Comparing The Efficacy Of An Anti-Human VEGF-A Neutralizing Antibody Versus Bevacizumab On A Laser-Induced Choroidal Neovascularization (CNV) Rhesus Monkey Model

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Purpose: To evaluate the efficacy of a therapy on improving characteristics of laser-induced choroidal neovascularization (CNV) via single intravitreal injection of a humanized anti-human VEGF monoclonal antibody (PRO-169) versus bevacizumab in a rhesus monkey model.

Methods: To induce experimental CNV, small high-energy laser spots were used to treat several areas, around the macula in the retinas of monkeys at Day −21. Eighteen rhesus monkeys were used for CNV induction. The efficacy endpoints were fluorescein leakage by FFA and retinal thickness by OCT. FFA examinations were performed 19 days after induction. Appropriate animals were enrolled for treatment and randomly divided into 3 groups: bevacizumab (n=5, 7 eyes), PRO-169 (n=5, 7 eyes), and vehicle controls (n=4, 7 eyes).

Results: In 25 of 36 (69.4%) eyes, CNV lesions were identified. The average percent change of retinal thickness in the eyes of bevacizumab group was −159.3±62.2% and −154.0±45.1% (p<0.01 vs Vehicle) at Day 14 and Day 28, respectively; in the eyes of PRO-169 group it was −131.6±68.7% and −131.5±63.8% (p<0.01 vs Vehicle), respectively. The average percent change of leakage area in the eyes of bevacizumab group was −75.3±49.4% and −78.0±42.6% (p<0.01 vs Vehicle) at Day 14 and Day 28, respectively; in the eyes of PRO-169 group it was −82.0±19.3% and −81.4±21.0% (p<0.01 vs Vehicle), respectively. There were no abnormalities found in behavior, skin and hair, excretion and overall eye appearance before and after treatment in all groups.

Conclusion: After photocoagulation, the eyes enrolled in this studio showed CNV related characteristics including increased retinal thickness, and fluorescein leakage at laser spots. PRO-169 (1.25 mg per eye) can reduce the retinal thickness and fluorescein leakage area after treatment for 14 and 28 days in this rhesus monkeys model, without toxic effect or adverse events. These findings suggested that PRO-169 can inhibit CNV.

Keywords: age-related macular degeneration, monoclonal antibodies, fundus fluorescein angiography, optical coherence tomography

Introduction
Choroidal neovascularization (CNV) comprises the invasion of new blood vessels from the choroid through breaks in Bruch’s membrane into the subretinal pigment epithelial and subretinal space.1,2 The CNV leads to the severely progressive decrease of visual acuity (VA) of neovascular/wet age-related macular degeneration (AMD).3,4 AMD is a highly complex disease that affects millions of people...
worldwide and is a leading cause of irreversible blindness among the elderly in developed countries.\textsuperscript{5,6} Over the past 20 years, it has become evident that antibody-based inhibitors of vascular endothelial growth factor (VEGF) present effective treatment for AMD, leading to clinically significant improvement of VA in a substantial number of patients.

Ocular neovascularization provides an ideal in vivo system in which to investigate the mechanism that control angiogenesis.\textsuperscript{7} There are several antiangiogenic molecules that target VEGF (also called VEGF-A) and are used in the management of AMD.\textsuperscript{8–10} Monoclonal antibodies (mAbs) can be used therapeutically in ophthalmology by binding to molecular targets with high specificity.\textsuperscript{11} Monoclonal antibodies inhibiting VEGF signaling (ranibizumab, bevacizumab) have provided to be effective in neovascular ocular disorders like AMD.\textsuperscript{1,2,7–14} Ranibizumab is an anti-VEGF antibody fragment, specifically designed for ophthalmic use, was developed for intraocular administration and born from the possibility that side effects might occur in an elderly population, such as those with AMD, and from toxicity in clinical trials with systemic administration of bevacizumab.\textsuperscript{15}

PRO-169 is a humanized-anti-VEGF mAb with a molecular mass of 149 kDa. PRO-169 had a target specificity similar to bevacizumab (Avastin, Genentech).\textsuperscript{9,14} Bevacizumab is a mAb approved by the United States Food and Drugs Administration for cancer therapy and used off-label to treat neovascular AMD and other ocular disease associated with macular edema and abnormal vessel growth.\textsuperscript{10,15} In head-to-head comparisons of ranibizumab with bevacizumab, the drugs had equivalent effects on VA throughout the first year of follow-up.\textsuperscript{13,14}

Recent studies evaluating the efficacy of bevacizumab and ranibizumab in rodent models of laser-induced CNV,\textsuperscript{12,16,17} demonstrated that these mAbs anti-VEGF offer no improvement in leakage or CNV in rat and mice. This absence of efficacy is supposed to be related to the structural differences between rodent and human VEGF (species-specific).\textsuperscript{1,16,18–20} Bevacizumab is human-specific and does not react with rat VEGF because of an amino acid substitution in the bevacizumab-binding site.\textsuperscript{20} Thus, the inhibition of corneal neovascularization of mAb is far from complete in murine models.\textsuperscript{17} However, the laser-induced CNV model in non-human primates (NHPs) has played a critical role in the development of new drugs for AMD.\textsuperscript{21} Although antibodies to VEGF have been shown to inhibit neovascularization in an experimental model of laser-induced CNV in rhesus monkeys.\textsuperscript{3,21} This model uses laser, to induce CNV lesions with significant angiographic leakage,\textsuperscript{1,12} and is a reliable and reproducible model that has been widely studied and is well characterized in the literature.\textsuperscript{1,3,5} The rhesus monkey model has relevance for preclinical pharmacologic studies due to the anatomic similarities between NHPs and human eye,\textsuperscript{22} and has become a model of choice for preclinical evaluation of the efficacy and safety of VEGF inhibitors.\textsuperscript{1}

The purpose of this in vivo preclinical study was to assess the efficacy and safety of intravitreal injection of mAb PRO-169 versus bevacizumab on improving characteristics of laser-induced CNV rhesus monkey model.

Materials And Methods

Animal Experiments

All animal studies were following the Animal Welfare Act, the Guide for the Care and Use of Laboratory Animals, and the Office of Laboratory Animal Welfare; they were approved by Institutional care and use committee (IACUC) of Sichuan PriMed Shines Bio-Tech Co., Ltd.

Monoclonal Antibodies And Vehicle

Positive control: Bevacizumab (Avastin; Genentech, South San Francisco, CA, USA) is a commercially available monoclonal antibody. Test article: PRO-169 is a mAb Anti-VEGF (Laboratorios Sophia, SA de CV, Zapopan, Jalisco, Mexico by KBI Biopharma), and placebo: the vehicle was a sterile, clear solution, pH= 6.2.

Induction Of CNV And Drug Delivery

A total of 18 rhesus monkeys of Chinese origin, obtained from Primed Non-human Primate Research Center (Ya’an, China), were used for CNV induction. The groups conformed are summarized in Table 1. Monkeys weighed 3.5–4.8 kg and aged from 3 to 5 years (9 males and 9 females). Baseline fundus photography (FP), fundus fluorescein angiography (FFA), and optical coherence tomography (OCT) were performed as part of animal and ocular health screening (Day –31). The exclusion criteria were serious inflammation in eye post laser-induction and having no point of grade III or IV of fluorescence leakage in both of the eyes. The CNV technique is based on previously published methods.\textsuperscript{1,3,12} On Day –21, after animals were anesthetized, both eyes of each monkey were treated with laser at 9 spots around the macula (laser wavelength 532 nm, diameter of facula 50µm, energy 600–775 mW, time of exposure 0.1 s). The laser (Vitra 532 nm, Quantel Medical) was mounted on a slit lamp with an...
adapter and the beam directed to the retina with a Volk Centralis diameter (Volk Optical, Mentor, OH). FFA was performed 19 days later. If there was more than 1 point with fluorescence leakage of grade III or IV lesion, the eye was considered clinically relevant and modeling was assessed as successful.\(^1\)\(^,\)\(^3\)\(^,\)\(^5\) According to the area of III+IV grade of fluorescence leakage spots, appropriate animals were enrolled for treatment and randomly divided into three groups: vehicle (n=4), bevacizumab (n=5) and PRO-169 (n=5). Importantly, the initial and only injection was delivered on postoperative Day 21 for all groups. After the animal was anesthetized with intramuscular injection of 1:1 ketamine–xylazine mix (10 mg/kg), a 5% povidone iodine solution (Chengdu Yogan Pharmaceutical Co, Ltd.) was placed in the conjunctival sac, eyelid and eyelash for disinfection. A 29-gauge needle attached to a syringe containing test or control article (50 \(\mu\)L bevacizumab 1.25 mg per eye), PRO-169 1.25 mg per eye or vehicle) was used for intravitreal (IVT) injection through an area 2–3 mm posterior to the limbus, aiming toward the center of globe. A broad-spectrum antibiotic eye drop was applied to the eyes (Oculentum Erythromycin, Xinxian Huaqing Pharmaceutical Co., Ltd).

**Clinical Observation**

Each animal was observed once daily by an examiner who was blinded to the treatment groups to minimize the observer bias. Each monkey was fed with 200g diet every day (protein 17%, fat 5%, carbohydrate 63%). Food consumption was assessed semi-quantitatively, abnormal findings were recorded. Animals got access to water all the time and 2 monkeys were pair-housed every day during the study. The body weight was weighed once before induction, once before administration, and once at Day 14 and Day 28.

**Fundus Photography (FP)**

FP was conducted once before induction (Day −31); once after induction (Day −21); once before test article administration (Day −2); once 2 weeks after administration (Day 14); and once 4 weeks after administration (Day 28), see Figure 1. Animals were anesthetized with 1:1 ketamine–xylazine mix

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### Table 1 Treatment groups

<table>
<thead>
<tr>
<th>Groups*</th>
<th>Number Of Animals Enrolled</th>
<th>Number Of Eyes Enrolled</th>
<th>Dose Conc. (mg/mL)</th>
<th>Dose Volume ((\mu)L)</th>
<th>Dose Amount (mg)</th>
<th>Route Of Dosing</th>
<th>Endpoints</th>
</tr>
</thead>
<tbody>
<tr>
<td>Vehicle</td>
<td>4</td>
<td>7</td>
<td>0</td>
<td>25</td>
<td>0</td>
<td>IVT</td>
<td>FP, FFA, OCT.</td>
</tr>
<tr>
<td>Bevacizumab</td>
<td>5</td>
<td>7</td>
<td>25</td>
<td></td>
<td>1.25</td>
<td></td>
<td></td>
</tr>
<tr>
<td>PRO-169</td>
<td>5</td>
<td>7</td>
<td>25</td>
<td></td>
<td>1.25</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Note:** Refer to section “Induction of CNV and drug delivery” for more information.

**Abbreviations:** IVT, intravitreal injection; FP, fundus photography; FFA, fundus fluorescein angiography; OCT, optical coherence tomography.

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**Figure 1** Study design.

**Abbreviations:** FP, fundus photography; FFA, fluorescein fundus angiography; OCT, optical coherence tomography.
(10 mg/Kg), 2 drops of tropicamide phenylephrine eye drop (Santen Pharmaceutical Co., Ltd) were applied to each eye for pupil dilation. The monkeys were placed in a dark room until pupil diameter was greater than 6 mm. FP and angiography were performed using a retinal camera (VX-20, Kowa Company, Ltd).

**Outcome Measurement**

**Retinal Thickness By OCT**

The retinal thickness of laser spots was assessed by Optical Coherence Tomography (OCT) once before induction (Day −31); once after induction (Day −21); once after test article administration (Day −2); once 2 weeks after induction (Day 14); and once 4 weeks after administration (Day 28), see Figure 1. OCT (Spectralis OCT Plus, Heidelberg Engineering GmbH) images were acquired after FP and FFA. To ensure high-quality OCT images, the eyes were evenly illuminated, and the camera was well focused on the macular fovea. The fast-macular scan procedure was applied to the eyes. The built-in software of Heidelberg OCT was used to measure the retinal thickness by determining the distance between the inner limiting membrane and base membrane BM automatically. Occasionally, manually locating the 2 membrane lines was necessary. The location with the maximal thickness around the burn spot was chosen for measurement. At 14 and 28 days after dosing, the same OCT examination was applied. Tracking mode was applied to keep the analysis consistent.

**Amount Of Grade III And IV Spots By FFA**

Fluorescein angiography was performed once before induction (Day −31); once after induction (Day −21); once after test article administration (Day −2); once 2 weeks after administration (Day 14); and once 4 weeks after administration (Day 28), see Figure 1. Rapid series of photographs were taken every 3 s during the first 1 min after fluorescein (Alcon Laboratories, Inc) injection to ensure that enough pictures centered on macular fovea at the early stage were collected for both eyes. 5 and 10 mins after fluorescein injection, two photographs were taken for both eyes centered on macular fovea. CNV was rated on an I to IV scale, where I= no hyperfluorescence; II= lesions exhibited hyper fluorescence early or mid-transit with late leakage; III= lesions showed hyperfluorescence without leakage; IV= lesions showed bright hyperfluorescence early or mild-transit with late leakage extending beyond the borders of the burned area. Lesions were graded systematically from the 12 o’clock position in a clockwise manner. The grade III and IV of fluorescein leakage spots were set as ROIs (region of interest). The area of ROI was measured by tracing the borders of fluorescein leakage using Image J software (Wayne Rasband, Research Service Branch, NIH, Bethesda, Maryland, US) with a fixed calibration.

The data that support the findings of this study are openly available in Open Science Framework (https://osf.io) at DOI 10.17605/OSF.IO/NA63J.

**Statistical Analyses**

Statistical analyses were carried out using SPSS 19.0 software for Windows (SPSS Inc., Chicago, IL, USA). Microsoft® Office Excel 2016 was used for data processing. Data for analysis consist of animal weights, the area of fluorescein leakage, amount of fluorescein leakage spots, and retinal thickness. The data of percentage change of area of fluorescein leakage spot, percentage change of retinal thickness and change of body weight were analyzed by one-way ANOVA. Data are expressed as means ± standard deviation (SD). The Tukey’s HDS test was applied for multiple comparisons. The amount of fluorescein leakage spots was analyzed using Chi-square test. All statistical analyses performed in this study were with P values ≤ 0.05 considered statistically significant.

**Results**

**Retinal Thickness**

A highly reflective light echogenic mass was found in monkey’s eyes 19 days after laser photocoagulation, which represented the formation of CNV (red arrows in Figure 2). The retinal thickness of laser spots increased significantly. The percent change of retinal thickness of the laser spot with the maximum retinal thickness was calculated as a sign of efficacy. In the eyes of vehicle group, the average percent change of retinal thickness was −13.4±46.4% and −5.7±76.2% at Day 14 and Day 28, respectively, see Table 2. In the bevacizumab group (1.25 mg per eye), the inhibition of CNV was found (blue arrows in Figure 2). The average percent change of retinal thickness was −159.3±62.2% and 154.0±45.1% at Day 14 and Day 28, respectively (see Table 2). Compared with vehicle, the group treated with bevacizumab had a significant decrease on the retinal thickness (p=0.001). Meanwhile, in the eyes of PRO-169 group (1.25 mg per eye), the inhibition of CNV was found (blue arrows in Figure 2). The average percent change of retinal thickness was −131.6±68.7% and −131.5±63.8% at Day 14 and Day 28, respectively, see Table 2. Compared with vehicle, PRO-169 decreased the retinal thickness significantly (p=0.004). No significant differences were observed between
Figure 2 Representative optical coherence tomography (OCT) images for vehicle, bevacizumab and PRO-169 on pre-laser induction, pre-treatment, Days 14 and 28 after intravitreal injection. The high reflect light echogenic mass (red arrows) was detected in the eyes on Day 19 after laser photocoagulation (Day −2). The light echogenic mass decreases after IVT injection (Day 14 and 28) compared with pre-treatment in bevacizumab and PRO-169 groups (blue arrows).
bevacizumab and PRO-169 groups at any time point ($p=0.668$ and $p=0.784$, respectively).

## Leakage Area Of Grade III And Grade IV Spots

Fluorescein leakage was found in monkey’s eyes 19 days after photocoagulation (pre-treatment), which represented the formation of CNV (red arrows in Figure 3). The average percent change of leakage area of grade III and IV spots was calculated as a sign of efficacy. In the eyes of vehicle, the leakage area slightly decreases caused by self-healing at Day 14 and Day 28. The average percent change of leakage area ± standard deviation was $-27.6\pm35.3\%$ and $-36.1\pm34.4\%$ at Day 14 and Day 28, respectively, see Table 3. In the eyes of bevacizumab group, the leakage area reduced significantly (blue arrows in Figure 3). The average percent change of leakage area was $-75.3\pm49.4\%$ and $78.0\pm42.6\%$ at Day 14 and Day 28, respectively, see Table 3. Compared with vehicle group, Bevacizumab decreased the leakage area significantly ($p=0.025$ and $p=0.040$). PRO-169 also decreased the leakage area significantly (blue arrows in Figure 3). The average percent

### Table 2 Change Of Retinal Thickness Of Lesion Spot (%)

<table>
<thead>
<tr>
<th></th>
<th>Vehicle, n=7 Eyes</th>
<th>Bevacizumab, n=7 Eyes</th>
<th>PRO-169, n=7 Eyes</th>
<th>p</th>
</tr>
</thead>
<tbody>
<tr>
<td>Day 14</td>
<td>$-13.4\pm46.4$</td>
<td>$-159.3\pm62.2^{**}$</td>
<td>$-131.6\pm68.7^{**}$</td>
<td>0.001</td>
</tr>
<tr>
<td>Day 28</td>
<td>$-5.7\pm76.2$</td>
<td>$-154.0\pm45.1^{**}$</td>
<td>$-131.5\pm63.8^{**}$</td>
<td>0.001</td>
</tr>
</tbody>
</table>

Notes: Data are presented as % mean ± SD. One-way ANOVA, **compared with vehicle group.

Figure 3 Bevacizumab and PRO-169 attenuate laser-induced CNV. Fluorescein fundus angiography (FFA) of vehicle (Veh), bevacizumab (Bevz) and PRO-169 (IVT delivery of 1.25 mg/eye, on post-operative day 21). Images provide representative examples of grade III and IV lesions identified by red arrows (pre-treatment phase), and the reduction of leakage area by blue arrows in Day 14 and Day 28 angiograms.
Table 3 Change Of Leakage Area Of Grade III+IV Spots (%)

<table>
<thead>
<tr>
<th></th>
<th>Vehicle, n=7 Eyes</th>
<th>Bevacizumab, n=7 Eyes</th>
<th>PRO-169, n=7 Eyes</th>
<th>p</th>
</tr>
</thead>
<tbody>
<tr>
<td>Day 14</td>
<td>−27.6 ± 35.3</td>
<td>−75.3 ± 49.4*</td>
<td>−82.0 ± 19.3*</td>
<td>0.025</td>
</tr>
<tr>
<td>Day 28</td>
<td>−36.1 ± 34.4</td>
<td>−78.0 ± 42.6*</td>
<td>−81.4 ± 21.0*</td>
<td>0.040</td>
</tr>
</tbody>
</table>

Notes: Data are presented as % mean ± SD. One-way ANOVA, *compared with vehicle group.

change of leakage area was −82.0±19.3% and −81.4±21.0% at Day 14 and Day 28, respectively, see Table 3. Compared with vehicle group, PRO-169 decreased the leakage area significantly (p=0.032 and p=0.05). No significant differences were observed between bevacizumab group and PRO-169 group at any time point (p=0.937 and p=0.980, respectively).

Amount Of Grade III And Grade IV Spots
Fluorescein leakage spots were found in monkey’s eyes 19 days after laser photocoagulation (pre-treatment Figure 2). The amount of grade III and IV spots was calculated as a sign of efficacy. In the eyes of vehicle group, the amount of grade III and IV spots did not change significantly compared with pre-treatment (31 vs 30 and 28, respectively).

In the eyes of bevacizumab group, compared with vehicle, the amount of grade III and IV leakage spots reduced significantly at Day 14 (30 vs 17, $\chi^2(1)=18.632$, $p=0.0001$) and Day 28 (28 vs 17, $\chi^2(1)=13.209$, $p=0.0001$). Meanwhile, in the eyes of PRO-169 group, the amount of grade III and IV leakage spots reduced significantly compared with vehicle at Day 14 (30 vs 21, $\chi^2(1)=9.908$, $p=0.003$) and Day 28 (28 vs 21, $\chi^2(1)=5.557$, $p=0.032$). No significant differences were observed between bevacizumab group and PRO-169 group at any time point ($p=0.218$).

Clinical Observations
Cage-Side Observations
There were no significant abnormalities in behavior, autonomic activity, skin and hair, excretion and overall eye appearance before and after administration in all groups.

Body Weight
After the injection of vehicle or bevacizumab, the body weight of animals decreased at Day 14 (−1.8±2.5% vs −1.9±5.0%) and Day 28 (−4.4±3.2% vs −1.1±5.4%). However, the body weight of animals that receiving PRO-169 increased compared with vehicle on Day 28 (−4.4±3.2% vs 2.3±1.5%), $p=0.05$. No significant differences were observed in body weight between bevacizumab group and PRO-169 group at Day 14 and Day 28 ($p=0.587$ and $p=0.360$, respectively).

Discussion
The process of angiogenesis is multi-factorial and complex and VEGF is considered playing a critical in angiogenesis role.3 Ocular neovascularization provides an ideal in vivo system in which to investigate the mechanism that control angiogenesis.7 Abnormalities or defects in Brunch’s membrane induce hypoxia, oxidative stress, and inflammation, which affect the balance of antiangiogenic factors in the direction of angiogenesis.23 Levels of angiogenic factors have shown to be related to the extent of CNV in vivo and antiangiogenic molecules that target VEGF are used in the management of AMD8,11,23 AMD is a prevalent and debilitating disease with a large demand for treatments. New drugs are continuously being designed and the importance of cost-effectiveness to keep health care sustainable is growing.9 VEGF inhibitors have been shown to slow down visual loss, and improve vision in AMD patients. Bevacizumab is a mAb used off-label to treat neovascular AMD and other ocular diseases associated with macular edema and abnormal vessel growth.10,12 Bevacizumab is a cost-effective, safe treatment option for neovascular age-related macular degeneration.9 In the current study, we designed a mAb PRO-169 structurally similar to bevacizumab but specifically for ophthalmic use. PRO-169 is a recombinant, humanized-anti-VEGF that having a target specificity like bevacizumab.9,14

Animal models of an ocular disease that is like the human condition would also be similar to humans in the efficacy, potency, and duration of action of drugs. These models are easier to validate for follow-on molecules in the same class of pharmacotherapy.24 However, bevacizumab and ranibizumab do not cross-react with mouse or rat VEGF; therefore, non-transgenic rodent models cannot be used.10,16–19,25 PRO-169 is specific to human VEGF and does not bind murine VEGF. Its use for studies in mouse models is rather limited.20

The anatomic similarity between eyes of rhesus monkey and human provides theoretical advantages of NHP CNV models over other neovascularization models.21,22 Antibodies to VEGF have shown to inhibit neovascularization in an experimental model of laser-induced CNV in rhesus monkeys. The penetration of bevacizumab through the monkey retina after the intravitreal injection has been demonstrated. The penetrating volume is sufficient to induce the therapeutic effects successfully in monkeys.26
We observed laser-induced CNVs in these monkeys’ eyes are similar to previous reports in other NHPs.$^{1,3,5,12}$ Several reports have defined clinically relevant laser-induced CNV of grade III and IV lesions. However, graded scoring of fluorescein angiograms still requires a somewhat subjective assessment by examiners.$^{1,26}$

Our findings demonstrated the inhibition effect of bevacizumab on laser-induced grade III-IV CNV was similar to previous studies. FFA was performed in all animals 19 days later to assess the severity and development of CNV following the laser treatment. There were no significant differences in the lesions diameter sizes or surface area in any eye of the vehicle or treatment groups before intravitreal injections. The change percentage of retinal thickness of the laser spot with the maximum retinal thickness was calculated as a sign of efficacy. The data from our study showed that the CNV induced by laser in vehicle group, which progressed during 14 and 28 days, was reduced after intravitreal injection of PRO-169. The average change percentage of retinal thickness was $-131.6\pm68.7\%$ and $-131.5\pm63.8\%$ on Day 14 and Day 28, respectively, without significant differences between bevacizumab and PRO-169 groups at any time point. IVT injection of PRO-169 can decrease the retinal thickness at CNVs for at least 28 days significantly. The effect is similar to ranibizumab from previous studies.$^{12}$ Also, PRO-169 can decrease the average percent change of leakage area of grade III and IV spots without significant differences between bevacizumab and PRO-169. A single IVT injection of PRO-169 (1.25 mg per eye) decreased leakage throughout the follow-up period suggesting that the duration of action on vascular permeability with one injection of PRO-169 is at least 4 weeks. In agreement with previous studies, the intravitreal administration of the current clinical doses of bevacizumab and ranibizumab would result in the effective VEGF inhibition of relatively similar duration, ~30 days post-injection.$^{12,19,27}$

A limitation of this study remains the low number of animal that can be ethically used. This limitation commonly results in studies with low statistical power.$^{4}$ However, using a model with anatomic similarity to the human eye clearly compensates this limitation. We carried out this study using a sample of 21 primate eyes and encountered no adverse event in any eye. There were no significant abnormalities in behavior, skin and hair, excretion and in appearance before and after intravitreal injections in all groups. We believe that the current study provides sufficient basis that PRO-169 appears to be safe like bevacizumab in NHPs and that clinical studies may be undertaken.

In summary, after laser photocoagulation, the eyes enrolled in this study showed CNV related characteristics including increased retinal thickness, and fluorescein leakage at laser spots. Our data indicate that intravitreal injection of PRO-169 appears to be safe and can reduce the retinal thickness and fluorescein leakage for 14 and 28 days without toxic effect or adverse events. These findings suggested that PRO-169 can inhibit CNV in the same manner as bevacizumab (Avastin) for CNV.$^{4}$

In conclusion, PRO-169 treatment should be carefully evaluated in future clinical trials to confirm the safety and efficacy in the treatment of CNV. After photocoagulation, the eyes enrolled in this study showed CNV related characteristics including increased retinal thickness, and fluorescein leakage at laser spots. PRO-169 (1.25 mg per eye) can reduce the retinal thickness and fluorescein leakage area after treatment for 14 and 28 days in this rhesus monkey model, without toxic effect or adverse events. These findings suggested that PRO-169 can inhibit CNV.

Abbreviations
AMD, age-related macular degeneration; CNV, choroidal neovascularization; FFA, fluorescein fundus angiography; FP, fundus photography; NHP, non-human primate; OCT, optical coherence tomography; VEGF, vascular endothelial growth factor.

Acknowledgments
We thank Sichuan Primed Shines for providing the experimental monkeys and conducting the monkey study and for their help during the experiment. This study was sponsored by Laboratorios Sophia, SA de CV (Zapopan, Jalisco, Mexico).

Author Contributions
All authors contributed to data analysis, drafting or revising the article, gave final approval of the version to be published, and agree to be accountable for all aspects of the work.

Disclosure
Olvera-Montaño O, Baiza-Duran L and Muñoz-Villegas P are employees of Laboratorios Sophia, SA de CV. Quintana-Hau JD and Quiñonez-Alvarado MG are employees of Centro de Investigación Sophia (CIS). The authors report no other conflicts of interest in this work.
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