

# Corneal abrasion

## Disclaimer

SEE ALSO: corneal foreign body, recurrent erosion syndrome, corneal laceration, penetrating eye injury (PEI), microbial keratitis

## DESCRIPTION

A corneal abrasion is an epithelial defect in the cornea.

## HOW TO ASSESS

### Red Flags:

- Exclude corneal laceration
- Exclude mechanism suggestive of penetrating eye injury, e.g. hammering, glass
- Corneal infection: corneal infiltrate/anterior chamber (AC) reaction (cells)

### On History:

- Symptoms: pain, foreign body (FB) sensation, redness, tearing, decreased vision (if central)
- Mechanism of injury:
  - High speed – increased risk of PEI (hammering, drilling)
  - Sharp object (fingernail, paper) – increased risk of recurrent erosion syndrome (RES)
  - Contaminated object (plant) – risk of infection
- Document details of protective eye wear

### On Examination:

Slit lamp examination may be facilitated by topical anaesthetic

- Conjunctival injection
- Evert upper lid to rule out subtarsal FB in all patients. Particularly important if vertical linear abrasions on cornea
- Position and size of abrasion: stain with fluorescein, measure/document size at slit lamp
- Assess depth of injury. If deeper injury, consult the AO or consultant as may be corneal laceration or penetrating injury. Perform Seidel test with fluorescein to rule out leak.
- Corneal infiltrate: if present indicates possible microbial keratitis
- AC reaction: may indicate presence of infection

## ACUTE MANAGEMENT:

- Antibiotic drops or ointment: chloramphenicol ointment QID (blurs vision for approximately 30 minutes) or eye drops QID for 3-5 days.
- Consider stat dose of cycloplegic (e.g. homatropine 2%) if patient has significant pain or photophobia.
- An eye pad is generally not used, as it can delay corneal healing. In the setting of large epithelial defect, a double eye pad may be used for 24 hours to reduce discomfort. Do not pad eye if abrasion caused by plant.
- Pain management: cool compresses, dim lights, rest, regular oral analgesia, e.g. paracetamol. Local anaesthetic drops should not be given to the patient to take home.

## FOLLOW UP:

Indications for follow up:

- Abrasion caused by plant/organic matter
- AC cells
- Symptoms not improving within 48 hours
- Children, if unable to evert upper lid and rule out subtarsal FB

If at increased risk of RES (plant/fingernail/sharp object injury), recommend paraffin based lubricant ointment (e.g. Refresh nighttime<sup>®</sup>, Polyvisc) nocte for 3 months

## DISCHARGE INSTRUCTIONS:

- Advise patient to return if increasing pain, photophobia or decreased vision
- Advise patient they will have FB sensation once local anaesthetic wears off
- Topical anaesthetic drops should never be prescribed on discharge
- Education regarding use of protective eye wear
- Advise patient regarding RES
- Contact lens wearer: discard previous lens and resume contact lens wear with a fresh contact lens once eye has been asymptomatic for 1 week.
- Give patient copy of [Corneal Abrasion Factsheet](#)

## REFERENCES:

Wills Eye Manual, 6<sup>th</sup> Edition 2012

Oxford Handbook of Ophthalmology, Oxford University Press 2006

Eye Emergency Manual, 2<sup>nd</sup> Edition 2009

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## Review Date:

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## Evidence Table

Author/s	Title	Source	Level of Evidence (I – VII)	Comments
Adam T. Gerstenblith Michael P. Rabinowitz	Wills Eye Manual, 6 <sup>th</sup> Edition 2012	-	VII	
Alastair Denniston Philip Murray	Oxford Handbook of Ophthalmology, Oxford University Press 2006	-	VII	
Dr Weng Sehu	Eye Emergency Manual, 2 <sup>nd</sup> Edition 2009	-	VII	

### The Hierarchy of Evidence

The Hierarchy of evidence is based on summaries from the National Health and Medical Research Council (2009), the Oxford Centre for Evidence-based Medicine Levels of Evidence (2011) and Melynck and Fineout-Overholt (2011).

- I** Evidence obtained from a systematic review of all relevant randomised control trials.
- II** Evidence obtained from at least one well designed randomised control trial.
- III** Evidence obtained from well-designed controlled trials without randomisation.
- IV** Evidence obtained from well designed cohort studies, case control studies, interrupted time series with a control group, historically controlled studies, interrupted time series without a control group or with case series.
- V** Evidence obtained from systematic reviews of descriptive and qualitative studies.
- VI** Evidence obtained from single descriptive and qualitative studies.
- VII** Expert opinion from clinician, authorities and/or reports of expert committees or based on physiology.

### **CPG Suite General Disclaimer**

These CPGs were written for use in the RVEEH speciality Emergency Department. They should be used under the guidance of an ENT or Ophthalmology registrar, and certain medications / procedures should only be undertaken by speciality registrars.

If you require clinical advice, please contact our admitting officer for assistance:

EYE: 03 9929 8033 ENT: 03 9929 8032