exceed 50%.

**Permanent vision loss may occur because of:**

1. Corneal ulcer and perforation secondary to exposure or neurotrophic keratitis.
2. Destruction of intraocular tissues following neovascular or inflammatory glaucoma, endophthalmitis, septic uveitis or retinitis and exudative retinal detachment.
3. Various other mechanisms affecting the globe or posterior orbit, such as secondary glaucoma due to elevated orbital pressure, infectious optic neuritis or inflammatory optic neuritis, pressure effects the optic nerve, thrombophlebitis of ocular veins and central retinal artery occlusion.

Blindness can result from elevated intraorbital pressure causing optic neuropathy or extension of the infection to the optic nerve from the sphenoid sinus.

### 3.6 Management

Prompt administration of appropriate antibiotics is the key for successful treatment of orbital cellulitis. After appropriate workup, all periorbital and orbital infections should be treated with broad-spectrum antimicrobial agents. There have been no controlled trials examining the required duration of antimicrobial therapy in orbital cellulitis. Treatment regimens are based upon coverage of the most likely organisms to cause infection in this setting and treatment of case series. The initial empiric antibiotic treatment should consist of parenteral broad-spectrum therapy. Infection due to methicillin-resistant *S. aureus* is best treated with vancomycin, clindamycin and cefotaxime.

Fungal orbital cellulitis occurs and is primarily due to mucor and aspergillums species. It requires antifungals, such as amphotericin. Corticosteroids may be helpful in bacterial infections, but they should not be started before surgery and until the patient has been on appropriate antibiotics for 2-3 days to ascertain eradication of the microbial agents.

If secondary glaucoma develops, ocular anti-hypertensive agents should be initiated promptly. Prompt diagnosis and therapy are important since delayed intervention can result in sustained vision loss.

#### 3.6.1 Antibiotic treatment

Due to increasing incidence of Methicillin-resistant *S. aureus*, empiric therapy with Vancomycin (Vancocin) (15 mg/kg IV every 12 hours in adults, 10 to 15 mg/kg IV every 6 hours in children, maximum daily dose of 4gr) is recommended. If susceptibility testing reveals Methicillin-sensitive *S. aureus*, Vancomycin should be replaced with Nafcillin (Unipen), or Oxacillin (Bactocill) - (both agents are dosed at 2gr IV every 4 hours in adults, and 200 mg/kg per day IV in 4-6 divided doses in children, maximum daily dose of 12gr) since these agents have better CNS penetration than vancomycin.
One of the following should be added:

1. Ampicillin-sulbactam (Unasyn) 3gr IV every 6 hours in adults, 300 mg/kg per day in 4 divided doses in children, maximum daily dose of 12 gr.

2. Ticarcillin-clavulanate (Ticar), which covers most of the Gram-negative bacteria as well as Gram-positive organisms, including atypical H. influenza, and has also excellent anaerobic coverage. The dosage is 3.1gr IV every 4 hours in adults, 200-300 mg of ticarcillin component per kg per day in 4-6 divided doses in children of less than 60 kg. The maximum daily dose of ticarcillin component is 18 gr.

3. Piperacillin-Tazobactam 4.5gr IV every 6 hours in adults, 240 mg/kg per day in 3 divided doses in children, with a maximum daily dose of 16gr of piperacillin component.

4. Ceftriaxone (Rocephin) is effective against penicillinase-producing S. aureus, most Gram-positive organisms, and most Gram-negative organisms except for Pseudomonas. Ceftriaxone also crosses the blood-brain barrier; therefore, it is an excellent choice if there is a suspicion of concurrent intracranial infection. 2gr IV every 12 hours in adults, 80-100 mg/kg per day in 2 divided doses in children, maximum dose of 4gr daily may be given.

5. Cefotaxime (Claven), a third-generation cephalosporin that covers most of the common sinus pathogens with the exception of Clostridium difficile, may be given 2gr IV every 4 hours in adults, 150-200 mg/kg per day in 3-4 equally divided doses in children, with a maximum daily dose of 12gr.

Patients allergic to penicillin and/or cephalosporins may be treated with a combination of vancomycin and a fluoroquinolone. For patients over 17 years of age, ciprofloxacin 500 mg twice a day or levofloxacin 500 mg daily may be prescribed. Fluoroquinolones are not recommended for use in pediatric patients as first-line therapy for any infection because of the musculoskeletal side effects - The mildest side effects include muscle pain, called fibromyalgia. A more serious side effect, though also less common, is tendon damage. Fluoroquinolones can, in high doses, cause tendon damage, which can ultimately lead to rupture of the Achilles tendon (in addition gastrointestinal effects predominating (nausea, vomiting, diarrhea, or abdominal pain in 1.0%-5.0% of the patients), followed by effects on the central nervous system (dizziness, headache, and/or insomnia in 0.1%-0.3% of the patients) and skin (0.5%-2.2% of the patients). Elevation in levels of hepatic enzymes occurred in 1.8%-2.5% of the patients, anemia in 0.2%-1.3%, and eosinophilia in 0.2%-2.0%. Initial antibiotic therapy should be administered IV under hospitalization. Generally switching to oral therapy is done after the patient is afebrile and skin findings have begun to resolve, which usually take 3-5 days.

The duration of treatment depends on the response. Patients should be treated with parenteral antibiotics until they show clear evidence of clinical improvement as manifested by decrease in orbital congestive signs such as proptosis, gaze limitation, cellulitis and edema. Intravenous therapy should continue for a minimum of 3 days. Then oral antibiotic therapy may be instituted for a total course of 10 days to 3 weeks, depending on the severity of infection. Associated bacteremia, however, should be treated with 7-10 days of IV therapy.

As aforesaid, the oral regimen should be tailored based upon the results of the cultures. If the results are not available, reasonable empiric oral antibiotic choices include 1. amoxicillin
clavulinate – 875 mg twice daily for adults or children over 40 kg, and 45 mg/kg/day divided every 12 hours for children over 3 months and under 40 kg or 2. fluoroquinolone – (levofloxacin 750 mg once daily in adults). Linezolid (ZYVOX) - (600 mg twice daily in adults and children over 12 years of age, 10 mg/kg three times daily for children under 11 years) should be added if MRSA is suspected.

Careful follow-up is indicated in all patients who present with orbital cellulitis. This should include twice-daily examinations with attention to visual acuity, confrontation visual fields, exophthalmometry, motility and pupillary examination.

3.6.2 Antifungals

(Amphotericin B- Ambisome) 1 mg/kg IV q24h or Voriconazole (VFEND, Pfizer) 6 mg/kg IV q12h for 2 doses, then 4 mg/kg IV q12h or Voriconazole 200-300 mg PO q12h

Antifungal is the treatment of choice for fungal orbital cellulitis. It is administered IV and may be appropriately administered before laboratory confirmation of fungal infection in cases of severe infection and debilitated patients (diabetes mellitus, drug abusers, human immunodeficiency disease, metastatic cancer, prolonged administration of antibiotics and/or corticosteroids).

3.6.3 Surgery

The timing for surgical intervention is critical. In cases of orbital cellulitis without abscess formation, in which visual acuity is 20/60 (Snellen notation), 6/15 (metric equivalent) or less, or declines with appropriate medical management, orbital exploration should be emergent. In cases in which the acuity is better than 20/60, the patient should be followed, expectantly, and frequently while more conservative management is initiated.

Orbital surgery is indicated if the patient:

1. Fails to respond
2. Deteriorates clinically despite treatment
3. Has worsening visual acuity or develops afferent pupillary defect
4. Develops an abscess, except selected pediatric cases with medial subperiosteal abscess, which may be successfully treated medically.

In patients older than 14 years of age, the author favours the latter approach because the risks of surgery are negligible compared with the visual and life-threatening risks of no intervention. In patients 9 years of age or younger, 25% of the subperiosteal abscesses are likely to resolve with antibiotic therapy alone. Several indications have been suggested for drainage of subperiosteal abscesses. These include age of 9 years or older, large abscess, frontal sinusitis, non-medial abscess, chronic sinusitis, dental infection, optic nerve involvement, suspicion of anaerobic infection and recurrence after drainage. All other cases may be managed conservatively by intravenous antibiotics.

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