28 The eye

28.1 Introduction

There are c. 285 million visually impaired people in the world, of whom >39 million are blind. WHO estimates that 43% of the visually impaired are so because of a lack of spectacles while 30% have cataract. Major causes of blindness in the world are cataract (50%), corneal infections (particularly trachoma, 25%), glaucoma, vitamin A deficiency, and onchocerciasis.

Washing the face regularly and the use of azithromycin every 3 months in children under 12yrs reduces the incidence of trachoma. In the industrial world 0-2% are blind, but in low-income countries blindness is ten times more common. You can treat cataracts, arrest glaucoma and prevent trachoma and vitamin A deficiency. It is unfortunate therefore that ophthalmology scares most doctors, who imagine that treating the eye must be impossibly difficult. This is not true: you can diagnose 90% of eye diseases with a torch and an ophthalmoscope.

Nonetheless, the eye may be difficult to examine, particularly in a child, and if the eyelids are swollen or the eye painful, the patient may forcefully resist examination. Do not give up, because the signs of serious trouble may be subtle. Danger signs are: haziness of the cornea, inequality of pupil size (especially if associated with reduced visual acuity), or circumcorneal redness.

Ideally someone in your district should be able to perform cataract removals. Surgery inside the eye, however, is difficult, so try to learn these operations by apprenticeship from an expert; they are not described here.

ANATOMY

The eye lies within the orbit, a V shaped enclosure, designed to protect the eye from trauma. Its blood supply comes from the ophthalmic artery, a branch of the carotid artery. Six muscles are attached to the eye and wall of the orbit; the IIIrd, IVth and VIth cranial nerves pass through a fissure in the superior part of the orbit to supply the muscles and the VIth cranial nerve gives sensation to the eye.

A septum is attached to the rim of the orbit and the eyelids are attached to this. The eyelids protect the corneal surface are made from skin, muscle, tarsal plate (28-18) and conjunctiva.

The eye itself can be divided into:
(a) the external eye: lids, conjunctiva, sclera,
(b) the anterior segment: cornea, iris and lens,
(c) the posterior segment: vitreous & retina, optic nerve.

Fig. 28.1 BASIC EYE ANATOMY.
A, B, flow of aqueous from the ciliary body (15) into the posterior chamber (6), through the pupil into the anterior chamber (3), then through the trabecular meshwork (19) via Schlemm’s canal (18) into the scleral sinus (17). C, The visual axis passes through the middle of the pupil (made by the iris (4) and through the centre of the lens (6), and the vitreous (7) to the fovea (8) which is at the centre of the macula (9). The optic nerve (11) enters the globe at the ‘blind spot’ and makes the optic disc (10). It is contiguous with the light-sensitive retina (14), bounded by the choroid (13), and sclera (12), which joins the cornea (2) anteriorly at the limbus (20), where the conjunctiva makes a groove known as the fornix. The globe rotates within a fascial layer, Tenon’s capsule, which covers the sclera and forms the sheaths of the extraocular muscles, the outer layer joining the conjunctiva at the limbus.

HISTORY

Always take a careful history; it may be critically important. Focus on how vision has changed and whether there is discomfort in the eye.

Vision can be divided into distance, near, peripheral, stereo double or single, colour, or night vision. Ask which type has been most affected.

Has vision been lost rapidly (specifically ask about trauma, resulting in retinal detachment, haemorrhage, or optic nerve damage), or slowly (cataract 28.4, presbyopia 28.8, diabetic retinopathy 28.9)? Is central vision lost (macular disease from diabetes, macular degeneration, cataract) or peripheral vision (glaucoma, retinal detachment, inherited eye disease)? Remember ‘double vision’ may actually be blurred vision. Ask about ‘floaters’ and ‘flashes’ in the vision.

Ask about ocular discomfort: conjunctival pain tends to feel like sand or hair in the eye, while very high pressure or inflammation of the eye can feel more like a deep ache or throbbing pain. Note any watering of the eye.

Optic nerve disease can present with pain on eye movements and loss of vision. Light sensitivity can be due to inflammation of the eye, or sometimes cataract and post trauma or surgery problems.

Ask about a family history of eye disease such as glaucoma, cataract, or night blindness.
THE STANDARD EXAMINATION OF AN EYE. The standard examination of an eye is time-consuming to do well, so train a nurse or medical assistant to test the visual acuity and examine the eyes. Your consulting room must be at least 6m long and you should be able to darken it. You must have a good pen light. Most examinations can be done while a patient sits in front of you.

ALWAYS test the visual acuity. Explain that you want to test the eyes. Begin by testing them separately (with distance glasses if worn); test them again on each subsequent visit; and record your results, so that you will know if vision is deteriorating or not.

If the patient can read, test each eye separately either with Snellen's or LogMAR charts. Stand him 6m from the well-lit chart (28-2), and close the left eye with a piece of paper or your left hand. Ask him to start at the top until he cannot read any more. If he is a young child or cannot recognize letters, use the tumbling E chart. It may be helpful to get him to point with fingers in the direction of the letters on the chart. Values are written with the top figure as the distance in metres to the test chart, the bottom one the distance at which a person with normal vision can read that line. The standard chart is calibrated: 6/6, 6/9, 6/12, 6/18, 6/60, and 3/60; these represent steadily deteriorating vision measurements. A value of 6/12 is normally required for driving a car. Counting fingers (CF) at 3m is equivalent to 3/60. If CF<1m, try hand movements (HM), and then test for the perception of light (PL). Get children to point at pictures of objects.

Visual acuity can be usefully divided into four groups:

1. good vision 6/6-6/18,
2. poor vision 6/24-6/60,
3. partially blind CF5m to PL,
4. totally blind to NPL.

The LogMAR charts have 5 standard shape letters in each row, varying logarithmically in size and spacing. The score is based on the number of smallest letters read.

If you shine a torch into each of the 4 quadrants of the visual field, can he tell you where it is coming from?

Blindness is ‘a loss of vision which results in the patient being unable to continue with normal life, and to walk unassisted’. It is usually equivalent to binocular visual of <3/60, which is the same as CF<3m. Before you decide that there is complete blindness, test with a very strong light. If an eye cannot see any light, and its pupil does not react to light, it is sure to be beyond help, so there is no point advising otherwise. If the vision is normal and remains normal and the eye is not inflamed, pathology is unlikely.

Here is some basic eye equipment: it does not include equipment for operating inside the eye. Many of the instruments are very fine: look after them with the greatest care:

CHARTS, visual acuity, (a) Snellen and (b) illiterate E charts, both for use at 6m. These are essential, and can usually be produced locally. They have patterns of Es of different sizes in different positions, and can be used by patients who cannot read.

TEST TYPE, reading pattern. Use this for examining older patients with presbyopia (28.8) who need glasses. If necessary, you can also use a book or newspaper.

TORCH, for focal illumination, local pattern, preferably pen type, with ‘lens bulb’. A locally available torch is adequate: it can be easily replaced, as can its bulb and batteries.

LOUPE (±2-8 magnifying spectacles), binocular, surgical, headband type. Some simple form of magnification is useful for examining the front of the eye, for removing superficial foreign bodies, and for other kinds of fine work, such as suturing nerves.

TONOmeter, Schiötz, (28-3). You must be able to measure the intraocular pressure (IOP) if you are going to diagnose glaucoma. Digital measurement is simple but unreliable. This instrument is not much seen nowadays, but is still very useful.

OPHTHALMOSCOPE, simple pattern, Keeler type, battery handle. An ophthalmoscope is very useful, but you can do much good eye work without one.

SLIT LAMP MICROSCOPE, on stand, simple pattern. You will find a slit lamp useful, although you can diagnose uveitis without one. If you need to do a lot of eye work, this is very useful: spend some time, though, with an experienced operator before you purchase one of these delicate instruments.

SPECULUM, ophthalmic, solid blades, hinged with screw adjustment.

SCISSORS, ophthalmic lid, blunt points. If necessary, you can use any fine scissors.

FORCEPS (clamp), tarsal cyst (chalazion, 28.12), 8mm ring, Lambert pattern. This has two blades, one with a ring and the other with a plate. Use it to hold an eyelid while you incise a tarsal cyst.

CURETTE, chalazion (tarsal cyst). CAUTERY, simple type, bulb pattern. Heat this on a spirit lamp, or get a battery-operated type.

CLAMP, eyelid, entropion, Desmarre’s or Snellen’s, (a) medium & (b) large. Use this to hold the eyelid when you operate for entropion.

SCISSORS, ophthalmic, spring pattern, Westcott’s or Castroviejo’s. These are very delicate instruments which need treating with special care.

BLADE, crescent type

FORCEPS, fine, toothed, St Martin’s.

RETraction, eye, Desmarre’s. Use this for examining children.

NEEDLE HOLDER, ophthalmic, curved with lock, Castroviejo pattern.

LENS LOOP for cataract extraction.

INTRAOCULAR LENS: standard PMMA 21 dioptre are sufficient.

CHEAP SPECTACLES can be made from malleable wire and polycarbonate instead of glass for the lens (www.omedolarglasses.org). Otherwise there are self-adjustable fluid-filled glasses available (28.8).

GLASSES, simple frames, second-hand if necessary, spherical lenses +1 to +3.50 - the most commonly needed glasses are ±2 and ±2.50. Collect unwanted glasses and allow patients to try for themselves: this way you can deal simply and effectively with the reading difficulties of many.

Do not operate with the large instruments of a basic set. Use special fine instruments listed above. For operations on the globe an operating loupe and a bright focal beam are almost essential, using preferably a 12V spotlight or LED source.
THE PIN-HOLE TEST is a useful way of screening for refractive errors. If there is poor vision, place a card with several 1mm holes (punched 5mm apart) in front of the eye. For repeated use, attach the card to the insides of each lens of a pair of spectacles. Cover each eye in turn for testing. If there is an uncorrected refractive error, the vision will be significantly improved to 6/9 or 6/12.

CHECK THE VISUAL FIELDS, in all 4 quadrants, while sitting face-to-face with the patient, covering his right eye and your left eye and vice versa, comparing them.

EXAMINING THE OUTER EYE

Start by looking at the whole face. Note any asymmetry of the position of the eyes, or any protrusion (best judged from above and behind the patient). Palpate for a lump, and for high intra-ocular pressure.

Ask the patient to look down and keep looking down, but not to actively close the eyes. Put the tips of both your index fingers on one of the globes, so as to feel the sclera through the upper lid up to the upper border of the tarsal plate. Gently press with alternate finger tips towards the centre of the globe:

1. Gently fluctuate it from one finger to another.
2. Indent it with one finger and estimate the sense of fluctuation imparted to your stationary finger.
3. Estimate the indentation of the sclera as you relax your indenting finger. You can judge the eye to be ‘soft’ (<10mmHg), ‘normal’ (10-40mmHg), or ‘hard’ (>40mmHg). This is a crude test, and there must be a significant rise of pressure (>40mm Hg) before you can detect a raised intra-ocular pressure.

Test the movements of both the eyes together, and then test each eye separately, in all directions, including convergence. Note any squint (28.9).

If there is much pain, and the eyelids are in spasm, a drop of sterile LA will make examination tolerable. This will allow also you to insert a speculum to examine the eye more easily. Whilst the patient is looking, grasp the top lid with your finger, and slip the top blade of the speculum under it. Then ask the patient to look up, grasp the bottom lid, and slip the lower blade of the speculum under that. Adjust the arm of the speculum until the eye is exposed, and then tighten the locking nut.

CAUTION! Beware that the speculum does not press on the eye or damage the cornea.

Note abnormalities of the lids, lacrimal apparatus, puncta and canaliculi, the lacrimal glands and sacs, and also any epiphora (tears running down the cheeks).

Do the eyelids open and close normally? You can see this best on blinking. Check the lids for swellings, and check that the lashes are in their normal position. Look at the conjunctiva. Note particularly the distribution of any redness. If it is maximal near the corneoscleral junction, this occurs in iritis and corneal ulcer. If it is maximal at the periphery but often extending all over, it is likely to be conjunctivitis.

To examine the conjunctiva of the upper lid, evert it (28-8H-K). This is necessary to exclude a foreign body.

Look for pus or mucopus in the inferior fornix. This is present in all cases of bacterial conjunctivitis, and in some cases of viral conjunctivitis. Look also for signs of vitamin A deficiency: dry-looking conjunctivae, or Bitot’s spots (white patches on the temporal side of the conjunctiva, produced by keratin mixed with gas-forming bacteria).

Look at the cornea of each eye. Is it shining and clear, reflecting the light of a torch, or its surface irregular? (corneal ulcer). Is there clouding superiorly (trachoma, 28.13), or a general haziness? (oedema from trauma, keratitis, or glaucoma). A bright light and a loupe can detect keratic precipitates and adhesions (synechiae, 28-9B, of iritis).

If you suspect the surface is injured or ulcerated, instil 1 drop of 2% fluorescein, or dip the end of a fluorescein impregnated filter paper inside the lower lid for a few seconds. Mop out the excess fluorescein with tissue paper. Shine a light on the eye at an angle. Gaps in the corneal epithelium (ulcers, abrasions) stain bright yellow-green.

Look at the anterior chamber and note its depth. Is there any blood (hyphaema), or pus (hypopyon, 28-9C) at the bottom of the anterior chamber?

Look at the pupils. Do they look black and do they react to bright light? Are the pupils grey or white? (opacities in the lens, cataract). Note their size, shape, and if their outline is irregular (synechiae, due to iritis, 28.5). If a pupil constricts incompletely when light is shone into that eye, and then constricts further when it is shone into the good eye, and when the light is shone back into the abnormal eye, both pupils enlarge, this means there is optic nerve damage, commonly caused by glaucoma, but you should exclude a stroke or brain injury. This is known as a relative afferent pupillary defect.

FUNDOSCOPY (OPHTHALMOSCOPY) examines the fundus and media of the eye. You must, either, dilate the pupils with a mydriatic such as cyclopentolate 1%, or do your examination in a dark room. This is however ineffective where the vitreous or cornea is opaque, or very unevenly curved (extreme astigmatism).

(1) Get the patient to keep both eyes open and look straight ahead.

(2) Start with the ‘0’ lens in the ophthalmoscope (unless you have a refractive error and are not wearing glasses; if so select the appropriate correcting lens and remember this as your starting point.

(3) Use your right hand and your right eye for the patient’s right eye and your left hand and your left eye for the patient’s left eye.

(4) Hold the sight hole of the ophthalmoscope close to your eye, resting against your nose and orbit, and move it with you as if it was attached to your head. To find this position, look through the sight hole at some distant object.
(5) With your thumb on the patient’s forehead, gently raise the upper lid clear of the pupil.
(6) Start with the ophthalmoscope 20cm from the eye, and shine the light into the pupil; it should glow uniformly red (the red reflex). This indicates the absence of a cataract.
(7) Move closer and watch for any opacities in the media silhouetted against the red reflex. Corneal opacities appear to move in the opposite direction to the ophthalmoscope; vitreous and posterior lens opacities appear to move with the ophthalmoscope. If you see a shadow, use the + lenses (+5 to +12) to see it more clearly.
(8) Ask the patient to look straight ahead, and move as close as you can to the eye without touching the eyelashes or cornea.
(9) Find and look at the optic disc: it is 15º to the nasal side of the optical axis of the eye.
(10) Turn the lens wheel in the ophthalmoscope with your forefinger from +6 down to zero to get the best view of the disc. Examine:
(a) the vertical cup/disc ratio (a ratio of >0.7 suggests glaucoma; 28.6, 28-11),
(b) the disc margins; if these are blurred all round (360º) it suggests papilloedema,
(c) the blood vessels, look for nasal displacement of the central retinal vessels and for haemorrhages and exudates suggestive of diabetic retinopathy,
(d) the macula (28-1C), by asking the patient to look directly into the light source, for black and white pigmentation which may suggest choroiditis (28.5) involving the macula (maculopathy).

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SCHIÖTZ TONOMETRY

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Fig. 28-3 SCHIÖTZ TONOMETRY.
The scale is merely an example; use the scale which is supplied with your instrument. 3 weights (5.5, 7.5, 10G) are usually supplied with each instrument.

SCHIÖTZ TONOMETRY
You may well never see this old-fashioned, but simple, instrument. If you happen to find one, clean the instrument with a pipe cleaner and ether. You’ll find it quite useful to diagnose glaucoma. It is however, a delicate instrument, so keep it carefully!
Using the standard 5·5g weight and the metal footpad, make sure the instrument is calibrated to zero.

Explain what you are going to do, lay the patient flat and instil LA into the conjunctiva. Ask him to open both eyes, and look straight up at a target placed on the ceiling.
With the 5·5g weight in place, put the tonometer plunger gently on the centre of the cornea with the eye open, and read the scale. If in doubt, repeat the reading 3 times. Use the tables provided with the instrument to calculate the IOP from the scale reading.
The normal IOP is 7-25mmHg. In practice, using the 5·5g weight, a scale reading of ≤2 (>28mmHg) indicates a raised IOP. A reading of ≥3 (>25mmHg) is 'normal'. If the IOP is >40mmHg, the cornea is likely to become oedematous (the characteristic 'hazy cornea' of glaucoma), and you can see this with a torch. This is usually a late sign of glaucoma.

**BINOCULAR INDIRECT OPHTHALMOSCOPE**

This allows good examination of the anterior & posterior segments at much less cost than a slit lamp: It is also portable (28-5).
(www.mercoframes.net/product/binocular-indirect-ophtalmoscope)

**INDIRECT OPHTHALMOSCOPY**

A binocular indirect ophthalmoscope (28-4) provides stereoscopic, wide angle, high resolution views of the entire ocular fundus. It is not hindered, as is the standard ophthalmoscope by a hazy media or scleral or central opacification. With the addition of a +20dioptre condensing lens, by varying the illumination and viewing angle, you can readily look at both the anterior and posterior segments.

1.1. Examine layer by layer: lid margin → conjunctiva → cornea → anterior chamber → lens → vitreous.
Lid margin: plugged orifices, lice, erosions?
Conjunctiva: foreign body?
Cornea: foreign body embedded in the cornea? Ulcer? Note its size and shape after instilling fluorescein and using the blue light. On the back of the cornea look for keratic precipitates (KP, these are clumps of white cells), indicating uveitis.
Anterior chamber: look for cells and flare, pus and blood; estimate its depth.
Lens: diffuse opacity, discoloration? Posterior synechiae from the iris? Focal opacities?
Vitreous: Particles from a recent posterior uveitis, or bleeding?
BINOCULAR INDIRECT OPHTHALMOSCOPY

Fig. 28-5 PRACTICE OF INDIRECT OPHTHALMOSCOPY. Use a +20D magnifying lens held close to the patient as shown.

SLIT LAMP MICROSCOPY

Fig. 28-6 SLIT LAMP MICROSCOPY (A) shines a narrow pencil of light illuminating the eye from an angle while it is examined with a low-power microscope. B, layers of the cornea and lens demonstrated. Particles in the aqueous and vitreous reflect light, like dust particles illuminated by a sunbeam in a darkened room. C, you may be able to see keratic particles directly with bright light.


SLIT LAMP MICROSCOPY. Use this for accurate visualization of the anterior part of the eye and its contents (28-6).

Position the head by placing the forehead and chin on the rest. Vary the angle of the light as convenient.

EXAMINING A BABY'S EYES

Fig. 28-7 EXAMINING A BABY'S EYES. Sit him on the mother's lap and hold the head between your knees.

BASIC EYE MEDICATION.

Drugs for use on the eye differ from other preparations: for topical use they come either as ointments or drops. The former are for longer-lasting effect, the latter for immediate and, usually, short-lasting effect.

Some drugs are toxic to the eye through systemic use: these include chloroquine and ethambutol. Others are locally toxic, such as penicillin, or dangerous if used for the wrong condition, such as steroids if used when there is a herpetic corneal ulcer present.

N.B. Many antibiotic eye preparations also contain steroids: avoid these!

Certain drugs are specifically used to help examination: cyclopentolate 1%, or phenylephrine 10%, will dilate the pupil for some hours only, so use these when you want a temporary effect for example when using an ophthalmoscope. Useful LA agents are: lidocaine hydrochloride 4% or amethocaine hydrochloride 1%. Remember that an anaesthetized eye is in great danger if an unnoticed foreign body gets into it, or that an abrasion injury is not felt; so shield it (28-8B) after appropriate examination.

To diagnose corneal injury, fluorescein papers are better than fluorescein drops, because you can more readily keep them sterile.

To use eye drops, pull the lower eyelid down so that you can see the conjunctiva. Ask the patient to look up. Put drops or ointment into the outer third of the conjunctiva.

Close the eye for 2mins to allow the drug to enter the eye. Do not let the dropper touch the eye, and do not put the dropper down on a surface, as it may become contaminated. If possible, each patient should keep his own drops, because of the danger of cross-infection.
TO MAKE YOUR OWN CHLORAMPHENICOL EYE DROPS
dissolve two 250mg capsules in 100ml of water. Filter the solution into
sterile 10ml dropper bottles. Screw the caps on loosely, and sterilize
them in a hot water bath or autoclave at 100ºC for 30mins, without
letting water splash over the necks of the bottles. Refrigerate them; their
shelf-life is 2months at 2-8ºC. The shelf-life of commercial drops is only
4months, so this is a useful method.

Subconjunctival antibiotics are indicated if there is a
severe corneal infection or ulceration, especially with
hypopyon (28.9A,C).

N.B. If you use 0·2ml of 2% lidocaine, the injection will
be almost painless, but be careful mixing antibiotics and
LA in the same syringe to maintain sterility.

If it is in the right layer, you will see its point under the
conjunctiva. Then inject 0·5mL (max 1mL).

If the infection is getting worse, repeat the injection the
next day. A severely infected eye is likely to improve,
or be lost within 6hrs, so use a subconjunctival injection
usually only once. If you fear the development of an
endophthalmitis, use IV antibiotics.

28.2 Operating on the eye

Try to get special training, especially for cataracts. Learn how to do the more important procedures: entropion (28.13), tarsal cysts (chalazion: 28.12), tarsorrhaphy (28.10), and evisceration, enucleation, and perhaps exenteration of the eye (28.14).

PREPARATION.
Prepare the face from the hairline to the chin and from ear
to ear, using iodine 10% in a non-alcoholic lotion which
will not harm the eyes, if it spills on them accidentally.
Make a special drape with a slit from the middle of one
end to the centre. Place this under the chin, and up each
side of the face. Fold it over the head and keep it there
with a towel clip. Place another drape across the forehead
over the eyebrows, and clip this to the first one.
If the patient is intubated, place a third drape over the
nose and the mount, which connects the patient’s
dontracheal tube.

If you are using LA, do not cover the nose or mouth.

ANAESTHESIA. You can usually use LA, using a
retrobulbar block with a 15mm 27G needle; otherwise use
ketamine or GA for a perforation (if LA is complicated by
retrobulbar haemorrhage, it may aggravate loss of eye
contents).

POSITION the table so that you can sit comfortably with
your knees underneath it. If necessary, put the head at the
foot end, or rest it on a board, or sheet of wood, pushed
under the mattress, and projecting beyond the table.
Sit your assistant on your left for a right eye, and on your
right for a left eye. Keep your own eyes on the patient.
This means you need to have trained your assistant to
hand you the right instruments properly.

Use a speculum, or lid stay sutures, to hold an eye open
while you operate on it. Keep your own eyes on the patient.
These sutures serve simply to hold the lids away from the eye while you
operate on it. They avoid the risk of a speculum
which may press on the eye, and perhaps scratch the
cornea.

In the upper lid insert two 3/0 silk or monofilament
sutures, just above the lash line and down to the tarsal
plate. In the lower lid insert one suture just below the lash
line. Do not penetrate the conjunctiva of either lid.
Hold these sutures with haemostats.
BLEEDING. The cornea is avascular and cannot bleed. If the conjunctiva or sclera bleed, apply a pad and very gentle pressure. Or irrigate the wound with saline from a syringe and an irrigating needle. The blood will stream in the clear saline, so that you can see the exact point where it is coming from, and control it with cautery. Heat a squint hook or a small cautery in the flame of a spirit lamp, until it is hot, but not red hot. Touch the bleeding point with this, through the stream of saline. This will cool its tip enough to prevent burning, but will leave it hot enough to seal the bleeding vessel. Do not use diathermy.

PAD THE EYE if there is had a minor injury with no suspicion of perforation (28-8A). An eyepad, with gentle firm pressure, will reduce discomfort, and promote healing by preventing the lids moving over the injured area. Close the eye, put a pad of gauze over it; place 2 pieces of adhesive strapping diagonally across the pad, from the forehead to the cheek, to hold the pad in place. Change the pad daily, and look for signs of ulceration or infection.

CAUTION! The great danger of an eyepad is that it may rub against an anaesthetized eye, and cause an abrasion. A layer of vaseline gauze on the pad will help to avoid this. So shut the eye when you apply the pad.

SHIELD THE EYE:
(1) after any severe injury, especially if there is a perforation.
(2) after any eye operation. Shielding it (28-8B) allows it to open and close, without anything extraneous touching the cornea, and perhaps scratching it. A shield is the safest way to protect an anaesthetized eye, and is very helpful for a painful inflamed eye with photophobia. Cut an 8cm diameter circle from cardboard, or an old X-ray film. Cut a radius in this, fold it into a cone, and maintain the cone with a piece of strapping. Hold the cone in place with two pieces of adhesive strapping, or plastic tape from the forehead to the cheek.

CAUTION! Never occlude the eye of a child <7yrs for >7 days, because this may cause amblyopia (28.9).

28.3 The painful red eye

Acute red painful eyes are due to:
(1) conjunctivitis (much the most common cause at any age). In the newborn this is often due to gonococcus, in children between 6 months and 6 yrs secondary to measles, and in adults in endemic areas, chlamydia.
(2) a corneal ulcer.
(3) acute iritis.
(4) acute glaucoma.
(5) trauma.

The problem in a busy clinic is that conjunctivitis is so much more common that these other causes are easily missed. So your first task in managing red eyes is to make sure that these rarer causes are recognized. The history, the visual acuity, and the examination of the eye with a torch should enable you to distinguish between conjunctivitis and something more serious.

Conjunctivitis can be infectious, allergic, or chemical. Bacterial conjunctivitis is common (especially from *neisseria, listeria and corynebacterium*) in the developing world, and may be mild, or so severe that the conjunctiva extrudes pus, and the lids swell so much that the eyes remain closed. Bacterial conjunctivitis needs an antibiotic. Viral conjunctivitis usually resolves spontaneously without, if the cornea is not involved. Allergic conjunctivitis rarely needs steroid treatment. Besides infecting the conjunctiva, bacteria can infect the lids (blepharitis), or the cornea, where they can cause changes in the stroma (keratitis and sometimes a corneal abscess), which may result in corneal ulceration, through which infection may spread inside the eye as an endophthalmitis, which may end in blindness.

A corneal ulcer may be due to:
(1) Bacteria.
(2) Herpes simplex virus.
(3) Fungi.
(4) Other conditions such as leprosy, causing incomplete eyelid closure (lagophthalmos) and exposing the cornea to trauma. Demonstrate a corneal ulcer with fluorescein. Bacterial infection can follow even a minor injury which damages the epithelium, or it can be spontaneous. Bacteria enter the eye through the anterior chamber. If pus gathers there, you will see a fluid level (hypopyon: 28-9C) when the patient stands upright.

Endophthalmitis may be the result of:
(1) a corneal ulcer, especially bacterial.
(2) a perforating injury of the cornea or sclera, especially if a foreign body has been left in situ, or if a wound is neglected, or after recent eye surgery. Once bacteria have entered the eye, the chance of total blindness is high. If presenting early, when the infection is fairly localized, some useful vision may remain. If you cannot control the infection, an evisceration is necessary (28.14).

DIAGNOSIS.
If there is conjunctivitis, the discomfort is of a gritty nature caused by rubbing of the conjunctivae on the cornea; pain varies from mild to severe:
(1) Both the eyes are usually involved.
(2) The visual acuity is normally good.
(3) There usually is a purulent discharge.
(4) Red conjunctivae, especially in the fornices (28-6C).
(5) The cornea is clear and does not stain with fluorescein (unless the conjunctivitis has produced a corneal ulcer).
(6) The pupils are normal.
(7) The tension in the globe is normal.
DIFFERENTIAL DIAGNOSIS.
Distinguish particularly between the redness of conjunctivitis, which is typically bilateral and maximal at the periphery, but is often uniform everywhere (very common), with redness which is most marked at the corneoscleral junction (less common).

CAUTION! Look for mucus in the inferior fornix (28-6C): it is always present in bacterial conjunctivitis; hesitate to diagnose conjunctivitis if you do not find any.

Suggesting acute iritis (28.5): one (sometimes two) moderately painful red eye(s) with no discharge. Pain is often only mild. Reduction in visual acuity is usually mild. A clear cornea is surrounded by redness at the corneoscleral junction. A small constricted pupil which becomes irregular on dilation, due to posterior synechiae (adhesions) is typical. An inflammatory exudate in the anterior chamber is visible most easily with a slit lamp: the aqueous is not as clear as it should be. The beam from the lamp shows a flare, like a beam of light shining across a dusty room. You may also see little lumps of cells (keratic precipitates or KP) sticking to the back of the cornea, and posterior synechiae between the iris and the front of the lens. The inflammatory cells in the anterior chamber may form a sterile hypopyon. The intraocular pressure (IOP) may be increased due to secondary glaucoma (28.6).

Suggesting acute angle closure glaucoma (28.6): one (seldom 2) very painful red eye(s) with severe unilateral headache, and slight watering. There is severely impaired visual acuity, often down to hand movements or perception of light only, with haloes, and sometimes even blindness. Circumcorneal hyperaemia is mild in the early stages. A hazy cornea (due to raised IOP) without its normal shine is associated with a shallow anterior chamber; this is best seen by shining a torch from the side. A vertically oval dilated pupil which does not react to light is classical. IOP is raised (28.1).

Suggesting a corneal ulcer: one severely painful red eye with reduced visual acuity (if the ulcer is central), scleral redness most marked round where the ulcer is situated, photophobia, swollen eyelids, and watering. Look for a grey-white spot (the ulcer) on the cornea, which stains with fluorescein. If it is not obvious, look for a defect in the smooth surface of the cornea in the reflection from a focused light. If the infection is severe, pus cells sediment at the bottom of the anterior chamber, with a fluid level (hypopyon). The pupil is usually regular.

Suggesting a foreign body: The signs of an abrasion, and a foreign body, are similar to those of a corneal ulcer: unilateral pain, photophobia, a watery discharge, sometimes impaired vision, and hyperaemia, which is marked near the lesion. Ask if there is a history suggesting trauma, and do not forget that contact lenses are foreign bodies and easily become infected if not kept scrupulously clean. Check underneath the upper eyelid!

ACUTE INFECTIVE CONJUNCTIVITIS
TREATMENT. Clean the eyes with a cotton swab and saline. Instil chloramphenicol or ciprofloxacin ointment hourly in severe infections, and 3hrly if less severe. Continue for 2 days after symptoms have resolved. Allow the exudate to escape, clean the eyes with a clean cloth and water, add an ointment at night to prevent the eyelids sticking together, and do not put a pad on the eyes.

If the conjunctivitis is severe, use subconjunctival antibiotics. Watch carefully for a corneal ulcer, and if necessary examine the cornea repeatedly with fluorescein. If the conjunctivitis is very severe, and especially if there is a corneal ulcer, instil chloramphenicol eye drops every min for 1 hr, every hour for 1 day, and then 3hrly.

If the cornea is not clear and the visual acuity is poor, there is a corneal ulcer and the eyesight is in danger.

If a neonate has severe conjunctivitis after birth (ophthalmia neonatorum), this may be gonococcal or chlamydial. This is an acute emergency, which may cause blindness. Treat with chloramphenicol drops as above, and add either ceftriaxone or gentamicin IV, and oral erythromycin.

If you are treating a child between 6 months and 6 yrs, check for a combination of malnutrition, vitamin A deficiency, and recent measles. Look for:
(1) Night blindness (inability to see in dim light).
(2) Bitot's spots (white foamy spots on the lateral conjunctiva).
(3) Xerosis (dryness of the conjunctiva with inability to produce tears, or a dry hazy cornea).
(4) Keratomalacia (corneal ulceration, softening of the cornea). Treat with vitamin A 200,000IU by mouth immediately, again after 24hrs, and again after 1 wk.
Also, use a topical antibiotic such as ciprofloxacin. Improve the nutrition especially with plenty of dark green leafy vegetables.

CHRONIC LOW-GRADE CONJUNCTIVITIS
characterised by yellow-grey dots (follicles) under the upper eyelid, in someone from an endemic area, is almost certainly TRACHOMA caused by chlamydia trachomatis.

N.B. Different strains of this bacteria cause 3 distinct diseases: urethritis & PID; lymphogranuloma venereum; and trachoma.

Trachoma passes through 4 stages (28.13). During the acute stage, make sure the patient actually puts tetracycline eye ointment 1% into the eyes b.d for 6wks. Add a single dose of 1g azithromycin orally. Advise thorough washing of the face and hands several times daily, avoiding rubbing of the eyes. Explain that the disease is due to the entry of dirt, often from flies, but also from sharing face towels with an infected person.
ALLERGIC CONJUNCTIVITIS

Suspect this if large gelatinous vegetations have formed on the upper tarsal conjunctiva, and look like cobblestones, or on the bulb conjunctiva surrounding the limbus. It is common in children and young adults. Their eyes may or may not be itchy but typically there is extreme watering. Suppress the inflammation with antihistamine drops or a very weak steroid. Beware of steroid glaucoma (28.6, 28.12D), because steroids, once started, may be needed for many years. Inject triamcinolone 1ml (40mg) IM into the upper eyelid for severe symptoms.

CORNEAL ULCER

This is an emergency needing admission. Start aggressive treatment with antibiotics urgently. A shield (28.2, 28-8B) or sunglasses will make life more comfortable. Do not use an eyepad or patch.

If the ulcer is severe, and particularly if there is a hypopyon, inject subconjunctival (28.1) gentamicin 20mg, or chloramphenicol 100mg and apply hourly chloramphenicol 1%, or ciprofloxacin 0-3% eye ointment.

If the ulcer is not so severe, and there is no hypopyon, treat as conjunctivitis.

Also, with any corneal ulcer, provided it has not already perforated, use atropine eye ointment bd or tid to keep the pupil dilated. This will prevent adhesions forming between the iris and the lens (posterior synechiae, 28-9A). Advise warm soaks: their use is effective for soothing a painful eye. Wrap a cloth round a spoon, dip this into very hot water, and let it cool till you can hold it as close to the eye as is bearable. Soaks are also useful for a stye (infected eyelash follicle, hordeolum).

Use vitamin A supplements if there is any suspicion that it may be deficient.

COMPLICATIONS of corneal ulceration include:
(1) Diffuse scarring of the cornea (28.4).
(2) A dense white scar (leucoma: 28.4).
(3) Perforation of the cornea, with adherence of the iris, and perhaps staphyloma (an opaque protrusion of the cornea, not related to staphylococci).
(4) Endophthalmitis.

If there is pain and watering without a history of a foreign body, look for a DENDRITIC ULCER (28-9E). Stain the cornea with fluorescein and look for a branching irregular pattern. This is due to infection by herpes simplex. Dendritic ulcers occur especially after fevers, particularly measles, malaria, and meningitis. If possible use an antiviral agent: idoxuridine ointment (x5 daily), trifluorothymidine drops (hourly), or aciclovir ointment (x5 daily).

If the lesion is severe, combine this with mechanical removal of the epithelium containing the virus. Apply a topical anaesthetic, and stain the cornea with fluorescein. Using a loupe, a good light, and a ball of cotton wool on the end of an applicator, gently scrub the surface of the cornea in the region of the ulcer to remove its epithelium. A chronic stromal keratitis with corneal scarring and blindness can complicate herpetic eye disease.

CAUTION! Never apply steroids, because these may spread the infection to the stroma of the cornea, and make the condition worse.

ENDOPHTHALMITIS (PANOPHTHALMITIS)
The anterior chamber is full of pus.

If the endophthalmitis is early, with some hope of vision, try to control infection and minimize pain. Use subconjunctival chloramphenicol and IV gentamicin for 7 days. The infection may settle.

If the endophthalmitis is due to a foreign body in the eye, remove it. It is usually superficial, so that it is possible to remove it through the wound by which it entered, which is usually in the cornea, even if this has to be enlarged. Remove any prolapsing iris, and leave the cornea unsutured. Use subconjunctival chloramphenicol and IV gentamicin.

If presentation is late, with no hope of vision and an anterior chamber full of pus, and the corneal ulcer has weakened, softened, and distorted the globe (phthisis bulbii, 28.4), especially with no improvement after 48 hrs antibiotic treatment, eviscerate the eye (28.14).

Be sure that the patient understands the necessity of removing the eye because of the mortal danger of orbital cellulitis and meningitis.

DIFFICULTIES WITH RED PAINFUL EYES

If a chemical has got into the eye, the conjunctiva is intensely red (more so than in infective conjunctivitis), the cornea may be opaque (from keratitis or an ulcer), and the vision impaired. Unlike infective conjunctivitis, mucopus is absent. Traditional medicine may have been inserted for a painful eye, which has made it worse. If the chemical is still present, wash it out with much water, making sure it does not spill over the other eye. Remember that if it is acid or alkali, a 1l water to alter the pH by a level of 3. Use an analgesic, and shield the eye. Instil an antibiotic ointment; its vaseline base will be soothing, and the antibiotic may prevent secondary infection.

If there is an acutely inflamed and oedematous lid or face, with a black slough, and surrounding thick oedema, this may be ANTHRAX, especially if there has been contact with animal carcasses or hides.

The eyelid may be completely destroyed, but the eye is normal. Use high doses of IV penicillin and sulphonamides. Anthrax responds rapidly to penicillin. Later, if necessary, toilet the slough and graft the raw area. If you leave raw lids ungrafted, severe scarring and a scar-induced ectropion (lid eversion) will follow.
THE IRIS AND THE CORNEA

A. Vertical section of the eye showing keratic precipitates floating in the aqueous (1), posterior synechiae (2), and a hypopyon (3). B. Iris bombé: the iris is adherent to the lens all round and is bulging forwards. C. Acute bacterial corneal ulcer with a hypopyon. D. Acute iridocyclitis. The pupil is small and irregular, because posterior synechiae have formed. E. Dendritic ulcer of the cornea, the result of herpes simplex infection.

Fig. 28-9 THE IRIS AND THE CORNEA.
A, vertical section of the eye showing keratic precipitates floating in the aqueous (1), posterior synechiae (2), and a hypopyon (3). B, iris bombé: the iris is adherent to the lens all round and is bulging forwards. C, acute bacterial corneal ulcer with a hypopyon. D, acute iridocyclitis. The pupil is small and irregular, because posterior synechiae have formed. E, dendritic ulcer of the cornea, the result of herpes simplex infection. After Parr J. Introduction to Ophthalmology, OUP 2nd ed 1982 with kind permission.

28.4 Loss of vision in a white eye

This is one of the common presentations of eye disease. Loss of vision in a white eye can be slow or fast. If there is slow loss of vision over months or years, there may be:
1. A corneal scar.
2. Cataracts.
4. A refractive error.
5. Disease of the retina due to:
   (a) Senile macular degeneration.
   (b) Retinitis pigmentosa (congenital photoreceptor deficiency)
   (c) Chloroquine maculopathy.
   (d) Old macular scars.
6. Optic atrophy.

If there is sudden loss of vision over minutes or days, the cause is usually inflammatory or vascular. If the complaint is simply that reading is difficult, especially in poor light, this is usually presbyopia (28.8).

Corneal scars cause 70% of blindness in children and 25% in adults in the developing world. They can be:
1. Diffuse.
2. A circumscribed white patch (leucoma).
3. A staphyloma, which is a bulging of the cornea forwards between the lids, due to its thinning, caused by previous ulceration (not staphylococci).
4. Phthisis bulbi, which is disorganization of the entire eye, leaving it small and shrunken. Bilateral scarring follows neonatal conjunctivitis (ophthalmia neonatorum), vitamin A deficiency, traditional eye medicine & trachoma (28.3, 13). Unilateral scars are more likely to be caused by corneal ulceration due to bacteria, herpes simplex, fungi or trauma.

Cataracts cause about half the blindness in the world, where the incidence is c.1:200. A large majority (85%) of cataracts occur in the elderly, and the rest are either congenital or familial, or due to trauma, iritis, or diabetes.

Vitamin D deficiency causes lamellar (flaky) cataracts in infants. Cataract presents with gradual loss of vision, in one or both the eyes. The corneas are clear, and there is an opacity in the pupil(s). A cataract can be immature (making the pupil grey), or mature, or hypermature (making it white). Sometimes a cataract swells, pushes the iris forwards, occludes the angle of the eye, and causes secondary glaucoma.

Removing cataracts is a standardized and repetitive task; it is also a skilled one but is rarely urgent. To learn this it is best to apprentice yourself to an expert for several months, and try to remove at least 50 under supervision. Or, better, send a motivated assistant to learn this skill. Cataracts can often be removed on a mass scale in special 'eye camps'.

In good hands the chance of success is >90%. If you operate on a patient for the right indications, even moderate success in one eye only will provide much sought-for independence.

Aim to insert an intra-ocular lens (IOL), which provides much better vision. The IOL can now be obtained for a reasonable price and is made in Eritrea & Nepal for example, so this is no longer an impossible dream in the developing world. It should be standard, as refractive errors are better corrected and waiting for maturity of the cataract is no longer necessary.

Manual small incision cataract surgery (MSICS) has become the preferred extracapsular method for removal of cataracts in low-income settings. It does not require sutures, can be done inexpensively, and produces high quality results. High cost cataract removal alternatives are not necessary. An operating microscope is especially valuable for reducing the incidence of complications; the less experience the surgeon has, the more important the microscope quality becomes. If you are more experienced you can use loupes, but if you are less experienced, you may cause problems which you may not even see. A cheaper special MSICS microscope is currently being developed.
Bad outcomes are related to:
(1) poor case selection, i.e. operating on patients who actually have a corneal scar or glaucoma,
(2) complications such as vitreous loss or infection,
(3) uncorrected refractive error,
(4) postoperative posterior capsule opacification.

COMMON CAUSES OF GRADUAL LOSS OF VISION IN A WHITE EYE

CORNEAL SCARS

If the cause of the scar is still present, and it is getting worse, remove the cause. This may include scratching of the cornea by the inwardly turned eyelashes of trachoma (trichiasis, 28.13). Vitamin A deficiency causes an acute ulcer in young children, and does not cause progressive scarring.

If there is still adequate vision in the other eye and disability is not severe, no treatment is indicated.

If there is no light visible at all, explain that nothing can be done.

If there is blindness, and a central leucoma which obscures the pupil, with an area of clear peripheral cornea, a peripheral iridectomy is necessary. This will provide an artificial pupil peripherally, behind the area of clear cornea, and should give enough vision for independent mobility. It is contraindicated if there is already enough vision for mobility, or if the peripheral cornea is opaque.

If there is blindness due to diffuse corneal scarring which has not made the eye perforate, a corneal graft is the only solution.

If the eye is blind and painful, consider evisceration or enucleation (28.14).

CATARACTS

(1) Measure the visual acuity accurately in both eyes. The pupils should react briskly to light. If they do not, suspect that there is also some other condition, such as optic nerve disease.
(2) Measure the IOP to make sure that the loss of vision is not due to glaucoma (28.6).
(3) Dilate the pupil and examine the red reflex with an ophthalmoscope to assess how dense the cataract is, especially if it is immature. If you can easily see the optic discs, the cataract may not yet be dense enough to be worth extracting.

CATARACT EXTRACTION (GRADE 2.5)

INDICATIONS.
(1) To improve sight.
(2) To treat complications, especially secondary glaucoma.

If there are bilateral cataracts, operate when the acuity in both eyes has fallen to worse than 6/60 (CF at 6m).

If there is a unilateral cataract, surgery is only indicated to treat or prevent secondary glaucoma, or uveitis. It will not improve sight significantly.

If there is already loss of sight in the other eye for any reason, and there is now a cataract in the remaining eye (cataract in an only eye), delay surgery until there is difficulty getting around independently and near blindness (CF <3m), because any complication will cause total blindness.

If the one cataract has already been successfully removed, you can schedule the second cataract at any time. But, this case now will be a lower priority.

If the cataract extraction is not possible, atropine ointment weekly, or minus (concave) glasses may improve eyesight.

CONTRAINDICATIONS.
(1) Unilateral cataracts with adequate sight in the other eye.
(2) Bilateral small immature cataracts with acuity above 6/60 in both eyes together; review the progress in 3-6months.
(3) Active uveitis: do not perform a cataract extraction at the same time as an iridectomy

METHOD

The principle is to make a self-sealing tunnel to extract the cataract and insert the new lens. It may be combined with a trabeculectomy (28.6). Dilate the pupil with cyclopentolate 1%.
Fig. 28-10 MANUAL SMALL INCISION CATARACT SURGERY
D, make the scleral tunnel trapezoid-shaped with a crescent blade. E, enlarge the tunnel. F, paracentesis: make a side port and inject dye to stain the lens capsule and air to protect the underside of the cornea.

G, capsulotomy: pierce the anterior capsule in a complete ring c.7-5mm in diameter with a keratome.
H, capsulorhexis: tear off the anterior capsule with a special hook or one made from a 27gauge needle. I, irrigate and dislodge the lens capsule. Avoid touching the inner surface of the cornea! J, introduce the lens loop within the capsular bag under the lens nucleus, and slowly ease both loop and lens out, whilst irrigating and pressing on the posterior lip of the tunnel. K, lens removed. Do not irrigate >30sec. Do not aspirate the posterior capsular surface!
L, instil 0·3ml viscoelastic (a cohesive substance usually of sodium hyaluronate 1·4%) and air to re-create the anterior chamber and seal the incision. M, introduce the intraocular lens into the capsular bag, holding onto the trailing haptic, which should curve to the right. Make sure the lens is the correct way up! Do not hold onto the lens itself!

Insert the eyelid retractor, or place stay sutures on the eyelids. Grasp the superior rectus at its insertion, about 7·5mm behind the limbus (28-1C) and rotate the eye inferiorly; insert a 4/0 or 5/0 stay suture through the conjunctiva and beneath the muscle (28-10A). Raise a flap by picking up the conjunctiva at the superior limbus (junction of sclera and conjunctiva) and buttonhole the conjunctiva with fine scissors. Extend this in the sub-Tenon’s space and lift the conjunctiva and Tenon’s capsule (28-1C) off the sclera for 1cm (28-10B). Ensure haemostasis with cautery and that the field is dry before you proceed to the next step.

Make a 6-7·5mm long curvilinear scleral partial thickness (0·3mm deep) shelved incision 3mm posterior to the limbus on the superior side of the eye (28-10C). Deepen this incision by advancing the crescent blade into the sclera and slowly cutting on either side, thus making further room for the blade. Judge the correct depth by making sure the crescent remains visible through the sclera (28-10D).

Keep the crescent flat on the globe during dissection, so that the tunnel depth remains uniform. Once you reach the limbus, extend the tunnel by forward and backward motion, cutting as you come out; this way, you create scleral pockets on either side of the tunnel which becomes trapezoid in shape with its inner margin 7·8-5mm, i.e. larger than the outer margin 6-7·5mm long, adapted to the size of the nucleus (28-10E).

Insert the retractor and push the conjunctiva back to expose the limbus. E, inject 0·2ml trypan blue dye through a sideport, using a 25gauge cannula and adding a little air which protects the underside of the cornea. Wait a full 30sec but no more, otherwise you will dye all the tissues blue. Then wash out the dye with balanced saline (0·9% saline made up to pH 7·3 with bicarbonate). Then deepen the chamber by injecting 0·3-0·5ml air (28-10F).

Now open the tunnel into the anterior chamber by advancing a keratome through the tunnel, tilting it downwards, and advancing into the anterior chamber (28-10G). Move the keratome medially and laterally the full length of the tunnel while keeping the tip of the blade in the anterior chamber. Insert a 27G needle with the tip bent slightly downwards like a hook.
Press the hook into the anterior capsular lens surface to create a circular 360° opening about 7.5-8.0mm cutting parallel to the limbus (28-10H). Irrigate this space to help free the nucleus (28-10I); rocking it side to side, or turning it round may help free it. Then introduce a lens loop into the tunnel, and pass this under the cataract in the capsular bag and slowly ease it and the nucleus out of the anterior chamber (28-10I), at the same time pressing gently on the posterior lip of the tunnel to help expel it. Avoid touching the inner corneal surface.

Continue irrigating as you manipulate the lens nucleus out, and once it is out, pick up the sclera edge and aspirate any cataract fragments, leaving a clear anterior chamber. Keep pressing gently on the posterior lip of the tunnel to allow debris to flow easily out. Do not irrigate for longer than 30sec. Do not aspirate the posterior capsule.

Inject 0.3ml viscoelastic (28-10L) and insert a 21dioptre polymethylmethacrylate (PMMA) intraocular lens into the capsular bag (28-10M). This is the standard size and is suitable for 80% or more of patients. Make sure the lens is the correct way (flat or concave surface) up! Hold the lens with long smooth (e.g. McPherson long-angled) forceps with the leading haptic (curved hook attached to the lens) sweeping to your left and the trailing haptic curving to the right. Advance the lens by holding the haptic with the forceps, but don’t hold the lens itself with the forceps.

Recreate the anterior chamber by injecting 0-3-0.5ml air through the sideport without applying any pressure on the tunnel (28-10N). No sutures are required since the properly formed tunnel acts as a one-way valve to prevent leaks. The conjunctival flap becomes covered by the eyelid, and needs no suture. Apply topical chloramphenicol or ciprofloxacin, and dexamethasone 0-1% 3hrly for 1wk and then 6hrly for 2wks, with or without LA.

N.B. Don’t forget to remove the stay suture in the superior rectus muscle!

POSTOPERATIVELY, watch for a leaking wound (with or without iris prolapse), infection, bleeding, and a raised IOP. Gently open the lids, and examine the eye with a torch.

If the patient is restless and expels the air bubble in the anterior chamber causing it to flatten, take him back to theatre and re-inject air through the wound. If this keeps leaking out, suture the tunnel with 10/0 nylon.

If the remaining lens matter is swollen and fluffy, keep the pupil dilated with atropine drops 1% bd.

If there is any iris prolapse, return to theatre, reduce the prolapsed, make sure the anterior chamber is filled with balanced saline, and suture the wound.

If the cornea is hazy with a striate pattern (striate keratitis): it will probably settle.
If there is blood in the anterior chamber (hyphaema), pad the eye and insist on bed rest.

If the anterior chamber is shallow and the pupil not round, the wound may be leaking. (You may prove this with a fluorescin test.) Return to theatre and wipe the wound with cellulose swabs, fill the anterior chamber with 0-3-0.5ml air, and close the wound properly.

If there is pus in the anterior chamber (hypopyon), there is infection (endophthalmitis). The eye is likely to be painful and the visual acuity very low. Use subconjunctival gentamicin or cefuroxime (28.1), and topical chloramphenicol or ciprofloxacin hrly. The eye may be lost in any case.

If the red reflex is absent after several months, there is some opacity in the media. This may be from re-growth of new fibres in the posterior capsule; perform a CAPSULOTOMY by holding the medial rectus tendon in forceps and pass a keratome through the cornea laterally backwards to cut the posterior capsule. Then keep the pupil dilated with 2 drops atropine 1% bd and add 2 drops chloramphenicol 0-5% 4hrly for 3days.

If there is much pain and the cornea is hazy, the IOP is probably raised (aphakic glaucoma) so measure it. The vitreous jelly may be blocking the pupil. Immediately dilate the pupil with cyclopentolate and phenylephrine drops, followed by atropine ointment for 6wks.

If visual acuity is not improved and there is no evidence of endophthalmitis, increase topical steroid to 2hrly and check for improvement in 1wk.

RARER CAUSES OF GRADUAL LOSS OF VISION IN A WHITE EYE
Examine the macula and the optic cup with particular care. For many of the following, there is no remedy.

If an old person has gradual loss of central vision, atrophy, and irregular pigment at the maculae, suspect senile macular degeneration.

If there are pale, white, flat optic discs (distinguish these from the pale cupped discs of glaucoma, 28-11C), and normal maculae, there is optic atrophy. Try to find the cause (there are many, including a space-occupying lesion around the optic chiasma).

If there is gradual loss of vision at any age, often starting with night blindness, a family history, and dark pigmentation which follows the retinal vessels and takes the form of ‘bone spicules’, suspect retinitis pigmentosa, a congenital disease of photoreceptor loss.
If there is gradual loss of central vision from excessive doses of chloroquine (>1.5g weekly for >1yr), or ethambutol, suspect maculopathy. The macula has a typical 'bull's eye' pattern with a dark centre and a paler surrounding ring. Stop the drugs.

If there are old macular scars (large white areas with black edges, often around the optic disc and the macula), they may be due to previous toxoplasmosis (treat with pyrimethamine 25mg od & cotrimoxazole 40mg/kg od for 3wks) or toxocariasis (treat with albendazole 5mg/kg bd for 5days)

SUDDEN LOSS OF VISION IN A WHITE EYE

Loss of vision can occur over minutes, hours or days, in one or both eyes, which are white.

If at any age there is steady loss of vision over 24hrs, in one eye or occasionally both eyes, suspect posterior choroiditis (28.5) due to toxoplasmosis or other causes. The important sign is inability to see the retinal vessels due to hazy vitreous caused by inflammatory cells. Treat as above.

If symptoms started with a flash of light followed by black objects floating in the field of vision, and then a curtain or cobweb across it, suspect retinal detachment. Part of the retina may look grey-green. Dilate the pupil and examine the fundus. You will see an abnormal red reflex in one part of the fundus, with elevation of part of the retina, and tortuosity of its vessels, which are difficult to focus on. Expert surgery may save eyesight.

If there is instantaneous loss of vision, suspect occlusion of the central retinal vein (a swollen disc with many haemorrhages all over the retina), or artery (a swollen disc, oedema of the retina, and often a cherry-red spot at the macula). Or, suspect a stroke (cerebrovascular accident).

Check the blood pressure. If it is not elevated, and visual loss is less than 6hrs old, start anticoagulants. There is otherwise no definitive treatment. If there is central retinal vein thrombosis, follow up to check for secondary glaucoma, which needs treatment.

If there is loss of central vision with an abnormal pupil response to light, suspect optic neuritis (any age, usually in the 3rd and 4th decades, and usually unilateral). The vitreous and optic disc are usually normal. This will usually improve over about 8wks. Bilateral optic neuritis following methyl alcohol or quinine is permanent. There is no specific treatment.

28.5 Anterior uveitis: iritis & iridocyclitis & posterior uveitis: choroiditis

Any part of the uveal tract can become inflamed: the iris (iritis), the ciliary body (cyclitis), or the choroid (choroiditis). More than one part may be involved at the same time (iridocyclitis).

Although iridocyclitis may be caused by bacteria invading the eye through a corneal ulcer (28.3), it and other forms of uveitis are more often due to a sterile inflammation, usually from an unknown cause.

Uveitis of several kinds is common. Iritis (more strictly iridocyclitis) has several consequences:

(1) The inflamed iris may stick to the lens by posterior synechiae (adhesions) or less often to the back of the cornea by anterior synechiae.

(2) If the entire margin of the pupil sticks to the lens, the iris balloons forwards (iris bombé: 28-9B), and causes secondary glaucoma (28.6).

(3) Abnormal proteins enter the aqueous, and cause an aqueous flare, which you can see with a slit lamp. You can also see leucocytes as tiny particles floating in the aqueous.

(4) These particles may stick to the back of the cornea as keratic precipitates (KP), and they may be numerous enough to gather at the bottom of the anterior chamber similarly to a hypopyon. Unlike the hypopyon that results from entry of bacteria through a corneal ulcer, the fluid in iridocyclitis is usually sterile. Untreated iridocyclitis eventually subsides spontaneously, typically in c.6wks, leaving the eye severely damaged. It may relapse, or it may be insidious and chronic, with few symptoms except progressive loss of vision.

Uveitis presents in 2 ways (or when in both ways together as panuveitis), as anterior uveitis (iritis) or posterior uveitis (choroiditis); the former presents as an 'acute red eye'; so being one of the important differential diagnoses of conjunctivitis (28.3), whilst the latter presents as progressive loss of vision in a white eye (28.4).

Iritis is usually a sterile reaction to one of the infections listed below. If onchocerciasis (28.7) is endemic, it will certainly be the most common cause. Usually, no cause is found, and iritis is presumed to be due to an autoimmune disease. Atropine will keep the pupils well dilated, and help break down synechiae. Steroid use is controversial: it probably hastens resolution, but do not use it if there is any sign of infection, especially a corneal ulcer.

Remember also that steroids:

(1) will make a red eye white, regardless of the cause, without necessarily curing it;
(2) will suppress the normal inflammatory response, without killing the causative agent;
(3) may raise intraocular pressure, and may rarely cause a secondary glaucoma that could produce blindness;
(4) may cause a cataract if used long-term, but this will not happen in the short time needed to treat acute iritis.
Uveitis may be unilateral, or bilateral, and presents in various ways.

**Acute anterior uveitis** (iritis, iridocyclitis: 28.3) presents as a red, painful eye, with photophobia, lacrimation; and often blurred vision. There is circumcorneal redness, and often general hyperaemia also. The pupil is constricted.

**Posterior uveitis** (28.4) mainly involves the choroid, and presents as fairly sudden loss of vision over 24-48hrs in a white and usually painless eye, due to damage to the retina and an exudate of cells and pigment into the vitreous. After dilatation, you can see these as a vitreous haze with an indistinct retina. At a later stage, when the haze has cleared, you may see foci of white depigmentation, surrounded by heaped up black pigment which results in impaired vision, especially if it involves the macula.

**Panuveitis** (quite common) is a combination of anterior and posterior uveitis, and causes loss of vision in a red, painful eye.

**CAUSES.** You will probably find no cause, but if any of these are present elsewhere, they may be responsible: syphilis, tuberculosis, leptospirosis, leprosy, herpes, toxoplasmosis, toxocariasis, onchocerciasis (28.7), HIV disease, trauma, or leakage of lens protein from a hypermature cataract.

**TREATMENT.**

Dilate the pupil with short-acting mydriatics (cyclopentolate and phenylephrine). When the pupil is dilated, maintain atropine ointment 1% tid, until the uveitis is no longer active, as shown by the absence of KP bodies and redness. This will prevent posterior synechiae, which would lead to the complication of secondary glaucoma and cataract, and so blindness. So, keep the pupil dilated until all the inflammation has subsided.

**If the disease is unilateral,** cover the eye with a shield (28.1) if it is severe, and a shade if it is not.

**If the disease is bilateral,** use eye shades. You can make these from exposed X-ray film.

**STEROIDS.**

Iritis will subside spontaneously, but topical steroids will hasten its resolution.

CAUTION! NEVER use steroids if:

1. there are signs of infection.
2. there is a corneal ulcer.
3. if there is anterior uveitis (iritis, cyclitis), instil hydrocortisone drops 1% into the conjunctival sac 3hrly.
4. if there is posterior uveitis, use oral prednisolone 20-30mg/day for 3-6wks. Do not continue beyond 6wks. Tail these off over 1wk at the end of the course.

If the IOP is raised, add oral acetazolamide 250mg qid, until the inflammation is under control. Double the dose if the IOP remains raised. If possible monitor the IOP weekly by tonometry (28.1).

**DIFFICULTIES WITH UVEITIS**

If posterior synechiae develop, a cataract may follow. The adhesions may occlude the pupil and cause pupil block glaucoma (iris bombé) with an increased IOP. Be sure to dilate the iris vigorously with atropine, so that it does not stick to the lens.

If there is secondary glaucoma, the pupil will not dilate. Try to arrange an iridectomy, but it is often too late even to recommend this.

**28.6 Glaucoma**

Glaucoma is a group of blinding diseases in which the intraocular pressure (IOP) is usually raised, causing damage to the optic nerve, and resulting in loss of vision.

There are 4 kinds:

1. **Primary (chronic) open-angle glaucoma (POAG).**
2. **Primary angle closure glaucoma (ACG).**
3. **Secondary glaucoma, as a complication of trauma, swollen cataract, iritis, etc.**
4. **Congenital glaucoma (buphthalmos).**

**Open-angle glaucoma** (POAG, chronic glaucoma) occurs in eyes in which the angle between the iris and the cornea is normal, and is probably due to a block in the drainage of intraocular fluid at the trabeculum (28.1). POAG causes most cases of glaucomatous blindness in Africans. Up to 1% of those >40yrs may be affected.

POAG is bilateral, but is often asymmetrical; it is insidious and progressive, and causes no symptoms until much eyesight is already lost. Glaucoma cannot be prevented, and even early treatment cannot restore lost vision. The best that can be done is to recognize it early, and to prevent vision getting worse.

For this to be possible, all health workers must be aware of the possibility of glaucoma in any patient who complains of loss of vision. The key to early diagnosis is to pick up early changes in the optic discs and a raised IOP, both of which can be recognized by eye assistants. The aim of medical and surgical treatment is to lower the IOP to a level which will stop further damage to the optic nerve, and therefore preserve vision at its present level.

Trabeculectomy is a relatively simple operation, with a reasonable chance of preserving what vision still exists; however, you may need to repeat the operation within 2yrs because recurrence is common. Learn it from an expert at the same time that you learn cataract extraction.
The symptoms of POAG are non-specific. There is slow loss of vision in one or both the eyes over months or years (28.4). Sometimes, there is marked loss of vision in one eye, while the other eye is normal, or nearly so. Occasionally, there is pain and headache, but this is late. Glaucoma is often familial.

**Angle closure glaucoma** (ACG, acute glaucoma) usually occurs >55yrs, in women more often than men, with an abnormally narrow angle between the iris and the cornea. If this angle should happen to close a little more than usual, it causes an abrupt rise in the IOP with resulting unilateral episodic attacks of pain, misty vision, and rainbow-coloured haloes round lights. Between attacks the eye is normal. Sooner or later, an episode of raised IOP does not resolve, causing classical acute congestive glaucoma (28.3). Acute glaucoma is relatively uncommon, and is rare in Africa. Its incidence is highest in Inuits and Mongolian peoples, in Burma, and in South East Asia.

The dangers of atropine in glaucoma result from its effect in dilating the pupil:
1. The iris is kept away from the lens, and prevents adhesions (synechiae) forming between them, which is valuable in iritis.
2. The iris is crowded into the angle of the anterior chamber, where it impedes the drainage of aqueous. This is never a desirable effect, but it does not matter in a normal eye or in iritis; it can however turn an eye blind if drainage is already impaired by glaucoma!

So, use atropine in iritis, but not in glaucoma!

**IF AN OLDER PERSON COMPLAINS OF POOR VISION, CHECK FOR GLAUCOMA.**

**PRIMARY OPEN-ANGLE GLAUCOMA,**
(POAG, CHRONIC GLAUCOMA)

DIAGNOSIS. Measure the visual acuity of any patient who presents with loss of vision. A hazy cornea or a pupil which does not respond normally to light should make you suspect glaucoma. If the IOP is >28, or the cup/disc ratio is >½, there may be glaucoma (28-12G,H,I). The end stage of glaucoma is a patient with a blind, or nearly blind eye, with a large pupil that does not react to light. Aim to diagnose it long before this with the following three tests.

Loss of visual field is an early sign, but is not easy to test with simple equipment.

CAUTION!
1. The level of IOP is useful in confirming glaucoma, but is not absolute, but is not always related to optic nerve loss.
2. Glaucoma can occur with a normal IOP (30% of patients with glaucoma have an IOP <22mmHg).
3. The IOP fluctuates, so if you are in doubt, repeat the measurements over a few days.
4. **LOSS OF VISUAL ACUITY is not an early sign in POAG.**

(5) **CUPPING OF THE OPTIC DISCS** is the important sign. Chronic glaucoma causes the discs to become deeper and wider, and the remaining rims of disc tissue to atrophy (28-11). One eye is commonly affected more than the other, so that a definite difference between the eyes is probably abnormal. Enlargement starts at the upper or lower margins, so that a vertically ovoid cup with a cup/disc diameter ratio of >½ is probably abnormal. Eventually, the margin of the cup approaches the margin of the disc, so that only a narrow rim of tissue remains. Its wall becomes steep, so that vessels bend abruptly as they reach the level of the surrounding disc. If the edge of the disc overhangs the cup, you may lose sight of the vessels until they appear over the edge of the cup.

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**SOME OPTIC DISCS OF THE LEFT EYE**

![Fundoscopic appearances: Normal](image1)
![Fundoscopic appearances: Normal large physiological cup](image2)
![Fundoscopic appearances: Glaucoma](image3)

**Fig. 28-11 SOME OPTIC DISCS OF THE LEFT EYE.**
A. normal optic disc with a moderately sized cup. The lamina cribrosa is a mesh-like structure of collagen in the sclera wall which maintains the pressure gradient. B, another normal optic disc. This has a large physiological cup and a temporal scleral crescent. C, optic disc of a patient with gross 'chronic' open-angle glaucomatous cupping. After Putt J. Introduction to Ophthalmology. OUP 2nd ed 1982, with kind permission

A large physiological cup can, however, be difficult to distinguish from that in early glaucoma. With practice, eye assistants should be able to distinguish 'normal discs', 'suspicious discs' and 'advanced glaucomatous cupping'. This is the most important sign to look out for in glaucoma: you usually need to dilate the pupils.

**Normal discs:**
1. (a) The discs are the same in both eyes.
2. (b) The ratio of optic cup to optic disc is ≤½ (28-12G).
3. (c) The cup is circular and the periphery of the disc (the optic nerve rim) is pink.
4. (d) The appearance of the disc remains constant over time.

**Signs suggestive of glaucoma:**
1. (a) A cup/disc ratio >½.
2. (b) A vertically oval cup, perhaps with notching at the upper or lower poles.
3. (c) An area of pallor >30% of the disc area.
4. (d) Asymmetry of the cup/disc ratio of the 2 eyes.
(6) ABNORMAL PUPIL RESPONSES are a useful way of testing for glaucoma, and only need a torch. Initially, one pupil does not react as briskly as the other. Finally, there is no response at all.

If you shine a light into a normal eye in a semi-dark room, its pupil will constrict (direct response), and so will also the other pupil automatically (consensual response).

**If the optic nerve is completely destroyed**, there will be no direct or consensual response (total afferent pupil defect).

**If the optic nerve is partly destroyed** (for example 90%), the pupil will constrict slowly when the light shines in it (partial afferent pupil defect), and the consensual response will be present.

The swinging torch test, is a useful test for early asymmetrical optic nerve damage, and does not need an ophthalmoscope. It is theoretically difficult, but is easy in practice. In a semi-dark room shine a light into the good eye, and then swing it across into the bad eye (the eye with reduced vision).

As the light shines in the good eye, the pupil of the bad eye will constrict. As you swing the light quickly across to the bad eye, its pupil, which was previously constricted, will now dilate. This indicates a relative afferent pupil defect, early optic nerve damage, and a difference in function between the two optic nerves.

The practical test is to swing the torch from one pupil to the other and back again in a semi-dark room. If one pupil consistently dilates as light shines on it, that eye has a reduced pupil response, relative to the better eye. You should investigate this for optic nerve disease, perhaps POAG.

**RISK FACTORS FOR POAG:**
1. Age >40yrs.
2. A positive family history in first-degree relatives.
3. A vertical cup/disc ratio >½.
4. An IOP of >28mmHg (<2 with a 5.5g weight).

**MANAGEMENT** will preserve what sight exists, but will not improve it.

**If there is any sight left**, perform a trabeculectomy. Even when sight is as poor as CF 1m (28.1), there may still be some benefit from treatment.

Medical treatment can lower the IOP, but it has to be constant, consistent and continue for life, which is usually impracticable, so that immediate surgery is better. Start with pilocarpine 4% qid or timolol 0.5% bd (expensive), and if this fails to maintain the IOP <20mmHg, add acetazolamide 250mg qid orally for short periods.

**N.B. Several side-effects may occur with continued use.** To treat glaucoma effectively, measure the intraocular pressure, and monitor the visual fields regularly. Do this in a specialist clinic.

Surgical treatment is a trabeculectomy which removes a piece of the filter (the trabecular meshwork), and so allows the intraocular fluid to drain under the conjunctiva; this increases drainage and reduces the IOP. The operation has an immediate success rate of >80%, and is the recommended treatment for most patients with POAG, but it does not have a permanent effect. You should follow up these patients long-term; they may need repeat surgery.
ANGLE-CLOSURE GLAUCOMA,
(ACG, ACUTE GLAUCOMA)

Most acute glaucoma is usually this type, but secondary glaucoma (see below) may occasionally present acutely.

DIAGNOSIS.
Presentation is with an 'acute red eye' at any age (one of its rarer causes, 28.3), severe unilateral headache in and around the eye, and sudden profound loss of vision.
Vision is reduced, the eye is red, the cornea is hazy from oedema, the anterior chamber is shallow, the pupil is usually dilated, and IOP usually >40mmHg.

CAUTION! Blindness is inevitable unless you treat this quickly.

TREATMENT.
Admit the patient as an emergency. If referral is delayed, start treatment before operating. Use analgesic to ease the pain, and treat both the eyes. Aim to:
(1) Lower the IOP by increasing the drainage of aqueous. Use acetazolamide 500mg orally followed by 250mg qid, as soon as the nausea has subsided. Apply timolol 0·5% immediately and repeat this twice.
(2) Keep the pupils constricted. This will keep the periphery of the iris away from the angle of the eye, where the aqueous flows out, and so help it to drain. Treat both eyes with drops of pilocarpine 1% every 15mins, for 2hrs. If this makes the pupil constrict, the angle will be opened.
(3) You can also lower the IOP by increasing the tonicity of the blood. Use 50ml of flavoured glycerine by mouth if nausea has settled. Alternatively, use 200ml of 20% mannitol IV over 20mins.

When medical treatment has reduced the IOP to normal, arrange a peripheral iridectomy soon, to prevent a future attack; recommend a prophylactic iridectomy on the other side also.

CAUTION!
(1) Blindness can occur in 12hrs, so treatment is urgent.
(2) Rainbow-coloured haloes round lights, and misty vision, are important prodromal signs, and need urgent investigation and treatment.
(3) Atropine can precipitate an attack in a patient with a shallow angle!

TRABECULECTOMY & PERIPHERAL IRIDECTOMY
GRADE 2.4
It helps to have a microscope to do this operation.
Insert the eyelid retractor. Secure the eye by means of a traction suture of 4/0 silk through the belly of the superior rectus muscle. Select the site of the incision cranially (usually at the 10-11 o'clock position) by injecting 0·2ml saline subconjunctivally. Incise the conjunctiva at the limbus (28-1B) for 5mm and undermine it to free it from Tenon’s capsule (28-1C). Control bleeding with a fine cautery. Mark a rectangular sclera flap 5x5mm of half the scleral thickness hinged at the limbus, dissecting this forward till you see the transparent cornea.

Make an incision through the cornea into the anterior chamber behind the hinge of the sclera flap and cut out a 4x2mm block of tissue containing the trabeculum and Schlemm’s canal (28-1B). Then grasp the peripheral iris with forceps, prolapsed it and excise it to make a peripheral iridectomy.

N.B. If you incise too much sclera, you may reach the ciliary body and cause a choroidal detachment.
Return the sclera flap on its hinge to its bed, and close wound, and then close the conjunctiva at each corner both with 9/0 or 10/0 nylon.

SECONDARY GLAUCOMA.
This complicates:
(1) trauma, including hyphaema (blood in the anterior chamber).
(2) swollen cataract (see below).
(3) some cases of iritis. Treat the primary condition, and try to arrange referral: in the meantime, treat with acetazolamide, mannitol, or glycerol as above.

DIFFICULTIES WITH GLAUCOMA

If there is an acute red painful eye, a fixed dilated white pupil, a shallow anterior chamber and a hazy cornea, this is a SWOLLEN CATARACT, causing secondary glaucoma. Use acetazolamide 500mg immediately followed by 250mg qid, and arrange cataract extraction.

If there is loss of vision from chronic use of steroid drops, suspect STEROID GLAUCOMA. Topical steroids cause a genetically determined rise in IOP in 30% of people. This is sometimes severe enough to cause glaucoma, exactly like POAG. Stop the steroids.

If a child has big eyes, which may be associated with photophobia, blepharospasm, and tears, suspect CONGENITAL GLAUCOMA (buphthalmos, ox-eye), due to maldevelopment of the angle of the anterior chamber. The sclera of a child are soft, so that the eyes enlarge when the IOP rises.

Other signs are: an increased IOP, a corneal haze (variable), sluggish reaction of the pupils to light, and enlargement of the cornea (>12mm), or of the whole globe. Try to arrange an urgent incision of the trabecular meshwork to allow flow of aqueous.

28.7 Onchocerciasis (River blindness)

Onchocerciasis is a parasitic infection of the skin and eyes caused by Onchocerca volvulus, which used to be endemic in parts of West Africa, with foci in East Africa and Latin America. WHO-led programmes have virtually eliminated this disease except in West & Central Africa.
In endemic areas 20% of the population are infected by a worm transmitted by the blackfly. Microfilariae invade all parts of the eye: the cornea (keratitis), the anterior chamber (iritis), the retina (chorioretinitis), and the optic nerve (optic neuritis). Blindness and irreversible eye lesions are most often found in people ≥30yrs.

A patient from an endemic area complains of itching, with or without a rash. There may be skin nodules on the hip or shoulder (in Africa) or scalp (in S. America), night blindness (enquire about this), gradual loss of vision in both the eyes, and sometimes tears and photophobia. Look for microfilariae in snips from the skin. There may also be huge adenolymphocoeles in the groin (18-5).

CORNEA. There is a sclerosing keratitis with opacification of the lower third of the cornea. Tongues of opacification invade the cornea from the 3 & 9 o’clock positions, or from anywhere in the lower ½, where they may form an apron across the cornea. If this is not treated, opacification slowly advances upwards over the pupil, until all that may be left is a clear area at 12 o’clock. Slit lamp microscopy shows 0·5mm linear and fluffy opacities at all levels in the stroma, and minute wriggling microfilariae in the anterior chamber.

IRIS. The final stage is a small, non-reactive, down drawn, pear-shaped pupil. Earlier stages are a loss of pigment in the margin of the pupil, exudation in front of, across, or behind it, posterior adhesions which turn it inwards, and small keratic precipitates (KP). A gelatinous exudate sometimes drags the lower margin of the pupil down and everts it. Posterior synechiae and peripheral anterior synechiae lead to secondary glaucoma.

RETINA AND OPTIC DISCS.
Look for:
(1) Diffuse white areas and black pigmented ones most marked temporal to the macula (the Ridley fundus).
(2) Optic atrophy, with sheathing of the vessels close to the nerve.
(3) Microfilariae wriggling in the anterior chamber (after putting the head between the knees for at least 1min)

TREATMENT. Use ivermectin once yearly as a single dose according to weight, for 15yrs (the lifespan of an adult worm):

<table>
<thead>
<tr>
<th>Weight (kg)</th>
<th>Dose (mg)</th>
</tr>
</thead>
<tbody>
<tr>
<td>15-25</td>
<td>3</td>
</tr>
<tr>
<td>26-44</td>
<td>6</td>
</tr>
<tr>
<td>45-64</td>
<td>9</td>
</tr>
<tr>
<td>65-84</td>
<td>12</td>
</tr>
</tbody>
</table>

If there is also loiasis, (28.18) with high quantities of microfilariae, add prednisolone 20mg od 3days before starting treatment and till 3days afterwards to prevent severe encephalopathy.

If there is an onchocercal iritis, dilate the pupil with atropine and use topical steroids (28.5). If secondary glaucoma (28.6) develops, arrange surgery.

28.8 Refractive errors: difficulty reading & presbyopia

In terms of comfort, increased efficiency and the number of people who benefit, the prescription of glasses is among the most valuable procedures in medicine. So do what you can to supply them. You may be able to get cheap self-tuning glasses (which contain an adjustable amount of fluid between the clear plastic making the lenses). However these are difficult to maintain and require skill to use.

The refractive errors are:
(1) Myopia (short sight).
(2) Hypermetropia (long sight).
(3) Astigmatism (the refractive mechanism is aspherical).
(4) Presbyopia (common long-sightedness of old age).
(5) Aphakia (‘no lens’) after old-style simple cataract extraction.

Presbyopia is part of the normal process of ageing: it appears earlier in the tropics than in higher latitudes, and is easily diagnosed and treated. It is responsible for 85% of the need for glasses.

Whereas a myopic child has to be specially fitted, because the eyes may not be the same, a presbyope can, if necessary, be left to choose the glasses which best suit him from a pile of second-hand ones. Astigmatism is more difficult to correct, but it is usually so mild that it needs no correction: its correction is often overemphasized.

N.B. The visual acuity of all patients with refractive errors improves when they look through a pin-hole, which uses only their central vision. This is the basis of the pin-hole test (28.1).

REFRACTIVE ERRORS
If there is astigmatism, due to irregularly curved lenses, the vision is blurred.
If there is hypermetropia (long-sightedness), a patient has difficulty reading or seeing things close at hand, e.g. threading a needle.
If there is myopia (short sightedness), a patient cannot see clearly at a distance, but near objects are clear.

Unless you can perform retinoscopy, test each eye alone by trial and error with spherical lenses. Use the charts (28-2).

For long-sightedness, start with +1 dioptries, the usual requirement at 43yrs, and by +0·5 dioptries increments for each 5yrs. The smallest number that gives the best acuity is the prescription for glasses needed.

For short sightedness, start -0·5 dioptries and continue adding -0·5 increments. Again the smallest number that gives the best acuity is the prescription needed.

Aphakia after old-style cataract extraction requires +10, 11 or 12 dioptries lenses for distance, and +13, 14 or 15 lenses for reading. These are easily broken, rendering the patient blind again!
Check that the eye is not red, and that there are no other serious eye problems.

A child with reduced visual acuity due to refractive errors needs glasses, whereas an adult probably does not.

A child <7yrs with a squint needs surgery (28.9)

Remember glasses for reading and other close work, are not useful for seeing at a distance.

If an adult has a squint and no double vision it may be the cause of reduced vision in the squinting eye (amblyopia). There is no treatment at this age.

If a child <7yrs presents with a squint:
(1) Dilate both the pupils, and use an ophthalmoscope to make sure that the squint is not due to a retinoblastoma in one of the eyes (an uncommon cause, 28.16: look for a yellowish mass on the retina).
(2) If the retinae are normal, try to assess the visual acuity in both eyes. This is relatively easy at >3yrs, but is difficult at younger ages (28.1).

If there is reduced vision in either eye, or a definite squint at >7yrs, there may be need for glasses, and perhaps surgery on the extraocular muscles.

Correct any refractive error and occlude the eye that is most used. Occlude it for ½-2hrs/day during close work, reading, or drawing. The duration of treatment depends on the duration of the amblyopia. If treatment is prompt, 6-8wks of intermittent occlusion treatment may be enough. If it is delayed you may need to continue for 1yr.

AFTER THE AGE OF 6 MONTHS DON’T IGNORE A SQUINT!

28.9 Disease of the neuromuscular system: squints, amblyopia, and diplopia

If the eyes do not look in the same direction, this is known as a squint (strabismus). Squints are common, and are usually accepted with resignation. Although treating a squint needs relatively simple technology, it is time-consuming and needs skill.

If the eyes do not look in the same direction, two images are generated. This causes confusion, and to avoid this, the brain suppresses one image. If this suppression continues for long enough, vision disappears in that eye. This is called amblyopia, which is a reduction in vision, due to lack of use of an apparently normal eye. If amblyopia is uncorrected in a child by 7yrs, it becomes permanent. So try to diagnose a squint and arrange surgery before this age.

CAUTION! Never occlude the eye of a child <7yrs for several days, because this may cause amblyopia.

SQUINTS AND AMBLYOPIA
DIAGNOSIS.

The corneal light reflex: Shine a pen torch directly in front of you, and ask the patient to look at it. If each of the eyes is properly fixing the torch, its reflection from the corneal mirrors will be the same, and more or less central on each cornea. Are the reflections from your torch equally centred on the pupils?

The cover test: Ask the patient to look straight ahead at some target in the distance. Cover the left eye with a piece of paper. If the right eye moves, in or out, to fix on the distant target, it was previously squinting. If it does not move, it was looking straight at the target.

Now put the paper in front of the right eye. If the left eye moves as you remove the paper, it was previously squinting in or out. If it does not move, there is no manifest squint, and both eyes look straight.

MANAGEMENT is limited.

If an adult has a squint and double vision, this suggests a serious recent disease of the extraocular muscles or their nerves, such as diabetic neuropathy, myasthenia gravis, or raised intracranial pressure. He needs a full medical and neurological examination.

If a child <7yrs presents with a squint:
(1) Dilate both the pupils, and use an ophthalmoscope to make sure that the squint is not due to a retinoblastoma in one of the eyes (an uncommon cause, 28.16: look for a yellowish mass on the retina).
(2) If the retinae are normal, try to assess the visual acuity in both eyes. This is relatively easy at >3yrs, but is difficult at younger ages (28.1).

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AFTER THE AGE OF 6 MONTHS DON’T IGNORE A SQUINT!

28.10 Diseases of the lids & nasolacrimal apparatus

Diseases of the eyelids and nasolacrimal system include tumours, deformities of the lids, and watering (epiphora). Globally, the most important disease of the lids is trachoma, which scars the lids, and causes them to turn inwards (entropion, 28.13). The commonest and usually the most harmless disease of the lid is a stye. The lid is involved in herpes zoster ophthalmicus, where the virus affects the VIIth cranial nerve, often associated with HIV infection. Cutaneous leishmaniasis causes the lid to scar and produce an everted eyelid (ectropion). Loiasis is found in equatorial rain forests, where the vigorously mobile loa loa worm is found under the conjunctiva, causing an acute inflammatory swelling.

THE LIDS

If a patient has a red swelling on the lid margin, with an eyelash coming out of it, this is a stye (hordeolum). It is a staphylococcal infection of an eyelash follicle. Pull the eyelash out of the swelling, using an analgesic. Only use antibiotics in recurrent styes, or if infection is spreading beyond the lid (cavernous sinus thrombosis is a rare complication, 6.6). Warm soaks (28.3) are useful.
If there is a swelling in either lid, some distance from its margin, pointing towards its conjunctival surface, it is probably a tarsal cyst, (chalazion, Meibomian cyst). Avoid an external scar by incising the conjunctiva wherever the cyst is about to burst (28.12).

If a few lashes turn in on the eye (trichiasis), remove them with electrolysis.

If most of the lashes or the margin of the lid are turned in (entropion), perform surgery (28.13).

If the upper eyelid, eyebrow and forehead are involved in a vesicular eruption with an abrupt demarcation in the midline, this is herpes zoster. Acyclovir given early will help. Later complications can result in ectropion, keratitis, uveitis and secondary glaucoma.

If there is an acute attack of localized oedema of the orbit in an endemic area, consider loiasis. If you can see the worm under the conjunctiva, instil lidocaine drops, make a small incision and remove the worm with forceps. Ivermectin annually is useful as a prophylactic.

If the lids do not close properly, as a result of a VIth cranial nerve palsy (from herpes, iatrogenic injury in parotid surgery, leprosy, or parotid malignancy), the cornea will remain exposed, especially at night, and exposure keratitis will result. To avoid this, do a tarsorrhaphy unless you can supply artificial tears.

If the margin of either lid is everted (ectropion), usually as the result of scarring, or long-standing palsy, release of lid contracture with skin grafting of the defect is necessary.

**TARSORRHAPHY (GRADE 2.2)**

**INDICATIONS**
Eyelids which permanently fail to cover the cornea properly, especially when the cornea is anaesthetic.

N.B. This is usually a temporary measure.

**METHOD.**
Stretch the lid by grasping one end of the lid margin and make an incision 2mm deep along the midline of the upper and lower margins, just beyond the eyelash follicles (28-13A). Then pass a 4/0 silk suture through the skin above the upper eyelash line and out through the incision, and similarly in through the lower lid incision and out below the lower eyelash line (28-13B). Turn the suture back, through 3mm rubber rings to take the tension, through the skin (28-13C) and tie it so that the lids are well approximated (28-13D).

You need as many sutures as required to get adequate eyelid closure; this may only be necessary on the lateral aspect, but do not make the tarsorrhaphy too small. Put chloramphenicol eye ointment or drops qid and make sure no eyelashes project back inside the suture line.

**RELEASE OF LID CONTRACTURES (GRADE 2.2)**

**INDICATIONS**
Contractures starting to expose the eyelids and expose the cornea. You should have done a tarsorrhaphy already.

**METHOD**
Put stay sutures through the lids (28-8F), so that you can move them up or down as necessary. Make the relaxation incisions (28-14A). When you have prepared a satisfactory bed for the graft, and controlled bleeding, stretch it, apply a piece of split skin graft, and hold it in place with tie-over sutures using 4/0 monofilament (28-14C). If the graft contracts after 12wks, repeat the release and apply another graft. Do not worry too much about what the patient’s eye looks like at this stage. What matters is that the cornea should not be exposed.

**DO NOT DELAY GRAFTING THE LIDS**

CAUTION! Stretch the lid first so that there will be some slack tissue when it contracts later. The thinner the graft, the more the shrinkage. If you are skilled, apply a full thickness graft. Primary skin grafting will not prevent ectropion, and you may need 2 or 3 operations to insert enough skin.
CONTRACTURES OF THE EYELIDS

Fig. 28-14 CONTRACTURES OF THE EYELIDS. A, retraction of scar tissue everting the eyelid and exposing the cornea. B, make relaxing incisions and get ready for grafting. C, hold grafts in place by the tie-over method. Kindly contributed by Randolph Whitfield II.

THE NASOLACRIMAL APPARATUS

If something interferes with the drainage of an adult’s tears, the eye waters (epiphora) even if there is no local irritation. Epiphora can occasionally be so severe that it needs surgery (a dacryocystorhinostomy).

If a mother brings you her young child saying that he has had a watering eye since birth, this is congenital atresia of the nasolacrimal duct. It will probably resolve spontaneously by the age of 18 months. Reassure her, and use a topical antibiotic if conjunctivitis develops.

If the eye is still watering at 2yrs, the nasolacrimal duct needs probing and syringing.

If there is a tender swelling between the eye and the side of the nose, this is probably acute dacryocystitis (an abscess in the tear sac). Use IV gentamicin, an analgesic, warm soaks, and incise the skin of the lower eyelid over the lacrimal sac (6.6).

28.11 Proptosis (Exophthalmos)

If there is a space occupying lesion in the orbit, it pushes the eye forwards. Proptosis is always serious, and it can be difficult to diagnose, but is uncommon. Some of its causes need medication (orbital cellulitis: IV antibiotics and Burkitt’s lymphoma: cytotoxics).

Most patients who need surgery are either going to die from malignant tumours anyway, no matter what is done, or they have slow-growing benign tumours, which you have time to try to refer. So your ability to help a patient with proptosis is limited; but you should try to make a diagnosis.

An adult may have:
(1) A retrobulbar haematoma following an injury (common). This is only an incident in a head injury, and the diagnosis is obvious.
(2) A mucocoele of the frontal sinus (the commonest cause), due to an infection followed by an obstruction, which prevents the sinus draining into the nose.
(3) Orbital cellulitis, or an orbital abscess, usually following frontal or ethmoid sinusitis (6.6), or occasionally trauma.
(4) A pseudotumour of the orbit due to a granuloma of unknown cause.
(5) An epidermoid or dermoid cyst, which may be of the ‘dumb-bell’ type, and extend into the anterior cranial fossa. Do not operate on these, unless you are skilled enough to dissect widely, and have made an accurate diagnosis.
(6) A lacrimal pleomorphic adenoma, palpable at the inferior orbital rim.
(7) A haemangioma; you may be able to empty a haemangioma temporarily by pressing it back into the orbit.
(8) A hydatid cyst, if this disease is endemic (15.10).
(9) A cavernous sinus thrombosis (6.6)
(10) A carotico-cavernous fistula, where there is an audible noise in the head and a bruit heard over the eye; this may follow trauma but is usually spontaneous.
(11) A metastasis.
(12) A malignant melanoma (34.6).
(13) A conjunctival carcinoma.
(14) The hyperophthalmopathic form of thyrotoxicosis (Graves’s disease, 25.2).
(15) A meningioma of the sphenoid.

A child may have:
(1) A retinoblastoma in the first 5 years of life (28.16).
(2) Acute ethmoiditis, commonly around 2yrs.
(3) Burkitt’s lymphoma (17.6), usually associated with a jaw tumour.
(4) A rhabdomyosarcoma (34.15).
(5) Some other kind of lymphoma (17.6).
(6) A neuroblastoma.
(7) A metastasis.

Proptosis can occur slowly over years, or rapidly over days. Its causes vary geographically, and with the age of the patient. The more common causes are listed first; the later ones are mostly very rare.

N.B. Thyrotoxicosis may cause unilateral proptosis, so check for weight loss, & tachycardia.
PROPTOSIS

A, B, Burkitt’s lymphoma, before and after treatment. C, proptosis, cause not yet established. D, carcinoma of the maxillary antrum extending into the orbit.


EXAMINATION. Sit the patient down, stand behind him, look down on the eyes from above, and observe the relative positions the globes. This will help to distinguish pseudoproptosis, due to the relative widening of one palpebral fissure.

Hold a ruler horizontally, and measure the position of each cornea from the midline. If there are 2 protruding globes, and they are both equidistant from the midline, this is probably thyrotoxicosis (the most likely cause of bilateral proptosis). If they are not equidistant, one globe has probably been pushed out of place by an orbital mass. Examine the fundi for papilloedema and optic atrophy. Search for signs of a primary malignant tumour.

CAUTION! Do not confuse proptosis with a staphyloma due to a neglected corneal ulcer (28.3). The normal intraocular pressure has caused the previously weakened cornea to bulge, in a manner which you can mistake for a tumour. The globe however remains in its normal position.

RADIOGRAPHS may demonstrate:
(1) Erosion of orbital bones.
(2) Sclerosis of orbital bones (typical of a meningioma).
(3) Calcification (sometimes in a retinoblastoma).

BIOPSY may be practical.
If there is a tumour palpable externally, take a biopsy, but if there is a swelling of the upper lateral quadrant of the orbit, pushing the eye downwards and inwards, which has grown slowly over many months or years, this is probably a lacrimal pleomorphic adenoma, which you should not biopsy for fear of spreading the tumour tissue. Do not be deceived by the small mass of tumour palpable externally: most of it will be inside the orbit behind the eye. It needs removing through a lateral orbitotomy.

DIAGNOSIS AND MANAGEMENT.
If the proptosis arose acutely, and the lids are red and swollen, perhaps with a fever and tachycardia, this is orbital cellulitis, or an orbital abscess. Use IV antibiotics. If there is an abscess drain it (6.6).

If a child c.2yrs has sudden unilateral proptosis, with swollen lids and conjunctiva, fever and tachycardia, suspect acute ethmoiditis. Use IV antibiotics.

If there is an acute pulsating proptosis, which may be unilateral initially, but soon becomes bilateral, with engorgement of the veins, and total inability to move the eye, with severe prostration or loss of consciousness, suspect cavernous sinus thrombosis (6.6).

If there is a swelling which has enlarged slowly (weeks or months) in the superior nasal quadrant of the orbit, pushing the eye downwards and outwards, this is probably a mucocele of the frontal sinus (common). By an approach between the periosteum and the frontal bone, keeping outside the orbit, enter the sinus and drain the mucopus. Place a drain from the sinus into the nose. Suture the skin in layers. Remove the nasal drain at 6wks.

If the patient is between 15-35yrs, and the proptosis occurred over several weeks or months, suspect idiopathic orbital inflammation. The diagnosis is largely made by excluding other causes. It will respond well to prednisolone 60mg od for 1wk, reducing slowly to 5mg od by the 4th wk. Maintain 5mg a day for several months, or it will recur.

If the proptosis of thyrotoxicosis does not respond to medical or surgical treatment (25.2), try high dose systemic steroids. If this fails, try to arrange surgical orbital decompression by removal of fat or bone.

TREATMENT FOR THE EXPOSED CORNEA.
Examine the cornea to make sure that it is not ulcerated. Apply antibiotic eye ointment qid, and especially at night. If necessary, protect it by tarsorrhaphy (28-13). Padding can be dangerous, because the pad may abrade and ulcerate the cornea.
28.12 Tarsal (Meibomian) cysts (Chalazions)

Meibomian glands secrete oil at each blinking of the eyelid; this prevents evaporation of the fluid film on the eye. Cysts may form in these glands on the conjunctival side of the tarsus (or eyelid). They present as a swelling in either lid, which may become chronically or acutely infected. Small asymptomatic ones need no treatment, and may resolve spontaneously. Incise an acute infection and curette a chronic one. These cysts are common everywhere, so that treating them is a common outpatient eye operation. Sometimes, they present as granulomas.

Fig. 28-16 CURETTING A TARSAL CYST.
A, chalazion close to the medial canthus. B, chalazion forceps. C, introducing LA. D, chalazion clamp in place ready to incise a chalazion in the centre of the lower lid. E, curetting the chalazion

ANAESTHESIA.
Anaesthetize the conjunctiva with drops of lidocaine 4%, or amethocaine hydrochloride 1%. Infiltrate the lid with lidocaine and adrenaline around the chalazion. Insert the needle at the upper margin of the upper tarsus, and the lower margin of the lower tarsus. Carry it forwards to the lid margin, on either side of the chalazion.

EQUIPMENT. Chalazion clamp, #11 scalpel blade and curette.

CURETTAGE. (GRADE 1.2)
Evert the lid slightly. Put the chalazion clamp over the cyst, so that the solid blade lies on the skin of the eyelid, and the ring lies on the conjunctiva over the cyst. Close it so that it holds the lid and the cyst. Insert the tip of a #11 blade, so that it cuts away from the eye. Always make the incision perpendicular to the lid margin so as to avoid cutting the levator muscle. Swab its contents clean.

CAUTION! Take care to curette away any pockets of granulation tissue, which may be hidden by a flap of conjunctiva, or have herniated themselves through the tarsal plate into the orbicularis muscle. If you do not do this, it may recur.

Remove the clamp and pinch the lid until it stops bleeding. If this is troublesome, wash it with warm saline. Place chloramphenicol ointment in the conjunctival sac for 1wk.

If the material you incise is hard, and not gelatinous, suspect a carcinoma. Send it for histological examination.

28.13 Entropion

Trachoma is the commonest eye infection in the tropics and also amongst Aborigines in Australia, and in its blinding hyperendemic form is worldwide the 2nd commonest single cause of blindness and impaired vision (cataract is the first). It is a chlamydial infection, which spreads from the eyes of one person to another, especially among children, in the poorest and most disadvantaged communities, particularly those in the Middle East and Africa.

Trachoma is a chronic follicular conjunctivitis; it scars the conjunctiva of the eyelids and the cornea, and goes through 4 stages (28.3). A single dose of 20mg/kg azithromycin (or 500mg bd erythromycin for 1wk in pregnancy) orally is effective in arresting the disease:
STAGE I. There is a mildly red watery, eye due to bilateral conjunctivitis, especially of the upper lids, but without any distinguishing features.

STAGE II. Under the upper lid there are dilated blood vessels and hyperaemic, oedematous epithelial tissue (papillae). There are also yellow-grey swellings (follicles). Look at the corneoscleral junction with a loupe. If the edge of the cornea looks mildly grey, owing to an arcuate (crescent-shaped) grey infiltration, and blood vessels go beyond the grey area into the cornea, there is pannus (meaning a curtain). This starts at the 12 o'clock position, and extends to 9 & 3 o'clock. Follicles and pannus indicate stage II trachoma. Follicles are not diagnostic, but pannus is.

STAGE III. The follicles in the lids become coarser and pannus spreads, sometimes across the pupillary area of the cornea. Scarring makes the margin of the upper lid irregular, and turns the upper tarsus inwards (entropion), taking the lashes with it, so that they scratch the cornea during every blink (trichiasis, 28-18C). This causes recurrent attacks of keratitis, which eventually results with a corneal opacity causing blindness.

STAGE IV. Fibrous tissue replaces the follicles in the lids. This is the stage of scarred, or healed trachoma. The cornea is grey and scarred, the vision severely impaired, and the lids are deformed.

If you work in an endemic area, you are likely to have many patients with entropion, so learn how to correct their eyelids yourself, and if necessary train an assistant to do this. The operation is always worth doing, even if the lids are severely scarred: sight may recover surprisingly.

Several operations are possible:
1. Splitting the eyelid margin
2. Radical eyelash excision
3. Tarsal eversion

If only a few lashes are turning in, try removing these with an electrolysis needle under LA. Epilation (pulling out the lashes) is ineffective as they will grow back. Do not cut the eyelashes short because this makes them sharp and even more irritant.

ENTROPION CORRECTION

INDICATIONS.
Trachoma which has distorted the upper tarsus, so that it has curled inwards and made the lashes scratch the globe. Operate as soon as possible after entropion occurs.

You can operate on both the eyes at the same time, but this will require admission for 3 days, to allow the oedema of the eyelids to subside. Absorbable sutures allow discharge without the patient needing to return.

EQUIPMENT. An eye set, a scalpel with #15 blade, 4/0 absorbable suture. A cautery will help to control bleeding.

ANAESTHESIA.
Anaesthetize the upper lids through the skin with 1 ml of 2% lidocaine with adrenaline. Anaesthetize the conjunctiva with 2 drops of amethocaine, or lidocaine.

EYELID MARGIN SPLITTING FOR TRICHIASIS (GRADE 1.3)

Split the eyelid into two parts along a ‘grey’ line defined by the orbicularis oculi muscle: just inside are the openings of the Meibomian glands and just outside are the eyelash roots. The divided eyelid will then have an inner part containing the conjunctiva and tarsal plate, and an outer part containing the orbicularis oculi muscle, the inverted eyelashes and their roots. Rotate this outer part outwards and fix it by an evertting suture over a gauze swab.

Fig. 28-17 RADICAL EYELASH EXCISION. A, direction of the incision. B, operation complete.

RADICAL EYELASH EXCISION FOR ENTROPION (Malcolm Phillips) (GRADE 2.3)

Removing the eyelashes completely will mean they cannot re-grow and scratch the cornea. This results in little cosmetic disability, especially with darker skin, and the relief that follows is dramatic.

If you are a right-handed operator, start with the right eye. Use a scalpel to incise the margin of the upper lid, at the lateral end of the lashes, to a depth of 3 mm (28-17A). Using small sharp scissors, remove the margin of the lid bearing the roots of the lashes. Cut towards the medial end and preserve the punctum. Evert the lid as you do this, by pressing it with a swab. Then repeat the procedure on the left eye.

Control the considerable bleeding that will result by suturing the conjunctiva to the skin of the eyelid with 3/0 absorbable sutures on a cutting needle. Insert about 5 sutures, 5 mm apart, knotting them, and use the same suture to hold little rolls of gauze. Apply an eyepad for 24 hrs. Remove the part of the sutures holding the gauze rolls after 3 days.

Fig. 28-17 RADICAL EYELASH EXCISION: A, direction of the incision. B, operation complete.
TARSAL EVERSION FOR ENTROPION (GRADE 2.4)

For this method, the tarsal plate must be stiff enough to take sutures. Place 3 stay sutures of black braided silk in the upper lid, evert it over a roll of gauze, and clamp the sutures to a drape.

Using a #15 scalpel, make an incision about 3mm from the inner margin of the lid, and parallel to it. Cut through the conjunctiva and the full length of the tarsal plate, at 90° to its surface, so as to free a strip from its edge (28-18E). Curve each end of the incision towards the free edge of the lid, so that you can evert the strip of lid that bears the lashes.

This operation gives the upper lid a new edge, and makes it c. 3mm shallower; but it will still meet the lower lid on shutting the eye.

Alternatively, if the tarsal plate is shrunked and degenerated, it will not take sutures, so remove it entirely.

A more complicated procedure is a tarsal plate rotation with a mucosal graft from the mouth, but this is for an expert.

If the eye is painful and blind, it may be better removed. This is one of the occasions on which the indications are more critical than the operation.

**CAUTION!** Take care not to buttonhole the skin.

You will now be able to rotate the distal fragment through 180° (28-18F). If you cannot, undermine the conjunctiva more widely. Insert 3 small mattress sutures of 4/0 absorbable, so that the knots are buried (28-18G). Put tetracycline eye drops into the conjunctiva tid for 1wk.
28.14 Destructive methods for the eye

Evisceration is the least radical procedure; scrape out the contents of the globe and leave the sclera intact. This is the only safe procedure if the eye is infected, because a sleeve of dura containing CSF surrounds the optic nerve. Other operations require that you cut it, and so open up a potential path of infection to the meninges. You may need to eviscerate the eye:

1. When antibiotic therapy fails to control a severe infection causing suppurative endophthalmitis, leading to orbital cellulitis, and oedema of the lids. If you do not eviscerate the eye and drain the pus from it, the infection may spread and cause cavernous sinus thrombosis and meningitis, and death.

2. When there is a chronic less urgent infection in a blind painful useless eye. Try to find a prosthesis to insert when the sepsis has settled: it vastly improves the patient’s appearance.

Enucleation (excision) removes the globe by dividing the conjunctiva, the extrinsic muscles of the eye, and the optic nerve. Do this only where there is no active infection; it is contraindicated when there is.

Exenteration is a bloody, mutilating operation. It removes the entire contents of the orbit, together with its periosteum, the globe, and all its extrinsic muscles. Consider doing this when there is a fungating malignant tumour of the eye or orbit. It will not prolong life, but the last days might at least be more comfortable. An empty orbital cavity will remain, which you can line with split skin, or allow to granulate.

**Before you start any destructive operation:**
(a) Get signed informed consent from a child’s parent or guardian or from an adult himself
(b) Make sure you operate on the correct eye!

**Do not rush in to perform these procedures:**
(a) The main indication for enucleation is persistent severe pain in a blind eye.
(b) If it has been injured, always try to repair it first, no matter how hopelessly injured it is. In some cases the eye may scar but remain stable for years.
(c) Prostheses may be difficult to find, and the best one is the natural eye, even if it is blind.

**CAUTION!**
(1) Unless you are operating for malignancy or acute infection to save life, the eye must be totally blind. Test this with a strong light.
(2) When pain is the main indication, it must be considerable. Pain is subjective, so make sure, if you can, that it is genuine. Review this on several occasions. If there is any sensation of light, do not do a destructive procedure. That little eyesight may be useful later.

**Evisceration of an eye (GRADE 3.2)**

**INDICATIONS.**
(1) The failure of antibiotics to control a suppurative endophthalmitis.
(2) A blind, painful eye, especially if it is infected.

**Do not take the eye out unless you are left with no other option.**

**ANAESTHESIA.** If there is no significant infection, you can use the combination of a facial and a retrobulbar block. Otherwise use ketamine or GA.

**METHOD.**
Incise the conjunctiva all round 360° at its junction with the cornea, using fine-toothed forceps and fine scissors (28-19A). Separate Tenon’s capsule (28-1C) bluntly from the underlying sclera in 4 quadrants. Cut through the corneoscleral junctional at the limbus with scissors (28-19B). Excise the entire cornea (28-19C). Scoop out all uveal tissue contents of the eye with a evisceration spoon, curette or a periosteal elevator (28-19D). Pack the sclera for a few minutes to control bleeding. Excise a 5mm triangle of sclera from each side (28-19E), to help make the globe collapse. Rinse the inside of the globe with hydrogen peroxide.

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**Fig. 28-19 EVISCERATING AN EYE.**
A, conjunctiva already incised and undermined. Stab a #11 scalpel blade through the cornea into the anterior chamber. B, continue the corneal incision with scissors. C, excise the cornea completely. D, scoop out the contents of the eye with a curette. E, cut a triangular section out of each side of the hole in the sclera. F, inspect the inside of the eye to make sure that no choroid remains. After Galbraith J.E.K. Basic Eye Surgery, Churchill Livingstone 1979 p.84 Figs 9.17-24 with kind permission.
Make sure that no black choroid remains to avoid the risk of sympathetic ophthalmitis. Close Tenon’s fascia and conjunctiva with 6/0 nylon.

If you are operating for acute infection, leave it open to drain.

If there is a chance of getting an artificial eye, insert a plastic conformer shell.

Postoperatively, control bleeding by bandaging 2 eye pads firmly over the socket. Leave the dressing on for 2 days. Clean the lids and lashes bd, and put 2 drops of chloramphenicol 0.5% into the socket. You can insert an artificial eye after 4-6 wks.

**ENUCLEATION OF AN EYE (GRADE 3.3)**

**INDICATIONS.**
(1) A malignant intraocular tumour (retinoblastoma or melanoma) is an absolute indication.
(2) A blind, persistently painful eye, which is not infected (evisceration is an alternative).
(3) A penetrating wound, especially in the circumcorneal region, complicated by iridocyclitis, and entanglement of the iris, lens capsule, and vitreous. If you leave an eye like this, sympathetic ophthalmitis may follow in the other eye.

**ANAESTHESIA.**
A retrobulbar block using ≥6ml of lidocaine, combined with a VIIth cranial nerve block. Otherwise use ketamine or GA.

**METHOD.**
Incise the conjunctiva at its junction with the cornea, using fine-toothed forceps and fine scissors (28-20A). Cut around it, and undermine it back to the insertion of the extraocular muscles, about 8mm from the edge of the cornea (28-20B).

Push closed scissors through the conjunctiva to open up the plane between the conjunctiva and the globe. Open them to expose the sclera, anterior to the insertion of the rectus muscles.

Snip Tenon’s capsule (28-1C) between the insertions of these muscles. Pass scissors through the incision, until you have defined the muscle insertions. Slip a squint hook under the medial rectus muscle (28-20C), and pull it into view.

If you are going to put in an implant, lift the muscle and insert a mattress suture of chromic catgut through its belly, about 3mm from its insertion. Clamp its insertion, remove the squint hook, and cut the muscle with scissors. Separate each rectus muscle in the same way. Leave the stump of the medial rectus tendon a little longer, so that you have something with which to hold the globe.

**Fig. 28-20 ENUCLEATING AN EYE.**
A, incise the conjunctiva. B, undermine the conjunctiva for about 8 mm. C, slip a muscle hook under each rectus muscle, bring it forwards into the wound, and cut it. D, draw the eye forwards by pulling on the insertion of the medial rectus muscle. E, cut the optic nerve from the medial side. F, cut any remaining adherent tissue. G, suture the conjunctiva with absorbable suture. After Galbraith JEK Basic Eye Surgery, Churchill Livingstone 1979 p.81-2 Figs. 9.9-15 with kind permission.
Make the globe prolapse forwards out of the orbit by closing the arms of the speculum behind it, and pushing them backwards. If the eye is so enlarged, that it will not fit between the blades of the speculum, pull it forwards by holding the stump of the medial rectus muscle with a haemostat (28-20D). Pass a pair of curved scissors, with their blades closed, down the medial side of the orbit, feel for the optic nerve behind the eye, open the scissors, and cut it (28-20E).

If you are excising it for a malignant tumour, cut it as far posteriorly as you can, because it may have been infiltrated by tumour. Pull the eye forwards, and cut any tissue that remains attached to it (28-20F). Put a hot wet pack into the orbit, and press on it until bleeding stops.

If you are going to put in an implant, it will probably be a simple glass globe. Place it in the muscle cone, and Tenon's capsule, and suture the conjunctiva over it.

If there is no possibility of an implant, close the conjunctiva and Tenon's capsule separately with 5/0 absorbable (28-20G). Irrigate the socket with 0·5% chloramphenicol.

EXENTERATION OF THE ORBIT (GRADE 3.4)

INDICATIONS.
A malignant tumour of the orbit, often an advanced conjunctival carcinoma (28.15) or a retinoblastoma, which has penetrated the globe and caused proptosis.

ANAESTHESIA.
Use ketamine or GA. Have blood for transfusion available.

METHOD.
If the lids have been involved by tumour, suture their margins together with 3/0 silk. Use a #15 scalpel blade to cut round the margins of the orbit. If the lids are not involved, incise closer to the scleral margins, so as to save all, or some, of the skin from the lids to line the empty orbit.

Control bleeding from the upper inner margin of the orbit with diathermy, adrenaline solution or hydrogen peroxide. Incise the periosteum round the margin of the orbit, and reflect it as far posteriorly as you can. It is firmly adherent at the bony suture lines of the skull.

CAUTION!
(1) The bone on the medial wall of the orbit is very thin, so elevate the periosteum here with special care.
(2) The tumour may have eaten through the wall of the orbit, into the brain. If so, you may find it difficult to be sure of the anatomy.

The periosteum should strip easily until you reach the orbital fissures, and the nasolacrimal duct (6-4). Cut this.

Separate the palpebral ligaments, the trochlea (the pulley structure for the tendon of the superior oblique), and the inferior oblique muscles from the bone with the periosteum. Use curved scissors to cut the structures entering through the orbital fissures. Pull the contents of the orbit forwards, and cut the tissues at its apex with strong scissors as far back as you can.

Bleeding may be profuse. Remove the contents of the orbit quickly, and then control bleeding.

Turn the skin at the edges of the orbit back into it. Graft its raw surfaces with split skin, either now or as a secondary procedure in 10-14days. Gently pack the orbit. If you are not grafting it, dress it with disinfectant such as acriflavine wool, and apply a firm bandage.

DIFFICULTIES WITH DESTRUCTIVE METHODS FOR THE EYE

If the patient refuses to have a painful eye enucleated or eviscerated, consider injecting absolute alcohol behind it to destroy its sensory nerves. You can use any strength of alcohol, provided it is more than 50%, but you may need to repeat the injection if pain returns. Permanent relief is uncertain.

Retrobulbar alcohol is very painful for about 30secs, so use a retrobulbar block of lidocaine 1ml. Remove the syringe and needle. When the block is effective, put another syringe on the needle and inject 2mL of alcohol. The orbit will become severely oedematous for 10days. Add chloramphenicol eye drops qid for 1wk.

29.15 Conjunctival carcinoma

The conjunctiva is a thin transparent mucous membrane which covers the inner aspects of the eyelids and is reflected over the anterior part of the eyeball and ends at the limbus. Conjunctival epithelium is non-keratinized. It is continuous with the cornea at the limbus and the skin at the lid margins.

Squamous carcinoma of the conjunctiva used to occur mostly in the elderly but now is commoner among young HIV+ve patients, in whom it acts much more aggressively. It may arise on its own or from a pre-existing lesion like a pterygium (28.18). When small, it appears as a whitish lesion surrounded by a hyperaemic area, whose surface is irregular with small nodular parts. Initially it is mobile but as it invades, it becomes fixed.

Advanced cases are easy to diagnose, and may be associated with metastases to pre-auricular and submandibular lymph nodes.
MANAGEMENT

Small lesions you can completely excise; this is advisable as it can then provide histological confirmation as well as information on the invasiveness of the tumour.

Large lesions need an exenteration (28.14) with removal of regional lymph nodes. Radiotherapy is useful, but chemotherapy needs the help of an expert, especially if HIV disease is present.

28.16 Retinoblastoma

This uncommon, malignant, radiosensitive tumour of the embryonic cells of the retina usually presents in children <2yrs. It is bilateral in 25% of cases, and occurs in 2 forms, familial and sporadic. At first, the tumour enlarges within the eye; later it grows through the sclera, or chambers of the eye, to perforate the cornea. It can also spread through the optic nerve (where it may cause ulcer leading to perforation, n the VII, exenteration type II reaction and is less common if lmic division of the V
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If the tumour has extended through the globe, but not through the wall of the orbit, you should remove the globe and refer for radiotherapy, but the prognosis is so poor that a long journey to a referral hospital is not worthwhile. Consider chemotherapy.

If there is proptosis and a fungating mass, exenteration of the orbit (28.14) will remove the mass, but is unlikely to prolong life.

CHEMOTHERAPY.

In 35% of cases the following regime causes a partial response, and in 65% it prevents progression for a while. Use vincristine 1-4mg/m², doxorubicin 50mg/m² and cyclophosphamide 1g/m². Administer all drugs IV once every 21days.

28.17 The eyes in leprosy

Leprosy causes:

(1) Paralysis of the facial nerve, affecting the orbicularis muscle so that the eye does not close (lagophthalmos).
(2) Loss of sensation in the ophthalmic division of the Vth cranial nerve, which makes the cornea anaesthetic.
(3) An acute iritis (uncommon), which is usually associated with a type II reaction and is less common if the patient is taking clofazimine.
(4) A chronic iritis (common) causing atrophy of the dilator pupillae and a small unreactive pupil.

The combination of these lesions can have a devastating effect on the sight. An anaesthetic cornea prevents a patient noticing that there is something in the eye, or that it is dry. The blink reflex disappears, so that, even if there is still enough power in the VIIth cranial nerve to blink, it does not wash and wet the conjunctiva automatically. As a result, the cornea may be unprotected, especially during sleep, so that it may develop exposure keratitis, and ulcerate. If the centre of the cornea becomes opaque, sight is damaged. So warn of the danger of an anaesthetic cornea, and examine the eyes regularly.

To decide if the cornea has been damaged, look for superficial scars, and use fluorescein drops, or papers, to search for central staining. If the cornea is anaesthetic, the eye is at great risk. If there is lagophthalmos, but the cornea is not anaesthetic, he may have enough sensation to complain of discomfort or burning.

To find out if there is significant lagophthalmos, examine the closed eyes. If the cornea is completely covered, all is well. But if any part of it remains exposed, you should do something to protect eyesight. Several operations are possible. Tarsorrhaphy aims to reduce, or remove entirely, the gap between the closed eyelids. It has cosmetic disadvantages, but it does save sight, and it is not difficult, so you should be able to do it if you care for leprosy patients. A more effective procedure is transfer of the temporalis muscle.
28.18 Other eye problems

Here are some of the other eye problems you may meet.

If opening the eye is impossible, this is PTOSIS. This may be congenital, traumatic, acute as the result of an oculomotor palsy, HIV disease, or the result of myasthenia gravis.

If there is a small yellowish-white lump adjacent to the cornea in the region exposed by the palpebral opening, this is a PINGUECULA. It is harmless so give reassurance, though it may develop into a pterygium.

If there is a wing-shaped vascular thickening of the conjunctiva which grows on to the cornea, usually from the medial side, this is a PTERYGIUM. It is a wedge-shaped benign fleshy growth on the scleral conjunctiva that may grow onto the cornea; this is more frequent in increased UV exposure from outdoor activities. It is seldom serious. If vision is good, leave it. If it is advancing over the centre of the cornea and impairing vision, excise the pterygium carefully under LA off the conjunctiva, and dissect it off the cornea. Leave the sclera bare, and gently cauterize it. Up to 50% recur. Scarring can result in a squint.

If there is a worm under the conjunctiva (e.g. in the rainforest belt of West Africa), this is loiasis. Use albendazole 5mg/kg bd for 3wks.

If an old person has a complete or incomplete white ring encircling the cornea about 1mm within the limbus, this is ARCUS SENILIS. It is a sign of high blood cholesterol levels.

If the sclera are deep yellow, this is jaundice. There may be surgical correction possible if the cause is cholestasis (15.9).