# Purpose

This evidence-based guideline provides clinical practice advice for clinicians involved in the emergency management of children with Peri-Orbital and Orbital cellulitis.

# Scope

This guideline applies to all Queensland Health Hospital and Health Services Staff involved in the care and management of children with Peri-Orbital and Orbital cellulitis.

# Related documents

**Procedures, Guidelines, Protocols**

CHQ Paediatric Antibiodcard: Empirical Antibiotic Guidelines (CHQ-GDL-01202)
**Guideline**

**Introduction**

Infection of the skin and other soft tissues, in and surrounding the eye, is sometimes referred to by the umbrella term, ‘Peri-Orbital Cellulitis’ \(^1,2,3,4,5\). This can be confusing, as the term encompasses a range of disease processes, each with differing aetiologies and prognoses. It is important to understand the anatomy of the orbit, in order to appreciate the pathophysiology of infection in this area. Arising from the orbital rim is a tough fibrous layer of fascia, the ‘Orbital Septum’. This attaches to the tarsal plates of the eyelids, and provides a physical barrier to infection between the superficial structures of the face, and the deep orbital structures \(^1,2,3,4,6,7,8,9,10\). The infection will therefore usually occur in either the pre-septal, or post-septal tissues. Infections in the pre-septal region are most often referred to as ‘Peri-Orbital Cellulitis’ \(^1,2,3,11,12\), whilst infections in the post-septal region, involving the orbit and its contents, are commonly referred to as ‘Orbital Cellulitis’ \(^1,2,3,11,12\). These terms will be used in this guideline.

![Figure 1: Sagittal Cross Section of Orbit](#)  

All children with either ‘Peri-Orbital (pre-septal)’ or ‘Orbital (post-septal)’ Cellulitis will present with erythema and swelling of the eye and/or surrounding skin. The underlying disorder can often be very difficult to distinguish clinically \(^2,3,6,13,14\).

Peri-Orbital Cellulitis occurs due to local infection of the skin of the face \(^4\). This can be due to a superficial infection of the eyelids, such as dacrocystitis, or a styel \(^1,2,8,14,15\). It can also occur following a break in the skin of the face, such as an insect bite, or wound \(^2,7,8,14,15\). The most common organisms causing these infections are *Streptococcus pyogenes*, *Staphylococcus epidermidis*, and *Staphylococcus aureus* \(^6,8,9,15,16\).

Orbital Cellulitis is usually a complication of sinus disease \(^1,2,4,5,8,9,11,12,13,14,15,17,18\), orbital trauma, or less often, occurs via direct haematological spread \(^1,6,9,11,15\). The infection spreads most frequently from the ethmoid sinuses \(^1,10,11,12,13,14\). These lie directly medial to the orbits, and are separated by a thin bone layer, the lamina papyracea \(^1,2,3,4,6,7,11,14,19\). The most common causative organisms are *Streptococcus pneumoniae*, *Staphylococcus aureus*, *Streptococcus pyogenes*, *Staphylococcus epidermidis*, and *Haemophilus species* \(^1,2,3,4,5,6,8,9,10,11,13,15,16,17,18,19\). Prior to immunisation against *Haemophilus influenzae type B (HIB)*, this was a common cause of Orbital Cellulitis in children \(^2,11,14,16,17\).

Orbital Cellulitis is a sight, and potentially life-threatening, disease \(^3,9,13,14\). Infection within the orbit can lead to direct compression of the optic nerve causing blindness \(^1,2,12,13,14\). The infection may also spread into surrounding tissues and cause a sub-periosteal, or orbital abscess. If the infection spreads posteriorly via the valveless veins of the orbit, it can cause cavernous sinus thrombosis, intra-cerebral abscess or meningitis \(^1,2,3,4,5,6,8,10,11,13,14,18,19\). As Orbital Cellulitis is primarily a disease of the sinuses, patients are cared for by an Ear, Nose and Throat (ENT) team \(^19\), with Ophthalmology team consultation, and involvement of other multi-disciplinary teams as required.
Assessment

A thorough clinical assessment is required in all patients presenting with redness and swelling around the eye, to help determine the severity, and location (pre or post-septal) of any infection.

Ensure **adequate analgesia** (may require opiates). A child in pain will be difficult to examine thoroughly.

### History
- Age of patient (Peri-Orbital cellulitis more common in younger patients <5yrs \(^{1,4,6,10,15}\))
- Recent infections (upper respiratory tract infection, sinus, teeth, ears)?
- Eye problems (nasolacrimal duct obstruction, dacrocystitis, sty, chalazion, watery eye)?
- Injury to eyes, face or skin (insect bites, penetrating injury, eczema)?
- Recent surgery to eyes, nasolacrimal ducts (probe/ syringe), teeth, or sinuses?
- Immunisation status (especially **HIB**)?
- Co-morbidities (immune-compromise, diabetes)?
- Risk for non-multi resistant Methicillin Resistant Staphylococcus Aureus (nmMRSA)
  - previous nmMRSA, history of boils, Aboriginal or Pacific Islander Descent?
- Personal or family history of boils?
- Symptoms:
  - Redness and swelling around eye
  - Eye pain
  - Headache
  - Fever
  - Neurological symptoms (drowsy, altered level of consciousness)

### Examination
- General appearance (toxic or shocked)
- Temperature and other vital signs
- Neurological examination in the presence of altered conscious level
- Evidence of skin lesion/ wound as source for skin infection
- Eye:
  - Eyelids and surrounding skin (extent of erythema and swelling)
  - Conjunctiva (injection, chemosis, discharge)
  - Sclera (injection)
  - Proptosis
  - Eye movements (reduced movement, pain, diplopia)
  - Visual Acuity (use Snellen or Lea chart, as age-appropriate)
  - Visual Fields
• Colour Vision, specifically perception of Red (can use Ishihara colour plates if available)
  Note: Loss of red-colour perception is an early sign of optic nerve injury.
• Pupil size and reaction (include checking for relative afferent pupil defect [RAPD])
• Fundoscopy

Investigations

If clinical assessment is sufficient to diagnose peri-orbital erythema and swelling of a non-infectious cause, such as allergy, then no investigations are required, and the patient should be managed according to the clinical diagnosis. A thorough clinical assessment may also be sufficient to diagnose mild cases of infectious peri-orbital cellulitis, in which case, any investigation requests can be tailored to the patient.

- **Swab of any conjunctival discharge** for Microscopy & Sensitivity (MC&S)\(^3,4,5,9,11,14,15,18\), include specific swabs for *Chlamydia* and *Gonorrhea* in neonates
  • May help direct antibiotic therapy if initial treatment is unsuccessful

- **Blood Tests**\(^5\): Full Blood Count (FBC)\(^4,9,11,14,16\), C-Reactive Protein (CRP)\(^4,14\), Blood Cultures\(^4,9,11,14,15,16\), Urea & Creatinine, Electrolytes, Liver Function Tests\(^1,3,18,19\)
  • FBC and CRP may help differentiate between moderate peri-orbital, and severe peri-orbital or orbital cellulitis
  • Blood cultures may help direct antibiotic therapy in more severe infections

- Consider lumbar puncture if clinical concern for meningitis (caution re: raised intracranial pressure)\(^9,11,16\)

- **Medical Imaging**
  Usually best to be considered in consultation with ENT team, in order to limit radiation exposure\(^2,17,18\). Medical imaging is used to help to delineate the source of the infection, as well as diagnose complications (such as abscess) that may require surgical intervention.
  *(Note: Some patients may require general anaesthetic, due to young age.)*
  • **Computerised Tomography (CT) of orbits, brain and sinuses, with contrast.**
    • Initial imaging choice in the majority of cases\(^2,3,4,5,6,8,9,10,11,12,13,14,15,16,18,19\).
    *(Note: Use of contrast is important to reduce need for further imaging later, when considering surgery.)*
  • Magnetic Resonance Imaging (MRI) brain and sinuses\(^2,3,9,11,12,13,14,17,18,20\).
    • Gives less definition of bony disease, better for assessing intracranial complications.
  • Bedside Ultrasound may be considered *(in experienced hands only)*
    • Can help delineate pre-septal, and post-septal infection\(^14,19,21\).
    *(Note: Will then require CT to confirm, and give further information.)*
Alert:
If high-risk features are present, will require urgent medical imaging: 2,3,4,6,8,10,12,18,19

- Altered level of consciousness/ seizure
- Gross proptosis (especially with marked conjunctival chemosis)
- Ophthalmoplegia (reduced eye movements)
- Altered visual acuity or loss of red-colour perception
- Abnormal pupil response, or afferent pupil defect
- No clinical improvement, or deterioration, after 24hrs of appropriate intravenous antibiotics

Diagnosis

Peri-orbital Cellulitis

Mild
- Age >3 months 2 (if age <3 months, treat at minimum as ‘moderate’)
- Not immune-compromised
- Minimal erythema and swelling around eye, not involving eyelid 2
- Patient able to fully open, and allow Doctor to examine eye 2
- White sclera, with non-injected conjunctiva
- Eye movements normal, with no pain 7,8,9
- Vision intact (red-colour perception, fields and acuity) 8,9
- No fever 2,7
- White Cell Count (WCC), if tested, is normal 7
- Obvious history of insect bite/ skin lesion/ stye 2,6

Moderate
- Moderate erythema and swelling around eye and involving eyelid
- Patient or Doctor able to fully open eye, and allow Doctor to examine eye
- White sclera, with non-injected conjunctiva
- Eye movements normal, with no pain 1,7,8
- Vision intact (red-colour perception, fields and acuity) 8
- No fever 7
- WCC normal 7
- Obvious history of insect bite/ skin lesion/ stye 6
SEVERE

- Extensive erythema and swelling around eye and eyelid
- Doctor able to fully open eye, and to examine eye

(Note: If not able to open eye for assessing eye movements and pupils, then assume diagnosis is ORBITAL Cellulitis) 2,9,14,15

- White sclera, with non-injected conjunctiva
- Eye movements normal, with no pain 1,7,8
- Vision intact (red-colour perception, fields and acuity) 8
- Possible fever
- WCC may be elevated
- Medical imaging confirming pre-septal infection

ORBITAL CELLULITIS

- Erythema and swelling around eye and eyelid (may not be extensive in early stage) 5,7,9,11
- Injection of sclera, and conjunctiva
- Chemosis of conjunctiva 1,2,5,7,9,15 (late sign)
- Eye movements decreased 2,5,6,7,11,15,16, and painful 9,10 (late sign)
- Vision change (late sign = optic nerve compression) 2,3,5,7,9,15,16
- Altered pupil response (late sign = optic nerve compression) 2,3,9,11
- Diplopia (late sign) 1,3,6,9,11
- Proptosis (late sign) 1,2,3,6,7,9,10,11,15,16.
- Fever 2,3,5,6,9,11
- WCC elevated 3,7,10,11,14,15
- Headache and Nausea 3,6,9,11,16
- Risk factors for Orbital Cellulitis (Eye surgery, Sinus disease 15, Non HIB-immunised, Immune-compromise 2)

**ALERT:**

Diagnosis based on clinical findings can be very difficult 2,3,6,8,13,14,19.

If in doubt, treat with intravenous antibiotics, and refer for ENT (and Ophthalmology) opinions immediately 2.
Management

**Antibiotics**

**DO NOT DELAY** starting intravenous antibiotics if considering Severe Peri-Orbital or Orbital Cellulitis. Early treatment with antibiotics may be sight, or life-saving.

<table>
<thead>
<tr>
<th>INFECTION</th>
<th>1ST CHOICE ANTIMICROBIAL</th>
<th>Alternative if hypersensitivity to penicillins and cephalosporins</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>MILD</strong> Peri-Orbital Cellulitis</td>
<td><strong>Cephalexin Per Oral (PO)</strong> 25mg/kg/dose four times a day (Max 1000mg/dose) <em>(For children unable to swallow capsules)</em> or <strong>Flucloxacillin PO</strong> 25mg/kg/dose four times a day (Max 1000mg/dose) <em>(For children who can swallow capsules)</em></td>
<td>Immediate - type hypersensitivity: <strong>Trimethoprim/Sulfamethoxazole PO</strong></td>
</tr>
<tr>
<td><strong>If at risk of nmMRSA</strong> (previous nmMRSA, history of boils or Aboriginal or Pacific islander descent), or if family/personal history of boils</td>
<td><strong>Clindamycin PO</strong> 7.5mg/kg/dose four times a day (Max 450mg/dose) <em>(Round to multiples of 150mg, as only available in capsules)</em> or <strong>Trimethoprim/Sulfamethoxazole PO</strong> 4mg/kg/dose twice daily (Max 160mg/dose Trimethoprim component)</td>
<td></td>
</tr>
<tr>
<td><strong>MILD</strong> Peri-Orbital Cellulitis</td>
<td><strong>Flucloxacillin Intravenous (IV)</strong> 50mg/kg/dose every 6 hours (Max 2g/dose) for 48 hours, then seek ID review</td>
<td>Delayed - type hypersensitivity: <strong>Cephazolin IV</strong> Immediate - type hypersensitivity: <strong>Lincomycin IV</strong> and seek Infectious disease team (ID) advice</td>
</tr>
<tr>
<td><strong>If &lt;5yrs of age and not HIB immune with MODERATE – SEVERE Peri-Orbital Cellulitis</strong></td>
<td><strong>Cefotaxime IV</strong> 50mg/kg/dose every 6 hours (Max 2g/dose) for 48 hours, then seek ID review</td>
<td>Immediate - type hypersensitivity: Seek ID advice</td>
</tr>
<tr>
<td><strong>ORBITAL</strong> Cellulitis (all patient groups)</td>
<td><strong>Add Lincomycin IV</strong> 15mg/kg/dose every 8 hours (Max 1.2 g/dose)</td>
<td></td>
</tr>
</tbody>
</table>
ALERT:
If ORBITAL Cellulitis, and signs of optic nerve compression:

- Inability to spontaneously open or close eyelids,
- Proptosis,
- External ophthalmoplegia,
- Decreased visual acuity/ red perception,
- RAPD,
- Increased intraocular pressure,

= Orbital Compartment Syndrome.

Requires URGENT surgical decompression (lateral canthotomy)\(^2,4,9,11,18\).
Request immediate Ophthalmology team advice +/- attendance.

Disposition

All children with a diagnosis of Peri-Orbital or Orbital Cellulitis, other than MILD disease, *must be admitted* to the hospital\(^3,4,19\). Most cases can be managed at a regional hospital, with local ENT (+/- Ophthalmology) consultation. If no local ENT service available, should be discussed with regional referral centre.

- **MILD Peri-Orbital** Cellulitis
  - Consider admission to Short Stay Unit (SSU), or under a General Paediatric team, for period of observation and clinical review (if any doubt about severity of infection).
  - Give first dose of oral antibiotic in the Emergency Department (ED)/ SSU, and write prescription for ongoing medication. Must complete *minimum of 7 days of oral antibiotic therapy*\(^1,6,8,9,14,16,23\).
  - Ensure sufficient volume of liquid antibiotics prescribed (calculate exact volume needed, may require prescription for multiple bottles), and parents aware of need to complete full course.
  - Must have thorough history and examination documented, and included on discharge paperwork\(^4\).
  - Advise parents of signs of deterioration, and reasons for urgent return to ED (including being unable to administer medication regularly).
  - All children must have review by a doctor within 24 hours of discharge. If the child is not able to attend own General Practitioner/ Local Medical Officer, then arrange for review in the ED\(^2,4,9,11\). If concerns about reliable follow-up, then admit to SSU/ ward overnight.
  - Discuss with ENT (and Ophthalmology) teams only if re-presentation to the ED despite antibiotic therapy.
**- MODERATE Peri-Orbital Cellulitis**

- Refer to a General Paediatric team for admission.
- Request inpatient ENT consultation (within 12 hours of admission), or discuss with regional referral centre if no inpatient ENT team available.
- Intravenous antibiotic therapy to be continued for minimum of 48 hours. 
- All children must have a minimum of twice daily medical review, with examination of optic nerve function (pupil reaction, visual acuity, colour vision), and eye movements. Any signs of deterioration require urgent review by ENT (and Ophthalmology) teams.
  
  *(Note: Should only be managed in outlying ward if medical staff able to attend easily for regular clinical review.)*
- If improving clinically (erythema and swelling decreased, fever and WCC improved), then discharge home on oral antibiotics (as per MILD) after completion of 48 hours of intravenous therapy.

**- SEVERE Peri-Orbital Cellulitis**

- Urgent referral to ENT +/- Ophthalmology teams.
- Consider need for medical imaging if clinical diagnosis (Peri-Orbital vs. Orbital) not clear.
- Admission under ENT team (may require inter-hospital transfer if no local service), with ongoing Ophthalmology team review.
- Intravenous antibiotic therapy to be continued for minimum of 48 hours, although may require longer intravenous course (liaise with ID team).
- All children must have a minimum of twice daily review with examination of optic nerve function (pupil reaction, visual acuity, colour vision), and eye movements.
- If improving clinically (erythema and swelling decreased, fever and WCC improved), then consider discharging home on oral antibiotics (as per MILD) to complete a total of 14 days of antibiotic therapy.

**- ORBITAL Cellulitis**

- Emergent referral to ENT +/- Ophthalmology teams.
- Organise medical imaging if high-risk features present, or on advice from ENT team.
- Admission under ENT team (may require inter-hospital transfer if no local service), with ongoing Ophthalmology team review.
- Inpatient treatment with a minimum of 72 hours of intravenous antibiotics, and eventual discharge with oral antibiotics to complete total of 14 days of antibiotic therapy. Seek ID review of antibiotic therapy after 48 hours.
- Steroid medication, nasal sprays and surgical management will be at the discretion of the treating ENT team, and dependant on the clinical situation.
Consultation

Guideline prepared by:

- Clinical Fellow, LCCH Emergency Department
- Registered Nurse, LCCH Emergency Department

Key stakeholders who reviewed this version:

- Infectious Diseases Consultant, CHQ
- Infectious Diseases Consultant, CHQ
- Antimicrobial Stewardship Pharmacist
- ENT Director, CHQ
- Ophthalmology Director, CHQ
- Ophthalmology Clinical Fellow, CHQ
- Quality and Safety Pharmacist, LCCH

Definition of terms

<table>
<thead>
<tr>
<th>Term</th>
<th>Definition</th>
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<tbody>
<tr>
<td>Dacrocystitis</td>
<td>Infection of the lacrimal sac, secondary to obstruction of the nasolacrimal duct.</td>
</tr>
<tr>
<td>Sty</td>
<td>Inflamed swelling on the edge of an eyelid, caused by infection of the gland at the base of an eyelash.</td>
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<tr>
<td>Chalazion</td>
<td>A cyst in the eyelid that is caused by inflammation of a blocked Meibomian gland, usually on the upper eyelid.</td>
</tr>
<tr>
<td>Chemosis</td>
<td>Swelling/oedema of the conjunctiva</td>
</tr>
<tr>
<td>Proptosis</td>
<td>Abnormal protrusion or displacement of an eye</td>
</tr>
<tr>
<td>Diploia</td>
<td>Double vision</td>
</tr>
<tr>
<td>RAPD</td>
<td>A Relative Afferent Pupillary Defect (Marcus Gunn Pupil) is observed during the swinging-flashlight test whereupon the patient's pupils constrict less (therefore appearing to dilate) when a bright light is swung from the unaffected eye to the affected eye.</td>
</tr>
<tr>
<td>Lateral Canthotomy</td>
<td>Emergent orbital decompression by incision of the lateral canthal tendon</td>
</tr>
</tbody>
</table>

References and suggested reading

5. Clinical Practice Guidelines: Orbital Cellulitis in Children. Boston Children’s Hospital, Boston, MA, USA. (Downloaded April 2016) http://www.childrenshospital.org/conditions-and-treatments/conditions/orbital-cellulitis/overview
16. Clinical Practice Guidelines: Peri orbital and Orbital Cellulitis. Royal Children’s Hospital, Melbourne, Australia. (Downloaded April 2016) www.rch.org.au/clinicalguide/guideline_index/Periorbital_and_ Orbital_Cellulitis/
18. Orbital Cellulitis – Guidelines on Best Management. Morris, S. Gold Coast University Hospital, Gold Coast Hospital and Health Service, Australia: Effective 25 May 2015 (Downloaded April 2016)


Guideline revision and approval history

<table>
<thead>
<tr>
<th>Version No.</th>
<th>Modified by</th>
<th>Amendments authorised by</th>
<th>Approved by</th>
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<tr>
<td>1.0</td>
<td>Director, Pediatric Emergency Medicine</td>
<td>Divisional Director, Critical Care</td>
<td>Executive Director Hospital Services</td>
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</tbody>
</table>

Keywords

- Peri-Orbital Cellulitis, Orbital Cellulitis, 00723

Accreditation references

- EQuIP and other criteria and standards: 12
- NSQHS Standard: 3, 4
Appendix 1:

Figure 2: Example of Snellen Chart (not to scale)

Figure 3: Example of Lea Chart (not to scale)

Figure 4: Example of Ishihara Colour Plate (not to scale)