The control of conjunctival fibrosis as a paradigm for the prevention of ocular fibrosis related blindness. “Fibrosis has many friends”

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Abstract
The processes involved in ocular fibrosis after disease or ocular tissue injury, including surgery play an important part in the development or failure of treatment of most blinding diseases. Ocular fibrosis is one of the biggest areas of unmet need in ophthalmology. Effective anti-scarring therapies could potentially revolutionise the management of many diseases like glaucoma worldwide. The response of a quiescent or activated conjunctiva to glaucoma surgery and aqueous flow with different stimulatory components and the response to different interventions and future therapeutics is a paradigm for scarring prevention in other parts of the eye and orbit. Evolution in our understanding of molecular and cellular mechanisms in ocular fibrosis is leading to the introduction of new and re-purposed therapeutic agents, targeting a wide range of key processes. This review provides current and future perspectives on different approaches to conjunctival fibrosis following glaucoma surgery and highlights the challenges faced in implementing these therapies with maximal effect and minimal side effects.

Introduction
Ocular fibrosis after disease or injury, including surgery, is a complex biological process and features in many blinding ocular diseases, which result from autoimmunity (e.g. pemphigoid and fibrosis in dysthyroid eye disease), angiogenesis and wound healing (e.g. glaucoma surgery and capsular opacification). Fibrovascular scarring features in numerous ocular conditions such as, diabetic retinopathy, macular degeneration, proliferative vitreoretinopathy, retinopathy of prematurity, and neovascular glaucoma, and is a consequence of the underlying inflammatory or hypoxia-driven neovascularization. Therefore, prevention of underlying drivers is the most appropriate therapeutic approach to preserve visual function. Furthermore, scarring is a major potential obstacle to regenerative therapies such as cellular transplantation [1].

Collectively, ocular fibrosis results in a huge amount of ocular morbidity worldwide. Many of these conditions were discussed during the XLIX Cambridge Ophthalmological Symposium. Discussion during the symposium drew on lessons learnt from aberrant wound healing pathogenesis and strategies to lessen scarring after GFS, and ocular fibrosis in general. Improved understanding of the molecular and cellular mechanisms of wound healing processes and the mode and mechanism of action of modulating agents, has led to the identification and modulation of potential therapeutic targets, and more effective use of existing agents. This paper aims to provide an overview of some of the approaches used to modulate conjunctival fibrosis after GFS and how these approaches can be applied to other forms of ocular fibrosis.

In glaucoma surgery the intent is to make a drainage fistula from the anterior chamber to the sub-Tenon space to allow aqueous outflow and lower the intraocular pressure (IOP). In this paper we will concentrate on glaucoma surgery involving fistulas draining externally to the sub-Tenon space. The healing response and the fibrosis that occurs are the primary determinants of the final IOP. This is because conjunctival fibrosis determines the hydraulic conductivity of the conjunctiva which controls the fluid flow through the tissue. Intraocular pressures between 10 to 14 mmHg are associated with minimal fibrosis of the conjunctiva, and slower rates of visual field deterioration [2].

Several different stimulatory components arise from glaucoma surgery which activate quiescent conjunctiva. Conjunctival fibrosis can result in the failure of conventional bleb-forming glaucoma filtration surgeries. New drainage implants employing different outflow pathways including subconjunctival, trabecular meshwork/Schlemm’s canal and suprachoroidal routes of drainage are used clinically. All of these drainage implants are eventually limited by fibrotic reaction around the device regardless of its mode of action. Despite advances in glaucoma surgical techniques, devices and post-operative care, fibrosis remains the principal impediment to long-term control of IOP following surgery.
Conjunctival wound healing responses after glaucoma surgery

Like all wound healing, conjunctival wound healing involves a complex interplay between growth factors, cytokines, receptors and enzymes and cross-talk between different cell types. The early phase of conjunctival wound healing following glaucoma surgery involves leakage of plasma proteins (fibrinogen, plasminogen and fibronectin) and blood cells (platelets, polymorphonuclear neutrophils and red blood cells) from disrupted vasculature. Conversion of fibrinogen to fibrin results in fibrin clot formation and blood coagulation. The combination of fibrin clot formation and released proteins, lead to migration and attraction of neutrophils, macrophages, and lymphocytes to the surgical site during the inflammatory phase [3].

Pleiotropic genes, involved in conjunctival fibrosis include, but not limited to, transforming growth factor-B (TGF-β), vascular endothelial growth factor (VEGF), platelet derived growth factor (PDGF), fibroblast growth factor (FGF) and connective tissue growth factor, which contribute to wound healing regulation. Aqueous contains a number of growth factors, including transforming growth factor (TGF)-β, which is a main mediator of fibrosis during wound healing [4, 5]. The later phases of conjunctival fibrosis can be subdivided into inflammation, proliferation, and tissue remodelling (see Fig 1). During the proliferation phase, activated fibroblasts and endothelial cells migrate to the site of injury, and granulation tissue formation and angiogenesis ensues. Activated Tenon fibroblasts proliferate and lay down extracellular matrix. During tissue remodelling, blood vessels regress and fibroblasts adopt a myofibroblast phenotype which induce cross-linking of collagen type 1 and elastin resulting in scar formation.
Fig 1. Sequence of events after glaucoma surgery. Patient risk factors should be considered and mitigated against to modulate conjunctival fibrosis after surgery (A). Surgical creation of the aqueous outflow pathway and suture material results in the release of blood constituents and release of cytokines and growth factors from the aqueous, Tenons and sclera (B). In addition to these factors, fibrin clot initiates infiltration of macrophages and polymorphic neutrophils (PMN) to the bleb site (D). Activation, elongation, and proliferation of Tenon fibroblasts ensues with significant deposition of extracellular
matrix (ECM), which persists due to stimulatory tractional forces and growth factors (E). Apoptosis of fibroblasts and vascular endothelial cells block outflow of the aqueous humour resulting in surgical failure due to scarring (F). (Adapted with permission).

The increase in the number of fibroblasts during the proliferative phase, starts as early as two days post-operatively, and provides an initial advantage to maintain the integrity of the conjunctiva and protect the eye against infection. This reflects an evolutionary pressure towards tissue fibrosis in response to injury. The degree of tissue damage and the host response in terms of severity and duration of inflammation, determines the extent of conjunctival scarring. Moorfields Safer Surgery System, is a modified surgical technique, which aims to minimise the insult and modulates the effects of antimetabolites [6]. The extent of conjunctival fibrosis after glaucoma filtration surgery (GFS) will vary between individuals, as it is influenced by risk factors such as, pre-existing inflammation, duration and type of eye drop therapy, and a previous history of surgery [7]. Other drivers of fibrosis after GFS include suture material, aqueous humour derived growth factors, mechanical transduction from aqueous fluid flow, signalling molecules released from extracellular matrix storage sites, myofibroblast trans-differentiation leading to further matrix deposition [8]. Post trabeculectomy, the Moorfields bleb grading system [9] is a useful tool to predict long term IOP outcome. The grading system assesses inflammation and effective application of anti-metabolites [10].

**The fibroblast: origins, activation, differentiation, and proliferation**

Following glaucoma filtration surgery, fibrosis is mediated primarily by Tenon fibroblasts and their precursors, myofibroblasts. Myofibroblasts are the key effector cells resulting in surgical failure. Myofibroblasts are derived from either one or more origins irrespective of the nature of the initial injury. The exact origin of myofibroblasts during conjunctival fibrosis is debated, and beyond the scope of this paper. Nevertheless the diversity of myofibroblast progenitor cells is relevant in the development of novel therapies. In brief, myofibroblasts involved in conjunctival fibrosis might originate from beyond locally residing cells in the scleral, conjunctiva and Tenons, such as circulating bone-marrow derived fibroblasts (see Fig 2). The relative contribution of the different origins of progenitor conjunctival myofibroblasts is unknown due to the lack of a specific myofibroblast cell marker. Myofibroblasts present microfilament bundles expressing α-smooth muscle actin (α-SMA), responsible for wound construction. Myofibroblasts deposit an abundant amount of extracellular matrix (ECM), produce crosslinking enzymes, and inhibitors of ECM, degrading metalloproteinases. Persistent activation of myofibroblasts and mechanical stress such as that found in a tight encapsulated bleb potentiates profibrotic factors. Myofibroblast contractility attracts macrophages, which modulate ECM composition and stiffening [11].
Figure 2. Different origins of myofibroblast precursors and stages of myofibroblast differentiation. 

SMC smooth muscle cell, TGF-β transforming growth factor-beta

**Therapies targeting fibrin clot**

Fibrin clot formation is an important step of the wound healing process. Fibrinolytic agents, such as tissue plasminogen activator, were thought to be useful to dissolve fibrin and blood clots after glaucoma surgery. Urokinase is a thrombolytic agent that could be used as an alternative to tissue plasminogen activator, and intracameral urokinase injection reduced IOP after glaucoma surgery in the short term [12]. Prolonged bleeding is the main side effect that deters the use of these agents, and in addition fibrin breakdown products can stimulate the proliferation of fibroblasts. Treatment with saratin, an anti-thrombotic and anti-fibrotic agent, in a rabbit GFS model, improved bleb survival when administered as two post-operative subconjunctival injections when compared to topical saratin treatment alone [13].

**Therapies targeting inflammation**

Reducing inflammation is a standard paradigm for preventing scarring after surgery or injury as chronic inflammation is associated with much more fibrosis. Broad-spectrum anti-inflammatory drugs are standard therapy post-operatively, and are usually administered as topical steroids. The use and tapering of topical steroids such as prednisolone acetate 1% are an important feature of the post-operative care to modulate inflammation and maintain adequate intra-ocular pressure control. Steroids can be started prior to surgery if the eye is inflamed. The use of topical non-steroidal anti-inflammatory agents (NSAIDs) after glaucoma surgery remains controversial, as these agents may be associated with a greater incidence of wound problems such as bleb leakage [14]. Nonetheless, where inflammation persists despite steroid use, NSAIDs may be a useful adjunct.

Studies have investigated other methods of administration to enhance steroid bioavailability and effectiveness in dampening the inflammatory response. A randomised controlled trial showed intra-operative sub-Tenon triamcinolone during augmented trabeculectomy surgery in patients with secondary glaucomas did not improve success rate or reduce complications [15].
In another study with a small sample size, intracameral triamcinolone showed no benefit after heterogenous glaucoma operations with or without use of mitomycin-C (MMC) [16].

Synthetically engineered macromolecules such as dendrimer glucosamine and cyclodextrins have been developed to enhance steroid delivery in experimental studies. Encapsulation of drugs in colloidal carriers liposomes and nanoparticles can enhance permeation across cell membranes and prevent enzymatic degradation [17]. Also, ocular implants are being investigated as alternatives to topical or injectable steroid drug delivery systems. Subconjunctival sustained-release dexamethasone (Ozurdex) have been piloted as an adjunct to trabeculectomy [18]. Ozurdex is comprised of a biodegradable co-polymer of lactic acid and glycolic acid so does not require surgical removal.

Anti-inflammatory drug loaded contact lenses have been developed to increase residence time of steroids in the tear film. Contact lenses fabricated from poly-2-hydroxyethyl methacrylate hydrogels with drug molecules entrapped into a nanoparticulate system can provide controlled drug release over days [19]. Due to increased residence time, drug permeation through the tissue can offer prolonged exposure. The delivery rate can be further modified by altering the loading dose in the hydrogel. However, this drug delivery system has yet to be tested in the context of glaucoma filtration surgery post-operative care.

One would assume favourable outcomes by targeting 'upstream' immune responses. However, broad-acting immunosuppressive therapies have proved sub-optimal to date, and have not entered routine use for glaucoma surgery. Calcineurin inhibitors such as cyclosporine, tacrolimus and sirolimus, are immunosuppressive agents that reduce the production of interleukin-2 (IL2) and IL2 receptor expression. Following GFS, topical cyclosporine, had no effect on IOP, although ocular surface disease improved [20]. A combination of therapies targeting several different pathways or the development of novel agents addressing multiple components of fibrosis following glaucoma surgery appear to be required.

**Therapies targeting proliferation**

Antimetabolites, 5-fluorouracil (5-FU) and mitomycin-C (MMC) are conventionally used at the surgical site to limit conjunctival scarring following GFS. While these agents have been shown to improve clinical outcomes, they do so in a non-selective manner inhibiting the proliferation of subconjunctival Tenons fibroblasts. Consequently, use of antimetabolites is associated with a significant side-effect profile that can lead to hypotony, bleb leakage, blebitis, endophthalmitis, and vision loss.

5-FU is an effective inhibitor of Tenons fibroblast growth by antagonizing pyrimidine metabolism inhibiting DNA synthesis and ultimately resulting in cell death [21]. Use of 5-FU at high concentrations, however, often leads to corneal epithelial toxicity and apoptosis, since 5-FU is toxic to all actively replicating cells. Intraoperative or post-operative 5-FU significantly reduces the risk of trabeculectomy failure but causes increased rates of wound leakage, hypotony and epitheliopathy compared to trabeculectomy alone [22-24].

MMC exhibits antiproliferative effects by acting as a DNA cross-linking alkylating agent that can interfere with cells at any stage of the cell cycle, inhibiting DNA replication, cell mitosis and protein synthesis. MMC increases the success rate of GFS, mainly through inhibition of proliferation of fibroblasts. MMC is more potent than 5-FU, inhibiting proliferation for a longer period [25, 26]. Meta-analysis of eleven studies provided some evidence that favoured MMC to 5-FU in lowering IOP at 1-year post trabeculectomy. Less epitheliopathy and hyphaema was observed with MMC [27]. The method of applying the MMC is also critical and has a huge bearing on the long-term bleb appearance and complications, and these general physical principles of drug applications are also very important in the prevention of scarring [28].
Beta-radiation only has superficial tissue penetration and can be used over the conjunctival bleb to arrest cell growth rather than cause widespread cell death. Tenons fibroblasts exposed to beta radiation express high levels of p53 to prevent cell replication in the presence of DNA damage, decreasing ECM production [29]. β-radiation can be applied locally, very rapidly (in less than one minute) and accurately, making it a useful anti-fibrotic option in GFS. The current 1000cGy β-radiation dose is based on cell culture studies [30] and is much lower compared to dosages historically used to treat pterygium, which varied from 2000 to 6000 cGy. The protocols to apply these higher doses of β-radiation had a poor safety profile in pterygium management [31]. In glaucoma filtration surgery β-radiation has proven to be an effective antifibrotic therapy in high risk patient groups such as patients from West African descent. A meta-analysis including four studies showed intraoperative beta irradiation lowered the risk of surgical failure compared to trabeculectomy alone [32]. However, the β-radiation group had a higher risk of cataract [32], but this may have been associated with lower pressures which is less often seen with newer GFS such as the Moorfields Safer surgery System [6]. When β-radiation was compared with intraoperative 5-FU, there was no difference in risk of failure or complication following phacotrabeculectomy [33]. More recently, randomised trials have shown that β-radiation with MMC are superior to MMC alone [34]. In a South African study, eyes receiving β-radiation were 5.55 times more likely to have an IOP <16mmHg on no medical treatment compared to eyes receiving MMC [35]. New and improved radiation delivery systems and sources should allow treatment times as short as 30 seconds.

The use of an adenoviral delivery system may also be considered feasible. The p21 gene is an important downstream effector of the p53 gene, which is a tumour suppressor gene that arrests cell cycle progression. A recombinant adenovirus carrying human p21 gene inhibited the proliferation of Tenon’s capsule fibroblasts, prevented conjunctival scarring, and significantly reduced IOP in animal models of GFS [36].

**Therapies targeting growth factors**

Stimulatory growth factors and cytokines, released at the site of surgery can both amplify and reduce inflammation during the wound healing process (Fig 3). Identifying therapeutic targets can be challenging due to the pluripotency and redundancy of these different signalling molecules [37]. TGF-β is instrumental in numerous processes of fibrosis, such as stimulating inflammation, migration of cells, loss of cell-cell adhesions, epithelial-mesenchymal transition (EMT) into myofibroblast phenotype and modulates ECM deposition.

**Transforming growth factor-β**

TGF-β2 is the predominant intraocular TGF- β isoform associated with ocular fibrosis such as proliferative vitreoretinopathy, posterior capsular opacification, and fibrosis after GFS [38-41]. Lerdelimumab, a monoclonal antibody to TGF-β2 was trialled to prevent the progression of fibrosis in primary trabeculectomy. The dose level and treatment regimen were derived from rabbit studies [42]. There was no difference between Lerdelimumab and placebo in preventing surgical failure in a phase III human trial [43]. The use of a monoclonal antibody directed solely against TGF-β2 may represent too narrow of an approach since other isoforms of TGF-β are also involved. The very short subconjunctival drug half-life and the mode of delivery may not have been optimal to neutralise TGF- β2, and the absence of neutralisation of the other isoforms may have also been a disadvantage. A pan-TGF-β antibody, Fresolimumab, which neutralises all three main isoforms of TGF-β, may have potential applications in the treatment of a broad range of medical conditions [44]. To our knowledge Fresolimumab has not been evaluated in the context of ocular fibrosis and may not be appropriate in the context of corneal-related fibrosis which requires a more targeted approach to downstream TGF- β signalling pathway mediators, in order to maintain normal corneal wound healing and corneal transparency [45].
RNA sequencing analyses have identified long non-coding RNAs (LncRNAs) and short RNA molecules (miRNAs) involved in conjunctival fibrosis, as new approaches to targeting the TGF-β signalling pathway. MicroRNAs are non-coding molecules and function to silence gene expression or targeted degradation of messenger RNAs. MicroRNAs regulate many biological processes, including tissue fibrosis and their function is likely to be tissue and species specific [46, 47]. A study performed to identify miRNAs that inhibit the transcription of profibrotic proteins in Tenon fibroblasts showed that overexpression of miRNA-29b, targets profibrotic TGF-β signalling pathway, inhibited production of collagen type I-α1 and protected the conjunctiva from fibrosis [48]. Data from other specialities suggest, inhibition of miRNA-21 and overexpression of miRNA-29 seem to be promising antifibrotic approaches [49, 50]. Potential drugs in the development pipeline which could be repurposed include an inhibitor of miRNA-21, RG-012 and is under evaluation for the treatment of Alport syndrome. There is also a miRNA-29 mimic, MRG-201, that is being assessed to treat cutaneous scleroderma [51].

LncRNAs do not code for proteins instead they interact with chromatin to modify gene expression. LncRNAs induced by TGF-β/Smad3 signalling have been identified in human tenon fibroblasts obtained from patients who had undergone GFS [52]. Although evidence is limited, lncRNAs are likely to have a function in conjunctival fibrosis and could present new areas for future investigation [53-55].

Vascular endothelial growth factor

Angiogenesis is a crucial process in wound closure by enabling inflammatory cells and fibroblasts to migrate to wound site, and is mediated by vascular endothelial growth factor (VEGF) [56, 57]. VEGF is one of the most potent vasodilators known and may facilitate the extravasation of blood proteins and cytokines. VEGF among other growth factors is found to be upregulated in the aqueous following GFS [58]. There remains uncertainty whether anti-VEGF agents such as bevacizumab and ranibizumab are as effective in preventing fibrosis following trabeculectomy as MMC or sham control [59]. The optimal dose and frequency of applications of anti-VEGF in the context of GFS remains unclear. Better bleb morphology and reduction in bleb vascularisation has been observed following intravitreal ranibizumab in combination with intraoperative topical MMC [60]. Subconjunctival bevacizumab has been used successfully to rescue failing filtering blebs that exhibit neovascularization [60, 61]. Anti-VEGF treatments have a role in the treatment of neovascular glaucoma, a disease where angiogenesis is the underlying pathology [62, 63].

Platelet-derived growth factor

Platelet-derived growth factor-B (PDGF-B) is another growth factor that can activate certain ocular fibroblasts. Aptamers are nucleic acid-based macromolecules that can recognise, bind, and block PDGF-B with high affinity and specificity. Two aptamers, ARC126 and ARC127, have been tested in a rabbit model of proliferative vitreoretinopathy [64], and might also be useful in glaucoma surgery.

There are also potential kinase inhibitors that can be used for antifibrotic therapy. Nintedanib inhibits PDGF receptor, VEGF receptor and FGF receptor tyrosine kinases. Nintedanib is FDA approved as an orally administered medicine to treat idiopathic pulmonary fibrosis (IPF). An in vitro study using human Tenons fibroblasts showed Nintedanib inhibits proliferation and motility induced by TGF-β1 [65]. Pirfenidone is also an orally administered medicine used to treat IPF. While the mode of action of Pirfenidone has not been firmly established, it appears there is reduced fibroblast proliferation and collagen production [66]. Both Nintedanib and Pirfenidone may be two of the few drugs licenced for fibrosis because of the multiple possible
mechanisms of action given the many cross covering biological systems responsible for wound healing and fibrosis.

**Tissue remodelling – therapies targeting matrix, collagen, and enzymes**

Mechanical stress induces the protomyofibroblast phenotype and expression TGF-β, which is critical for the final myofibroblast transformation. In addition to mechanical stress and TGF-β, integrin signalling are also necessary for the maintenance of the myofibroblastic phenotype [67]. TGF-β upregulates connective tissue growth factor (CTGF) expression, which is another important fibrogenic cytokine that facilitates a persistent pro-fibrotic state. Drug targeting either CTGF or TGF-β signalling can maintain filtering bleb function by minimising scarring [68-70]. Blocking of CTGF using an antibody against TGF-β2 receptors reduced conjunctival scarring in ocular cicatricial pemphigoid [71].

The expression of matrix metalloproteinases (MMPs) varies throughout wound healing as they contribute to ECM remodelling by degrading extracellular matrix proteins. MMPs mediate inflammation by stimulating fibroblasts, altering endothelial permeability, and activating VEGF [56, 72, 73], and are known to be upregulated in the fibroblasts of the bleb walls surrounding glaucoma implants [74]. Ilomastat, a broad-spectrum MMP inhibitor, can reduce subconjunctival scarring after experimental GFS [75]. A cyclodextrin-Ilomastat-complex suspension eye drop could permeate through conjunctival tissue and therapeutic concentrations were present in the anterior chamber, conjunctiva and sclera in a preclinical in vivo study [76]. Doxycycline was found to inhibit MMPs but to a lesser extent than MMC in a rabbit model [77]. Sustained drug delivery is crucial to maintain beneficial drug effect throughout the ECM remodelling phase of wound healing.

Other enzymes such as lysyl oxidase (LOX) and lysyl oxidase-like (LOXL), crosslink collagen and elastin in the ECM, which leads to fibrosis [78]. A monoclonal antibody to LOXL-2 reduced both inflammation as well as fibrosis in vivo [79]. Elevated TGF and LOXL-2 levels in Tenon’s tissue at the time of GFS strongly correlated with surgical failure at one year, suggesting LOX-2, could be a promising therapeutic target for reducing scar formation after glaucoma surgery [79, 80].

Rho and Rac1 are small GTPases that play a critical role in the regulation of actomyosin cytoskeletal organisation and cell motility. A Rac1 inhibitor, efficiently inhibited fibroblast-mediated matrix contraction, reduced MMP-1 expression in conjunctival tissue and did not cause any significant toxicity [81]. The Rho-associated protein kinase (ROCK) inhibitor, also reduced fibroblast-mediated matrix contraction in vitro, and improved bleb survival in experimental GFS [82].

Activated myofibroblasts are the key effector cell in conjunctival fibrosis and scar formation as previously described. Myocardin-related transcription factor/serum response factor (MRTF/SRF) transcription pathway plays an important role in myofibroblast activation in fibrosis and is a key upstream regulator of MMP expression in ocular fibrosis. MRTF inhibition can be achieved through siRNA-mediated silencing or pharmacological inhibition. A few small molecule inhibitors, have been reported to be more effective in reducing MRTF/SRF-regulated gene transcriptional signalling than ROCK inhibitors [83]. Sustained release liposome MRTF/SRF pathway inhibitors have shown efficacy in preventing conjunctival fibrosis in an established rabbit model of GFS [84, 85].
Fig 3. The phases of conjunctival wound healing include clotting, vascular response, inflammation, fibroplasia and tissue remodelling/scarring [37]. Some of the mediators and factors involved in each phase are listed in the balances. PDGF platelet-derived growth factor, VEGF vascular endothelial growth factor, TGF-β transforming growth factor-beta, SAP serum amyloid P, anti-COX2 anti-cyclooxygenase 2.

Delivery of anti-fibrotic therapies

Application of antifibrotic agents at the bleb site is hampered by rapid clearance from the subconjunctival space due to aqueous outflow resulting in conjunctival absorption and systemic clearance. Mediating the localised healing response after surgery for extended periods of time may best be achieved by prolonging the presence of a wide-spectrum anti-fibrotic agent in the subconjunctival space. Developing long acting dosage forms for subconjunctival administration require that material-tissue interactions are optimised to avoid additional localised inflammation due to foreign body responses. Sustained release implants and in situ forming gels have been described for intraocular and subconjunctival use which may potentially be developed for administration after glaucoma filtration surgery or drainage device implantation. Strategies for the administration of anti-fibrotic therapies including sponges, hydrogels, injections, implants, gene therapies, cell-based therapies are shown in Fig 4 and Table 1.

Local accumulation in the sub-tenon space can be achieved with the use of particulate associated formulations (i.e. nano or micro sized colloids). Particulate carriers have been described that bind to cell-surface receptors on myofibroblasts and facilitate drug internalisation which is triggered by binding to the receptor. Theoretically such strategies may increase focal conjunctival antifibrotic activity with lowered unwanted side effects and toxicity to other ocular tissues. Following glaucoma surgery, there is overexpression of LDL receptor in activated Tenons fibroblasts [86]. Engineered LDL-MMC-nanoparticles may provide a novel target for drug delivery systems that specifically bind to LDL receptors mainly in activated fibroblasts. This may achieve highly selective targeting, smaller drug dose requirement, better bioavailability, and reduced cellular toxicity to quiescent cells [86].
Fig 4. Localised treatment options, some which are based on the development of subconjunctival specific dosage forms that can prolong the local exposure of an anti-fibrotic agent at the bleb site.

<table>
<thead>
<tr>
<th>Therapeutic agent</th>
<th>Mode of delivery</th>
<th>Effect</th>
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<tbody>
<tr>
<td>Mitomycin-C, 5-Fluorouracil</td>
<td>Sponge, Subconjunctival injection</td>
<td>Decrease fibroblast proliferation, apoptosis</td>
</tr>
<tr>
<td>Beta-radiation</td>
<td>Local irradiation of tissues</td>
<td>Decrease fibroblast proliferation</td>
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<tr>
<td>Local anaesthetic</td>
<td>Sub-Tenon’s injection, Subconjunctival injection</td>
<td>Might decrease pain and local inflammation</td>
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<tr>
<td>Steroid</td>
<td>Topical, Ozurdex implant</td>
<td>Decrease IOP, Prolong bleb survival</td>
</tr>
<tr>
<td>Nonsteroidal anti-inflammatory drug (NSAID)</td>
<td>Topical</td>
<td>Decrease IOP but greater incidence of wound leaks and conjunctival retraction</td>
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<tr>
<td>Anti-VEGF (Bevacizumab)</td>
<td>Subconjunctival injection, Intracameral injection</td>
<td>Controversial: some studies report improved bleb survival and lower IOP. Decrease needling interventions after trabeculectomy.</td>
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Table 1. Currently available treatment modalities to modulate wound healing during and after glaucoma surgery

**Surgical principals and modifications**

Simple changes in surgical techniques and the application of antimetabolite treatment have significantly improved the success rate and reduced the complication rate of glaucoma surgery [6] such as minimising tissue damage and application of tailored dose of antifibrotic agent. The principle of using physical spacers to attenuate mechanical stress might also help to maintain the subconjunctival pocket after GFS. Spacers using human amniotic membrane, perfluoropropane gas, or sodium hyaluronate, have been reported to increase bleb survival and to lower intraocular pressure [87]. More research is needed to develop better biocompatible materials that would lead to less scarring and better post-operative results after glaucoma surgery.

Use of ocular anti-hypertensives after surgery may play a role in conjunctival fibrosis and tissue remodelling. After glaucoma drainage device surgery, the use of aqueous suppressants is preferable to prostaglandin analogues to treat raised intraocular pressure [88, 89]. In experimental models prostaglandin analogues induced collagen contraction [90], upregulated
interleukin-2 and MMP-9, while aqueous suppressants decreased expression of α-SMA, reducing the transformation of fibroblasts to myofibroblasts [91]. Aqueous suppressants may attenuate growth factors derived from the aqueous and reduce the mechanical stress from aqueous pressure in the bleb.

Needling with 5-FU or MMC augmentation, can be performed to rescue failing blebs particularly in high risk cases. Bleb failure can occur early or late following glaucoma filtration surgery. Fibrosis can occur at the level of the conjunctival, episcleral or internal ostium. Needling was most successful soon after trabeculectomy, but rescue of late failed trabeculectomy bleb has a lower likelihood of success [92]. Therapies targeting early tissue remodelling phases may influence long term post-operative outcomes, as it is the overall accumulation of fibrotic tissue that leads to pressure rise and ultimate surgical failure.

Concomitant surgery may worsen fibrosis such as the combination of phacoemulsification with glaucoma drainage surgery. A greater amount of postoperative inflammation and flare have been observed after cataract surgery compared to trabeculectomy for up to 6 months surgery. Inflammation associated with cataract surgery is likely to be as a result of crystallin lens protein, ultrasound energy, and the speed and volume of intraocular fluid irrigation during surgery. The consequence is an upregulation of pro-fibrotic cytokines and growth factors in the aqueous humour and increased risk of bleb failure. The extent of time these profibrotic proteins are upregulated should be borne in mind when prescribing post-operative anti-inflammatory agents in cases of phacotrabeculectomy and sequential surgeries, (cataract surgery after trabeculectomy), if excessive scarring is to be reduced [93]. The earlier cataract surgery is performed within 6 months, the greater the risk of bleb failure [94].

**Can fibrosis be reversed?**

Direct *in vivo* reprogramming of myofibroblasts into fibroblasts has potential as a strategy for regeneration in fibrosis-related conditions. Hepatic fibrosis has been reversed by using transcription factor induction, directly inducing myofibroblasts to transform into hepatocytes using a lentiviral vector in mouse models of chronic liver disease [95]. For this regenerative activity to be achieved in glaucoma surgery, transcriptional mechanisms underlying myofibroblast phenotype would need to be fully elucidated, and the development of lineage tracers that are tissue/cell specific. Presently, there is no evidence demonstrating reversal of conjunctival fibrosis but the absence of evidence does not make it impossible. Prevention of ECM deposition is more likely to result in the restitution of conjunctival architecture. Therefore, it is important to identify patients who may be more susceptible to aberrant fibrosis at an early stage.

**Summary and perspective**

Conjunctival wound healing modulation continues to receive interest as it holds the key to glaucoma filtration surgery attaining and sustaining optimal intraocular pressure (10 – 14 mmHg) and preventing glaucoma progression associated with higher pressures. New treatments will arise from a better understanding of tissue specific associated molecular, cellular, and biomechanical processes. Innovative solutions that can directly and indirectly modulate inflammatory and fibrotic processes at the optimal time, dosage and locality will result in more favourable outcomes. A combination of treatments or sequential treatments may be more efficacious than a single agent, particularly in individuals with a high risk of scarring. There are many areas for future investigation yet to be explored, in terms of novel or repurposed anti-fibrotic agents and sophisticated drug delivery systems. Genetic manipulation of cells involved in wound healing or *in vivo* reprogramming of myofibroblast may be future methods used to treat ocular fibrosis. Ultimately, the prize is great. Glaucoma is increasing exponentially with the rapidly ageing populations around the world, and a management plan that can cope with the
scale of demand should involve surgical therapy as accessible and successful as cataract surgery.

We know from clinical evidence that pressures around the 10 mmHg mark are associated with minimal optic nerve disease and minimal visual field progression over a decade. Micro-devices now allow a degree of surgical control and pressure control to achieve low pressures around 10 mmHg safely and rapidly. This means the 10 10 10 target we have proposed for glaucoma treatment (10 minutes to conduct the treatment to achieve 10 mmHg that lasts 10 years) would revolutionise the treatment of glaucoma could become a possibility as long as we can control wound healing and fibrosis over the period. With the considerable advances that have been made in understanding and modulating fibrosis, this exciting prospect is in sight.

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