Natural latex graft in lamellar and penetrating sclerectomies in rabbits

Enxerto de látex natural na cicatrização de esclerectomias lamelar e penetrante em coelhos

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ABSTRACT

The objective of study was to investigate the effects of natural latex with 0.1% of polylysine on lamellar and penetrating sclerectomies in rabbits. Two groups of twelve rabbits each (lamellar GI and penetrating GII) were studied. Scleral square incisions near the limbus were performed on the left eye of each animal. The latex biomembrane was fixed to the recipient sclera and it covered with a conjunctival flap. The clinical evaluations were followed for 60 days. Aplanation tonometry, binocular indirect ophthalmoscopy and slit-lamp biomicroscopy were performed during evaluation. Bright field microscopy and polarization microscopy were employed. Blepharospasm, graft infection, mucoid ocular discharge and chemosis were not observed in either treatment group. The conjunctival hyperemia varied from moderate to hardly noticeable. The postoperative IOP was not statistically significant, comparing to the preoperative IOP, for GI and GII. The histopathology by polarization microscopy showed that the neoformed tissue was primarily dependent on adjacent vascularized tissues and was constituted by collagen type III. Both groups presented optimum graft adhesion to the receiving sclera. The natural latex biomembrane with 0.1% polylysine constitutes a new alternative for scleral reconstruction. Furthermore, this is a durable material, easy to obtain and manipulate.

Key words: biomaterial, graft, healing, sclera, scleroplasty.

RESUMO

Este trabalho teve como objetivo investigar os efeitos do látex natural com polilisina a 0,1% na cicatrização de esclerectomias lamelar e penetrante em coelhos. Foram estudadas dois grupos de 12 coelhos (GI - lamelar e GII - penetrante). As esclerectomias foram realizadas no olho esquerdo de cada animal. A biomembrana de látex foi fixada à esclera receptora e foi recoberta com conjuntiva bulbar. As avaliações clínicas foram realizadas durante 60 dias. Para tal, empregaram-se a tonometria de aplanação, a oftalmoscopia indireta binocular e a biomicroscopia em lâmpada de fenda. Realizou-se análise histopatológica das escleras em microscopia de luz e sob luz polarizada. Não houve blefarospasmo, secreção ocular mucóide, quemose ou sinais de infecção do enxerto em ambos os grupos. A hiperemia conjuntival variou de moderada a tênue. Não houve diferença estatisticamente significativa entre a pressão intraocular do período pós-operatório e do período pré-operatório, para o GI e o GII. A histopatologia por microsopia de polarização demonstrou que o tecido neoformado foi oriundo de tecidos vascularizados adjacentes e constituído por colágeno do tipo III. Os dois grupos apresentaram ótima adesão do enxerto de látex à esclera receptora. A biomembrana de látex natural com polilisina a 0,1% representa uma alternativa para a reconstrução escleral, além de apresentar fácil obtenção, manuseio e durabilidade.

Palavras-chave: biomaterial, enxerto, cicatrização, esclera, escleroplastia.
INTRODUCTION

A new biomaterial (Biocure, Pele Nova Biotecnologia S/A, Ribeirão Preto, Brazil – PCT/BR 97/00054) was developed at the Ribeirão Preto Medicine College of the University of São Paulo, using natural latex extracted from the *Hevea brasiliensis* rubber tree. The membrane was obtained through a cold polarization of the latex constituents. In experimental studies, latex biomembrane showed to be useful for repairing defects of the esophagus, peritoneum, pericardium (SADER et al., 2000), bulbar conjunctiva (PINHO et al., 2004) and dental alveolus (BALABANIAN et al., 2006). It was successfully used for the reconstruction of the tympanic membrane perforation (OLIVEIRA et al., 2003) and for the treatment of chronic phlebopatic cutaneous ulcers (POTÉRIO-FILHO et al., 2005).

Disorders of the sclera constitute a group of congenital and acquired diseases. Most of them result in scleral stretching or necrosis and exhibition of the underlying uveal tract leading to surgical treatment (GREEN & BRANNEN, 1995; SKOROBOHACH & HENDRIX, 2003; TI & TAN, 2003).

In 1934, Van der Hoeve (NGUYEN & FOSTER, 1999) first recommended the use of grafts in *scleromalacia perforans* in human, using mucous membranes. Several biological or synthetic materials were studied in the scleroplasties: allogenous sclera (CARPINETO et al., 2002; GOPAL et al., 2003; ÖZCAN et al., 2005), autogenous sclera (MA et al., 2002), corneosclera (SRIDHAR et al., 2002), allogenous corneal lamellar (TI & TAN, 2003; ÖZCAN et al., 2005), fresh cadaveric limbus (SONY et al., 2005), amniotic membrane grafting (TI & TAN, 2003; MA et al., 2002; SRIDHAR et al., 2002; FAIRBANKS et al., 2003), fascia lata (ÖZCAN et al., 2005), elastic layer of the aorta (KANIUKOV et al., 1999), autogenous perioisteum (NGUYEN & FOSTER, 1999), autogenous buccal mucosa and conjunctiva (MURDOCH et al., 1997), dura mater (TI & TAN, 2003; KANIUKOV et al., 1999), auricular cartilage (GREEN & BRANNEN, 1995; NGUYEN & FOSTER, 1999) and autogenous dermal grafts (NGUYEN & FOSTER, 1999).

In animals, several tissues were successfully used: porcine small intestinal submucosa (LEWIN, 1999), autogenous temporalis fascia (SKOROBOHACH & HENDRIX, 2003), equine pericardium (BARROS et al., 1996) and equine renal capsule (ANDRADE et al., 2004) and frozen canine amniotic membrane (BARROS et al., 2005). There are reports of the use of synthetic grafts, as the polytetrafluoroethylene (Gore-Tex™ Soft Tissue Patch), the polymethylacrylate (Miragel™) and the polyethylene terephthalate (Dracon™) (WHITMORE et al., 1990). The scleral graft should be accompanied by conjunctival flap, because it provides vascularization, avoids epithelial loss, maintains the graft integrity and acts in the prophylaxis of the infection (LEWIN, 1999; TI & TAN, 2003; ÖZCAN et al., 2005).

In this study we investigated the healing of the sclera in rabbits presenting experimental lamellar and penetrating lesions using the natural latex biomembrane with 0.1% polylysine, according to bright field microscopy and polarization microscopy.

MATERIALS AND METHODS

Twenty-four adult male and female albino New Zealand rabbits were used. They were submitted to slit-lamp biomicroscopy (Portable Slit Lamp/SL-14, Kowa), indirect ophthalmoscopy (Heine – Omega 180, Heine), aplation tonometry (TonoPen XL, Mentor O & O), and fluorescein test (Fluorescein Strips, Ophthalmos Ltda). The rabbits were divided into two groups of twelve members each. Healing events were investigated in the lamellar scleroplasties with natural latex biomembrane in GI and in the penetrating scleroplasties in GII.

The protocol used for anesthesia was acepromazine (0.5mg/kg SC - Acepran 1%, Univet), xylazine (1.0mg kg⁻¹ - Rompun, Bayer) and tiletamine / zolazepam (15mg kg⁻¹ 1M - Zoletil, Virbac). Scleroplasty was performed on the left eye. Tetracaine and epinephrine eyedrops (Colírio Anestésico, Allergan) were administered before the preparation of the surgical bed.

The microsurgical procedures were performed with the aid of an operating microscope (Surgical Microscope MC – M 900, DF Vasconcellos) of 16X magnification. Bulbar conjunctiva was cut with scissors and retracted for scleral beds exposure. Scleral square incisions (lamellar – GI and penetrating - GII) of 4.0 x 4.0mm were performed in a 9-12 clockwise position, near the limbus. The depth of the lamellar sclerectomy corresponded to half of the stromal thickness (Figure 1A). In the penetrating sclerectomy, the scleral defect was removed up to the lamina fusca (Figure 1B).

Units of natural latex biomembrane with 4.0 x 4.0mm and 0.5mm thick and sterilized in ethylene oxide were used. The biomembrane was fixed to the recipient sclera using an interrupted pattern with 9-0 mononylon (Mononylon 9-0, Cirumédica) (Figure 1C), and then it was covered with conjunctival flap sutured in the

periphery of the limbus and in the adjacent conjunctiva (Figure 1D). The conjunctival flap was kept during the whole experiment.

After the surgery, the rabbits were treated with flunixin meglumine (1mg kg⁻¹ SC - Banamine, Schering-Plough Veterinária) and meperidine (12mg kg⁻¹ SC - Dolosal, Cristália), twice a day for the first three postoperative days. Topically, flurbiprofen (Ocufen, Allergan) and tobramicin (Tobrex, Allergan) eyedrops were applied twice a day for seven days.

Clinical evaluations started on the first day of the postoperative period and were performed at 24-h intervals for 60 days, using slit-lamp biomicroscopy in dark room. The evaluated ocular signs were blepharospasm, ocular discharge, chemosis, conjunctival hyperemia, reflex uveitis, as well as the viability and the time length of the graft.

The intraocular pressure was measured by applanation tonometry in left eye on the preoperative day and on the 1st, 3rd, 7th, 12th, 15th, 22nd, 30th, 35th, 45th, 55th and 60th days of the postoperative period. The results were submitted to variability tests in order to verify if there were differences between the groups and throughout the days, with a significance level of 5%. The Dunnett test was used to compare the differences between the means and the preoperative and postoperative IOP.

Indirect ophthalmoscopy was performed in the operated eye after induction of mydriasis with tropicamide eyedrops (Mydriacyl, Allergan) on the 3rd, 7th, 15th, 30th, 45th and 60th postoperative days.

Two animals of each group were evaluated to histopathology for each period (1st, 3rd, 7th, 15th, 30th and 60th days). The samples were stained with either hematoxylin-eosin, Masson trichrome or picrosirius.

RESULTS

Clinical evaluation

Conjunctival hyperemia was moderate on the 1st day, hardly noticeable by the 5th day, becoming tenuous on the 30th day in both groups. Mucoïd ocular discharge was observed in only two animals of each group, being moderate on the 1st day and modest on

the 3rd day. Chemosis was modest on the 1st day, fading completely on the 5th day in two rabbits of each group. Flare and miosis were not observed in any of the groups.

On the 3rd day, two rabbits of GII presented a small dimension clot, permeating the superior medial portion of the vitreous, close to the penetrating sclerectomy. On the 15th day, there was a decrease in the size of the clots in both animals and by the day 30th, they were permeating the inferior medial of the vitreous. On the day 60th, they were no longer observed. Regarding intraocular pressure there was no significant difference between the groups (P>0.05) and throughout the days (P>0.05) as shown in table1.

Histopathology

In order to better describe the results, the area on the graft in both groups was denominated “conjunctival face”. The area below the graft was denominated “scleral face” in GI, and “choroidal face” in GII.

Hematoxilin and eosin

In both groups, the inflammation was predominantly polymorphonuclear on the 1st day in the conjunctival and scleral faces (GI) and choroidal face (GII), and also between the choroid and the sclera. From the 3rd day on, there was a decrease of the polymorphonuclear infiltrate. Mononuclear cells became evident on the 3rd day, increasing gradually through the end of the evaluation. On the 30th day, the inflammation became chronic granulomatous. These inflammatory cells, originated from the uvea and the adjacent episclera, were observed in the choroidal and conjunctival faces of the graft. In some areas, they crowded around many fibroblasts and few collagen fibrils, forming granulomas in both faces of the graft. In GI, on the 60th day, the chronic granulomatous inflammation persisted moderately, including the biomembrane (Figure 2A). Giant cells were observed around the graft from the 7th day on in GI and from the 15th day on in GII, increasing on the 60th day.

In both groups, the blood vessels of the choroid were intensely swollen on the 1st day, fading on the 15th day in GI and remaining up to the 60th day in GII. In five animals of GII, retinal edema and sub retinal hemorrhage focuses were observed in every period of the evaluation, except on the 60th day.

In the GII the fibroblastic migration to the lesion bed, in the scleral and conjunctival faces began on the 3rd day of the evaluation, formed by a row of cells. On the 7th day, in the peripheries of the graft and adjacent to the borders of the lesion, fibroblasts were found in moderate amount, migrating to both faces of the graft. The neoformed tissue started to involve the whole graft on the 15th day of the evaluation and on the 30th day it became thicker. In the neoformed tissue of both faces of the graft, neovessels were observed firstly on the 15th day in the periphery of the grafts, reaching their largest values on the 60th day.

As the sclera was absent due to penetrating sclerectomy, the fibroblasts migrated from the choroid and the episclera (Figure 2B) to the choroidal and conjunctival faces, respectively, through a fibrin net. At the end of the evaluation, a great number of fibroblasts, few fibrocytes and great amount of collagen fibrils were observed, characterizing the maturation of the healing tissue (texto removido). Neovessels increased from the 7th day throughout the end of the evaluation.

Masson trichrome

Fibroblastic migration (Figure 2C) and the neoformation tissue were observed as for the hematoxilin-eosin. Fibroblasts were predominant compared to collagen fibrils in both faces of the graft in GI (texto removido). Collagen fibrils, marked in light blue, permeated the fibroblasts (texto removido). In GII, on the 30th and 60th days, neoformed tissue with a larger amount of collagen fibrils was observed in the choroidal face of the graft (Figure 2D). Neoformed vessels were observed in both groups.

On the 60th day of the evaluation, the neoformed tissue showed to be more mature in both groups. In GI, a larger amount of collagen fibrils and

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Significant difference (P<0.05) is indicated with *
Natural latex graft in lamellar and penetrating sclerectomies in rabbits.


A few fibroblasts were observed in the scleral face of the graft (Figure 2C). In the choroidal face of the graft of GII, however, neoformed tissue with young collagen fibrils was observed in light blue, and also mature collagen in dark blue, exhibiting organization.

Picosirius

It was observed that the neoformed tissue of both groups has green pigments, from the initial periods of the evaluation, increasing gradually in thickness in the following healing phases. The green
color suggests tissue formed by type III collagen fibers (Figures 2E). In GI, on the 60th day of evaluation, it was particularly observed that the neoformed tissue presented type I collagen fibers (Figure 2F).

**DISCUSSION**

In terms of convenience and availability, the synthetic materials constitute excellent alternatives to biological grafts. Moreover, the donor collagen can be involved with the sclera disease events, being digested by enzymes or being substituted for receiver’s fibroblasts, leading to scleral weakness.

The choice for the study of biomembrane with polylsine is due to the results obtained in previous researches (POTÉRIO-FILHO et al., 1999). The property of promoting angiogenesis and the capacity of promoting cellular adhesion and formation of extracellular matrix are relevant for the process. Their physical properties, such as elasticity and molding, are also useful in reparative surgery, besides allowing sterilization in ethylene oxide and being easily obtained (FRADE et al., 2001; OLIVEIRA et al., 2003). It was observed that the addition of 0.1% amino acid polylsine strengthened the healing effects when applied to native tissue.

This study is based on positive results of conjunctival flap in scleroplasties. Several studies show that it promotes vascularization of the area, maintains the integrity of the graft and acts in the prophylaxis of epithelial losses or infections (LEWIN, 1999; TI & TAN, 2003).

Blepharospasm was not observed in any of the groups, due to the attention paid to the conjunctival flap, which protected the suture and the areas of the sclera bare (ANDRADE et al., 2004).

The conjunctival hyperemia, more intensely manifested in the initial stages in both groups, is admitted to be a result of the surgical trauma. Unlikely, some severe ocular inflammation has already been found, characterized by blepharospasm and accentuated conjunctival hyperemia, by the use of synthetic graft of polytetrafluoroethylene (HUANG et al., 1994). The researchers observed suture dehiscence and poor adhesion between graft and surrounding tissue. It is believed that such intercurrences were a result of the non-use of conjunctival covering.

Chemosis was observed in some animals, especially in those of the group submitted to the penetrating scleroplasty. Supposedly, this is the result of manipulation and of direct surgical trauma on the ocular surface, which were certainly superior in that group.

Clinical signs of reflex uveitis such as flare, miosis and iris edema were expected, mainly in GI. However, those signs were not observed in neither of the two groups in despite of the histopathology had showed blood vessels of the choroid intensely swollen on GI and GII. In addition, retinal edema and sub retinal hemorrhage focuses were observed in every period of the evaluation, except on the 60th day in five animals of GI. Certainly, the manipulation in GI justify these histopathological alterations.

In regard to intraocular pressure, data show that the surgical trauma in both groups was not enough to produce uveitis. In the GI, the biomembrane acted quickly as a stamp in the sclera, resisted the intraocular pressure and was sufficiently rigid to avoid distortion of the eye.

It is known that the healing of the sclera depends, excessively, on vascularized adjacent tissue, as the episclera and the uveal tract (DUKE-ELDER & LEIGH, 1977). Such discoveries were confirmed by Masson trichrome stain.

The latex biomembrane, in both groups, was involved by a conjunctive-vascular capsule. Fibroblasts, collagen fibers and giant cells did not invade the graft, which acted as a support so that the healing tissue could develop around it. The fact of the graft been encapsulated by neoformed tissue instead of incorporated, had already been reported (WHITMORE et al., 1990; WILKIE & WOLF, 1991; PEZZUTTI et al., 2001) in those scleroplasties in which polytetrafluoroethylene was used.

In the present research, blood vessels were observed in the neoformed tissue. These results were in accordance with the indication that the latex biomembrane is capable of stimulating the angiogenesis of the receiver’s tissue, as it had already been pointed out (OLIVEIRA et al., 2003; PINHO et al., 2004). Moreover, it has already been admitted the interference of a vascular growth factor present in the rubber tree *Hevea brasiliensis*, which would be the inductor of the angiogenesis (OLIVEIRA et al., 2003). It is believed that the fact of been craft made by a simple polymerization method, might have preserved the biological activity of angiogenic substances.

In this study, strong birefringency of the collagen fibers was evident when the picrosirius method was used. The results were in accordance with other researchers’ results, who studied the kinetics of the fibroplasia for the distinction among the types of collagen (JUNQUEIRA et al., 1979). It is admitted that in the neoformed tissue, initially, type III collagen fibers prevailed, although with the evolution of the repairing, type I started to prevail. Predominance of types I and III collagen fibers in sclera had already been notified (WATSON & YOUNG, 2004).
The results obtained through this study agree with other researchers’ results regarding the advantages of using latex biomembrane grafts. The utilization of these grafts in scleral defects deserves attention and it should be advisable to conduct another study with a longer follow up in order to observe the position, size and life span of the latex biomembrane in the sclera.

CONCLUSION

This research showed that the natural latex biomembrane maintains properties that induce to a fast and regular healing. Among these properties the main ones are cellular adhesion that is processed on its surface; induction of angiogenesis and formation of extracellular matrix and good adherence that maintains the native tissue, serving as support for the growth of fibrovascular tissue, without infection signs or rejection.

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