Prospective trial of intraoperative mitomycin C in the treatment of primary pterygium

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Abstract

Aims—A prospective, randomised, double blind, placebo controlled study of intraoperative mitomycin C as adjunctive treatment of primary pterygium was conducted.

Methods—A total of 66 eyes of 54 patients with primary pterygium were treated with excision, with or without a single intraoperative application of mitomycin C (0·1 mg/ml for 5 minutes) to evaluate the efficacy and toxicity of this adjunctive treatment. The mean follow up was 14·1 months (range 12–23 months).

Results—Of the 36 eyes that underwent simple excision, 14 (38·8%) exhibited recurrences whereas only one of 30 eyes (3·3%) treated with excision and intraoperative application of mitomycin C had recurrence (p=0·0006). Neither serious ocular complications nor systemic toxicity were noted in the mitomycin C treated group.

Conclusion—Intraoperative mitomycin C appears to be an effective and safe adjunctive treatment of primary pterygium.


Pterygium is a triangular sheet of fibrovascular tissue that invades the clear cornea from the bulbar conjunctiva, and has a high recurrence rate after simple excision1–3 unless any adjunctive treatment is applied. The use of postoperative β irradiation renders a lower incidence of recurrence but may lead to complications such as scleral ulceration, infection, and cataract.4 Other effective treatments such as the use of topical thiotepa can cause depigmentation of the lids,5 6 and postoperative topical mitomycin C has been associated with serious ocular complications.7

We studied the efficacy and toxicity of intraoperative mitomycin C in the treatment of primary pterygium in a randomised, prospective, double blind, placebo controlled study comparing standard excision with excision and a single intraoperative application of mitomycin C.

Patients and methods

A total of 66 consecutive primary pterygia were surgically excised in 54 patients between September 1992 and August 1993. A complete ocular examination, photographic documentation of the pterygium, and haematological examination (complete cell counts with differential, sedimentation rate, electrolytes, prothrombin time, and prothrombin thromboplastin time) were performed for each patient. The inclusion criteria were (1) age older than 20 years, (2) primary pterygium which invaded more than 2 mm into the cornea. Exclusion criteria were (1) external ocular diseases such as Sjögren syndrome and ocular rosacea, (2) abnormal cell counts. The protocol was approved by the ethics and clinical trials committee of La Fe University Hospital. Informed consent was obtained from all patients.1

All the surgical excisions of the pterygium were performed on an outpatient basis by two different surgeons (MJM, EV) using the same technique under an operating microscope. The surgical technique was as follows:

1. conjunctival topical anaesthesia with 0·1% tetracaine chlorhydrate and 0·1% oxybuprocaine chlorhydrate eyedrops twice before entering the operating theatre,
2. placement of rigid lid speculum,
3. subconjunctival injection of 0·5 ml of 0·4% meipivacaine chlorhydrate into the body of the pterygium with a 25 gauge needle, 4. dissection of pterygium from cornea with Baird Parker No 15 surgical blade with partial superficial keratotomy,
5. dissection with spring action scissors onto conjunctiva and Tenon's capsule,
6. complete resection leaving 3 mm or more of bare sclera exposed with occasional light bipolar cautery of the bleeding vessels,
7. at this moment, patients were randomised in a masked fashion to receive a 5 minute scleral application of a 4×5 mm fragment of surgical sponge (k sponge No K 20-5000, Katena Products Inc, Denville, USA) soaked in a solution of 0·1 mg/ml of mitomycin C or in distilled water in the control group,
8. the site was irrigated copiously with 10 ml of a saline solution after removal of the sponge,
9. no conjunctival sutures were used,
10. instillation of topical eyedrops of 0·1% dexamethasone and gentamicin ointment followed by eye patching. Oral analgesics (paracetamol every 8 hours) were given to relieve postoperative pain.

Mitomycin C was present in a blue-violet crystalline powder that was reconstructed and diluted in sterile water just before surgery. All the pterygia were evaluated on postoperative days 1, 7, 15, and monthly thereafter by one of us (JCP) who was masked to the patient's treatment status. All patients had a repeat haematological examination on postoperative day 7, and periodic ocular photographic documentation was obtained. A total of 12 patients with bilateral pterygium were
Table 1  Demographic and surgical data of 54 patients with primary pterygium

<table>
<thead>
<tr>
<th>Treatment group</th>
<th>No of patients</th>
<th>Mean age (SD)</th>
<th>No of eyes</th>
<th>Sex (M-F) (%)</th>
<th>No of eyes with recurrence</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>at treatment (yrs)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Excision</td>
<td>29</td>
<td>50 (6)</td>
<td>36</td>
<td>22:7</td>
<td>14 (38:8)†</td>
</tr>
<tr>
<td>Excision + mitomycin C</td>
<td>25</td>
<td>54 (3)</td>
<td>30</td>
<td>20:5</td>
<td>1 (3:33)†</td>
</tr>
<tr>
<td>Total</td>
<td>54</td>
<td>51.8 (5)</td>
<td>66</td>
<td>42:12</td>
<td>15</td>
</tr>
</tbody>
</table>

*12 patients with bilateral pterygium were treated by different procedures. Each eye of these patients was entered into each group. tp=0.0006, χ² test.

Table 2  Postoperative complications

<table>
<thead>
<tr>
<th>Complication</th>
<th>Mitomycin C treated eyes</th>
<th>Control eyes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Delayed wound healing (1–2 weeks)</td>
<td>30</td>
<td>0</td>
</tr>
<tr>
<td>Conjunctival granuloma</td>
<td>5</td>
<td>14</td>
</tr>
<tr>
<td>Astigmatism (+2 D)</td>
<td>1</td>
<td>12</td>
</tr>
<tr>
<td>Superficial punctate keratitis</td>
<td>8</td>
<td>2</td>
</tr>
<tr>
<td>Anterior chamber reaction</td>
<td>2</td>
<td>0</td>
</tr>
</tbody>
</table>

These patients complained of severe discomfort. A mild anterior chamber reaction developed in the first postoperative day in two mitomycin C treated eyes (6-8%). Neither conjunctival nor corneal infection was observed in this group. All the mitomycin C treated eyes showed characteristic areas of sclera with a relatively avascular surface, however, we did not observe any sceral thinning of these eyes during the follow up. Neither significant haematological changes nor systemic toxicity was recognised in any of the treated patients with intraoperative mitomycin C. These eyes are being followed up further to study possible long term complications.

Discussion

Primary pterygium is one of the most common corneal disorders seen in our community because the inhabitants have a high exposure to ultraviolet light. This causative factor has been established as the most important risk factor in its development. Excision with the bare sclera technique as described by Ombrain is the most widely used procedure of treatment of pterygium. Nevertheless this technique is accompanied by a recurrence rate of 30–50%. Different adjunctive treatments to excision of the pterygium have been shown to diminish the recurrence rate; however, variable complications have been reported. The use of conjunctival autograft also has a recurrence rate between 5-3% and 21%. The recurrence of pterygium appears not to be associated with ultraviolet light exposure and would be due to an accelerated fibroblast proliferation produced by the trauma of operation in the same way as the production of keloid tissue.

Mitomycin C is an antineoplastic antibiotic agent isolated from the fermentation filtrate of Streptomyces caesius. Its action is similar to those of alkylation agents, alkylates and crosslinks DNA and, in addition, may generate superoxide and hydroxyl radicals in solution. It also inhibits DNA synthesis and RNA and protein synthesis. These combined effects may result in long term effects on cellular proliferation.

In a previous experimental study of the effects of mitomycin C on cultured rabbit subconjunctival fibroblasts, the authors found that its antiproliferative effect is both dose and time dependent. Other in vitro studies have shown that the effect of 5 minute exposure with mitomycin C at a concentration of 0.1 mg/ml to human Tenon’s capsule fibroblasts results in a significant long term inhibition of fibroblast proliferation. At this dose, the number of cells did not increase more than 2-5 times to the density at day 0 during a 36 day period of the experiment, despite stimulation throughout this period with 10% fetal bovine serum. In contrast, at a concentration of 1 mg/ml more than 65% of the cells died.

However, there are limitations in applying results of in vitro studies to the in vivo situation. The action of a drug over a mono-layer of cells with relatively small amounts of
extracellular matrix, and where most of the cells are proliferating may be different from the in vivo situation where cells are surrounded by connective tissue, which may impede drug penetration. Nevertheless, at the time of conjunctival surgery there are several growth factors, such as platelet derived growth factor (PDGF), that are released and might play an important role in pterygium recurrence. If the process is impaired even at this early stage, the ability of the fibroblast to proliferate as well as the capillaries to grow may be impaired.

The use of mitomycin C eyedrops in the postoperative period of pterygium surgery in both doses, 0.2 and 0.4 mg/ml, four times daily for 5–15 days has been effective in reducing the recurrence rate of pterygium between 0%–9%. However, in a recent report this postoperative treatment has been related to serious ocular complications such as secondary glaucoma, corneal edema, corneal perforation, iritis, sudden onset mature cataract, and scleral calcification. In contrast, no serious complications have been noted in another study with the postoperative use of 0.1 mg/ml mitomycin C twice daily for 5 days (with a mean follow up period of 15–3 months) or in the intraoperative mitomycin C treated eyes of our current study. A common element in toxicity with mitomycin C is a relatively large cumulative dose. Therefore, we consider that a single intraoperative exposure to mitomycin C would reduce the complication rate of mitomycin C eyedrop regimen.

Although serious complications with low concentrations of mitomycin C are rare, we noted in our study only minor complications. Delayed epithelial closure as a side effect of mitomycin C may predispose to postoperative infectious scleritis and endophthalmitis, particularly in tropical countries. None of the complications of mitomycin C treated eyes presented in Table 2 significantly troubled the patients. Moreover, conjunctival granulomas were found to be less frequent among the mitomycin C group than among the control group.

Recurrence of pterygium commonly occurs within 6 months after the initial surgery and the minimum follow up in our study has been 12 months. However, there are late complications such as scleral malacia related to the use of topical mitomycin C. Therefore, it is necessary to continue to assess the eyes treated with mitomycin C after this period.

We have shown that the single intraoperative exposure to mitomycin C (0.1 mg/ml) reduces the recurrence rate of primary pterygium without serious complications over a mean follow up of 14–1 months. We suggest that the single intraoperative exposure of mitomycin C appears to be a safe, simple, effective, and useful form of adjunctive therapy to the surgical treatment of the primary pterygium.

The authors have no financial interest in this drug.