ABSTRACT.- Milani J.F., Barros P.S.M., Guerra J.L. & Brooks D.E. 2008. Effects of topical 0.2% Cyclosporine A on corneal neovascularization induced by xenologous amniotic membrane implantation into a corneal stroma micropocket of rats. Pesquisa Veterinária Brasileira 28(8):379-386. Laboratório de Investigação em Oftalmologia Comparada, Faculdade de Medicina Veterinária e Zootecnia, Universidade de São Paulo, Av. Prof. Orlando Marques de Paiva 87, São Paulo, SP 05508-900, Brazil. E-mail: pauloeye@usp.br

The objective of the study was to evaluate the topical effects of 0.2% Cyclosporine A (CsA) on corneal neovascularization of rats following surgical implantation of equine amniotic membrane into a corneal stroma micropocket. The implantation of xenologous amniotic membrane was performed bilaterally in 90 rats. In the same day of the surgery each right eye started receiving topical CsA twice a day. The left eye received no medication and served as a control. The evaluation of corneal neovascularization was performed by computerized image analysis and histopathological evaluation at 1, 3, 7, 15, 30 and 60 days postoperatively. For the image analysis 10 animals were used per time period, and for the histopathological examination, five animals were used per time period. Image analysis found that corneal neovascularization began on the 3rd postoperative day, reached its peak on the 7th day, and then progressively and rapidly decreased. Statistic analysis indicated that neovascularization of the CsA treated eye on the 7th day was significantly higher than that observed in untreated eyes. On the 30th day, however, this pattern was reversed with the neovascularization observed in the CsA treated eyes declining to the low levels observed on the 3rd day. The degree of neovascularization on the 30th day declined to the baseline level obtained on day 3 at the 60th day. Histopathological analysis indicated that deposition of collagen in the implanted tissue was completed by the 15th day. Therefore, we concluded that (1) equine amniotic membrane in rat corneal stroma produced an intense neovascularization until the 15th day postoperatively and then regressed, (2) deposition of collagen of the implanted tissue was completed on the 15th day postoperatively, and (3) use of CsA was associated with increase in the corneal neovascularization initially, followed by a quick and intense regression.

INDEX TERMS: Amniotic membrane, cyclosporine, cornea, veterinary, ophthalmology.
CsA iniciou-se no mesmo dia da cirurgia, nos olhos direitos dos animais, duas vezes ao dia. Os olhos esquerdos não receberam nenhum tratamento e serviram de controle. A avaliação da neovascularização corneana foi feita por análise de imagem computadorizada e por exame histopatológico aos dias 1, 3, 7, 15, 30 e 60 de pós-cirúrgico. Para a análise de imagem foram utilizados 10 animais por período, e para o exame histopatológico, 5 por período. A análise de imagem demonstrou que a neovascularização iniciou-se no 3° dia pós-cirúrgico, alcançou seu pico no 7° dia e então regrediu rápida e progressivamente até o 60° dia. A análise estatística indicou que a neovascularização no 7° dia nos olhos tratados com CsA foi significativamente mais acentuada do que aquela observada nos olhos não tratados. Entretanto, no 30° dia deste fato se reverteu, e a neovascularização observada nos olhos tratados com CsA diminuiu a níveis baixos comparáveis àquela do 3° dia. Já nos olhos não tratados, o grau de neovascularização somente pode ser comparado ao nível básico encontrado no 3° dia aos 60 dias de pós-operatório. A análise histopatológica demonstrou que a deposição de colágeno no tecido implantado se completou no 15° dia. Desta maneira, foi possível concluir que (1) a membrana amniótica em estroma corneano de ratos produz intensa neovascularização até o 15° dia de pós-operatório com posterior regressão, (2) a deposição de colágeno do tecido implantado foi completa ao 15° dia de pós-operatório, e que (3) o uso de CsA esteve associado com aumento inicial da neovascularização corneana, seguido de rápida e intensa regressão.

TERMOS DE INDEXAÇÃO: Membrana amniótica, ciclosporina, córnea, veterinária, oftalmologia.

INTRODUCTION

Biologic membranes grafts such as amnion, pericardium, renal capsule, and conjunctiva are being used for the repair of corneal diseases such as deep ulcerative keratitis, descemécone, staphyloma, iris prolapse, or after excision of cornea, sclera or limbal tumors. The problem with such membranes is that they induce varying amounts of neovascularization and/or opacity of the cornea to result in decreased vision of the patient (Hakanson & Merideth 1986, Barros et al. 1995, Andrade et al. 1999, Godoy et al. 2002).

The angiogenic producing capacity of biologic membranes can vary. A study comparing xenogenous amniotic membrane with xenogenous pericardium using micropocket implants in rat corneal stroma demonstrated that the pericardium induced more corneal neovascularization than amniotic membrane (Safatle et al. 2002). Another study showed that angiogenesis was effectively retarded by cyclosporine in rat models of corneal angiogenesis induced by xenotransplantation and/or by chemical cautereization (Benelli et al. 1997). Many drugs have been used to suppress angiogenesis such as cytokine inhibitors interleukin-1 receptor antagonist (Coxon et al. 2002), FGF inhibitors as rapamycin (Kwon & Kim 2006), FGF-2 inhibitors as hypercin (Lavie et al. 2005), doxycycline and triancinolone (Riazi-Esfahani et al. 2006), anti-VGEF antibodies as ranibizumab (Lucentis®) (Heier et al. 2006), pegaptanib (Macugen®) (Macugen Diabetic Retinopathy Group 2006), bevacizumab (Avastin®) (Manzano et al. 2006, Barros & Belfort Jr 2007, Hosseini & Nejabat 2007) or VEGF trap drugs (Liu et al. 2006).

This study evaluated the topical effect of 0.2% cyclosporine A by histopathological and computerized image analysis on corneal neovascularization of rats with implantation of equine amniotic membrane in a micropocket of corneal stroma.

MATERIALS AND METHODS

Animals. Ninety female, adult, healthy Wistar rats without any corneal injury with an average weight of 200g were used. The animals were kept in appropriate boxes and wood shavings, and fed with *ad libitum* water and animal feed.

Anesthesia. Animals were anaesthetized by intraperitoneal injection of 0.1ml/100g of ketamine chloral hydrate (50mg/ml) and xylazine chloral hydrate, associated at the rate of 2:1.

Implant of the amniotic membrane. Equine amniotic membrane, stored in 98% glycerin was used. The implant was hydrated with a solution of sodium chloride 0.9% for 15 minutes prior to implantation. With the animal anaesthetized and in lateral recumbency, an operating microscope with a number 11 needle without a bevel was used to perform a longitudinal incision about 0.3mm from the limbus followed by dissection of the corneal stroma towards the center of the cornea. A microcavity or pocket was formed between the collagen layers up to ¾ of the cornea radius. Fragments (1mm X 1mm) of amniotic membrane were then inserted inside each micropocket in both eyes. This pocket-surgery has been previously proved not to cause any significant neovascularization alone (Safatle et al. 2002).

Treatment with 0.2% Cyclosporine A (CsA). Right after the implantation of the amniotic membrane in the rat corneas, the treatment with 0.2% Cyclosporine A (ointment presentation - Optimmune®/Shering-Plough) was initiated twice a day in right eyes only (the left eyes serving as controls).

Filling of the vascular bundle with Nankeen Ink. Animals were deeply anaesthetized on the 1st, 3rd, 7th, 15th, 30th and 60th day postoperatively (10 animals per period). The animals were catheterized and given intravenous (femoral vein) heparin (500 UI). Five minutes later, the thoracic region skin was incised and the thorax was opened through a mediastinal incision. Animals were then euthanized by exsanguination through an incision in the left ventricle, and a cannula was put in the brachiocephalic trunk to slowly infuse 10ml of Nankeen Ink (Faber-Castell®) in order to fill the vasculature. The eyes were enucleated and fixed in Formaldehyde 10%.

Dehydration and diaphanous rendering (Spalthölz Method). After 48 hours, the eyes were washed for one hour in running water, dehydrated in crescent alcoholic solutions (50°GL, 60°GL, 70°GL, 80°GL, 95°GL and 100%, 24 hours in each solution), rendered diaphanous in two passages of 24 hours each of benzol, and preserved in a solution of methyl salicylate and benzyl benzoate (mixed at the rate of 5:3). The corneas were then removed from the eyes, divided into two equal parts, and put between two microscope glass slides fixed by staples (paper staple of grampomol type), in such a way 48 hours later the corneas were totally widened and could be fixed between a microscope glass slide and a microscope covers lip with resinoid.
Quantifying of corneal neovascularization by computerized image analysis. The neovascularization density in a defined area was used to quantify the neovascularization of the corneas after diaphanous rendering. The area utilized was a rectangle of defined area \((x)\) in the most vascularized region between the limbus and the implant, as close as possible to the limbus. To choose the most vascularized area a computer system (Bio Scan Optimas®) indicated the degrees of vascularization with different colors. The software also gave the area of vascularization inside the rectangle \((y)\), which was different for each sample. The neovascularization density for each sample was obtained by dividing \((y)\) by \((x)\). The rectangular area \((x)\) was kept constant for all samples.

Statistic analysis. Statistic analysis was performed at a 5% level of significance by means of four comparisons of the neovascularization: analysis of the averages and their standard deviations; analysis of variances with repeated measures; Bonferroni’s and Tukey-Kramer’s multiple comparisons between the eyes; and Bonferroni’s and Tukey-Kramer’s multiple comparisons between the periods.

Histopathological analysis. We analyzed corneas from the eyes of 1st, 3rd, 7th, 15th, 30th and 60th days postoperatively (five per period). For this, the eyes were preserved in formaldehyde 10% for 24 hours. Afterwards the corneas were removed (preserving the cornea-sclera region) and sectioned into four parts. The parts with the implant were dehydrated in alcoholic solutions of crescent concentrations, included in paraffin and sectioned into a microtome (5mm of thickness). Thereafter, the fragments were stained with haematoxylin-eosin (HE) and analyzed under a light microscope.

RESULTS

Computerized image analysis

Computer image analysis indicated that there was obvious neovascularization beginning the 3rd day postoperatively, reaching its peak on the 7th day postoperatively, and then decreasing until the 15th day postoperatively (Fig.1, 2). A progressive decrease until
the 60th day postoperatively was later observed (Fig.3, 4). Such analysis did not indicate that there was an individual difference in neovascularization between treated and untreated eyes.

Statistical analysis

Analysis of averages and their standard deviations indicated there was more obvious neovascularization the 3rd day postoperatively and reaching its peak in treated and no treated groups on the 7th day postoperatively. The neovascularization decreased rapidly until the 15th day postoperatively, and then more regressed until the 60th day postoperatively. Analysis of variances with repeated measures indicated that there was a significant difference in the interaction term (p=0.0067), i.e., neovascularization densities vary in a different way between treated and untreated eyes in each period.

Bonferroni's and Tukey-Kramer's multiple comparisons between eyes found that the variation indicated in analysis of variance occurred on the 7th day, and that treated eyes showed more neovascularization than untreated eyes.

Bonferroni's and Tukey-Kramer's multiple comparisons between periods were performed for both treated and untreated eyes, and indicated:

**Treated eye.** There was neovascularization equivalence among days 1, 3, 30 and 60, which differed from days 7 and 15, which are equivalent between themselves.

**Untreated eye.** There was neovascularization equivalence among days 1, 3 and 60, but not between day 30, which was equivalent to day 15. There was also equivalence between day 7 and 15.

Therefore, statistical analysis found that the peak of neovascularization occurred on the 7th day postoperatively, and that eyes treated with CsA had a larger peak than untreated eyes. However, on the 30th day postoperatively, such a difference was inverted, and untreated eyes showed more neovascularization than treated eyes (Fig.5).

**Histopathologic analysis**

Histopathologic analysis indicated that on the 1st day postoperatively there was intense inflammatory reaction at the limbus both in treated and in untreated eyes. On the 3rd day, there was a small amount of neovascularization in the implant region. On the 7th day, neovascularization became more obvious, and deposition of collagen in the implant begins. On the 15th day, there was total deposition of collagen in the implant area (Fig.6, 7, 8). On the 30th day, neovascularization displayed clear signs of regression and amniotic membrane becomes confounding with stroma, especially in eyes treated with CsA. On the 60th day postoperatively, there was a difference in the deposition of collagen in treated eyes compared to untreated eyes (Fig.5).
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The cornea is almost normal without neovascularization or signs of eyes implant (Fig.9).

DISCUSSION


In veterinary medicine its utility has been growing. For instance, it has been already used to repair full-thickness defects of the cornea of dogs (Barros et al. 1998), generalized keratomalacia, ankyloblepharon, and after fibrous histiocyteoma removal in two dogs and a cat (Barros et al. 2005), to reconstruct corneal surface after excision of corneolimbal squamous cell carcinomas in nine horses (Oliver et al. 2006), and to repair corneal ulceration and keratomalacia in three horses (Lassaline et al. 2005).

It has also been used in human patients with deficient production of limbal stem-cells, not solely for eye reconstruction, but also for the ex vivo culture of such cells (Tsai et al. 2000, Anderson et al. 2001, Meller et al. 2002, Ti et al. 2002).

The amniotic membrane is a thin tissue, semi-transparent with a thick and continuous basal membrane, and an avascular stromal matrix (Van Herendael et al. 1978, Modesti et al. 1984). It has a great immunological advantage, since its cells do not express leukocyte antigens of class I, and has some immunoregulatory factors rendering it less susceptible to transplant rejection (Adinolfi et al. 1982). The main advantages of its use in the reconstruction of the eye surface include: viability of the amnion, convenience and facility of its use, and facility of forming a new epithelium with decreased inflammation, vascularization and eye secretion (Tseng 2001).

However, its use leads to corneal neovascularization which has an important role in reabsorbing the implanted amniotic membrane and in deposition of stromal collagen, besides being responsible for injured cornea’s cicatrization (Rehany & Waisman 1994, Gris et al. 2002). Notwithstanding, such neovascularization also leads to undesired effects, such as loss of corneal transparency with decreased visual acuity and intense inflammatory response, and with loss of the immunological advantage and possible rejection of the transplant tissue (Rehany & Waisman 1994).

Cyclosporine A is a cyclic polypeptide from fungus Tolypocladium inflatum Gans, and presents the ability to inhibit the activation and proliferation of T-helper lymphocytes by blocking the release of lymphokines (Diasio & Lobuglio 1996). It has been systemically used since 1979 in preventing transplant and for immunological diseases (Calne et al. 1979). In ophthalmology, it is used especially for decreasing cases of rejection of corneal implant and transplant (Calatayud et al. 2001). Since its systemic use has a high cost and results in several collateral effects, and its topical use leads to concentration levels in corneal epithelium higher than when it is used systemically, ophthalmologists began to use eye drops or ointments for the application of cyclosporine (Ponn et al. 2001, Theng et al. 2002).

Our study indicated that amniotic membrane implant led to corneal neovascularization which reached its peak on the 7th day postoperatively, with a slight decrease until the 15th day and then total regression at 60 days. On the 15th day at the implant site, the corneal epithelium regenerated, and deposition of collagen in the stromal implant site was total. Topical application of CsA twice a day led to an increase in neovascularization in the
beginning of the treatment, with a quicker and more intense regression of neovascularization as of the 15th day when comparing treated to untreated eyes.

While one study detected Beta Transforming Growth Factor (TGF-β) in vascularized corneas, another study demonstrated that TGF-b is related to the tridimensional structure of capillary structures "like" in vitro. (Cursiefen et al. 2000, Darland & D’Amore 2001). Since Cyclosporine stimulates the expression of TGF-B in humans, in vivo and in vitro, this may be one of the mechanisms that led to an increase in neovascularization at the beginning of the experiment (7 days postoperatively) (Suthanthiran & Strom 1994, Khanna et al. 1997, Goodman et al. 2001, Stabellini et al. 2002).

One study evidenced that after a corneal injury, neovascularization process begins with a non-proliferative stage in the first 48 hours, followed by a proliferative stage with a high rate of vascular growth and, thereafter, with a regression stage with decrease in density of the new vessels (Edelman et al. 1999). It is hypothesized that in the first two stages there is an increase in expression of the Vascular Endothelium Growth Factor (VEGF), whilst in regression stage there is a decrease of VEGF, thus suggesting that the increase in expression of VEGF is important for inducing and maintaining the neovascularization in cornea, and that its decrease leads to a regression of neovascularization (Edelman et al. 1999). Another study demonstrated that VEGF needs activation of Nuclear Factor of Activated T-Cells (NFAT) for inducing the genetic expression of Cyclo-Oxygenase-2 (COX-2) in endothelial cells (Hernandez et al. 2001). Since CsA inhibits NFAT, the angiogenesis stimulated and maintained by VEGF also is inhibited by CsA (Chreiber & Crabtree 1992, Erlanger 1992, Mascarell & Truffa 2002). Therefore, we demonstrated the possible mechanism through which CsA stimulated regression of neovases in our study.

CsA also inhibits Interleucin-2 (IL-2) and Interferon-Gamma (INF-g) that are associated to activation of macrophages (Hingorani et al. 1999). Macrophages are directly related to the angiogenesis, liberating proteolytic enzymes responsible for the discontinuance of vascular stability (Auerbach & Auerbach 1994). Hence, the use of CsA may inhibit the activation of macrophages and decrease the neovascularization by this mechanism. Therefore, the use of Cyclosporine may be indicated in the postoperative period for implant/ transplantation of amniotic membranes in cornea in a way to decrease neovascularization led by its initial stage.

Our study concluded that equine amniotic membrane in rat corneal stroma led to an intense neovascularization until the 15th postoperative day followed by later regression; collagen deposition of the implanted tissue was entirely achieved on the 15th postoperative day, and use of CsA was linked with increase in the corneal neovascularization at first, followed by a rapid and intense regression.

However, many other factors are involved in corneal neovascularization, as well as in cyclosporine’s mechanism of action; so several studies in this area will still be necessary to elucidate the way Cyclosporine acts on transplant angiogenesis.

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REFERENCES


transforming growth factor beta modify the pattern of extracellular glycosaminoglycans without causing cytoskeletal changes in human gingival fibroblasts. Transplantation, Baltimore, 73(10):1676-1679.


