A descriptive study of primary pterygium surgery with fibrin glue

Estudo descritivo de cirurgia de pterígio primário com adesivo de fibrina

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ABSTRACT

Objective: To evaluate the surgical technique for excision of primary pterygium with fibrin glue, as the symptoms and epidemiological data. 

Methods: Clinical prospective non-controlled and randomized study with 30 patients of the Hospital Universitário Lauro Wanderley, consistent with the term informed consent, approved by the Ethics in Research. The surgeries were evaluated at 1, 7 and 21 postoperatively (PO).

Results: Of the 30 operated patients, 16 (53.3%) were female, aged 21-67 years (42.2). As for the degrees of pterygium (1-3): grau 1-10 (33.3%), grade 2-16 (33.3%) and grade 3-10 (33.3%), 16 (53.3%) in the right eye and 16 (44.4%) in the left eye. 56.7% of patients had intense sun exposure during their lifetime and cases in the family. In the trans-operative period, there were complaints of pain 1-10 (33.3%), grade 2-16 (33.3%) and grade 3-10 (33.3%), 16 (53.3%) in the right eye and 16 (44.4%) in the left eye. 56.7% of patients (PO).

Conclusion: Autologous conjunctiva is currently the most effective alternative for the treatment of pterygium. Associated with the use of fibrin adhesive, which is composed of fibrinogen and thrombin and has the advantage of being totally absorbable, has advantages, among them a reduction in surgical time. In this study, corroborating recent studies, the mean operative time was 19.05 (± 6.1) minutes and there was a high incidence of family pterygium patients and UV exposure. Complaints of dry eye increased progressively after surgery, revealing a possible change in blink rate. Rarely addressed in the literature, the symptoms assessed in this study, showed that complaints Intraoperative mild and marked the first POD, regressed from the 7th POD, justifying the higher patient satisfaction and excellent aesthetics that using this technique.

Keywords: Pterygium; Transplantation, Autologous; Fibrin foam; Epidemiology; Local symptoms

Resumo

Objetivo: Avaliar a técnica cirúrgica de exérese de pterígio primário com adesivo de fibrina, quanto à sintomatologia e dados epidemiológicos. Métodos: Ensai clínico prospectivo, não-controlado e aleatório com 30 pacientes do Hospital Universitário Lauro Wanderley, concordantes com o termo de consentimento livre e esclarecido, aprovado pelo Comitê de Ética em Pesquisa. As cirurgias foram avaliadas nos 1º, 7º e 21º pós-operatório (PO).

Resultados: Dos 30 pacientes operados, 16 (53.3%) eram do sexo feminino, com idade variando de 21 a 67 anos (42.2). Quanto à graduação do pterígio (1 a 3): grau 1-10 (33.3%), grau 2-16 (33.3%) e grau 3-10 (33.3%), sendo 16 (53.3%) no olho direito e 14 (46.7%) no olho esquerdo. 56.7% dos pacientes afirmaram intensa exposição solar durante a vida e possuíam casos semelhantes na família. No transoperatório, houve queixas de dor (43.3%) e sensação de corpo estranho (46.7%). O tempo cirúrgico variou de 11 a 32 minutos (17.7). As queixas nos 1º, 7º e 21º PO, respectivamente, foram: dor (60%, 26.6% e 6.66%), hiperemia (93.3%, 66.6% e 36.6%), sensação de corpo estranho (53.3%, 46.6% e 20%), epífora (83.3%, 43.3% e 6.66%), secreção (33.3%, 36.6% e 6.66%), ardência (53.3%, 36.6% e 16.6%) e olho seco (6.66%, 26.6% e 23.3%). 43.3% afirmaram estar muito satisfeitos com a cirurgia e 63.8% consideraram o aspecto estético excelente. 3 (10%) pacientes não compareceram ao 7º DPO e 5 (16.6%) ao 21º DPO. Não houve perda do enxerto nos casos estudados. Conclusão: O transplante autólogo de conjuntiva é atualmente a alternativa mais eficaz para o tratamento do pterígio. Associado ao uso do adesivo de fibrina, que é composto de fibrinogênio e trombina e tem a vantagem de ser totalmente absorvível, apresenta vantagens, dentre elas a redução do tempo cirúrgico. Neste estudo, corroborando com estudos recentes, o tempo cirúrgico médio foi 19.05 (±6.1) minutos e houve alta incidência de familiares portadores de pterígio e exposição aos raios ultravioleta. A queixa de olho seco aumentou progressivamente após a cirurgia, revelando uma possível alteração do ritmo do piscar. Pouco abordada na literatura, a sintomatologia avaliada neste estudo, mostrou que as queixas levam transoperatoriamente e acentuadas no 1º DPO, regrediram a partir do 7º DPO, justificando a maior satisfação dos pacientes e o excelente aspecto estético referido com o uso desta técnica.

Descritores: Pterígio; Transplante autólogo; Espuma de fibrina; Epidemiologia; Sintomas localis

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**INTRODUCTION**

Pterygium excision associated with autologous conjunctival grafting is currently the most effective method to treat pterygium due to its low rates of recurrence(11). Sutures can be used in the procedure, usually polyglactin (Vycril™) or nylon 10.0. However, pterygium excision with sutures often causes postoperative discomfort, as despite the procedure’s high rates of success, the use of sutures increases tissue manipulation, requiring constant repositioning for adequate fixation. Furthermore, using a needle and suture produces increased tissue trauma, thus increasing the incidence of granuloma. In such situations, loose sutures need to be removed, thus increasing the chance of infection and, more importantly, the surgical time(3).

In order to minimise the adverse effects of sutures, a compound with excellent adhesiveness and functionality has been developed: the fibrinogen adhesive or biological glue, used as an alternative to skin sutures(3). Fibrin glue mimics the final reactions of the coagulation cascade, resulting in a firm fibrin clot resulting from the polymerisation of fibrinogen by thrombin. Thus, it aims to provide adhesion between tissues during surgical procedures as an alternative to sutures. Additionally, it provides several benefits in terms of tissue regeneration, reducing surgical time, the friction and tension applied to tissues, tissue necrosis, inflammation, the risk of infection, and the formation of foreign body granuloma, thus providing greater patient comfort(4,5). However, the surgeon needs to be experienced in the technique and has to handle the glue swiftly due to its fast precipitation(6).

In ophthalmology, fibrinogen adhesives have been tested in various surgical procedures, such as: amniotic membrane fixation and autologous conjunctival grafting for pterygium, limbal transplantation, lamellar keratoplasty, reconstruction of the ocular surface, glaucoma surgery, cataract surgery, phacoemulsification with scleral tunnel, corneal perforations, oculoplastic surgery, conjunctival surgery, radial keratotomy, lamellar grafting, penetrating keratoplasty, refractive surgery, and even surgery for superficial burns(5,6).

Thus, we aimed to conduct a prospective analysis of patients submitted to pterygium surgery using fibrin glue for autologous conjunctival graft fixation, assessing factors such as: surgical time, symptoms, and intra- and postoperative visual acuity; patient satisfaction and postoperative aesthetic appearance; the need for symptom-relieving medication; and the time needed to readapt to daily activities. This surgical technique is still a matter of debate due to the effective and well-established use of sutures; however, many ophthalmologists are still unfamiliar with the multiple benefits of fibrinogen adhesives(6,7).

The study’s inclusion criteria were: symptomatic patients with primary medial pterygium grade 1 (up to 2 mm), 2 (2 to 4 mm) or 3 (more than 4 mm) who voluntarily opted for surgical treatment. Exclusion criteria were: associated corneal or surface disorders, poor general health, and mental or physical inability to collaborate with the study.

Patient charts included the following information: date, name, age, sex, group, eye, start and finish time of the procedure; symptoms reported during surgery (pain, foreign body sensation, other); postoperative complaints (pain, redness, foreign body sensation, epiphora, discharge, burning, and dry eye); postoperative discomfort rated from 0 to 10 on the Visual Analogue Scale (VAS); patient satisfaction on postoperative (PO) days 1, 7 and 21, rated as very poor, poor, fair, good, and excellent; pre- and postoperative visual acuity (PRE VA, POST VA); aesthetic appearance on PO day 21; and rehabilitation time to perform daily activities. The VAS measures the intensity of ocular discomfort and is an important tool to reliably assess the patient’s condition during treatment: the patient is asked to rate their degree of ocular discomfort, with 0 being no discomfort and 10 the maximum tolerable level of discomfort.

All procedures were performed by the same surgeon. After preparation of the surgical field, asepsis and topical anaesthesia with 1% proparacaine eye drops, the operative time was measured and the following surgical steps were performed: infiltration with 2% lidocaine in the body of the pterygium; detachment of the pterygium head with a cold blade number 15; dissection and excision of the pterygium body from the limbus with conjunctival scissors; and excision of Tenon’s capsule, avoiding cauterisation whenever possible. The ipsilateral upper conjunctival graft was marked and dissected, being 2 mm larger in size than the excised conjunctiva. The two components of fibrin glue were then applied separately, with fibrinogen on the base of the sclera and thrombin on the graft’s inner surface. Then, within 1 minute, the graft was placed on the desired position, the donor limbus was positioned on the host limbus, the conjunctival epithelium was exposed, and excess glue was removed.

Postoperative assessments were conducted by an independent, previously-calibrated observer. After surgery, an occlusive dressing with an ointment of tobramycin plus dexamethasone was applied. After removal, patients were prescribed gatifloxacin plus prednisolone acetate (ZyprerX™, Allergan) eye drops, 1 drop every 6 hours until PO day 21, with lubricating eye drops (Optive™, Allergan) every 6 hours or more if necessary.

Statistical analysis was done using SPSS (Statistical Package for Social Sciences) version 18.0. The Shapiro-Wilk test was used to assess the normality of the quantitative variables. The chi-square test and the Fisher exact test were used to compare the frequency of categorical variables among the categories of the variable “grade of pterygium”. The Friedman test and the Wilcoxon test were used to compare the progression of quantitative variables over time postoperatively. The Kruskal-Wallis test and the Mann-Whitney test were used to compare the surgical times of the three grades of pterygium. Differences with p<0.05 were considered statistically significant.
RESULTS

The study sample consisted of 30 participants. Mean age was 45.2 years (SD=12.8), ranging from 22 to 67 years, with a predominance of females (53.3%). Participants were separated into groups 1, 2 and 3 according to the grade of pterygium, with 10 patients in each group. Sixteen (53.3%) patients had pterygium in the right eye and 14 (46.7%) in the left eye. As regards their personal background, 56.7% of patients reported a history of intense exposure to sunlight, and 56.7% of patients had a family history of pterygium.

The Shapiro-Wilk test was used to verify if the data to be analysed were normally distributed. In other words, if the p-value in the table below is <0.05, then the data have a normal distribution (Table 1).

<table>
<thead>
<tr>
<th>Variable</th>
<th>p-Value</th>
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<tbody>
<tr>
<td>Age</td>
<td>0.391</td>
</tr>
<tr>
<td>Time</td>
<td>0.004</td>
</tr>
<tr>
<td>PRE VA</td>
<td>0.011</td>
</tr>
<tr>
<td>POST VA</td>
<td>0.003</td>
</tr>
<tr>
<td>VAS PO 1</td>
<td>0.041</td>
</tr>
<tr>
<td>VAS PO 7</td>
<td>0.003</td>
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<tr>
<td>VAS PO 21</td>
<td>0.000</td>
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</table>

Significant differences between the three grades of pterygium were found in the variable “surgical time” (p=0.033). The average surgical times of grades 1, 2, and 3 were 14.8, 17.8, and 20.5 minutes, respectively. A significant difference was found between grades 1 and 3, with longer surgical times for grade 3 (p=0.019).

As regards intraoperative symptoms, 43.3% of patients reported pain, 46.6%, reported foreign body sensation, and 23.3% reported burning. No statistically-significant differences were found between the study groups regarding the frequency of postoperative symptoms, except for the variable “discharge” on PO day 7 (p=0.048). A higher frequency of discharge was observed in subjects with pterygium grade 3. A gradual and progressive regression of pain, redness, foreign body sensation, burning, and epiphora was observed on PO days 1, 7, and 21 (Figure 1). However, groups 2 and 3 showed an increase in complaints of dry eye over time.

Table 1

Assessing the normality of quantitative variables using the Shapiro-Wilk Test

<table>
<thead>
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<td>VAS PO 21</td>
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VAS: Visual Analogue Scale for ocular discomfort; PRE VA: Preoperative visual acuity; POST VA: Postoperative visual acuity; PO: Post-operative day.

To compare PRE VA vs. POST VA and VAS, as the data were not normally distributed, we used the nonparametric Wilcoxon and Friedman tests (PRE VA vs. POST VA) followed by the Wilcoxon test (VAS PO 1 vs. PO 7 vs. PO 21). Statistically-significant differences were found between PRE VA and POST VA for the sample as a whole (overall p=0.005) and more specifically for pterygium grade 3 (p=0.027). An improvement in POST VA in comparison with PRE VA was observed.

Figure 2 shows the analysis of the degree ocular discomfort according to VAS using the Wilcoxon test, where pterygium grades 1, 2 and 3 were assessed on PO days 1, 7, and 21. For pterygium grade 1, the mean and standard deviation were 2.7 (3.4), 1.8 (2.8), and 0.5 (0.8), respectively, with p=0.043; for grade 2, 4.2 (2.9), 2.0 (1.9), and 1.8 (1.8), p=0.028; for grade 3, 4.3 (3.1), 2.3 (2.6), and 1.7 (2.7), p=0.026. Significant differences were found between PO days 1, 7 and 21 for the sample as a whole (overall p=0.001) and more specifically for grades 2 (p=0.023) and 3 (p=0.037). For the total sample and for grade 2, significant differences were found between the PO days 1 and 7 and between days 1 and 21. For grade 3, a significant difference was found only between PO days 1 and 7. A progressive decrease in postoperative VAS scores was observed in all situations.
Fibrin monomer. The preparation of autologous tissue adhesives using fibrin glue causes less postoperative discomfort and pain (e.g. by preventing the need to remove sutures postoperatively, as by preventing the need to remove sutures postoperatively, fibrin glue causes less postoperative discomfort and pain (e.g. Figure 3). No studies were found in the literature that assess these variables.)

The topics covered in this study are part of an ongoing project which will follow-up participants for a period of 3 months using biomicroscopy and will perform a comparison with surgical procedures that used nylon 10.0 or vycril 8.0 sutures. The results of this project will be published upon its conclusion.

Figura 3: Aesthetic appearance preoperatively and on postoperative day 21.
CONCLUSION

O uso da cola de fibrina, como opção terapêutica do pterígio primário, mostra resultados satisfatórios quanto aos sintomas per e pós-operatórios e benefícios na estética, proporcionando satisfação aos pacientes já nos primeiros dias após a cirurgia, assegurando ser uma técnica rápida e eficaz.

REFERENCES