Intravitreal bevacizumab combined with infliximab in the treatment of choroidal neovascularization secondary to age-related macular degeneration: case report series

Abstract

Purpose: To evaluate the feasibility of the combined use of bevacizumab (Avastin®) and combined with infliximab (Remicade®) in the treatment of naïve choroidal neovascularization due to age-related macular degeneration eyes.

Methods: Intravitreal injections of bevacizumab combined with infliximab in 6 neovascular age-related macular degeneration eye. All patients underwent complete ophthalmologic examination on the initial visit and at days 1, 30, 60, 90, 120, 150 and 180 following the first injection. Optical coherence tomography and fluorescein angiography were performed during at initial visit and monthly during the 6 months follow-up period. Electroretinography was performed before and 30 days after initial injection, in order to evaluate retinal toxicity induced by such treatment.

Results: Thirty days after the first injection, 5 eyes (83%) shown decrease in macular thickness. No change was seen in electroretinogram in any eyes compared to initially performed electroretinogram. All phakic eyes developed cataract. One patient developed vitritis and was submitted to medical treatment successfully. At the end of the 6 months follow-up period, 4 patients showed significant improvement in the exudative process of choroidal neovascularization. One eye had mild persistent subretinal fluid without active choroidal neovascularization, and another eye had persistent amount of intraretinal fluid due to active choroidal neovascularization.

Conclusion: The combined use of bevacizumab with infliximab in eyes with neo-vascular age-related macular degeneration was effective in reducing leakage and improving the macular thickness. However, it is not possible to assert that the results were related to synergic effects of the combination therapy. A controlled study with more cases is necessary to precisely define the complication rates; however the dosage and/or association of drugs studied in this research should not be recommended in clinical practice due to inflammatory reaction.

Keywords: Retina, Macular degeneration/complications; Choroidal neovascularization/etiology; Intravitreal injections; Optical coherence tomography; Angiogenesis inhibitors/therapeutic use

Resumo

Objetivo: Avaliar a viabilidade do uso combinado do bevacizumabe (Avastin®) e do infliximabe (Remicade®) no tratamento da degeneração macular relacionada à idade neovascular em pacientes sem tratamentos prévios.

Métodos: Foram realizadas injeções intravitreas de bevacizumabe combinado com infliximabe em 6 pacientes portadores de degeneração macular relacionada à idade neovascular. Todas foram submetidas ao exame oftalmológico completo, no primeiro dia de consulta, no dia seguinte a cada injeção e mensalmente até completar seis meses após a primeira injeção. Foram realizados tomograma da coerência óptica e angiografia fluoresceínica na primeira consulta e mensalmente, até completar 6 meses após o primeiro procedimento. Electroretinografia também foi realizada antes da injeção e 30 dias após, no intuito de avaliar toxicidade retiniana.

Resultados: Ao final de 30 dias da primeira injeção, 5 (83%) pacientes apresentaram diminuição na espessura macular. Não foi visualizada alteração à eletroretinografia em relação ao exame inicial em 100% os pacientes. Cinco pacientes (100% dos fá­cios) desenvolveram catarata. Um paciente desenvolveu vitreíte e foi tratado com sucesso. Ao final dos 6 meses, 4 pacientes apresentaram melhora significativa da neovascularização de coroide, porém ainda com foco de neovascularização em atividade, um paciente apresentava discreta persistência de fluido submacular sem neovascularização ativa e 1 paciente persistia importante quantidade de fluido intraretiniano com neovascularização em atividade.

Discussão: Avaliou-se o uso combinado do bevacizumabe com infliximabe em pacientes portadores de degeneração macular relacionada à idade neovascular e a associação mostrou-se eficaz na redução do vazamento da neovascularização de coroide e da espessura macular ao tomograma de coerência óptica. Não é possível, no entanto, afirmar se os resultados apresentam efeitos sinérgicos pela associação entre as duas drogas. Um estudo com maior número de casos é necessário para definir exatamente as taxas de catarata e vitreíte da associação entre as drogas, no entanto, ao menos na dosagem estudada no presente trabalho, a associação não deveria ser recomendada na prática clínica.

Descritores: Retina; Degeneração macular/complicações; Neovascularização retiniana/etologia; Injeções intravitreas; Tomografia de coerência óptica, Inibidores da angiogênese/uso terapêutico
INTRODUCTION
Age-related macular degeneration (ARMD) is the leading cause of central vision loss in individuals over 60 years of age. In the United States, it is related to 54.4% of visual impairment cases and 22.9% of legal blindness in Caucasian population[1,2]. The neovascular form of ARMD has more aggressive and devastating effect, accounting for 80% of the cases of legal blindness by the disease[3].

The treatment for neovascular ARMD with drugs inhibiting vascular endothelial growth factor (VEGF) such as ranibizumab and bevacizumab are currently the most widely used[4,5]. The efficacy and safety of the use of ranibizumab was proven in studies such as MARINA and ANCHOR, which determined the benefit of using monthly injections of the drug. In both studies patients showed significant visual gain after 3 applications of medication[6,7].

Other studies such as PIER and PRONTO assessed alternative schedules for the use of ranibizumab involving fewer injections. Both studies showed the benefit of using ranibizumab but with results for visual acuity slightly worse than the MARINA and ANCHOR studies[8,9].

Whether ranibizumab or bevacizumab, a factor of this therapeutic modality is the necessity for multiple injections to maintain the visual benefit achieved in the first applications. The limitation of this therapeutic modality is the need for multiple injections to maintain the visual benefit achieved in the first applications.

One way to try to extend the anatomic and visual benefits would be to try to block another agent of the angiogenic pathway. An important molecule in the formation of neovascularization is the tumor necrosis factor (TNF). It is released in the early stages of the disease and participates in the cyclic inflammatory process in the formation of choroidal neovascularization (CNV)[10].

The blockage of TNF expression can be achieved through the use of monoclonal chimeric antibodies such as infliximab. This drug is used in the treatment of rheumatoid arthritis, spondyloarthropathies and Crohn's disease. Intravenous infusion of the drug in the treatment of arthritis patients who also carried the neovascular form of ARMD, led to improvement in visual acuity and regression of the CNV. Neither ocular nor extraocular effects were observed in these patients[11,12].

Giganti et al., in a pilot study evaluating the safety of intravitreal infliximab in rabbits, showed no electrophysiological or histological damage in dosage up to 1.7 mg[13]. Theodossiadis et al., also demonstrated the safety of intravitreal infliximab in New Zealand white rabbits, which received intravitreal injections of up to 2 mg of infliximab, and Rassi et al., demonstrated safety in the use of 2 mg of 2 to 3 serial applications in rabbits of the same breed[14,15].

The intravitreal use of infliximab in humans was first reported in 2009, when three selected eyes with active CNV previously and unsuccessfully treated with bevacizumab received intravitreal injections of 0.05 ml of infliximab in each eye and resulted in improvement in best-corrected visual acuity (BCVA) as well as the resolution of central macular thickness[16].

The purpose of this study is to evaluate the feasibility of the combined use of bevacizumab (Avastin®, Roche, Brazil) and infliximab (Remicade®, Schering-Plough, United States) in the treatment of neovascular ARMD.

METHODS
The study was approved by the Ethics Committee of Federal University of Goias. It was conducted as a prospective, interventional treatment study in the period from March to October 2011, where patients with neovascular ARMD were selected.

Inclusion criteria: Eyes with CNV secondary to ARMD diagnosed by means of fluorescein angiography (FA) (Topcon TRC-50DX®, Topcon, Japan) and by optical coherence tomography (Stratus - OCT®, Carl Zeiss Meditec Inc, USA); central macular thickness ≥ 300 µm by Thickness Map Report on OCT®, only one eye in each patient could be enrolled in the study; eyes should not be previously treated with antiangiogenic drugs; and the patient should provide a signed consent form.

Exclusion criteria: Eyes with cataracts or corneal opacities that precluded adequate visualization of the retina; cataract surgery performed less than three months before the possible intravitreal injection; eyes with retinal diseases other than ARMD; vitreoretinal surgery or previous treatment for neovascular AMD at any time; and refusal by the patient to participate in the study.

All intravitreal injections were performed by the same surgeon, with 30-gauge needle positioned at 3.5 mm from the limbus. Two separate injections were made with measurements at 2 different sites. After the injections, anterior chamber paracentesis was performed.

Infliximab (Remicade®, Schering-Plough, United States) is available in a bottle containing 100 mg. Its aliquotation was performed in a sterile environment (Laminar Flow AHC-2D1, ESCO, USA) using Eppendorf vials containing 2 mg of infliximab. After aliquotation, the vials were kept under refrigeration at 2-8°C.

Patients enrolled in the study underwent a complete ophthalmologic examination, on the first day of medical appointment, the day after each injection and monthly until completion of six months from the first injection.

The FA and optical coherence tomography (OCT) examinations were performed at the first visit and then monthly until completion of the six months after the first procedure. Full field electroretinography (ERG) examinations were performed in all patients before the injection and 30 days after each injection in order to evaluate the possibility of retinal toxicity. To classify the lens status, the LOCS III[17] classification system was used.

RESULTS
Six eyes (6 patients) treated with combined bevacizumab and infliximab were evaluated. Regarding gender, 5 patients (83%) were female and all of them aged over 70 years.

On biomicroscopic examination, one patient was pseudophakic and five patients had NO3 NC3 cataract.

Two consecutive injections were performed in 5 patients and 3 injections in 1 patient in the follow-up period of 6 months.

Thirty days after the first injection, 5 (83%) patients showed decrease in macular thickness. In one patient a significant presence of macular fluid was present, but the FA has shown less leakage in late phases when compared to the pre-injection (Figures 1 and 2).

Thirty days after each injection, no patient had abnormal electroretinogram findings compared to initial examination.

Two months after the first injection, two eyes developed NO4 NC4 cataract and 3 eyes developed NO5 NCS cataract, these last, it was necessary to perform phacoemulsification with implantation of intraocular lens for better retinal visualization and performance of FA and OCT exams (Table 1).

Anterior chamber cells were observed in 3 patients (50%) in the first days after each injection. One patient showed vitritis and vasculitis 1 month after the second application. Treatment was conducted, showing improvement of the retinal vascular process after 30 days (Figure 3). Treatment was performed with prednisolone eye drops 6/6h for 15 days. After 6 months, 4 eyes showed significant improvement of CNV fluid and retinal edema. However focusing on active CNV, one eye had mild persistent submacular fluid without active CNV, and another eye had persistent amount of intraretinal fluid due to active CNV.

The intraocular pressure remained within normal limits during the study (Table 1).
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DISCUSSION

There are many advances in the treatment for neovascular ARMD involving monoclonal antibodies. The results of treatments using anti-VEGF drugs have been proven through various studies, although, therapies with drugs that inhibit TNF are still being investigated.

The tumor necrosis factor is a very active pro-inflammatory cytokine and can stimulate specific and nonspecific immune systems. It activates B and T lymphocytes, and macrophages, increases the levels of pro-inflammatory cytokines such as interleukin 1, 6 and 8 (IL-1, IL-6 and IL-8), increases the attraction of neutrophils to the inflammatory...
Table 1. Clinical eye exams before and after injections

<table>
<thead>
<tr>
<th></th>
<th>BCVA Before treatment</th>
<th>BCVA 30 days after first injection</th>
<th>BCVA 90 days after first injection</th>
<th>IOP Before treatment</th>
<th>IOP 90 days after first injection</th>
<th>Anterior chamber inflammatory reaction by biomicroscopy</th>
<th>Cataract Before treatment</th>
<th>Cataract 2 months after first injection</th>
</tr>
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<tbody>
<tr>
<td>Eye 1</td>
<td>0.2</td>
<td>0.4</td>
<td>0.5</td>
<td>16</td>
<td>14</td>
<td>+</td>
<td>NO4</td>
<td>NC4</td>
</tr>
<tr>
<td>Eye 2</td>
<td>HM</td>
<td>CF 3’</td>
<td>CF 3’</td>
<td>14</td>
<td>13</td>
<td>-</td>
<td>NO4</td>
<td>NC4</td>
</tr>
<tr>
<td>Eye 3</td>
<td>0.1</td>
<td>0.1</td>
<td>0.15</td>
<td>14</td>
<td>18</td>
<td>+</td>
<td>NO5</td>
<td>NCS</td>
</tr>
<tr>
<td>Eye 4</td>
<td>CF 4’</td>
<td>CF 4’</td>
<td>CF 4’</td>
<td>14</td>
<td>12</td>
<td>-</td>
<td>NOS</td>
<td>NCS</td>
</tr>
<tr>
<td>Eye 5</td>
<td>CF 7’</td>
<td>CF 7’</td>
<td>CF 1’</td>
<td>12</td>
<td>13</td>
<td>-</td>
<td>NOS</td>
<td>NCS</td>
</tr>
<tr>
<td>Eye 6</td>
<td>CF 4’</td>
<td>CF 4’</td>
<td>CF 4’</td>
<td>13</td>
<td>12</td>
<td>+</td>
<td>Pseudophakic</td>
<td>Pseudophakic</td>
</tr>
</tbody>
</table>

BCVA= best-corrected visual acuity; IOP= intraocular pressure; HM= hand motion; CF 1’= count finger at one foot; CF 3’= count finger at three feet; CF 4’= count finger at four feet; CF 7’= count finger at seven feet.

Electrophysiology was performed before injection and at 30 and 60 days. All results were normal. For these findings it is supposed that this association may be non-toxic to the human retina controlled. Studies with more patients should be performed to improve the reliability of results.

Arias et al., reported the use of intra-vitreous infliximab (2 mg/0.05 ml) in patients with neovascular ARMD does not respond to serial injections of bevacizumab and ranibizumab. In this series of cases, reaction was observed in the anterior chamber and vitritis in 50%(17). In this study, three patients had anterior chamber reaction (1+) with flare and cells in the first postoperative day.

Some authors in 2011 evaluated the use of infliximab in rabbits (2 mg). In the study, the presence of few lymphocytes in the retina of two rabbits was seen and had an important inflammatory membrane on the surface of the vitreous and retinal posterior cortex. Despite inflammatory findings, the electroretinogram study in the rabbits was normal, as in this current study(18).

Thus, despite the findings of inflammation in the eyes treated with infliximab, it did not alter the electroretinography (ERG) suggesting no specific toxicity to the retina and its operation.

In one patient uveitis and retinal vasculitis was detected, during a visit of 30 days. The patient was treated with steroids. After 15 days of treatment, the patient had regression of angiographic signs of vasculitis.

Inflammatory reactions were also observed by Wu and colleagues in 8 patients with diabetic macular edema (20% of total patients) who underwent intravitreal injection of infliximab (2 mg)(19) similar to 4 cases reported by Giganti and colleagues who injected infliximab (0.5 mg) in 2 patients with macular edema and 2 patients with neovascular ARMD where 100% of the patients showed uveitis(20).

Both infliximab and bevacizumab are monochlonal humanized antibodies. Bevacizumab is comprised of 5% murine to 95% human framework while infliximab is 25% murine and 75% human framework.

One might suppose that the greater presence of murine components in infliximab would be an important factor for the greatest potential to generate vitritis in treated eyes, in the current series. Other anti-TNF drugs such as adalimumab are 100% humanized and is believed that it might have have fewer inflammatory components than infliximab.

Other adverse effect was observed. All phakic eyes (5 patients) had progression of cataracts in the two months following the injection and in 3 additional eyes (50% of patients) the phacoemulsification was necessary in order to provide the possibility of fundus evaluation.

In none of the animal studies and case series in humans the progression of cataract was demonstrated(12,21). As in studies with site when it stimulates the synthesis of adhesion molecules and the activation of neutrophils(16,17). Oh et al., in 1999, investigated the distribution of inflammatory mediators such as IL-1, TNF and angiogenic cytokines and as well as vascular endothelial growth factor (VEGF), in the choroidal neovascularization process. Results showed that IL-1 and TNF alpha, secreted by macrophages can promote at least in part, angiogenesis in CNV by stimulating the production of VEGF by retinal pigment epithelium (RPE) cells(18).

Studies have shown that TNF alpha plays an active role in the pathogenesis of inflammatory eye diseases and neurodegenerative eye diseases, macular edema, retinal neovascularization and proliferative vitreoretinopathy (PVR). Intravitreal injection of TNF alpha in mice causes inflammation and abnormal permeability in inner blood-retinal barrier, manifesting an inflammatory process, as well as increased levels of this cytokine in the eye(19-22).

The blockade of TNF can be obtained by use of chimeric monoclonal antibodies such as infliximab. This drug has been used to treat rheumatoid arthritis, spondyloarthopathies and Crohn’s disease. Intravenous infusion of this drug in the treatment of arthritis patients who also carried the neovascular form of ARMD, led to improvement in visual acuity and regression of choroidal membrane. No adverse ocular or extraocular effect was presented in these patients(10,13).

The safety of intravitreal infliximab has been demonstrated by studies in experimental animal models indicating that it can be safely administered at a dose of 2 mg. In these studies electrophysiological tests were performed which showed normal results in all rabbit eyes that were injected with 1 or 2 mg infliximab(14). In this current study, we selected the dose of 2 mg of infliximab to be injected into the vitreous cavity of these eyes.
anti-VEGF (MARINA, ANCHOR and CATT) there is no cataract progression. It could be presumed that the association between infliximab and bevacizumab could accelerate cataract progression.

CONCLUSION

The combination was effective in reducing the leakage of CNV and macular thickness in OCT. However, it is not possible to assert that the results are due to synergic effects of both drugs combination. Despite the small number of patients and not evidence of functional changes by the ERG, the findings of vitritis and the progression of cataracts in eyes submitted to this treatment were evident.

A controlled study with more number of eyes is necessary to precisely define the rates of cataract and vitritis development as well as therapeutic effect of the bevacizumab/infliximab association; however, due to the clinical findings observed, the dosage analysed in this study and, should not be recommended in clinical practice. By an unknown; however this hypotheses should be evaluated by futures studies designed for this purpose.

REFERENCES


