The patient needed the trabeculectomy and you have done a great surgical job, but the pressure is now above the pre-operative level and you are feeling a failure. What are you going to do? What follows is a personal suggestion of questions to ask in developing a strategy.

**How long is it since the operation?**
If within three months of the operation, then routine post-operative management should be employed. During this period elevated intraocular pressure (IOP) develops in some patients. This occurs as early as four weeks in some African ethnic groups but is more typically at about six to eight weeks post-operatively. It is part of the wound remodelling process and resolves with good subsequent operative results. The difficulty lies in telling it apart from frank scarring and failure which has to be addressed with every tool available (page 73).

After three months, it is safe to say you have a fully-developed failure on your hands!

**Is there residual function in the trabeculectomy?**
It is always crucial to remember the purpose of the trabeculectomy, namely to prevent ongoing glaucomatous optic nerve damage. Occasionally the recorded IOP post trabeculectomy differs little from the pre-operative pressure and yet progression of glaucoma is halted. Nothing need be done in this case! All-day pressure readings post-trabeculectomy frequently show remarkably stable pressures, which is generally not the case when glaucoma is progressing. This may partly explain the apparent paradox.

Another option is to simply restart ocular hypotensive therapy. This is frequently sufficient, as evidenced by the ‘partial success’ figures in all trabeculectomy surgical trial outcomes. To optimise the topical therapy, remember to carefully explore the past records for the drug group that was best tolerated and most effective prior to the procedure.

‘As with any procedure, the results in your hands are what matter, not the minutaie of the technique compared to someone else’

**Is the environment working against success?**
The environment may be local concurrent ocular disease such as uveits, inflamed conjunctiva from allergy or blepharitis, lid pathology, or past ocular surgery. All of these should be treated first or a suitable strategy developed to minimise any subsequent procedure being at risk of failure from this secondary pathology.

The adverse environment may equally include external factors. Poor social situations for post-operative care include living alone and being unable to instil eye drops, or personal well-being having lower priority than child care and other responsibilities. Facilitate an improved environment wherever possible. Ask relatives to administer therapy. Help the patient to find a more satisfactory post-operative management environment. Consider depo-steroid use rather than intensive eye drop regimens. Consider review times that match the other time constraints for the patient.

**Is the trabeculectomy amenable to ‘resuscitation’?**
This involves careful examination with gonioscopy to ensure the sclerostomy is patent and free from internal obstruction. The conjunctival mobility, inflammation, and vascularisation should be noted. When examining the bleb, the most frequent site of failure is either scarring at the edge of the scleral flap with flat overlying tissues, or else encapsulation of the trabeculectomy flap with a raised profile.

Needling is not an exact science, but reports of outcomes suggest that greater success is achieved with a lower IOP immediately after needling, performing a course of needlings (i.e. more than one if required), and use of sub-conjunctival anti-scarring medications.2,3

**If needling is appropriate, how am I going to do it?**
As with any procedure, the results in your hands are what matter, not the minutaie of the technique compared to someone else. You want minimum complications and maximum success!

A good operative field is vital. Use of adequate anaesthesia (I use topical anaesthetic followed by sub-conjunctival lignocaine 2%), a speculum, vascular constriction (I use phenylephrine because it is handy in the clinic), and povidone iodine (or a similar preparation) make a world of difference.

Needling at the slit lamp has the advantage of immediate assessment of pressure effect and easy review at regular intervals after the procedure (more difficult when done in the middle of a busy operating list). Needling in the operating theatre allows you to proceed immediately to a more complex surgical intervention if necessary. Needles of size 25–30 g have been reported, as have MVR blades; some practitioners report being more adventurous in their choices. (I use a 30 g needle at the slit lamp and micro MVR in theatre.)

Most practitioners enter the conjunctiva at a distance from the area of scarring to be perforated and use a ‘slicing’ action to open a good-sized drainage channel (Figure 1). Plan your approach carefully and only work on the area of obstruction to flow.

The most common complication when
Medical treatment of open-angle glaucoma

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Most guidelines that exist for the treatment of primary open-angle glaucoma (POAG) — such the National Institute for Health and Clinical Excellence (NICE) guidelines (page 47) — recommend medicine as initial therapy, though laser treatment may also play a role.

None of these guidelines are based on research done in Africa, however, and there also is no evidence yet on the relative effectiveness of different glaucoma medications in African populations.

For medicines to be effective at controlling POAG, they must be used every day for the remainder of the patient’s life.

This means the medicine must not only be clinically effective, it must also be available, of good quality, affordable, and well tolerated by the patient.

Keep the following in mind when choosing medical treatment for your patient.

1. **The cost:** can the patient afford the recommended medication in the long term?
2. **Availability:** will the drug(s) always be in stock, and what are the consequences to the patient of any stock-outs?
3. **Quality:** for drugs such as latanoprost, will the cold chain be maintained? Can the patient refrigerate the medicine?
4. **Discomfort:** is there any discomfort associated with the medicine that will discourage the patient from continuing with treatment?
5. **Follow-up:** will the patient be able to attend regular follow-up appointments?

Choosing the right drugs

There are five main groups of glaucoma drugs, each acting in a different way to reduce IOP:

- **prostaglandin analogues** (bimatoprost, latanoprost, and travoprost) increase uveoscleral outflow
- **beta-blockers** consist of two main groups: selective (betaxolol) and non-selective (timolol, levo-bunolol), both of which decrease aqueous production
- **alpha-2 adrenergic agonists** (apraclonidine, brimonidine) decrease aqueous production and increase uveoscleral outflow
- **carbonic anhydrase inhibitors** decrease aqueous formation and can be applied topically (brinzolamide, dorzolamide), or systemically (acetazolamide, methazolamide)
- **parasympathomimetics** (pilocarpine, carbacol) increases aqueous outflow

Continues overleaf ➤

References


A selection of glaucoma drugs available in Nigeria