Retinal Biocompatibility of Brilliant Blue G with Deuterated Water for Chromovitrectomy

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Purpose: To investigate the retinal biocompatibility of Brilliant Blue G with deuterated water (BBG-D2O) as a vital dye for chromovitrectomy.

Methods: In this animal study, 0.05 mL of 0.25 g/L Brilliant Blue G (BBG) associated with 0.13 mL/mL of deuterium oxide (D2O) was injected intravitreally in the right eye and the same amount of balanced salt solution (BSS) was injected similarly in the left eye of rabbits. Clinical examination and histology with light microscopy were performed after seven days. Retinal cell layers were evaluated for morphologic alterations. Electroretinographic (ERG) changes were also assessed at baseline and 7 days after the injections.

Results: A total of 6 rabbits were included in the study. The gross histopathologic appearance of the retina, choroid, sclera and optic nerve was within normal limits without any sign of severe retinal necrosis or cystic degeneration. Light microscopy showed that BBG-D2O caused no substantial alterations in retinal layers as compared to control eyes. The injection of BBG-D2O did not induce considerable functional ERG alterations.

Conclusion: Intravitreal injection of BBG-D2O 0.25 g/L seems to induce no retinal toxicity as documented by lack of functional and histological changes.

Keywords: Brilliant Blue; Chromovitrectomy; Deuterated Water; Retinal Toxicity

INTRODUCTION

Chromovitrectomy refers to staining of pre-retinal membranes and tissues to facilitate their intraoperative visualization and removal.1 Indocyanine green (ICG) was initially shown to avidly stain the internal limiting membrane (ILM).2 However, numerous articles have raised concerns over retinal toxicity after intravitreal injection of ICG.3–5 Later, newer vital dyes emerged as alternatives, such as trypan blue (TB), triamcinolone acetonide (TA) and brilliant blue G (BBG).6–8 However, the blue dyes may stain epiretinal membranes (ERMs) rather than ILM elements, while TA deposits substantially in the acellular vitreous.9–13 Furthermore, the exact risk of BBG and TB retinal toxicity is not well defined but under investigation.14–22 Thus, the
best vital dye in terms of retinal compatibility and high affinity for the acellular ILM and ERM is yet to be determined.

BBG is approved for intraocular use and commercially available at a concentration of 0.25%. The dye should be injected into the fluid-filled globe and washed out immediately to prevent ocular damage. However, efforts are made to achieve higher dye concentrations directly on the surface of the ILM and prevent uncontrolled distribution of the dye within the vitreous cavity by adding heavy water (D2O, deuterium oxide) to the solution.

In the current study, BBG-D2O was tested in a rabbit model for retinal toxicity. The goal was to investigate the in vivo retinal biocompatibility of this dye injected intravitreally into rabbits eyes by clinical examination, histology using light microscopy (LM) and electroretinography (ERG).

METHODS

Dye Solutions, pH, Osmolarity, and Light Absorption Properties

Portions of 5 mg of BBG, in powder form, were weighed with an analytical balance (Mettler-Toledo Inc., Columbus, USA) and dissolved in 10 mL of the solvent, balanced salt solution (BSS), to obtain a concentration of 0.25 g/L. The mixtures were shaken for 5 minutes and sonicated (Unique Ind., Idaiautuba, Brazil) to obtain a complete solution. The pH was measured with a pH meter (Quimis Inc., Diadema, Brazil), while osmolarity was determined with an osmometer (Advanced Instruments Inc., Norwood, USA). Heavy brilliant blue G was created by adding D2O (water containing a higher-than-normal proportion of the hydrogen isotope deuterium) at a concentration of 0.13 mL/mL. Spectral absorption was measured with a spectrometer between 200 and 1,000 nm immediately after preparation (Spectronic Genesys 5 spectrophotometer, Milton Roy, Warminster, PA, USA).

Dye preparations that were well mixed and sonicated produced complete solutions for injection. The dye was prepared in a solution with BSS to achieve a concentration of 0.25 g/L associated with 0.13 mL/mL of D2O. The osmolarity of selected dyes for experiments ranged from 267 mOsm to 315 mOsm, while pH values were within 6.84 and 7.33.

Animals and Surgical Technique for Intravitreal Injection

A total of 6 male Dutch-belted rabbits, weighing 1.8 to 2.5 kg, were used in accordance with the ARVO statement, “Principles of Laboratory Animal Care” (NIH publication No. 85-23, revised 1985), the OPRR Public Health Service Policy on the Humane Care and Use of Laboratory Animals (revised 1986) and the US Animal Welfare Act, as well as university and national guidelines for research on animals.

After placing a lid speculum, a drop of 5% povidone–iodine was instilled followed by BSS washout. The vitreous cavity was accessed through the superotemporal sclera 2 mm posterior to the limbus using a 27-gauge needle connected to a 1 mL syringe containing dye or BSS. The animals were given an injection of 0.05 mL of 0.25 g/L BBG associated with 0.13 mL/mL of D2O into the right eye vitreous cavity, while the left eyes were injected with 0.05 mL of BSS (290 mOsm) serving as the control group. Eyes were immediately examined by indirect ophthalmoscopy; one drop of antibiotic–steroid solution (0.5% moxifloxacin and 0.1% dexamethasone) was then instilled in each eye.

Clinical Evaluation

Biomicroscopy and indirect ophthalmoscopy (Keeler Instruments, Broomall, USA) was performed immediately after the injection and at seven days to evaluate corneal transparency, conjunctival reaction, appearance of the lens and retina, and vitreous haze.

In vivo Retinal Functional Toxicity

ERG recordings were taken at baseline and 7 days after the intravitreal injections. The rabbits were kept in a dark room for 30 minutes and anesthetized with an intramuscular injection of
a solution containing 1 mL of ketamine (50 mg/mL) and 0.4 mL of xylazine (10 mg/mL). The pupils were dilated with 1% tropicamide eye drops and the cornea was anesthetized with 1% proparacaine drops. The rabbits were placed on a heating pad during the experiment and unipolar contact lenses with ERG jet electrodes (Universe SA, La Chaux-de-Fons, Switzerland) were placed on both corneas with 2% methylcellulose (Ophthalmos, São Paulo, Brazil).

A reference electrode filled with electrolytic gel was placed at the temporal canthus, while the ground electrode was also filled with gel and placed on the earlobe. They were then presented in a Veris System Ganzfeld stimulator (Electro-Diagnostic Imaging Inc., San Mateo, USA). After 30 minutes of dark adaptation, the procedure was performed according to two types of responses: scotopic rod response and scotopic maximal response. The responses one week after injection were compared to baseline levels and a decrease in post-injection values exceeding 50% was considered as remarkable.

Data were presented as median (range) values. Pre- and postinjection comparisons were performed using the Wilcoxon signed rank test. P-values less than 0.05 were considered as statistically significant. All analyses were performed employing Stata software version 11 (College Station, Texas, USA).

**RESULTS**

**Clinical Evaluation**

Promptly after intravitreal injection, the dye appeared in the vitreous cavity as a blue floating mass. At clinical examination immediately and 7 days after dye injection, all eyes were negative for cataracts, hemorrhage, retinal detachment and intraocular opacities.

**Morphologic Retinal Toxicity with Light Microscopy**

The histology of both BBG-injected and control eyes showed no major anatomical signs of toxicity. The histopathologic appearance of the retina, choroid and sclera was within normal limits without any sign of severe retinal necrosis or cystic degeneration.

The nerve fibre layer, retinal pigment epithelium (RPE) and choriocapillaris appeared normal after 7 days in both groups; both groups showed vacuolization and edema only in sparse regions of the retina (Figure 1).
Evaluation of Retinal Function with Electroretinography

No considerable alterations in ERG were found during the follow-up period. Median B-wave amplitude of the maximal response in the right eyes was 184.25 (range, 162-192.5) µV at baseline and 168.5 (range, 144.5-263) µV after 7 days (P=0.916). Corresponding values for the left control eyes were 164.25 (145-195.5) µV and 140.25 (71-176.5) µV, respectively (P=0.173).

DISCUSSION

The use of vital dyes in vitrectomy facilitates intraoperative surgical maneuvers such as ILM or ERM peeling. Some dyes available for chromovitrectomy such as ICG and TB may cause complications in macular surgery such as migration into the subretinal space and alterations in the RPE, and perimetric and papillary defects. In this rabbit model, the retinal biocompatibility of BBG with deuterated water was determined to evaluate the use of this dye as an alternative in chromovitrectomy. Our study showed that BBG-D2O at a dose of 0.25g/L caused no severe functional or morphologic retinal toxicity in this animal model. Another study using cell culture found that BBG-D2O is safe based on ERG values. Therefore, BBG-D2O may be useful and should be considered for application in chromovitrectomy. We used ERG B-wave amplitude as a criterion for evaluating retinal function in rabbits as it is the most relevant parameter for drug toxicity.

Compared to cell culture alone, animal models are usually considered to provide better evidence of pharmacological activity since it better simulates the human ocular and retinal environment. However, our results may not exactly correspond to the results in human chromovitrectomy due to interspecies variations. In our study, we did observe considerable anatomical or functional damage to the retinal layers. Considering the longer duration of dye exposure by the retina in the current study as compared to the shorter time of application during vitrectomy, we may assume that toxicity is even less likely during surgical procedures in humans. However one should consider the possibility of enhanced toxicity due to light exposure.

Drawbacks to our study include: (1) testing only one concentration of the dye in contrast to the large number of concentrations that can be examined in cell culture studies. (2) We did
not evaluate the affinity of the dye for the ILM. (3) Lack of cell viability analysis.

In summary, intravitreal injection of BBG-D2O did not induce significant anatomical or functional toxicity in the retina in this animal model. However this agent should be studied further to prove its capacity to stain preretinal membranes and the vitreous with tolerable toxicity. Future surgical experience will help improve the use and interpretation of new dyes in chromovitrectomy.

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Conflicts of Interest

None.

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